Probing fibronectin adsorption on chemically defined surfaces by means of single molecule force microscopy

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Atomic force microscope (AFM) based single molecule force spectroscopy (SMFS) and a quartz crystal microbalance (QCM) were respectively employed to probe interfacial characteristics of fibronectin fragment FNIII8–14 and full-length fibronectin (FN) on CH3–, OH–, COOH–, and NH2-terminated alkane-thiol self-assembled monolayers (SAMs). Force-distance curves acquired between hexahistidine-tagged FNIII8–14 immobilised on tris-NTA-Ni2+ functionalized AFM cantilevers and the OH and COOH SAM surfaces were predominantly ‘loop-like’ (76% and 94% respectively), suggesting domain unfolding and preference for ‘end-on’ oriented binding, while those generated with NH2 and CH3 SAMs were largely ‘mixed type’ (81% and 86%, respectively) commensurate with unravelling and desorption, and ‘side-on’ binding. Time-dependent binding of FN to SAM-coated QCM crystals occurred in at least two phases: initial rapid coverage over the first 5 min; and variably diminishing adsorption thereafter (5–70 min). Loading profiles and the final hydrated surface concentrations reached (~950, ~1200, ~1400, ~1500 ng cm−2 for CH3, OH, COOH and NH2 SAMs) were consistent with: space-filling ‘side-on’ orientation and unfolding on CH3 SAM; greater numbers of FN molecules arranged ‘end-on’ on OH and especially COOH SAMs; and initial ‘side-on’ contact, followed by either (1) gradual tilting to a space-saving ‘end-on’ configuration, or (2) bi-/multi-layer adsorption on NH2 SAM.

Key to the successful integration of all medical ‘biomaterials’ in human body, in contact with tissue and blood, be they synthetic or natural, is their ‘biocompatibility’ defined by Ratner1 as the ‘ability of biomaterials to: (1) locally trigger and guide wound healing and tissue regeneration; and (2) reside in the body for long periods of time with only low degrees of inflammatory reaction’. Their contact with proteins is particularly important in determining the response of the surrounding tissues, given that they in turn mediate the attachment of cells2. The conformation adopted by surface adsorbed proteins has been identified as a primary determinant of cell binding; while certain conformations promote cell binding and successful integration of the biomaterial, others inhibit it, inducing an immune response, which ultimately leads to implant rejection3,4. The form that given proteins assume on interacting with biomaterials is strongly influenced by the surface chemistry of the latter5–7. It is for this reason that medical devices are commonly modified with a coating that is chemically and biologically compatible with the intended target tissue. For example, coating a medical grade material with hydroxyapatite enhances its osseointegration, i.e. accelerates and increases bone contact with the implant surface8. Continued advance in surface coatings for biomaterials is required however; in the case of bone and dental implants to improve osseous integration by booting osteoblast adhesion9–11. The development of medical grade materials with improved biocompatibility requires comprehensive understanding of the effects of surface chemistry on
protein adsorption. Less obvious, though nevertheless important, is the need to quantify molecular interactions between differently functionalized surfaces and adsorbing proteins. With the growing availability of powerful non-destructive quantitative methods such as atomic force microscope based single-molecule force spectroscopy (SMFS)21–23, and high resolution mass sensing on a Quartz Crystal Microbalance, QCM41,42, it is now possible to analyze/measure interactions between proteins and biomaterials at the molecular level.

One of the main proteins mediating the interaction of cells with surfaces is fibronectin (FN), a major component of the extracellular matrix in many tissues. FN is a complex high molecular weight (470–500 kDa) glycoprotein dimer that binds other extracellular matrix proteins (e.g. collagen, fibrin, syndecans, tenacins), and importantly, integrin adhesion molecules, a large family of cell membrane spanning receptor proteins.16,18,19,37. The nearly identical subunits are linked by a pair of C-terminal disulfide bonds, each comprising multiple homologous modules (termed FNI, FNII and FNIII) arranged into many distinct functional and binding domains20,21. FNIII domains 9 (FNIII9) and 10 (FNIII10) are responsible for cell binding. The crucial attachment site for cell surface anchored FN receptors (e.g. integrins α,β1 and α,β3) is the RGD (Arg-Gly-Asp) sequence22,23 in FN10, while the synergy site PHSRN (Pro-His-Ser-Arg-Asn)24–26 located in FNIII9, modulates the association of FN with integrins. SMFS has been applied in the study of FN’s mechanical stability and interactions with other molecules, notably integrins. Collectively, these investigations reveal FNIII modules containing the cell-binding site as weakest in the mechanical sense27–30 and that force-induced dissociation of the complex formed between α,β1 integrin and fibronectin fragment FNIII9–10 requires overcoming two activation barriers, indicating that integrin activation involves cooperative interaction of the FNIII9 RGD and FNIII10 PHSRN synergy sites31.

SMFS has also been used, albeit rarely to date, to interrogate the interaction of FN with homogeneous chemically-defined surfaces. In a notable example, Meadows and Walker32 employed SMFS to investigate the adsorption of full-length FN supplied at two different concentrations on a wide range of defined surfaces (including mica, gold, polyethylene glycol and SAMs terminated with methyl, hydroxyl or carboxyl functions). The authors reported that at high bulk phase FN concentration, i.e. high FN challenge to the surface (aggregate state), adsorbed FN adopted a looser conformation and exhibited weaker binding on hydrophilic cf. hydrophobic surfaces, whereas at low solution phase concentration (low FN challenge to the surface, semi-aggregate state) the larger number of substrate contacts available to each adsorbed FN molecule resulted in their adopting more rigid conformations cf. the corresponding aggregate state. However, because FN was pre-adsorbed onto the target surfaces rather than anchored onto AFM cantilevers the observed interactions were non-specific in nature, with only 15–20% of the acquired data displaying pulling events. In earlier work from the same laboratory, Meadows et al.33 investigated desorption of full-length FN from a negatively charged mica surface as a function of prior FN loading; here too FN was pre-adsorbed on the test surface not attached to the AFM tip. At high FN challenge to the surface force-distance curves exhibited successive rupture events indicative of protein unfolding, while at low FN challenge FN partially denatured upon adsorption, generating force-distance curves lacking dominant pulling events.

QCM has also been used to study the adsorption of proteins including fibronectin on chemically defined surfaces. Dickerson et al.34 employed QCM to investigate the two-stage adsorption under physiological conditions of a recombinant FN-binding outer membrane protein fragment, rTp0483, on four differently terminated SAMs, followed by FN. They reported that initial binding of rTp0483 was greater on the negatively charged COOH SAM cf. all others tested, i.e. NH2, OH and CH3 SAMs (a finding consistent with the fragment’s net positive charge under experimental conditions); and further that rTp0483 adsorption was also more uniform on the negatively charged SAM cf. other surfaces (revealed by AFM). These two factors combined explained the much higher FN loadings on rTp0483 primed’ COOH SAM cf. other fragment-primed SAMs. In a different study, using COOH and NH2 SAMs blended in different ratios to control the surface chemistry of gold-coated QCM crystals, Lin et al.35 reported that despite carrying a small net negative charge (−5.7 mV) under experimental binding conditions (phosphate buffered saline) FN adsorbed on both negatively and positively charged surfaces. Interestingly, the authors’ observed greater adsorption on strongly negatively charged SAMs cf. moderately negative and weakly positive SAMs, which they attributed to a fine balance of surface charge induced polarization of the protein, short-range holding forces and simple electrostatic interactions.

In this study we have employed SMFS and QCM to respectively probe the interactions of fibronectin domain FNIII9–10 and the full-length FN molecule with the same four chemically defined surfaces (i.e. NH2-, COOH-, OH- and CH3-terminated SAMs) to derive information on unfolding and denaturation, preferential binding orientation in the adsorbed state, surface loading, and binding kinetics. Specifically, we have used SMFS to extract information on the forces developed between FNIII9–11 immobilized in various forms on gold AFM tips and the different SAMs, QCM to extract complementary information (kinetics, loading, changes in binding orientation as function of time/loading) on the binding of free FN to gold-coated QCM crystals modified with the same SAMs. Note, a full-length human FN was used in QCM studies in place of FNIII9–11, given Michael et al.36 finding of very similar adsorption of fibronectin domain FNIII9–10 and the full-length molecule and because the costs of using FNIII9–11 in QCM were prohibitive.

Materials and methods

Materials. Full-length human plasma fibronectin (fc010) and recombinant human fibronectin fragment 3 protein, CF (FN-050; Carrier free histidine-tagged FNIII8–14 fragment) were obtained from Millipore Limited (Hertfordshire, UK) and R&D systems (Minneapolis, Minnesota, USA) respectively, and the trisNTA-EG3-C16-SH linker was synthesized as described previously37–39. Gold-coated QCM crystals were supplied by open-QCM (Novatech S.r.l., Napoli, Italy) and gold-coated AFM chips (NPG-10) were purchased from Bruker (UK) Limited (Coventry, UK). The chemicals/reagents, ethanol (>99.5% for HPLC), hydrogen peroxide solution (30% w/w), concentrated sulfuric acid (99.999%), 1-hexanethiol (HS–(CH2)5–CH3), 6-mercaptopentanoic
Preparation of self-assembled monolayers (SAMs). Prior to their use as substrates for SAMs gold-coated QCM crystals were cleaned with freshly prepared piranha solution (7:3 mix of concentrated sulfuric acid and 30% w/w hydrogen peroxide). Hydroxyl, methyl, amine and carboxyl terminated SAMs were prepared according to Sigma Aldrich Technical Bulletin AL-266. Briefly, this involved immersing gold-coated QCM sensors in ethanolic solutions of different 2 mM alkanethiol (HS–(CH₄)₅–CH₃, HS–(CH₄)₅–COOH, HS–(CH₆)₆–NH₂, HS–(CH₆)₆–OH) solutions for 24 h. Self-assembly was terminated by rinsing the modified QCM sensors for 120 s in ethanol in a sonicating bath, before drying under nitrogen. Static contact angle measurements of the resulting SAMs, hereafter identified by the terminal function they present (CH₃, COOH, NH₂ and OH SAMs) were made using a contact angle goniometer (Biolin Scientific, Manchester, UK). Values of 101 ± 1°, 50 ± 1°, 35 ± 1° and 11 ± 1° (n = 3, mean ± SD) found for the CH₃, NH₂, OH and COOH, terminated SAMs respectively are in agreement with previous reports40–43.

Zeta potential measurement. A Zetasizer Nano ZS (Malvern Panalytical Ltd, Malvern, UK) was used to determine the zeta potential, ζ, of FNIII8–14 in 1 M HEPES buffer at pH 7.4, and value of −14 ± 1 mV was found.

Functionalization of AFM cantilevers. FNIII8–14 fragment derivatized AFM cantilevers were prepared in stepwise manner using a variation of Gruber’s protocol46, exploiting sulfur–gold bonding and immobilized metal chelate–histidine binding interactions37,38,45. Briefly this involved: (1) cleaning gold-coated AFM chips with UV/ozone; (2) immersing the cleaned AFM chips in an ethanolic solution containing 2 mM of 11-mercapto-1-undecanol, 0.02 mM of trisNTA-EG₃-C₁₆-SH linker, 2 mM EDTA and 2 mM TCEP for 24 h; (3) successive rinsing of the ‘trisNTA-EG₃-C₁₆/undecanol SAM’ modified AFM chips with absolute ethanol followed by 40 mM HEPES pH 7.4 buffer supplemented with 40 mM EDTA and 85 mM NaCl, before drying under nitrogen; (4) incubating in 200 µL buffer containing 0.5 µM FNIII8–14, 200 µm nickel (II) chloride at 37 °C for 1.5 h; and finally, rinsing the FNIII8–14 with 1 M HEPES solution in readiness for AFM experiments.

Force spectroscopy. Force measurements were performed in 1 M HEPES buffer pH 7.4 at room temperature in the fluid cell of a Nanowizard II atomic force microscope (JPK Instruments AG, Berlin, Germany). The AFM was loaded with fully functionalized cantilevers (featuring chemically attached linker and FNIII8–14 fragment designated ‘gold-linker’); FNIII8–14 physisorbed on bare cantilever (‘gold-FNIII8–14’); and cantilevers functionalized with trisNTA-EG₃-C₁₆-SH linker (‘gold-linker’).

Prior to experiments, AFM cantilevers were calibrated as detailed by Hutter and Bechhoefer46. A force of magnitude 1 nN was applied to the tip as it approached the SAM surface with a velocity of 1 µm s⁻¹. Once the tip had reached the surface, contact was maintained for 3 s before it was retracted and the process repeated. The force-distance (FD) curves were acquired from 16 different positions (4 × 4 grid) on the surface used, while the distance between each position was 10 µm. The 10 measurements were repeated four times with different AFM cantilevers and surfaces to ensure the reproducibility of the measurements.

Surface adsorption. A quartz crystal microbalance (OpenQCM, Novaetech S.r.l, Italy) was employed to determine the adsorption of full-length human plasma fibronectin on the different SAMs surfaces. SAM-coated QCM crystals were loaded onto the QCM before introducing the fibronectin solution (25 µg mL⁻¹) in PBS pH 7.4 at a flow of 1 µL s⁻¹ using an Ismatec IPC-N 4 peristaltic pump (Ismatec, Wertheim, Germany). Masses of adsorbed hydrated protein were calculated using the Sauerbrey equation47 (Eq. 1):

\[ \Delta f_m = -\frac{2\mu_q}{A\sqrt{\rho_q}} \cdot \Delta m = -C_f \cdot \Delta m \]  

(1)

here \( \Delta f \) is the measured resonant frequency (Hz), \( \mu_q \) is the intrinsic resonant frequency of the unloaded crystal (Hz), \( \Delta m \) is the mass change (g), A is the mass sensitive area of the electrode (cm²), \( \rho_q \) is the density of quartz (2.648 g cm⁻³), \( \mu_s \) is the shear modulus of the quartz crystal (2.947 × 10¹¹ g cm⁻¹ s⁻²) and \( C_f \) is the quartz sensitivity factor or constant (Hz g⁻¹). The negative sign in the formula implies that an increase in mass on the surface of the crystal results in a reduction of the measured resonant frequency. A total of seven measurements were performed for each SAM surface.

Data handling. All data is presented as mean values ± the standard error of the mean. An analysis of variance (ANOVA) was used to evaluate the significance of the measured parameters.
Results and discussion

Force spectroscopy measurements. The functionalization of AFM tips with the FNIII8–14 fragment is a two-step process, involving: (1) initial attachment of a heterobifunctional linker (trisNTA-EG3-C16-SH) via its thiol end to the gold-coated AFM tip; followed by (2) anchoring of the FNIII8–14 to the Ni(II)-loaded chelate moiety at the end of the linker via the fragment’s His-tag. To identify the contribution from each chemical species, force measurements were performed between the AFM tip and the SAMs surfaces at every step of the functionalization. Prior to examining FD curves of the chemically attached FNIII8–14 (‘gold-linker–FNIII8–14′) with the different SAM modified surfaces a hierarchical series of tests were conducted using bare (‘gold’), physisorbed FNIII8–14 (‘gold-FNIII8–14′), and trisNTA-EG3-C16-SH linker (‘gold-linker’) AFM cantilevers.

Interaction between ‘gold’ AFM tips and SAMs. Figure 1a,b respectively show representative FD retraction curves of bare gold-coated AFM chips contacting the four different SAM surfaces, and the mean forces that are required to separate them. Adhesion was noted to two of the four SAMs, i.e. methyl and amine terminated, but was not detected for the OH- and COOH-terminated surfaces. The absence of extra peaks in the retracting part of the FD curves (indicative of the presence of surface contaminants on the AFM tip and/or surface being interrogated) for gold AFM cantilevers interacting with CH3- and NH2-terminated SAMs confirm the effectiveness of the protocol employed for cleaning the AFM tips and SAM surfaces used in this study. The weak adhesion between the hydrophilic gold AFM tip and the hydrophobic CH3 SAM likely arises from interfacial tension48, whereas the considerably stronger interaction with NH2-SAM probably reflects electrostatic attraction between the opposing surfaces. In principle, the gold surface is electrostatically neutral, but in aqueous environments the adsorption of anions onto its surface lends it an overall negative charge49–52. In the present example, the adsorption of zwitterionic 4-(2-hydroxyethyl)-1-piperazineethanesulfonate buffer anions53 onto gold likely renders the latter surface negatively charged; which would explain (1) the greater force required to separate a bare gold AFM tip from the NH2 cf. CH3 SAM (i.e. 580 ± 107 cf. 143 ± 25 pN; Fig. 1b), and (2) the absence of any interaction between the bare AFM tip and the OH- and COOH-functionalized SAMs.

Interaction between ‘gold-FNIII8–14′ AFM tips and SAMs. In contrast to the single peaks observed in the FD retraction phase between bare gold and the CH3 and NH2 SAMs, multiple peaks were observed in the FD retraction curves between the physisorbed FNIII8–14 fragment and all four SAMs. For adsorbed proteins, the shape of retraction part of the FD curve is known to be dependent on: (1) the velocity of the AFM tip retracting from the surface; (2) the nature of the interactions between the protein and surface; and (3) protein conformation in the adsorbed state54. When multiple peaks arise (Fig. 2) the traces may exhibit one of three general forms, i.e. either: (1) a saw-tooth (loop-like) pattern, characteristic of unfolding of protein domains55,56; (2) an extended plateau (train-like) shape associated with desorption of proteins from charged or hydrophobic surfaces57,58, which has been attributed to denaturation of the adsorbed proteins53; or (3) some combination (mix) of (1) and (2). In physisorbed state, FNIII8–14 adopts different conformations at the gold AFM tip surface. A direct consequence of the conformational heterogeneity of FNIII8–14 in interactions of the gold-FNIII8–14 AFM tips with all four SAMs was no preference for a particular type of retraction force curve (Table 1). For three of the SAMs (CH3-, NH2-, COOH-terminated) the largest number of retraction FD curves (obtained with gold-FNIII8–14) were of the mixed type (41–64%). This aside, there were notable differences in the distribution of retraction curve shapes against the different SAM surfaces. For FD retraction curves measured against CH3 SAM, ~53% exhibited mixed events, ~41% showed train-like events and just 6.3% displayed loop-like curves. By contrast, none of the curves obtained with the OH SAM were train-like, 59% were loop-like, and 41% were of the mixed type. Given Meadows et al. correlation of train-like events with denaturation upon adsorption59, it follows that the extent of denaturation induced on FNIII8–14 by interaction with the SAM surfaces assumes the order CH3 > NH2 > COOH > OH. The desorption forces developed between gold-FNIII8–14 AFM tips and SAM surfaces are shown in Fig. 3a, and the corresponding desorption distances are presented in Fig. 3b.

![Figure 1](https://example.com/fig1.png)
Figure 2. Characteristic FD curves illustrating examples of: (a) a ‘loop-like’ retraction event indicating protein unfolding; (b) a ‘train-like’ retraction event implying protein desorption; and (c) a ‘mixed’ retraction event inferring desorption and unfolding. Desorption force is determined from the last peak in the retraction curve at which point (↑), the ‘desorption distance’, the protein detaches from the surface.

Table 1. Distributions of ‘loop-like’, ‘train-like’ and ‘mixed’ conformations calculated from FD curves generated between differently functionalized AFM tips and SAM substrates.

| Substrate | Gold-FNIII<sup>8–14</sup> AFM tip (physisorbed) | Gold-linker AFM tip | Gold-linker-FNIII<sup>8–14</sup> AFM tip (chemisorbed) |
|-----------|-----------------------------------------------|---------------------|----------------------------------------------------------|
|           | % loop | % train | % mix | % loop | % train | % mix | % loop | % train | % mix |
| CH<sub>3</sub> SAM | 6.3 | 40.6 | 53.1 | 0.0 | 0.0 | 100 | 0.0 | 14.3 | 85.7 |
| COOH SAM | 20.0 | 16.0 | 64.0 | 26.7 | 0.0 | 73.3 | 75.8 | 0.0 | 24.2 |
| NH<sub>2</sub> SAM | 34.8 | 21.7 | 43.5 | 3.6 | 39.3 | 57.1 | 0.0 | 18.6 | 81.4 |
| OH SAM | 58.8 | 0.0 | 41.2 | 23.1 | 0.0 | 76.9 | 94.3 | 0.0 | 5.7 |

Figure 3. Plots of mean adhesion force (top) and desorption distance (bottom) for gold-FNIII<sup>8–14</sup> (a, b), gold-linker (c, d) and gold-linker-FNIII<sup>8–14</sup> (e, f) AFM tips against four different SAMs in 1 M HEPES buffer pH 7. The data represent the mean values ± standard error of n = 64 force curves. Statistical tests: P < 0.05 for (a, d–f), and P > 0.05 for (b, c).
Interactions between gold-linker AFM tips and SAM surfaces. Force spectroscopy experiments performed with AFM tips pre-functionalized with a 10:1 mixture of 11-mercapto-1-undecanol and trisNTA-EG3-C6-SH linker yielded FD curves featuring multiple pulling events (Table 1). By far the greatest number of interactions (57.1–100.0%) between the gold-linker AFM tips and SAMs were of the mixed type. Roughly 25% of the retraction curve shapes against the OH and COOH terminated surfaces were loop-like, whereas FD curves for the NH2 surface were the only ones to suggest a train-like conformation (40%). The magnitude of the desorption forces developed against all types of surfaces were of the order of 150 pN (Fig. 3c), but non-specific events likely arising from the presence of aggregates resulted in there being no significant differences (P > 0.05) between them in terms of force. By contrast, significant differences (P < 0.05) in desorption distances for the various surfaces were found (Fig. 3d), i.e. desorption distances for the CH3 and NH2 surfaces were roughly two and three fold greater than those for COOH and OH SAMs, respectively. Collectively, the above observations suggest that the pulling events detected in FD curves might arise from the presence of aggregates (derived from 11-mercapto-1-undecanol and/or trisNTA-EG3-C6-SH) with both hydrophobic and negatively charged regions sitting on top of the mixed ‘OH/linker’ SAM of the AFM tip post functionalization. The greater desorption distances observed with CH3 and NH2 terminated SAMs may reflect the presence of larger aggregates on these surfaces and/or greater unfolding of the entangled aggregates.

Interaction between gold-linker-FNIII8–14 AFM tips with SAM surfaces. In contrast to findings with physisorbed FNIII8–14, most (i.e. > 82 to 100%) of the FD retraction curves for chemisorbed FNIII8–14 were loop-like or mixed varieties (Table 1), highlighting a specific interaction arising from successful functionalization. The probability of acquiring a force curve with meaningful feature, as defined in Fig. 2, was 57, 52, 69, and 55% for CH3, COOH, NH2, and OH-terminated SAMs, respectively. Predominantly loop-like curves were observed with COOH and OH SAMs (i.e. 76 and 94%, respectively) inferring domain unfolding. In contrast, CH3 and NH2 SAMs surfaces favored curves of mixed type, indicative of ‘denaturation and unfolding’. The presence of train-like curves, plus complete absence of loop-like ones, in interactions between chemisorbed FNIII8–14 and the CH3 and NH2 SAMs, infer that these two surfaces induce greater denaturation of the fibronectin fragment cf. the COOH and OH terminated varieties; the extent of denaturation induced assumes the order: NH2 > CH3 > COOH > OH.

Of particular importance were striking differences between physisorbed and chemisorbed FNIII8–14. Whereas physisorbed FNIII8–14 showed a preference for CH3 and NH2 terminated surfaces (Fig. 3a), chemisorbed FNIII8–14 exhibited a contrary bias, i.e. substantially higher affinity for COOH and OH functionalized SAMs. For example, Fig. 3e shows that 2–3-fold greater desorption forces were required to release FNIII8–14 from COOH and OH cf. CH3 and NH2 terminated varieties, indicating that adsorption affinity for FNIII8–14 follows the order: OH ≥ COOH > NH2 > CH3. The exact opposite trend was noted for desorption distance, i.e. CH3 > NH2 > COOH > OH (Fig. 3f). Maps of desorption force versus distance (Fig. 4) show loose clustering of the data obtained for each surface (from left to right as follows: OH, COOH, NH2, CH3), and serve to highlight the greater spread in desorption force cf. distance for OH and COOH surfaces, and the opposite for NH2 and CH3 SAMs, i.e. wider distribution in desorption distance cf. force. It is worth noting that previous study confirmed that the linker, tris-NTA-His6, remains intact over a range of adhesion forces up to 600 pN, which provides reassurance that the pulling events observed were not compromised.

Given the differences in interaction types noted between the four SAMs, it follows that the conformations of FNIII8–14 adopted on these surfaces are likely different. The strength of interaction between a given protein and surface typically correlates with extent of unfolding/denaturation and ‘protein-surface’ contact area, i.e. the greater the interaction strength, the greater the unfolding, the larger the contact area. The contact area will be larger when FNIII8–14 is orientated ‘side-on’ to the binding surface (long axis parallel to surface), and lowest when adsorbed in a ‘head on’ orientation (long axis perpendicular to the surface).

Under the experimental conditions applied, the surface of FNIII8–14 exhibits a net negative charge (ζ = −14 ± 1 mV). FNIII8–14 binding to the positively charged surface of NH2 SAM is likely driven by large numbers of strong electrostatic attraction forces over a high contact area. A large footprint is also expected for FNIII8–14 on CH3 SAM the most hydrophobic surface (displaying a contact angle of 101 ± 1°) promoted by unfolding, exposure of and strong interaction with previously buried hydrophobic residues. Weaker forces steer the adsorption of FNIII8–14 on COOH and OH SAMs, i.e. van der Waals in the case of the OH SAM, and a combination of electrostatic and van der Waals interactions for the COOH SAM. Despite carrying a net negative charge positively charged regions within FNIII8–14 are attracted and eventually adsorbed on the COOH SAM surface. It is expected that OH and COOH surfaces have fewer contact points with the protein cf. CH3 and NH2 surfaces. We therefore envisage that a large fraction of FNIII8–14 molecules adsorb ‘side-on’ (i.e. with their long axis approximately parallel to the surface) on CH3 and NH2 SAMs, and ‘head-on’ (i.e. long axis approximately perpendicular) on the OH and COOH SAMs. The lower mean adhesion forces on CH3 and NH2 cf. OH and COOH SAMs (Fig. 3e) imply that these two surfaces induce less denaturation of FNIII8–14 (careful inspection of FD retraction curve data in Table 1 shows the reverse is true), and they also appear at odds with the above assumptions on the forces governing FNIII8–14 binding and its orientation in the adsorbed state. It should be stressed that numbers for the desorption force derive from the last peak in the retraction curve (Fig. 2). As FNIII8–14 molecules adhering to CH3 and NH2 surfaces are desorbed and unfolded over larger distances cf.
COOH and OH SAMs, smaller numbers of them will remain on the surface before the final desorption force is determined, resulting in lower measured values.

**QCM studies.** Figure 5a illustrates the time-dependent adsorption of full-length human FN onto gold-coated QCM crystals homogeneously functionalized with SAMs. At the relatively high FN concentration employed (25 µg mL⁻¹), at least two kinetic steps can be discerned from the curves. Hydrated mass loads on the sensors increased rapidly over the first 5 min at broadly similar rates, i.e. ~ 130 ng cm⁻² min⁻¹ for the hydrophobic CH₃ SAM and ~ 150 ng cm⁻² min⁻¹ for the three hydrophilic SAMs. Thereafter, adsorption rates diminished variably with increasing time, as surface occupancy increased and quasi-steady-state saturation was approached. At the very final stage (i.e. after 70 min of adsorption), surface concentrations (Fig. 5b) reached levels of 945 ± 22, 1158 ± 41, 1427 ± 32 and 1495 ± 45 ng hydrated mass per cm² for the CH₃, OH, COOH and NH₂ SAMs respectively.

FN is an extremely flexible molecule with a conformation strongly dependent on its local environment, and ability to adsorb on very different surfaces and adopt different conformations at the interface. Comparative studies of FN binding to hydrophilic and hydrophobic surfaces show that under normal physiological conditions, adsorption on hydrophobic, but not hydrophilic, substrates induces significant unfolding and footprint spreading. The degree to which these events occur depends on: (1) the hydrophobicity of the surface, i.e. the more hydrophobic the greater the effects; and (2) the bulk phase FN concentration and packing density (low surface coverage promotes strong unfolding and molecular expansion, whereas high packing density suppresses both effects). It follows that the lowest hydrated mass/area loading observed in this work for CH₃ SAM (945 ng cm⁻²) likely reflects monolayer binding dominated by a space filling 'side on' orientation, with the additional possibility of an increase in FN's molecular footprint (Fig. 6a). The significantly raised loadings...
on the OH (1157 ng cm\(^{-2}\)) and especially COOH (1427 ng cm\(^{-2}\)) SAMs \(\text{cf.}\) that on the methyl terminated SAM could be explained by greater numbers of FN molecules binding ‘end-on’ (Fig. 6b), in keeping with our findings in this work (Table 1, Fig. 4) of the single FNIII8–14 molecule’s observed preference for this orientation on these two surfaces.

Whilst binding to CH\(_3\) and OH SAMs involves short-range intermolecular forces, long-range electrostatics steer initial binding of FN to hydrophilic charged SAMs\(^{73}\). At first glance, the similar maximum loadings of FN on the negatively charged COOH and positively charged NH\(_2\) SAMs (1427 ± 32 and 1495 ± 45 ng cm\(^{-2}\), respectively), may be difficult to reconcile with the molecule’s small net negative charge (\(\zeta = -5.7\) mV) in PBS pH 7.4\(^{35}\). They are however, consistent with previous studies; indeed Lin et al.\(^{74}\) observed higher mass loadings on COOH \(\text{cf.}\) NH\(_2\) SAMs. Whereas FN binding to the amine-terminated surface is driven by electrostatic attraction, its adsorption on the carboxyl-terminated surface appears more complex, probably involving one or more positively charged residues, e.g. Lys 1469 located in a positive patch around the cell binding region\(^{73}\), augmented by shorter range holding forces\(^{74}\). Though similar surface concentrations were reached on NH\(_2\) and COOH SAMs the shape of the NH\(_2\) SAM binding profile implies a more complex adsorption behavior \(\text{cf.}\) the other surfaces. Here we

**Figure 5.** Plots of (a) change in surface concentration vs. time and (b) maximum surface concentration attained for the adsorption of fibronectin (25 µg/mL) on CH\(_3\)–, COOH–, NH\(_2\)– and OH-terminated SAMs in PBS buffer pH 7.4. The dip in the traces preceding the onset of binding (↓) reflects a momentary halt in pumping on switching from PBS to the protein solution. The data represent the mean values ± standard error of n = 7 measurements.

**Figure 6.** Schematic illustrations (a–d) showing possible changes in FN orientation and shape upon surface adsorption as a function of increasing time/surface coverage. The white and gray ovals represent free and bound FN respectively. (a) ‘Side-on’ binding with possible spreading forming a sparse monolayer; (b) ‘end-on’ monolayer binding; (c) initial ‘side-on’ binding switching to ‘end-on’ oriented monolayer; and (d) initial ‘side-on’ monolayer binding to multilayer.
envisage two possibilities. In both, FN binds initially in space-filling side-on manner commensurate with the single molecule's preference. With increasing supply to the surface, adsorbed FN molecules either rearrange, tiling to a space-saving end-on configuration to maximize packing density (Fig. 6c), or alternatively multi-layer binding of FN (Fig. 6d).

Conclusions
Knowledge on the adsorption behavior of FN molecules against a quartet of chemical defined surfaces was acquired by AFM-SMFS and QCM investigations. The following general conclusions can be drawn from the SMFS studies under physiological conditions: that extent of the unfolding and denaturation, strength of adsorption and binding orientation of FNIII15–18 correlated closely with one another. While the single molecule's preference on more strongly adhering and denaturing positively charged NH3 and hydrophobic CH3 surfaces appears to be binding in a 'side-on' orientation, an 'end-on' configuration is favored on the comparatively gentle (i.e. less denaturing) more weakly interacting hydrophilic neutral OH and negatively charged COOH surfaces. The preferred binding orientation of the single tethered FNIII15–18 molecule was also inferred for the initial binding of free full-length FN on gold-coated QCM crystals modified with the same SAMs, i.e. ‘end-on’ on OH and COOH surfaces, and ‘side-on’ on CH3 and NH3 SAMs; in accord with Michael et al. findings36. In the context of cell binding, the orientation of surface adsorbed FN molecules is of key import, given that it determines the accessibility or otherwise of RGD motifs, which in turn influence cell surface interactions on exposure to the biological environment and during wound healing7,4,7,8,11,75–80. The approaches detailed herein on the interrogation of FN binding to chemically defined SAMs could be usefully employed as quantitative tools in much broader context including but not limited the design/development of surfaces that control cell adhesion and platelet activation, bacterial colonization, or limit fouling38,16,62.

References
1. Ratner, B. D. The biocompatibility of implant materials. In Host Response to Biomaterials (ed. Badylak, S. F.) (Academic Press, 2015).
2. Ratner, B. D. Biomaterials Science: An Introduction to Materials in Medicine 3rd edn. (Elsevier/Academic Press, Amsterdam, 2013).
3. Anderson, J. M., Rodriguez, A. & Chang, D. T. Foreign body reaction to biomaterials. Semin. Immunol. 20, 86–100 (2008).
4. Ozboyaci, M., Kokh, D. B., Corni, S. & Wade, R. C. Modeling and simulation of protein-surface interactions: Achievements and challenges. Q. Rev. Biophys. 49, e4 (2016).
5. Denis, E. A. et al. Protein adsorption on model surfaces with controlled nanotopography and chemistry. Langmuir 18(5), 819–828 (2002).
6. Sigal, G. B., Mrksich, M. & Whitesides, G. M. Effect of surface wettability on the adsorption of proteins and detergents. J. Am. Chem. Soc. 120, 3464–3473 (1998).
7. Scotchford, C. A., Gilmore, C. P., Cooper, E., Leggett, G. J. & Downes, S. Protein adsorption and human osteoblast-like cell attachment and growth on alkylthiol on gold self-assembled monolayers, J. Biomed. Mater. Res. 59, 84–99 (2002).
8. Mandracci, P., Mussano, F., Rivolo, P. & Carossa, S. Surface treatments and functional coatings for biocompatibility improvement and bacterial adhesion reduction in dental implantology. Coatings 6, 7 (2016).
9. Auernheimer, J. et al. Titanium implant materials with improved biocompatibility through coating with phosphonate-anchored cyclic RGD peptides. ChemBioChem 6, 2034–2040 (2005).
10. Hersel, U., Dahmen, C. & Kessler, H. RGD modified polymers: Biomaterials for stimulated cell adhesion and beyond. Biomaterials 24(24), 4385–4415 (2003).
11. von Salis-Soglio, M. et al. A novel multi-phosphate surface treatment of titanium dental implants: A study in sheep. J. Funct. Biomater. 5, 135–157 (2014).
12. Lee, G. U., Kidwell, D. A. & Colton, R. J. Sensing discrete streptavidin-biotin interactions with atomic-force microscopy. Langmuir 10, 354–357 (1994).
13. Janshoff, A., Neitzert, M., Oberdorfer, Y. & Fuchs, H. Force spectroscopy of molecular systems: single molecule spectroscopy of polymers and biomolecules. Angew. Chem. Int. Ed. Engl. 39, 3212–3237 (2000).
14. Höök, F. et al. A comparative study of protein adsorption on titanium oxide surfaces using in situ ellipsometry, optical waveguide lightmode spectroscopy, and quartz crystal microbalance/dissipation. Colloid Surface B 24, 155–170 (2002).
15. Stadler, H., Mondon, M. & Ziegler, C. Protein adsorption on surfaces: Dynamic contact-angle (DCA) and quartz-crystal microbalance (QCM) measurements. Anal. Bioanal. Chem. 375, 53–61 (2003).
16. Parkov, R. & Yamada, K. M. Fibronectin at a glance. J. Cell Sci. 115, 3861–3863 (2002).
17. Pearlstein, E., Gold, I. L. & Garcia-Pardo, A. Fibronectin: A review of its structure and biological activity. Mol. Cell Biochem. 29, 103–128 (1980).
18. Rombherger, D. J. Fibronectin. Int. J. Biochem. Cell Biol. 29, 939–943 (1997).
19. Schwarzaurber, J. E. & DeSimone, D. W. Fibronectins, their fibrillogenesis, and in vivo functions. Cold Spring Harb. Perspect. Biol. 3, a005041 (2011).
20. Man, Y. & Schwarzaurber, J. E. Fibronectin fibrillogenesis, a cell-mediated matrix assembly process. Matrix Biol. 24, 389–399 (2005).
21. Potts, J. R. & Campbell, I. D. Structure and function of fibronectin modules. Matrix Biol. 15, 313–320 (1996).
22. Pierschbacher, M. D. & Ruoslahti, E. Cell attachment activity of fibronectin can be duplicated by small synthetic fragments of the molecule. Nature 309, 30–33 (1984).
23. Yamada, K. M. & Kennedy, D. W. Dualistic nature of adhesive protein function: Fibronectin and its biologically active peptide fragments can autoinhibit fibronectin function. J. Cell Biol. 99, 29–36 (1984).
24. Aota, S., Nomizu, M. & Yamada, K. M. The short amino acid sequence Pro-His-Ser-Arg-Asn in human fibronectin enhances cell-adhesive function. J. Biol. Chem. 269, 4756–4761 (1994).
25. Kimizuka, F. et al. Role of type-III homology repeats in cell adhesion function within the cell-binding domain of fibronectin. J. Biol. Chem. 266, 3045–3051 (1991).
26. Obara, M., Kang, M. S. & Yamada, K. M. Site-directed mutagenesis of the cell-binding domain of human fibronectin: separable, synergistic sites mediate adhesive function. Cell 53, 649–657 (1988).
27. Rief, M., Gautel, M., Schemmel, A. & Gauld, H. E. The mechanical stability of immunoglobulin and fibronectin III domains in the muscle protein titin measured by atomic force microscopy. Biophys. J. 75, 3008–3014 (1998).

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28. Oberhauser, A. F., Badilla-Fernandez, C., Carrion-Vazquez, M. & Fernandez, J. M. The mechanical hierarchies of fibronectin observed with single-molecule AFM. J. Mol. Biol. 319, 433–447 (2002).

29. Li, L., Huang, H. H., Badilla, C. L. & Fernandez, J. M. Mechanical unfolding intermediates observed with single-molecule force spectroscopy in a fibronectin type III module. J. Mol. Biol. 345, 817–826 (2005).

30. Erickson, H. P. Reversible unfolding of fibronectin type III and immunoglobulin domains provides the structural basis for stretch and elasticity of titin and fibronectin. P. Natl. Acad. Sci. USA 91, 10114–10118 (1994).

31. Li, F. Y., Redick, S. D., Erickson, H. P. & Moy, V. T. Force measurements of the alpha(3)beta(1) integrin-fibronectin interaction. Biophys. J. 84, 1252–1262 (2003).

32. Meadows, P. Y. & Walker, G. C. Force microscopy studies of fibronectin adsorption and subsequent cellular adhesion to substrates with well-defined surface chemistries. Langmuir 21, 4096–4107 (2005).

33. Meadows, P. Y., Bemis, J. E. & Walker, G. C. Single-molecule force spectroscopy of isolated and aggregated fibronectin proteins on negatively charged surfaces in aqueous liquids. Langmuir 19, 9566–9572 (2003).

34. Dickerson, M. T. et al. Fibronectin binding to the Treponeuma pallidum adhesion protein fragment rPp0483 on functionalized self-assembled monolayers. Bioconjug. Chem. 23, 184–195 (2012).

35. Lin, J. H. et al. Effect of surface potential on extracellular matrix protein adsorption. Langmuir 30, 10328–10335 (2014).

36. Michael, K. E. et al. Adsorption-induced conformational changes in fibronectin due to interactions with well-defined surface chemistries. Langmuir 19, 8033–8040 (2003).

37. Lata, S., Reichl, A., Brock, R., Tampe, R. & Pielker, J. High-affinity adaptors for switchable recognition of histidine-tagged proteins. J. Am. Chem. Soc. 127, 10205–10215 (2005).

38. Tinazli, A. et al. High-affinity chelator thiolis for switchable and oriented immobilization of histidine-tagged proteins: A generic platform for protein chip technologies. Chemistry 11, 5249–5259 (2005).

39. Valiokas, R. et al. Self-assembled monolayers containing terminal mono-, bis-, and tris-nitrilotriacetic acid groups: Characterization and application. Langmuir 24, 4959–4967 (2008).

40. Hoypierryes, J. et al. Two methods for one-point anchoring of a linear polysaccharide on a gold surface. Langmuir 31, 254–261 (2015).

41. Ostuni, E., Yan, L. & Whitesides, G. M. The interaction of proteins and cells with self-assembled monolayers of alkanethiols on gold and silver. Colloid Surf. B 15, 3–30 (1999).

42. Petrovykh, D. Y. et al. Alkanethiols on platinum: Multicomponent self-assembled monolayers. Langmuir 22, 2578–2587 (2006).

43. Wang, Z., Nania, S. L. & Shaw, S. K. Structure of aqueous water films on textured –OH-terminated self-assembled monolayers. Langmuir 31, 2382–2389 (2015).

44. Gruber, H. J. Functionalisation of AFM tips with tris-NTA for binding and His6-tagged proteins. https://www.jku.at/fileadmin/grupp/en/21608_AFM_tip_with_tris-NTA_2016_05_06.pdf (2016).

45. Tinazli, A., Pielker, J., Beuttler, M., Guckenberger, R. & Tampe, R. Native protein nanolithography that can write, read and erase. Nat. Nanotechnol. 2, 220 (2007).

46. Hutter, J. L. & Bechhoefer, J. Calibration of atomic-force microscope tips. Rev. Sci. Instrum. 64, 1868–1873 (1993).

47. Sauerbrey, G. Verwendung von schwingquarzen zur wagung dunner schichten und zur mikrowagung. Z. Angew. Phys. 156, 202–222 (1959).

48. Papastavrou, G., Akari, S. & Mohwald, H. Interactions between hydrophilic and hydrophobic surfaces on microscopic scale and the influence of air bubbles as observed by scanning force microscopy in aqueous and alcoholic mediums. Europhys. Lett. 52, 551–556 (2000).

49. Biggs, S., Mulvaney, P., Ziaoski, C. F. & Griesser, F. Study of adion adsorption at the gold-aqueous solution interface by atomic force microscopy. J. Am. Chem. Soc. 116, 9150–9157 (1994).

50. Kasuya, M. et al. Anion adsorption on gold electrodes studied by electrochemical surface force measurements. J. Phys. Chem. C 120, 15986–15992 (2016).

51. Kumal, R. R., Karam, T. E. & Haber, L. H. Determination of the surface charge density of colloidal gold nanoparticles using second harmonic generation. J. Phys. Chem. C 119, 16200–16207 (2015).

52. Thompson, D. W. & Collins, I. R. Electrical properties of the gold-aqueous solution interface. J. Colloid Interf. Sci. 152, 197–204 (1992).

53. Al-Ghobashy, M. A. Electrophotorec behavior of charge regulated zwittrer ionic buffers in covalently and dynamically coated fused silica capillaries. Bull. Fac. Pharm. Cairo Univ. 52, 71–78 (2014).

54. Zhang, Z., Tomlinson, M. R., Golestanian, R. & Geoghegan, M. The interfacial behaviour of single poly(N,N-dimethylacrylamide) chains as a function of pH. Nanotechnology 19, 035505 (2008).

55. Carrion-Vazquez, M. et al. Mechanical and chemical unfolding of a single protein: A comparison. P. Natl. Acad. Sci. USA 96, 3694–3699 (1999).

56. Fisher, T. E., Oberhauser, A. F., Carrion-Vazquez, M., Marszałek, P. E. & Fernandez, J. M. The study of protein mechanics with the atomic force microscope. Trends Biochem. Sci. 24, 379–384 (1999).

57. Horinek, D. et al. Peptide adsorption on gold surface results from an interplay of solvation, surface, and intrapeptide forces. P. Natl. Acad. Sci. USA 105, 2842–2847 (2008).

58. Kienle, S., Pirzer, T., Krysiak, S., Geisler, M. & Hugel, T. Measuring the interaction between ions, biopolymers and interfaces—One polymer at a time. Faraday Discuss 160, 329–340 (2013).

59. Verbelen, C., Gruber, H. J. & Dufrène, Y. F. The NTA-His, bond is strong enough for AFM single-molecule recognition studies. J. Mol. Recognit. 20, 490 (2007).

60. Lhoest, J. B., Detrait, E., de Aguilar, P. B. & Bertrand, P. Fibronectin adsorption, conformation, and orientation on polystyrene substrates studied by radiolabeling, XPS, and ToF SIMS. J. Biomed. Mater. Res. 41, 95–103 (1998).

61. Perndorf, N. et al. Fibronectin fibrillogenesis on sulfonated polystyrene surfaces. J. Biomed. Mater. Res. A 64, 684–692 (2003).

62. Erickson, H. P., Carrell, N. & McDonald, J. Fibronectin molecule visualized in electron microscopy: A long, thin, flexible strand. J. Cell Biol. 91, 673–678 (1981).

63. Osterlund, E. The secondary structure of human plasma fibronectin: Conformational changes induced by acidic pH and elevated temperatures; a circular dichroic study. Biochim. Biophys. Acta 955, 330–336 (1988).

64. Baujard-Lamotte, L., Noinville, S., Goubard, F., Marque, P. & Pauthe, E. Kinetics of conformational changes of fibronectin adsorbed on model surfaces. Colloid Surf. B 63, 129–137 (2008).

65. Benecky, M. J., Kolvenbach, C. G., Wine, R. W., DiOrtio, J. P. & Mosesson, M. W. Human plasma fibronectin structure probed by steady-state fluorescence polarization: Evidence for a rigid oblate structure. Biochemistry 29, 3082–3091 (1990).

66. Sjoberg, B., Eriksson, M., Osterlund, E., Pap, S. & Osterlund, K. Solution structure of human plasma fibronectin as a function of NaCl concentration determined by small-angle X-ray scattering. Eur. Biophys. J. 17, 5–11 (1989).

67. Grinnell, F. & Feld, M. K. Fibronectin adsorption on hydrophilic and hydrophobic surfaces detected by antibody binding and analyzed during cell adhesion in serum-containing medium. J. Biol. Chem. 257, 4888–4893 (1982).

68. Iwamoto, G. K. et al. Fibronectin adsorption detected by interfacial fluorescence. J. Colloid Interf. Sci. 106, 459–464 (1985).

69. Wolff, C. & Lai, C. S. Fluorescence energy transfer detects changes in fibronectin structure upon surface binding. Arch. Biochem. Biophys. 268, 536–545 (1989).
70. MacDonald, D. E., Markovic, B., Allen, M., Somasundaran, P. & Boskey, A. L. Surface analysis of human plasma fibronectin adsorbed to commercially pure titanium materials. *J. Biomed. Mater. Res.* **41**, 120–130 (1998).

71. Wolff, C. E. & Lai, C. S. Inter-sulfhydryl distances in plasma fibronectin determined by fluorescence energy transfer: Effect of environmental factors. *Biochemistry* **29**, 3354–3361 (1990).

72. Klotzsch, E. et al. Conformational distribution of surface-adsorbed fibronectin molecules explored by single molecule localization microscopy. *Biomater. Sci.* **2**, 883–892 (2014).

73. Liamas, E. et al. Adsorption of fibronectin fragment on surfaces using fully atomistic molecular dynamics simulations. *Int. J. Mol. Sci.* **19**, 3321 (2018).

74. Lin, J. H. et al. Effect of surface potential on extracellular matrix protein adsorption. *Langmuir* **30**(34), 10328–10335 (2014).

75. Campillo-Fernandez, A. J. et al. Analysis of the biological response of endothelial and fibroblast cells cultured on synthetic scaffolds with various hydrophilic/hydrophobic ratios: Influence of fibronectin adsorption and conformation. *Tissue Eng. Part A* **15**, 1331–1341 (2009).

76. Gossart, A., Battiston, K. G., Gand, A., Pauhe, E. & Santrerre, J. P. Mono vs multilayer fibronectin coatings on polar/hydrophobic/ionic polyurethanes: Altering surface interactions with human monocytes. *Acta Biomater.* **66**, 129–140 (2018).

77. Sun, Y., Welsh, W. J. & Latour, R. A. Prediction of the orientations of adsorbed protein using an empirical energy function with implicit solvation. *Langmuir* **21**, 5616–5626 (2005).

78. Arima, Y. & Iwata, H. Effect of wettability and surface functional groups on protein adsorption and cell adhesion using well-defined mixed self-assembled monolayers. *Biomaterials* **28**, 3074–3082 (2007).

79. Denis, F. A. et al. Protein adsorption on model surfaces with controlled nanotopography and chemistry. *Langmuir* **18**, 819–828 (2002).

80. Hersel, U., Dahmen, C. & Kessler, H. RGD modified polymers: Biomaterials for stimulated cell adhesion and beyond. *Biomaterials* **24**, 4385–4415 (2003).

81. See, J. H. et al. Designing dynamic surfaces for regulation of biological responses. *Soft Matter* **8**, 5477–5485 (2012).

82. Gorth, D. J. et al. Decreased bacteria activity on Si(3)N(4) surfaces compared with PEEK or titanium. *Int. J. Nanomed.* **7**, 4829–4840 (2012).

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**Author contributions**

E.L., R.A.B., P.A.M, O.R.T.T., and Z.J.Z. designed the experiments. R.T. and R.W. designed and synthesised the linkage molecule. E.L. performed SMFS and QCM measurements. E.L., O.R.T.T., and Z.J.Z. wrote the main manuscript. All authors reviewed and approved the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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