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New mechanism of fibronectin fibril assembly revealed by live imaging and superresolution microscopy — Source link ☑

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Published on: 10 Sep 2020 - bioRxiv (Cold Spring Harbor Laboratory)

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30 Abstract

31• The regulation of cell fate decisions, morphogenesis, and responses to injury are intimately 32 linked to the process of Fn1 fibrillogenesis. Live imaging and super-resolution microscopy revealed that Fn1 fibrils are not continuous. Instead, Fn1 fibrils arise from nanodomains 33 34 containing multiple Fn1 dimers. As they move toward cell center. Fn1 nanodomains become 35 organized into linear arrays with a spacing of 130 nm between the nanodomains, with little Fn1 in between: Fn1 nanodomain arrays are resistant to deoxycholate treatment demonstrating 36 that these beaded assemblies are indeed mature Fn1 fibrils. FUD, a bacterial peptide that 37 disrupts Fn1 fibrillogenesis, does not disrupt nanodomain formation; instead, it interferes with 38 39 the organization of nanodomains into arrays. The nanodomain composition of Fn1 fibrils is 40 observed in multiple contexts: in three-dimensional ECM in vivo, on substrata of different composition and stiffness, and is retained in the absence of cells. The modular architecture of 41 42 Fn1 fibrils bears important implications for mechanisms of ECM remodeling and signal transduction. 43

44 Introduction

45 Fibronectin (Fn1) is a requisite component of extracellular matrix (ECM) necessary for 46 embryogenesis and homeostasis (1). It is noteworthy that in the absence of Fn1 fibrillogenesis, 47 the binding of Fn1 to cells is not sufficient to regulate key biological processes including those 48 governing embryonic development, angiogenesis, vascular remodeling, or cartilage 49 condensation (2-5). Therefore, understanding the mechanisms by which Fn1 proteins 50 assemble into macromolecular fibrils is essential to gain insights into in vivo functions of Fn1. 51 Fn1 fibrillogenesis occurs following the binding of secreted Fn1 homodimers to cell-surface 52 integrins. Following integrin binding, intracellular cytoskeletal forces such as actomyosin 53 contractility acting through integrins generate pulling forces on Fn1 dimers, exposing epitopes 54 that promote Fn1 fibrillogenesis (6-10). At the cell biological level, the process of Fn1 55 fibrillogenesis is correlated with the formation of fibrillar adhesions, whereby mobile adhesions 56 containing Fn1 and integrin α 5 β 1 translocate toward the nucleus resulting in elongated linear 57 arrays termed focal and fibrillar adhesions, containing both Fn1 and intracellular cytoplasmic 58 effectors linking Fn1 and actin cytoskeleton (11-16).

59 It has been thought that Fn1 fibrils arise following partial unfolding and alignment of Fn1 dimers in a periodic end-to-end fashion of alternating N- and C-termini, forming continuous 60 fibers (17-19). However, this model is based on the analyses of fixed samples and electron 61 62 microscopy wherein the exact composition of the fibrous material imaged could not be easily 63 determined. To evaluate the process of fibrillogenesis in real time, we adopted a 64 CRISPR/Cas9-mediated mutagenesis approach to generate fluorescently-labeled Fn1, subject to physiological regulation of expression and splicing. This approach has enabled us to 65 66 visualize the initiation and progression of Fn1 fibrillogenesis over an extended period of time. 67 Using live imaging and super resolution microscopy, we uncovered an unexpected mechanism 68 of Fn1 fibrillogenesis. Our data demonstrate that Fn1 fibrils are composed of centripetally-

69 moving Fn1 nanodomains originating at cell periphery. As Fn1 nanodomains move toward the 70 nucleus, they become arranged into progressively longer and longer arrays wherein nanodomains containing a high number of Fn1 dimers alternate with regions containing low or 71 72 no Fn1. We show that the N-terminal Fn1 assembly region is not required for the formation of 73 Fn1 nanodomains or their centripetal translocation. Instead, the N-terminus of Fn1 regulates 74 the organization of Fn1 nanodomains into linear arrays. This model of fibrillogenesis integrates 75 the process of fibrillogenesis with adhesion maturation and provides significant new insights 76 into the mechanisms of ECM formation, remodeling, and signaling.

- 77
- 78 Results

79 While examining Fn1⁺ ECM by confocal immunofluorescence microscopy in mouse 80 embryos, we observed that Fn1 fibrils appeared discontinuous (Fig. 1A-B, arrows), containing 81 regularly-spaced regions of high and low fluorescence intensity (Fig. 1C, Movie 1). This suggested that Fn1 fibrils consisted of regions with a high number of Fn1 dimers separated by 82 regions containing a low number of Fn1 dimers (Fig. 1C). To test this hypothesis and to 83 84 determine mechanisms regulating the formation of Fn1 fibrils, we employed a CRISPR/Cas9 85 knock-in strategy to modify the endogenous Fn1 locus by replacing the termination codon of 86 Fn1 with a sequence encoding a fluorescent protein. This strategy has allowed to generate 87 fluorescently-labeled Fn1 proteins subject to endogenous regulation (Sup. Fig. 1A-B), and 88 enabled extended live imaging of Fn1 fibrillogenesis. Using this strategy, we obtained multiple 89 independent lines of mouse embryo fibroblasts (MEFs) expressing Fn1-mEGFP, Fn1mScarlet-I, Fn1-Neon Green, or Fn1-tdTomato fusion proteins (FP) generated by 90 91 CRISPR/Cas9-mediated insertion. Western blots showed that FP fusions to Fn1 were specific: 92 FPs were only fused to Fn1 as no other FP fusions were detected either by western blotting or 93 immunofluorescence (IF) (Sup. Fig. 2A-B).

94 Deoxycholate (DOC) insolubility of Fn1 ECM is a classical biochemical assay for 95 proteins stably incorporated into the assembled ECM (20-23). DOC assays demonstrated that the incorporation of Fn1-FPs into ECM was indistinguishable from wild-type, untagged Fn1 96 (Sup. Fig. 2C). To determine whether Fn1-FP proteins carried out the physiological functions 97 98 of Fn1, we generated Fn1^{mEGFP} knock-in mice. Fn1^{mEGFP/mEGFP} homozygous knock-in animals 99 are viable and fertile (Sup. Fig. 1B panels 4 and 5). Furthermore, Fn1-mEGFP proteins were 100 expressed in the same pattern as total Fn1 protein (Sup. Fig. 2d) (24). Together, these studies 101 demonstrated that Fn1-FP fusion proteins are suitable reagents for investigating mechanisms 102 of Fn1 fibrillogenesis.

To visualize the process of fibrillogenesis in real time, we plated Fn1^{mEGFP/+} MEFs on 103 104 gelatin-coated cover glass and imaged cells 16 hours after plating using total internal reflection 105 (TIRF) microscopy at the critical angle of incidence. These experiments showed that Fn1 106 fibrillogenesis initiated at cell periphery as distinct bright Fn1 densities that moved centripetally 107 in parallel with F-actin and aligned into linear arrays of "beads" (arrows in Movie 2). TIRF 108 imaging also showed that the domains of higher fluorescence intensity of Fn1 co-localized with integrin α 5 β 1 both in non-fibrillar adhesions (arrows in **Fig. 2A-A2**) and in fibrillar adhesions 109 110 (arrows in **Fig. 2B-B2**), and that Fn1 and α 5 β 1 fibrillar adhesions are beaded (**Fig. 2B-B2**). 111 arrows). We also observed the beaded architecture of Fn1 fibrils using an independent 112 imaging method by employing Zeiss Airyscan (Sup. Fig. 3). Staining using monoclonal and 113 polyclonal antibodies recognizing distinct and multiple epitopes in Fn1 resulted in the 114 discontinuous appearance of Fn1 fibrils (Fig. 3A1-C1). Fn1 fibrils formed by cells plated on 115 glass, gelatin, laminin 111 or vitronectin were beaded Fn1 fibrils (Fig. 3, and Sup. Figs. 3-4), 116 and so were Fn1 fibrils in cell-free areas (Fig. 3A, A1) and between cells (Fig. 3B-C). In addition, the beaded appearance of Fn1 fibrils was observed when cells were plated on soft 117 118 substrata such as hydrogels of variable stiffness (Sup. Fig. 4C-D). In the latter experiment,

119 Fn1 was detected by imaging the native fluorescence of Fn1-mEGFP, indicating that the

120 beaded appearance of Fn1 fibrils was independent of antibody staining. Taken together, these

121 studies indicated that the beaded appearance of Fn1 fibrils was a general feature of Fn1 ECM

seen in 3D ECM *in vivo* and under different conditions *in vitro*.

123 To test the hypothesis that the "beads" in Fn1 strings were contiguous, cultures were 124 treated with 2% DOC. 2% DOC treatment dissolves cell membranes and cytoplasmic 125 components, leaving insoluble ECM devoid of cell contact (see Movie 3 for time-laps of 126 dissolution of cellular components, F-actin and DNA). This experiment showed that Fn1 fibrils in assembled ECM retained their beaded architecture in the absence of cell contact (Fig. 3D). 127 Together, these data indicated that the beaded topology of Fn1 fibrils is a feature of 128 129 physiological three-dimensional Fn1 ECM in vivo and suggested that the beads are contiguous. 130

131 To determine the relationship between the beaded architecture of Fn1 seen by 132 diffraction-limited microscopy with Fn1 nanoarchitecture, we plated Fn1^{mEGFP/+} MEFs on glass 133 for 16 hours, then fixed and stained cells using a monoclonal antibody to the central region of 134 Fn1. Fn1-mEGFP fluorescence was imaged in the TIRF mode at the critical angle of incidence, 135 while the binding of the monoclonal α Fn1 antibody was detected with Alexa Fluor-647conjugated secondary antibody and imaged by Stochastic Optical Reconstruction Microscopy 136 (STORM). STORM was performed by illuminating samples at the critical angle of incidence, as 137 138 described in (25). Thin beaded fibrillar adhesions (arrow in Fig. 4A) were resolved by STORM 139 (Fig. 4B-B2) to be arrays of regularly-spaced nanodomains that were symmetrical in x, y, and 140 z dimensions (Fig. 4B1, B2, B2' and Movie 4). The space between nanodomains contained a few or no Fn1 localizations (Fig. 4B2 and Movie 4). 141

Fn1 is a large, multi-domain, ~250 kDa glycoprotein secreted as a homodimer, wherein
 Fn1 subunits are linked in the anti-parallel orientation by two di-sulfide bonds at their C-termini

144 (1, 26). To investigate the relationship between the domain structure of Fn1 protein and the 145 nanodomain architecture of Fn1 fibrils, we used antibodies to distinct parts of Fn1 protein 146 (depicted in Model 1 at the bottom of Fig. 5,) and STORM. For these experiments, wild-type and Fn1^{mEGFP/+} MEFs were sparsely-plated in Ibidi 8-well glass-bottom chambers overnight. 147 148 allowing to observe the structure of thin fibrils. Cells were then fixed and stained with 149 antibodies recognizing different Fn1 epitopes: polyclonal rabbit antibodies raised to recognize 150 the first six type III repeats of Fn1 (Fn1 III₁₋₆) (27) (Fig. 5 column 1), a monoclonal antibody 151 recognizing an epitope within the central region of Fn1 (Fig. 5 columns 2, 4-5), polyclonal anti-152 serum 297.1 raised against the entire Fn1 protein (28) (Fig. 5, column 3), or polyclonal antibodies to GFP, recognizing the C-terminus of Fn1-mEGFP protein (Fig. 6, column 5). To 153 154 maximize the labeling density, the polyclonal anti-Fn1 II_{1-6} and 297.1 antisera were used at a 155 40- and 4-fold higher concentrations than for routine immunofluorescence microscopy. 156 respectively. Together with the use of excess secondary antibodies each conjugated to 3-6157 molecules of Alexa-647, this approach maximizes the chance that all the epitopes recognized by the 1° antibodies will be localized by STORM (25). 158 159 We first focused on the analyses of thin Fn1 fibrils like those marked by the arrows in 160 (Fig. 4A-B) to study the initiation of fibril formation rather than fibril bundling or branching. 161 Each antibody resulted in the detection of long characteristic Fn1 fibrils (Fig. 5A-C). 162 Successive magnifications show that independent of the antibody used for staining, Fn1 fibrils

163 contained linear arrays of nanodomains (thin arrows in **Fig. 5D**, **magnified in 5E**) separated

by regions containing a small number or no Fn1 localizations (as in **Fig. 4, Movie 4**).

165

166 Surprisingly, staining with the polyclonal anti-serum 297.1 raised to the entire Fn1 and,

presumably, recognizing multiple epitopes along Fn1, resulted in the same pattern of regularly-

spaced nanodomains as staining with antibodies recognizing distinct parts of Fn1 molecule

169 (Fig. 5, column 3), suggesting that each Fn1 localization contained the entire Fn1 sequence,

and that each nanodomain contained multiple Fn1 dimers.

171

172 These data are not consistent with previous models suggesting that Fn1 fibrils are composed 173 of periodically aligned Fn1 dimers arranged in an end-to-end fashion of alternating N- and C-174 termini (17-19) (**Model 1** at the bottom of **Fig. 5**). Such periodic alignment of Fn1 dimers 175 necessitates that staining using polyclonal antibodies would result in a uniform labeling of thin 176 Fn1 fibrils, as depicted in **Model 1**. The dimers in the **Model 1** are aligned in an end-to-end 177 fashion, according to the current Fn1 fibrillogenesis model, with the predicted overlap between 178 the N-terminal Fn1 assembly domain (blue) and the first six type III repeats of Fn1 (red) (21)). 179 If Fn1 fibrils, were indeed composed of continuous, linear arrays of dimers, the use of all four 180 depicted in **Model 1** would be predicted to uniformly label Fn1 fibrils.

181

To test this prediction, we used Fn1^{mEGFP/+} MEFs and a cocktail of four antibodies recognizing 182 the beginning (α Fn1 III₁₋₆), middle- (α Fn1 monoclonal), and the end (α GFP) of Fn1-mEGFP 183 protein in addition to all the epitopes recognized by 297.1 polyclonal antibody. The binding of 184 185 all the antibodies in the cocktail was detected by a cocktail of secondary antibodies that were each conjugated to 3 – 6 molecules of Alexa-647 (Fig. 5, column 7). The nanoarchitecture of 186 187 thin Fn1 fibrils, the nanodomain spacing (124 + 25 nm between nanodomains in fibrils, on 188 average), nanodomain size (average diameter 77 + 18 nm), and the number of Fn1 189 localizations per nanodomain (average of 80 + 43 localizations) detected by the cocktail of four 190 antibodies were indistinguishable from those produced by each type of the antibody 191 individually (Fig. 5D – E, quantified in Fig. 5F-H and Tables 1 – 3). These results indicate that 192 Fn1 fibrils are not uniform. Instead, our data show that Fn1 fibrils are arrays of nanodomains 193 containing multiple Fn1 dimers separated by areas containing a few or no Fn1 molecules (Fig.

5, Model 2). The nanoarchitecture of Fn1 fibrils following the treatment with 2% DOC which
removes cells and cellular components, was similar to untreated fibrils (Fig. 5, columns 4 and
8, Fig. 5F – H). These data indicate that the nanodomain architecture is a feature of mature
Fn1 fibrils.

198

199 Fn1 nanodomains were also present outside of fibrils, we term them non-fibrillar nanodomains (e.g. notched arrowheads in Fig. 4). About 3 – 5 such nanodomains are seen as a "bead" in 200 201 conventional diffraction-limited microscopy. Non-fibrillar Fn1 nanodomains contained a similar 202 number of Fn1 localizations per nanodomain and were of similar size compared with Fn1 nanodomains in fibrils (Fig. 5H, Tables 2 – 3). Staining using antibodies to endosomal and 203 204 lysosomal markers (Rab5 and LAMP1) showed no appreciable co-localization with Fn1 205 nanodomains (data not shown). Together with imaging using TIRF microscopy, these findings 206 indicated that Fn1 nanodomains in fibrils and non-fibrillar nanodomains are on the cell surface. 207 Non-fibrillar Fn1 nanodomains were present at cell periphery and throughout the cell surface, but they were not organized into linear arrays and were spaced at an median distance of 329 208 209 nm (Table 1).

210 Live imaging experiments suggested that Fn1 fibrils form from centripetally-translocating 211 Fn1 nanodomains originating at cell periphery (**Movie 2**). To understand the relationship 212 between the observed nanodomain architecture of Fn1 fibrils and the process of fibrillogenesis, we adopted a live imaging approach using Fn1^{mEGFP/+} MEFs and inhibitors of fibrillogenesis. 213 214 Fn1 fibrillogenesis critically depends on the interactions mediated by the N-terminal assembly 215 domain of Fn1 (domains shaded in blue in **Model 1, Fig. 5**), and inhibitors that interfere with these interactions block the formation of Fn1 fibrils (29-34). One such inhibitor is a 49-amino 216 acid peptide derived from Streptococcus pyogenes adhesin F1, termed the functional 217 upstream domain (FUD) (31). FUD binds the N-terminal assembly domain of Fn1 and 218 functions as a competitive inhibitor of Fn1-Fn1 interactions (31, 35). To determine how N-219

terminal interactions regulate Fn1 fibrillogenesis, Fn1^{mEGFP/+} MEFs were plated on glass for 4 220 221 hours, and then imaged for 15 - 18 hours either in the imaging medium alone, or in the medium containing either 225 nM FUD or 274 nM 11-IIIC, a 68 amino-acid control peptide that 222 223 does not interfere with Fn1 fibrillogenesis (32, 36). Untreated cells or cells treated with the 224 control peptide developed and accumulated long Fn1 fibrils (Movie 5). In contrast, treatment 225 with FUD led to dismantling of the pre-existing Fn1 fibrils and inhibited the formation of new 226 Fn1 fibrils (Movie 6). Instead of fibrils, cells cultured in the presence of FUD mainly contained 227 centripetally-moving Fn1-mEGFP fluorescent "beads" that only rarely formed strings (Movie 6). These experiments suggested that FUD inhibits fibrillogenesis by interfering with the process 228 229 by which Fn1 "beads" become arranged or connected into linear arrays. To test this 230 hypothesis, Fn1^{mEGFP/+} MEFs were plated for 16 hours in the continuous presence of either 231 225 nM FUD or 274 nM III-11C control peptides, or were left untreated. Cell were then fixed 232 and stained without permeabilization using monoclonal anti-Fn1 antibodies which were 233 detected with Alexa Fluor 647-conjugated secondary antibodies, and imaged at the critical angle of incidence by STORM at the excitation wavelength of 640 nm (25). This approach 234 235 abolishes the detection of intracellular Fn1-mEGFP and maximizes the detection of cell-236 surface Fn1 due to the following: a) the absence of a detergent during fixation, staining and 237 washing, b) detecting the emission of Alexa 647-conjugated antibodies at > 670 nm, c) the use 238 of oxygen scavengers in the STORM buffer that inhibits GFP fluorescence (37), and d) 239 imaging at the critical angle of incidence to detect fluorescence in close proximity to the 240 plasma membrane. These experiments demonstrated that the organization of nanodomains 241 into linear arrays was lost upon incubation with FUD (compare Fig. 6A, B, A1, B1 with Fig. 242 6C, C1). Non-fibrillar Fn1 nanodomains in cells treated with FUD had a similar number of Fn1 243 localizations per nanodomain, and were of similar sizes compared with fibrillar or non-fibrillar 244 Fn1 nanodomains in untreated cells or cells incubated with the control peptide (Fig. 6A2, 6B2, and 6C1-1, guantified in Fig. 6D and Tables 2-3). Taken together, these data indicate that 245

FUD does not interfere with the formation of Fn1 nanodomains but inhibits the organization of Fn1 nanodomains into linear arrays. Since Fn1 proteins lacking the N-terminal assembly domain do not form fibrils, our experiments suggest that interactions mediated by the Nterminal assembly domain of Fn1 are critical for the linking of Fn1 nanodomains into strings.

250

251 Discussion

252 In this manuscript, we describe the discovery of a novel mechanism underlying the process of 253 Fn1 fibrillogenesis. We found that the three-dimensional beaded architecture of Fn1 ECM fibrils in vivo and in two-dimensional cell culture is due to the presence of Fn1 nanodomains, 254 255 and showed that each Fn1 nanodomain is composed of multiple Fn1 dimers. Time-laps 256 imaging by confocal and TIRF microscopy showed that Fn1 fibrillogenesis initiated at cell periphery, as bright fluorescent "beads" of Fn1-mEGFP moved centripetally in parallel with F-257 258 actin and became organized into linear arrays. Interestingly, the beaded appearance of focal 259 and fibrillar adhesions has been noted before (13, 16), and the beaded architecture of cell adhesions can be seen in micrographs from multiple studies (e.g. Fig. 1C in (16), Fig 7a and 260 261 Sup. Figs 1a and b in (38); Fig. 8B in (39); and lower left cell in Fig. 3A (40). However, the 262 significance of these observations has not been investigated to the best of our knowledge. Fn1 263 domains and beaded strings in our movies resemble mobile, Fn1⁺ adhesions similar to those 264 described previously (13-16). The centripetal movement of these adhesions was dependent on the rearward actin flow, and the linkage of Fn1 to actin was mediated by integrin α 5 β 1 and 265 tensin (13, 15, 16). Our TIRF microscopy experiments are consistent with these studies and 266 267 show that integrin $\alpha 5\beta 1$ co-localizes with the regions of higher Fn1 intensity in fibrillar and nonfibrillar adhesions, and that $\alpha 5\beta 1^+$ focal and fibrillar adhesions are beaded. Taken together, our 268 studies suggest that Fn1 fibrils arise from small mobile nanodomains containing Fn1⁺ and 269 integrin $\alpha 5\beta 1^+$ that move toward the cell's center. This centripetal translocation of Fn1 270

271 nanodomains is coordinated with their organization into linear arrays, which become longer

and longer as more nanodomains are added.

273

274 The number of Fn1 localizations in nanodomains is independent of the antibody type or 275 amount used for staining, indicating that we are using saturating amounts of antibodies. 276 Together with the preservation of the nanodomain architecture upon the treatment with DOC, 277 our studies indicate that the non-uniform, beaded architecture is a salient feature of mature 278 Fn1 fibrils. Two pieces of evidence suggest that Fn1 nanodomains in fibrils are contiguous: 1) 279 the preservation of the linear organization and the nanoarchitecture of Fn1 fibrils after the treatment of cells with DOC which dissolves cell membranes, and 2) the presence of fibrous 280 281 material between immunogold densities in electron micrographs (18, 19, 41). The sparsity of 282 Fn1 localizations between Fn1 nanodomains in fibrils indicates that molecules other than or in 283 addition to Fn1 participate in the linking of Fn1 nanodomains into strings.

284

285 FUD peptide specifically binds to the N-terminal assembly domain of Fn1 (35, 42) and acts as 286 a competitive inhibitor of Fn1-Fn1 interactions (31, 43). In the absence of FUD, individual fibrils 287 in an established matrix are stable and can be tracked for over 16 hours (data not shown). 288 When FUD is added to cells, it specifically co-localizes with Fn1 fibrils and dismantles the 289 mature Fn1 ECM (31, 34), suggesting that the linear arrangement of Fn1 nanodomains in the 290 fibrillar ECM is maintained through dynamic interactions mediated at least in part by the Fn1 N-291 terminal assembly domain. Our live imaging experiments demonstrated that in addition to 292 dismantling pre-existing fibrils, FUD effectively blocks their de-novo formation. STORM 293 showed that FUD does not affect the formation of Fn1 nanodomains, instead, it blocks the 294 organization of Fn1 nanodomains into linear arrays. Together, these data suggest that FUD may block the dynamic interactions between the N-terminal Fn1 assembly domain and the 295 296 factor(s) linking Fn1 nanodomains into fibrils.

297

298 The beaded architecture of Fn1 ECM has important implications for the mechanisms of ECM 299 formation, remodeling and signal transduction. The tensile strength of knotted strings is 300 significantly lower than that of strings with uniformly-aligned fibers (44, 45), thus the beaded 301 architecture of Fn1 fibrils may facilitate their rupture under strain (46). The non-uniform, 302 nanodomain architecture of Fn1 may facilitate the accessibility of Fn1 fibrils to matrix 303 metalloproteases. In this model, degradation of Fn1 fibrils by metalloproteases may be 304 accomplished by cleaving between Fn1 nanodomains facilitating ECM remodeling. Finally, Fn1 305 is known to bind growth factors (47-49), and cell adhesion to ECM is known to orchestrate growth factor signaling (50). Thus, Fn1 nanodomains could serve as platforms for the binding 306 307 and presentation of concentrated packets of growth factors to cells, the organization of Fn1 308 nanodomains into closely-spaced arrays could further facilitate clustering and signaling by 309 growth factor receptors.

310

311 Acknowledgements

We thank Richard Hynes and Nathan Astrof for insightful discussions and careful reading of the manuscript, Sydney Astrof for help with data entry and encouragement, Patrick Murphy for endothelial cells, Richard Hynes for the gift of 297.1 antibody, and Tung Chan for help with setting up Western Blotting using ProteinSimple.

316

Sources of Funding This work was supported by the funding from the National Heart, Lung, and Blood Institute of the NIH R01 HL103920 and R01 HL134935 to SA, by the NIH Office of the Director R21 OD025323-01 to SA, by the pre-doctoral fellowship F31HL151046 to BEA, by the National Institute of General Medicine R35GM122505 to AK, by the National Institute of Arthritis and Musculoskeletal and Skin Diseases R01 AR073236 to JES, by the Faculty Seed Grant from the Center for Engineering MechanoBiology (CEMB), an NSF Science and

- 323 Technology Center, under grant agreement CMMI: 15-48571. Any opinions, findings, and
- 324 conclusions or recommendations expressed in this material are those of the authors and do
- 325 not necessarily reflect the views of the National Science Foundation.

326

327 Materials and Methods

328 Generation Fn1-fluorescent protein targeting constructs

- 329 Sequences of monomeric (m) green fluorescent protein (GFP), mNeonGreen, mScarlet-I, and
- tdTomato were obtained from FPbase (<u>https://www.fpbase.org</u>). The sequence encoding one
- 331 of the above fluorescent proteins (FPs) was knocked into the Fn1 locus following the last
- coding exon of mouse Fn1, and separated from the last coding amino acid by a flexible,
- 333 proline-rich linker, PPPELLGGP (51). Targeting was achieved by CRISPR/Cas9 (52). The
- 334 sequence of the guide RNA was chosen and off-target sites were identified using GuideScan
- and Off-Spotter software (53, 54). The guide RNA (gRNA) sequence 5'-AGC GGC ATG AAG
- 336 CAC TCA AT-3' targeting the last coding exon of *Fn1* was subcloned downstream the U6
- 337 promoter into the PX459 vector (Addgene, cat # 62988) encoding the Cas9-2A-Puromycin
- 338 cassette (52). The homology-directed repair (HDR) template was constructed using pBS-KS
- vector (Sup. Fig. 1a). The sequence of the last coding exon of *Fn1* 5'-
- 340 AACGTAAATTGCCCCATTGAGTGCTTCATGCCGCTAGATGTGCAAGCTGACAGAGACGAT
- 341 TCTCGAGAG-3' was modified to 5'-
- 342 AACGTAAATTGCCCCAT<u>c</u>GA<u>a</u>TGCTTCATGCCGCTAGATGTGCAAGCTGACAGAGACGATT
- 343 CTCGAGAG-3' in the HDR template by introducing silent mutations (underlined) to prevent
- targeting of the template by the gRNA. Homology arm 1 contained 677 bp encoding exon #45,
- intron, and a portion of the last exon (#46), of the transcript *ENSMUST00000055226.12*.
- Homology arm 2 encoded 1739 bp immediately downstream of the *Fn1* termination codon and
- included the unmodified 3'UTR of Fn1. Knockin Fn1^{mEGFP/+} mice were generated by

Biocytogen using the same HDR construct and a longer gRNA, 5'-TAG CGG CAT GAA GCA

349 CTC AAT <u>G</u>G-3',

- 350 targeting the same sequence in the last coding exon (differences between the two gRNAs are
- underlined). Targeting was confirmed by sequencing and Southern Blotting
- 352 (Sub. Fig. 1b). 500 bp around each of the top ten predicted off-target sites were sequenced
- and no mutations were found in the founder mice. Mice containing correctly-targeted *Fn1* locus
- were used to establish living colonies of Fn1^{mEGFP/mEGFP} animals. Mice were housed in an
- 355 AAALAC-approved barrier facility. All experimental procedures were approved by the
- 356 Institutional Animal Care and Use Committee of Rutgers University and conducted in
- 357 accordance with the Federal guidelines for the humane care of animals.
- 358

359 Generation of Fn1-FP-expressing cell lines

360 Mouse embryonic fibroblasts (MEFs) were isolated from embryonic day (E) 13.5 embryos

derived from the C57BL/6J strain (Jackson Labs, stock # 664) according to established

- 362 protocols (55) and cultured in high-glucose Dulbecco's Modified Eagle Medium (DMEM,
- 363 Corning, cat # 10-013-CV) supplemented with 10% v/v fetal bovine serum (Gemini

Biosciences, cat # 100-106), 1% v/v penicillin/streptomycin solution (GE Healthcare, cat

- 365 #SV30010), 1% v/v L-glutamine (Gibco, cat # 35050-061). We refer to this as complete
- 366 medium henceforth in the methods. CRISPR targeting was performed by transfecting the
- 367 PX459 plasmid encoding Fn1 gRNA and the HDR template using lipofectamine 3000, as
- described (52). For all other experiments, MEFs expressing Fn1-mEGFP proteins were
- 369 generated from E13.5 Fn1^{mEGFP/+} embryos. Wild-type MEFs were isolated from the littermates
 370 of Fn1^{mEGFP/+} embryos.
- 371

372 Analysis of Fn1 matrix assembly

373 Matrix Assembly was performed according to the established protocols (22). MEFs were plated in 6-well dishes (9 cm² growth area) at a density of 2×10^5 cells per well for 48 h, in complete 374 medium and incubated under sterile conditions at 37°C, 5% CO2. Cells were washed twice 375 376 with ice cold PBS (supplemented with Mg²⁺ and Ca²⁺), scraped with a cell scraper and lysed with either 500 ul RIPA lysis buffer pH 8.0 (50 mM Tris-Cl. 150 mM NaCl. 2 mM EDTA. 1% v/v 377 378 NP-40, 0.5% w/v sodium deoxycholate, 0.1% w/v SDS, 1X protease inhibitor cocktail (Cell 379 Signaling Technology, cat # 5871), or DOC lysis buffer, pH 8.8 (20 mM Tris-CI, 2 mM EDTA, 380 2% w/v sodium deoxycholate, 1X protease inhibitor cocktail (Cell Signaling Technology, 5871). Extracts were carefully transferred to Eppendorf tubes containing 1 µl (250 units) Benzonase® 381 Nuclease (Sigma-Aldrich, E1014), mixed by inverting a few times and incubated at 37 °C for 382 15 mins. The samples were then centrifuged at 16,000 × g for 15 min at 4 °C. For cells lysed 383 384 with DOC lysis buffer, the supernatant containing DOC-soluble material was carefully 385 removed, and the pellet containing the DOC-insoluble material was resuspended in 100 µl SDS solubilization buffer, pH8.8 (20 mM Tris-Cl, 2 mM EDTA, 1% w/v SDS, 1X protease 386 387 inhibitor cocktail (Cell Signaling Technology, 5871). The DOC-insoluble pellet was thoroughly 388 dissolved by heating the sample to 95 °C and vortexing. All samples were aliquoted and stored 389 at -80 °C until further use. Prior to guantification of Fn1 in the samples, the total protein 390 concentration of the RIPA and DOC lysates was determined using the BCA protein assay 391 (Pierce[™] BCA Protein Assay Kit, 23225). Fn1 and Fn1-FP fusion proteins were resolved using 392 66-440 kDa Wes separation module (ProteinSimple, SM-W007). Primary antibodies were used 393 at the following dilutions: anti-total Fn1 – 1:1000 (Abcam, ab199056), anti-GFP – 1:1000 394 (Roche, 11814460001), anti-mCherry – 1:1000 (Abcam, 167453). Primary antibodies were 395 detected using horseradish peroxidase-conjugated secondary antibodies (anti-Rabbit 396 Detection Module ProteinSimple, DM-001), and chemiluminescence was guantified using the

397 Compass for SW software (v3.1.8). Prior to running experimental samples, care was taken to 398 optimize the dilutions of lysates to be within the linear range of the detection.

399

400 Chemicals and reagents

401 Cells were grown in complete medium consisting of high-glucose Dulbecco's Modified Eagle 402 Medium (DMEM, Corning, cat # 10-013-CV) supplemented with 10% v/v fetal bovine serum 403 (Gemini Biosciences, cat # 100-106), 1% v/v penicillin/streptomycin solution (GE Healthcare, 404 cat #SV30010), 1% v/v L-glutamine (Gibco, cat # 35050-061). During live imaging cells were 405 incubated in FluoroBrite DMEM (Thermo Fisher Scientific, catalog # A1896701) supplemented with 2% v/v fetal bovine serum (Gemini Biosciences 100-106), 1% v/v penicillin/streptomycin 406 407 solution (GE Healthcare, SV30010), 1% v/v L-glutamine (Gibco 35050-061). We refer to this as 408 imaging medium in the methods henceforth. The pH of the imaging medium was 8.14. FUD 409 and III-11C peptides were generated as described (31, 36) and stored in PBS at -80° C. 4% 410 DOC solution was prepared by dissolving 0.4 g deoxycholate salt (Sigma, catalog # D6750) in 10 ml of imaging medium; the solution was then vortexed and filter sterilized. The pH of the 411 412 final solution was 8.01. 16% paraformaldehyde (PFA) (Electron microscopy Sciences; catalog 413 # 50-980-487) was diluted in 1x PBS to prepare 4% PFA. The 4% PFA solution was aliguoted 414 into 1 ml microfuge tubes, stored at -80° C, and thawed at 37° C immediately before use. 415 Triton X-100 (100X stock, Sigma-Aldrich, catalog # T-8787) was used to prepare 1X PBST by 416 diluting in 10X PBS (VWR, catalog # 76180-740). Blocking buffer was prepared by adding 5% 417 Donkey serum (Sigma-Aldrich, catalog # D9663) to 1X PBST. 5 mg/ml stock of DAPI (Fisher 418 Scientific, cat #D3571) was prepared in water and used at 1:300 dilution. Stain Buffer (cat # 419 554656 BD Pharmingen) was used for antibody dilutions and washing of cells that were 420 stained without permeabilization. Hoechst 33342 Trihydrochloride, Thermo Fisher, catalog # 421 H1399, Stock- 10mg/ml, was used for labelling live MEFs at 1:300 dilution. In live MEFs, F-422 actin was labelled using SiR actin (cat# CY-SC001 used at 1 μ M final concentration).

423 mCardinal-Lifeact-7 was a gift from Michael Davidson (Addgene plasmid # 54663 ;

424 http://n2t.net/addgene:54663 ; RRID:Addgene_54663). Vectashield antifade mounting medium

- 425 (Vectorlabs, catalog # H-1000), was used for cover slipping. STORM buffer was prepared
- 426 using 50 mM Tris-HCI (fisher scientific, catalog # T-395-1), pH 8.0, 10 mM NaCI (Sigma-
- 427 Aldrich, catalog # S-7653), 10% glucose (Sigma- Aldrich, catalog # G8270), 0.5 mg/ml glucose
- 428 oxidase (Sigma- Aldrich, catalog # G2133), 40 μg/ml catalase (Sigma- Aldrich, catalog # C40),
- 429 10 mM mercaptoethylamine (MEA, Sigma-Aldrich, catalog # 30070), according to (56).

430 Cell plating and chambers

All live imaging and STORM experiments were performed using Ibidi glass bottom 8-well

432 chambers (catalog # 80827) or MatTek round glass bottom dishes (catalog # P35G-1.5-14-C).

433 For imaging fixed cells, cells were plated on #1.5 round glass coverslips (Electron Microscopy

434 Sciences. Catalog # 72230-01). Coverslips were used either without coating or were coated

435 with the following ECM proteins: gelatin (Sigma Aldrich, catalog # G2500) (stock 0.1% in

436 distilled water), vitronectin (Sigma Aldrich, catalog # SRP3186; stock solution was prepared as

437 200µg/ml in 0.1% BSA and water) and laminin (R&D systems, catalog # 3400-010-02, stock 1

438 mg/ml was pipeted into 10ul aliquots and stored at -80°C). To coat with gelatin, glass surfaces

- 439 were incubated with the 0.1% gelatin solution for 5 min at room temperature (rt). To coat with
- 440 vitronectin or laminin, glass surfaces were incubated at 37° C for 1 hr in 20 μg/ml of either

vitronectin or laminin, excess liquid was removed, cover slips were rinsed once with 1X PBS,

- and blocked with 10 μ g/ml heat denatured BSA for 30 min before plating cells (14). Cells were
- grown in complete medium. During live imaging cells were incubated in imaging medium.

444 Antibodies

All primary antibodies were checked for specificity on genetically-null tissues: Fn1-null tissue
 sections obtained from Fn1-null embryos were used to assay the specificity of each of the anti-

- 447 Fn1 antibodies; Tissues isolated from GFP-null, Itga5-null, and mCherry-null embryos were
- 448 used to check the specificity of anti-GFP, anti-Itga5, and anti-mCherry antibodies. For each of
- the antibodies, staining of control tissues resulted in no more fluorescent signal than the
- 450 background fluorescence produced by the use of secondary antibodies only.
- 451

452 Antibodies

Primary Antibodies (Ab)	Source, catalog #,	Dilution / ng used per
	concentration	staining
tFn monoclonal Ab	Abcam, cat # 199056,	1:300 for STORM, 429 ng
	0.429 mg/ml	
tFn polyclonal Ab, 297.1	Richard Hynes lab	1:2000 for IF, 1:500 for
serum		STORM
Fn1 N-term, R184 serum	Jean Schwarzbauer's lab	1:2000 for IF, 1:25 for
		STORM
GFP	Aves lab, cat# GFP-1010,	1:300 for STORM / 30 ng
	30 μg/ml	
ltgα5	BD biosciences, cat #	1:100
	553319	
mCherry	Abcam, cat # ab167453	1:100

Secondary Antibodies	Source, catalog #,	Dilution/ μ g of antibody in
	concentration	staining solution)
donkey anti-rabbit Alexa	Thermo Fisher Scientific,	1:300 / 2 μg
Fluor 647	A-31573, 2 mg/ml	

donkey anti-mouse Alexa	Thermo Fisher Scientific,	1:300 / 2 μg
Fluor 555	A-31570, 2 mg/ml	
donkey anti-rat Alexa Fluor	Jackson Immunoresearch,	1:300 / 1.5 μg
555	712-166-150, 1.5mg/ml	
donkey anti-chicken Alexa	Jackson Immunoresearch,	1:300 / 1.5 μg
Fluor 488	703-546-155, 1.5mg/ml	
donkey anti-chicken Alexa	Jackson Immunoresearch,	1:300 / 1.5 μg
Fluor 647	703-606-155, 1.5mg/ml	

454

455 Cell culture and treatments

MEFs were maintained by plating on 25 cm² dishes (25 cm² growth area) in complete medium 456 457 and incubated under sterile conditions at 37°C, 5% CO₂. For FUD and III-11C treatment Fn1^{mEGFP/+} MEFs were plated in 8-well glass Ibidi dishes (1 cm² growth area) without coating at 458 459 a density of 0.6x10⁴ cells/well in complete medium. After 5 hours, DMEM was removed and 460 cells were rinsed once with 1X PBS. Subsequently, the medium was changed to imaging 461 medium. For FUD experiments, imaging medium was supplemented either with 225 nM FUD 462 or 274 nM of control III-11C peptide. Untreated wells contained cells incubated with imaging medium. Following the addition of the imaging medium (with or without the peptides), the 463 chamber was immediately set up for live imaging under in the humidified Tokai Hit stage-top 464 465 incubator at 37°C, 5% CO₂.

466

In order to enrich for non-fibrillar nanodomains, Fn1^{mEGFP/+} MEFs were plated in 8 well glass
Ibidi dishes (1 cm² growth area) without coating. Cell were plated at the density of 0.6x10⁴
cells/well in imaging medium with or without FUD (225 nM) or III-11C (274 nM), and incubated
in at 37°C, 5% CO₂ for 1 hr. Subsequently, MEFs were rinsed once in warm 1X PBS and fixed

using pre-warmed 4% PFA for 20 min. After fixation, wells were rinsed three times, 5 min each
with Stain Buffer (cat # 554656 BD Pharmingen), blocked for 30 min at room using 5% Donkey
serum prepared in Stain Buffer, and incubated with the monoclonal anti-Fn1 (Abcam, cat #
199056) overnight at 4° C. Cell were then rinsed with Stain Buffer three times, 10 min each,
and incubated with anti-rabbit antibodies conjugated with Alexa-647 for 1 hour at rt. Cell were
then rinsed again with Stain Buffer three times, 10 min each, and stored at 4° C in 1X PBS for
imaging later.

- 478
- 479 Hydrogels

Methacrylated Alginate Synthesis: Methacrylated alginate (MeAlg) was synthesized 480 according to a previously established protocol (57). In brief, alginic acid sodium salt from brown 481 482 algae (Sigma-Aldrich, USA) (3% w/v) was fully dissolved in Dulbecco's phosphate buffered 483 saline (dPBS, Sigma-Aldrich, USA). Then, methacrylic anhydride (Sigma-Aldrich, USA) (8% v/v) was added drop-wise to the alginate solution and stirred for 12 h at 4°C, using 2M NaOH (Sigma-484 Aldrich, USA) to ensure that the pH remained between 8 and 9 for the duration of the reaction. 485 486 The resulting solution was passed through filter paper (GE Whatman) and poured into 487 Spectra/Por dialysis membrane with a 6-8 kDa molecular weight cutoff (Fischer Scientific) and 488 kept in DIW under stirring for 7 days to eliminate the unreacted MA and salts. Dialyzed solution 489 was then freeze-dried for 4 days to obtain MeAlg foam.

Fabrication of the Hydrogel Substrates: MeAlg substrates were fabricated using a previously established protocol (58). Briefly, petri dishes with glass bottoms were treated with UV/ozone (UVO) for 30 minutes, immediately followed by a coating of 3-(trimethoxysilyl)propyl methacrylate (TMS) (Sigma-Aldrich, USA) to methacrylate the glass surfaces (59). The dishes were left in a desiccator overnight. The hydrogels were fabricated using Michael-type addition polymerization. First, 2-hydroxy-4'-(2-hydroxyethoxy)-2-methylpropiophenone (I2959) (Sigma496 Aldrich, USA), a photoinitiatior (0.5% w/v) was completely dissolved in Dulbecco's PBS (dPBS), 497 followed by the lyophilized MeAlg (3% w/v) synthesized previously. This was kept at room temperature until a clear solution was achieved. Crosslinking occurs with the introduction of DL-498 499 Dithiothreitol (DTT) (Sigma-Aldrich, USA) to the solution, along with 0.2M triethanolamine 500 (Sigma-Aldrich, USA) at pH 10. To form 3kPa and 12 kPa gels, 20% and 30% (w/v) DTT are 501 used, respectively. To promote cell adhesion, GRGDSPC peptide (1% w/v) (Genscript) was 502 added to the solution. After all contents were thoroughly mixed, 5µL of MeAlg solution was 503 pipetted onto the surface of the dish before being covered with a glass coverslip in order to 504 create gels less than 30µm thick. These were left at room temperature for an hour to crosslink 505 before being submerged in dPBS to remove the coverslip.

Atomic Force Microscopy: For stiffness measurements, hydrogel samples were submerged in dPBS and placed in a Dimension Icon AFM with ScanAsyst (Bruker). Using the PeakForce-QNM mode, hydrogel samples were indented using an MLCT-Bio probe tip with pyramidal geometry (Bruker, CA) and a nominal spring constant of 0.03 N/m, checked by thermal calibration.

511

512 Treatment of cells with Deoxycholate (DOC)

10⁴ Fn1^{mEGFP/+} MEFs were plated for 48 hrs in 8-well glass bottom lbidi dishes in complete 513 514 medium and incubated at 37° C and 5% CO₂. Two hours before imaging SiR-actin was added 515 at 1 µM final concentration. SiR-actin contains a far-red dye, silicon rhodamine, conjugated to 516 iasplakinolide that labels F-actin in live and fixed cells (60). Just before imaging, complete medium was replaced by 150 µl imaging medium containing 33 µg/ml of Hoechst 33342. 517 Positions were marked in each well and live imaging was initiated at 37°C and 5 % CO₂ 518 519 humidified chamber. After 15 min, 150 µl 4% DOC solution prepared in imaging medium 520 containing 33 µg/ml Hoechst was added to the experimental well (final pH 8.01) and 150 µl

- imaging medium containing 33 μg/ml Hoechst but without DOC was added to the control well.
 Cells were imaged at 50 sec intervals until F-actin and DNA disappeared (see Movie 4). The
- 523 medium was then removed, cells were rinsed for 1 min with 1X PBS pre-warmed to 37°C, fixed
- 524 with 4% PFA pre-warmed to 37°C, and stained to detect Fn1, as described below.
- 525

526 Immunofluorescence staining of permeabilized cells

- 527 MEFs were grown either on #1.5 round glass coverslips in 24-well dishes or in 8-well glass 528 Ibidi dishes depending on the experiment for the times indicated in figure legends. MEFs were then rinsed with 1X PBS (warmed to 37° C) for 5 min, fixed with freshly thawed 4% PFA pre-529 warmed to 37^o C for 20 min, and washed three times with 1X PBS (warmed to 37^o C) with mild 530 shaking. All subsequent washing steps were done with shaking. For permeabilization, cells 531 532 were washed once in 1X PBS containing 0.1% Triton- X 100 (PBST). Blocking was done for 30 533 min in 5% Donkey serum prepared in PBST (blocking solution). After blocking, cells were incubated in primary antibodies were diluted in blocking solution overnight at 4^o C, as specified 534 535 in the table above. This was followed by 3 washes in PBST for 10 min each. Cells were then 536 incubated with secondary antibodies diluted in PBST for 60 min at rt. Finally, cells were 537 washed three times with PBST for 10 min each. DAPI (1:300) was added to the second wash. 538 Cells were mounted using Vectashield.
- 539

540 **Imaging**

Fixed samples on coverslips and 8-well Ibidi dishes were imaged using Nikon A1-HD25
inverted confocal microscope with the DUG 4-Channel Detector and 2 GaAsP, 2 highsensitivity PMTs, and a motorized XYZ stage with Nikon's Perfect Focus 4 system. Plan Fluor
40x Oil (numerical aperture 1.3, cat # MRH01401) was used for live imaging, and CFI

545 Apochromat TIRF 100xC Oil objective with numerical aperture 1.49 (cat # MRD01905) was

546 used with the enhanced resolution protocol, TIRF, and STORM.

547

548 Confocal Settings

549 Confocal images of fixed samples were recorded using Nikon A1-HD25 inverted confocal
550 microscope equipped with CFI Apochromat TIRF 100xC Oil objective with the pinhole set to

- 551 0.8 Airy units, and imaged through 2 4 microns with step size of 0.125 μ m 0.15 μ m at a
- sampling of 40 nm per pixel. Crop function was used to reduce imaging time and sample
- 553 bleaching. Deconvolution was done using Nikon 3D deconvolution software (v5.11.01).
- 554 Airyscan imaging was performed using Zeiss LSM 880 fitted with a 32 array AiryScan GaAsP-
- 555 PMT detector and the Plan Apochromat 63X Oil (NA 1.4) objective. Deconvolution and pixel
- reassignment were done using Zeiss LSM software.
- 557

558 Live imaging

Ibidi 8-well glass-bottom slides or MatTek glass bottom dishes were placed into humidified Tokai Hit stage-top incubator maintained at 37° C and a 5% CO₂ atmosphere. mEGFP was excited using 488 nm laser at 1% power and pinhole set to 1 Airy unit. An optical zoom of 2 and Z step size of 0.5 µm were used, and stack size was set to 10-15 microns allowing to image the entire cell. For overnight movies, each position was filmed every 1.5 min – 4 min, as noted in Movie legends, for the DOC assay, imaging was performed at 54 sec intervals.

565

566 Movies

567 Movies in the mp4 format were generated using Imaris 9.5.1 (Bitplane), titles and arrows were 568 added using Adobe Premiere Elements Editor 2020.

570 **TIRF imaging**

TIRF microscopy was performed using Nikon A1-HD25 inverted confocal microscope equipped with 4 laser lines of 100mW per line at 405, 488, and 561nm and 125mW at 640nm, and motorized TIRF illumination. CFI Apochromat TIRF 100xC Oil objective and EMCCD camera were used. Before imaging lasers were aligned and the critical angle of incidence for imaging was determined by the software. The exposure time was 20 ms and readout speed was set at 10 MHz.

577

578 STORM imaging

579 Following IF staining, primary antibodies were detected using secondary antibodies coupled 580 with Alexa-647 fluorescent dve. Samples were washed and stored in PBS at 4°C. Prior to 581 imaging, freshly prepared STORM buffer was added and the chamber was immediately sealed 582 using parafilm. STORM was performed using Nikon A1-HD25 Ti2E microscope equipped with motorized TIRF illumination, 125mW 640 nm solid-state laser, Perfect Focus, and a 583 584 100x/1.49NA objective. Images were acquired at the critical angle of incidence and recorded 585 using a 512 x 512 EMCCD camera (Princeton Instruments). Calibration, drift correction, and z-586 rejection were based on the calibration file obtained by imaging of 100 nm Tetraspeck beads 587 (Life technologies, catalog # T-7279) using the same glass surface and buffer conditions. To 588 drive Alexa-647 into the dark state, samples were pre-bleached by the illumination at 640 nm 589 for 10 seconds at 100 % laser power. Images were acquired for 40,000 frames at 8.4 ms 590 exposure. Blinking events were fitted using the Nikon N-STORM localization software. Images 591 in which the Gaussian distribution of spot sizes was centered at 2 – 5 nm were used for further 592 analyses. Localization events with fewer than 800 or more than 50000 photons were filtered 593 out to remove blinking evens that were either too faint or too bright. In addition, blinking events 594 were filtered out if they occurred in more than 3 consecutive periods or where outside the z-

range determined by the calibration using 100 nm Tetraspeck beads. Images in which z-

rejection was below 50% were used for the analyses.

597

598 Analysis of the number of molecules within non-fibrillar and fibrillar Fn1 nanodomains 599 The free-hand ROI tool in the STORM window (Nikon Elements AR Software v5.11.01) was 600 used to draw ROI around nanodomains in fibrils or non-fibrillar nanodomains in a fibril to get a 601 count of molecule numbers. Fn1 nanodomains were analyzed in 5 random regions from 3 602 independently acquired images (a total of 15 fields) for each sample/antibody type. To 603 determine the molecule number in Fn1 nanodomains within fibrils, we analyzed more than 20 fibrils per antibody, from 3 or more independently acquired images. All the counts were plotted 604 605 in Prism 8.2.1 (GraphPad Software, USA), and compared using either one-way ANOVA test 606 with Tukey's correction or Kruskal-Wallis test with Dunn's correction for multiple testing.

607

608 Analysis of distance between fluorescence Fn1 nanodomains in fibrils Distances between nanodomains were quantified by measuring the distance between the centers of 609 610 nanodomain within fibrils using Nikon image analysis software and by Fiji peak analysis plugin, 611 which was done in the following manner: In order to quantify distance between nanodomains in 612 a fibril, a snapshot of the STORM window was generated. Rectangular ROI box was drawn 613 around a fibril and plot profiles were generated. We analyzed more than 20 thin fibrils from 3 or 614 more independently acquired images for each antibody type. Plot profiles were analyzed using 615 Find Peak function in the BAR module of Fiji (Tiago Ferreira et al., (2016) 616 10.5281/zenodo.28838). Default settings were used to generate lists of maxima and minima 617 for each plot. Lists were extracted to Microsoft Excel and distances between peak maxima 618 were computed. Distances measured manually and automatically were comparable and all

619 were plotted using Prism 8.2.1. See **Supplemental Figure 5a** for the automatic peak finding

620 workflow.

621

622 Analysis of distance between non-fibrillar nanodomains In order to quantify distance between non-fibrillar nanodomains in an image, a snapshot of the STORM window was 623 624 generated. Rectangular ROI box used for analysis of distance between density peaks in a fibril 625 was reloaded and positioned to capture as many non-fibrillar nanodomains as possible in a 626 rectangle. More than 5 regions from 3 or more independently acquired images were analyzed for Fn1^{mEGFP/+} MEFs immunostained with a cocktail of four antibodies, anti-N-terminal Fn1 627 628 antibody, monoclonal anti-tFn1, polyclonal anti-tFn1, and anti-GFP antibodies. Plot profiles were analyzed using Find Peak function in the BAR module of Fiji ((Tiago Ferreira et al., 629 (2016) 10.5281/zenodo.28838). Default settings were used to generate lists of maxima and 630 631 minima for each plot. Lists were extracted to Microsoft Excel and distances between peak 632 maxima were computed. Distances were plotted using Prism 8.2.1. See Supplemental Figure 633 **5B** for the workflow.

634

Analysis of area and diameter of non-fibrillar and fibrillar nanodomains Snapshots of 5 635 636 regions were generated from 3 independently acquired images and saved as .png files with 637 scale bars. Images were converted to 8-bit files and thresholded in Fiji. "Analyze particle" 638 function was used to extract the area for all nanodomains in the image. These values were 639 copied into an excel file where they were sorted from largest to smallest. Areas smaller than 10⁻⁴ µm² (~11.9 nm in diameter) were excluded from the analyses (see **Supplemental Figure** 640 641 6A-C). There were no statistical differences between non-fibrillar nanodomain areas of 642 untreated, FUD- or 11-IIIC treated cells without filtering (Supplemental Figure 6D-E). The 643 values for nanodomain diameters were extracted from the area measurements and confirmed 644 manually for a smaller sample, by measuring nanodomain diameters using the Nikon software.

645

647 Figure Legends

648 Figure 1. Beaded architecture of Fn1 fibrils in embryonic ECM. Wild-type E9.5 mouse 649 embryos were fixed and stained with the monoclonal antibody to Fn1 (white) and DAPI, and 650 imaged using 100x oil objective, N.A. 1.49, pinhole 0.8, and sampling rate of 40 nm/pixel. A -651 A1. Sagittal optical section through the first pharyngeal arch and b. the cardiac jelly, an ECM-652 rich region between the myocardial and endocardial layers of the outflow tract of the heart. 653 Large arrowheads in **A** – **A1** point to the ECM at the ectoderm-mesenchyme boundary of the 654 1st pharyngeal arch. The box in (A) is expanded in A1 to show the beaded Fn1 655 microarchitecture. Arrow in A1 points to Fn1 fibril within the arch mesenchyme; B. Beaded architecture of Fn1 fibrils in cardiac jelly, e.g., arrow. **C.** Intensity profile plot of a Fn1 fibril 656 657 shows a regularly-undulating profile, with peaks corresponding with Fn1 "beads"; a.u. arbitrary 658 units. 659 660 **Figure 2.** Integrin α 5 β 1 and Fn1 co-localize in beaded adhesions. Wild-type MEFs were

plated for 16 hours on glass coverslips, then fixed, stained with antibodies to Fn1 and integrin
α5 (Itga5), and imaged at the critical angle of incidence using 100x oil objective, NA 1.49. A –
A2 cell periphery. Arrows in A – A2 point to examples of non-fibrillar Fn1 adhesions ("beads")
at cell periphery. B - B2 medial portion of a cell containing beaded fibrillar adhesions (arrows).
Note that both Itga5 and Fn1 stainings are beaded. Magnifications in all panels are the same.

Figure 3. Beaded architecture of Fn1 fibrils is present in fibrils between cells and is retained in the absence of cell contact. A – B. Fn1 secreted by wild-type MEFs and deposited A) on glass or B) between cells. C. Fn1 fibrils between endothelial cells. Boxes in A– C are magnified in A1-C1. Arrows point to fibrils left behind on substrata (A1) or between cells (B1-C1). d. Fn1^{Neon Green} MEFs were treated with 2% DOC pH 8 until cells were dissolved

(see Movie 3). Following DOC treatment, Fn1 was imaged using 100x oil objective, NA 1.49,
pinhole size 0.8 Airy units, and sampling resolution of 40 nm/pixel in x,y. Note beaded
architecture (arrows).

675

676 Figure 4. Fn1 fibrils are composed linear arrays of Fn1 nanodomains. Fn1^{mEGFP/+} MEFs 677 were plated on glass for 16 hrs, then fixed and stained with antibodies to GFP followed by 678 secondary antibodies conjugated to Alexa-647. A. Native GFP fluorescence imaged by TIRF 679 using 100x objective NA 1.49. Arrow in **A** points to a thin, beaded Fn1 fibril resolved by 680 STORM in **B1**, **B1**' and **B2**. **B**. The entire field of view in **A** is imaged by STORM detecting Alexa-Fluor 647-conjugated antibody (see Methods). The box in **B** is expanded in **B1**. The box 681 682 in **B1** is expanded in **B2**. The arrow in **B1** points to the same region as the arrow in **A**. Arrows 683 in **B2** point to Fn1 nanodomains; arrowheads point to Fn1 localizations between the 684 nanodomains, notched arrowheads point to Fn1 nanodomains that are not in fibrils and wide arrows point to Fn1-free areas between the nanodomains in the fibril marked by the arrow in **A**. 685 **A. B. B1. B1**' and **B2** are in x-v planes: **B2**' shows the nanodomains in **B2** in the x-z plane. 686 687 Images in B1' and B2' are depth-coded according to the scale in B1'. See Movie 4 for 3D 688 rotation of the fibril underlined in **B1**' around the x-axis.

689

Figure 5. Fn1 fibrils are composed of nanodomains containing multiple Fn1 dimers.

691 Wild-type or Fn1^{mEGFP/+} MEFs were plated on glass for 16 hrs, fixed and stained with different

antibodies to Fn1 followed by Alexa 647-conjugated secondary antibodies. Columns showing

693 cells treated with DOC prior to fixation are marked. Cells were imaged using STORM. A.

2004 zoom-out views to show the overall appearance of Fn1 fibrils. **B-E.** Successive magnifications

of fibrils shown in (A). Arrows in D point to nanodomains magnified in E; arrowheads in D-E

point to Fn1 molecules between nanodomains, wide open arrows point to Fn1-free zones

697 between nanodomains in a fibril. F. distances between nanodomains within fibrils or non-

fibrillar (NF) nanodomains, **** p<10⁻⁴, Kruskal-Wallis test, with Dunn's correction for multiple 698 699 testing. Inset in **F** is a plot profile of the fibril marked by the box in the column **6D** showing regularly-spaced peaks of intensity. DOC-treated samples are marked. 4-antibody cocktail 700 701 contains four antibodies as in Model 1. G. Diameter of nanodomains in fibrils. H. number of 702 Fn1 localizations per nanodomain in fibrils. **Model 1**. Two Fn1 dimers in a fibril. Depiction of 703 antibody coverage of Fn1 fibrils if dimers were oriented in end-to-end fashion with alternating 704 N- and C-termini. Model 2. Our data show that Fn1 fibrils consist of a linear array of 705 nanodomains (black arrows) containing multiple Fn1 dimers (Fn1 dimers are depicted as balls 706 and color-coded according to the antibody scheme in **Model 1**. Arrowheads point to Fn1 and 707 open arrows point to Fn1-free areas between nanodomains.

708

709 Figure 6. The N-terminal Fn1 assembly domain regulates the organization of Fn1

nanodomains into linear fibrillar arrays. Fn1^{mEGFP/+} MEFs were plated on glass and were 710 711 either left untreated, or were incubated with the control 11-IIIC peptide, or the FUD peptide for 16 hrs. Cell were then fixed and stained with the monoclonal antibody to Fn1 followed by Alexa 712 713 647-conjugated secondary antibodies. Cells were imaged at the critical angle of incidence by 714 STORM. A – A2. Untreated, unpermeabilized cells. B – B2. Cells incubated with controls 11-715 IIIC peptide. C – C1, C1-1 FUD-treated, unpermeabilized cells. Boxes marked 1 in A-B were 716 expanded in A1-B1. Boxes marked 2 in A-B, were expanded in A2 – B2. The box in C is expanded in C1 and C1-1. Arrows in A1-B1 point to Fn1 nanodomains (NDs) in fibrils. Arrows 717 718 in **C1** points to non-fibrillar nanodomains expanded in **C1-1**; **D.** Quantification of the number of 719 Fn1 localizations in NDs in fibrils and in non-fibrillar NDs after various conditions. Red lines 720 mark medians. Differences are not statistically significant, Kruskal-Wallis test with Dunn's 721 correction for multiple testing. E. Model of fibril formation: Fn1 dimers assemble into small 722 nanodomains containing integrin α 5 β 1 at cell periphery, move rearward with actin flow, and

- 523 become organized into linear arrays of nanodomains. Joining of the additional Fn1
- nanodomains to these arrays leads to the generation of longer fibrils as the assembly moves
- toward the cell center. FUD does not interfere with the formation of Fn1 nanodomains. But
- instead, it blocks the organization of Fn1 nanodomains into linear arrays.
- 727
- 728 Legends for Movies
- Movie 1. Rotational views through the Fn1+ ECM in the cardiac jelly. Whole E9.5 embryo
 was stained using rabbit monoclonal anti-Fn1 antibody and imaged using 100x objective, N.A.
 1.49, with the pinhole set at 0.8 Airy units, and sampling of 40 nm per pixel in x, y. The movie
 shows 3D reconstruction through 3.4 µm of tissue sampled every 0.121 µm in z. Fn1 is in
- white, DAPI is in blue. Arrows point to examples of beaded Fn1 fibrils.
- 734

Movie 2. Fn1 fibrillogenesis imaged by TIRF microscopy. Fn1^{mEGFP} MEFs were transiently
transfected with mCardinal-lifeact, plated on gelatin-coated glass cover slips, and imaged 48
hours later. Filming was done every 2 min for 30 min using TIRF and 100x objective, N.A.
1.49. The first set shows Fn1-mEGFP channel. Yellow arrows point to centripetally-moving
Fn1 nanodomains that appear to be organizing into an elongating linear fibril. The second set
is an overlay between Fn1-mEGFP and mCardinal-lifeact.

741

Movie 3. 2% DOC dissolved cytoplasm and nucleus in under 13 min leaving Fn1 fibrils.
MEFs expressing Fn1-mEGFP were plated on glass-bottom slides and labeled with SiRActin
(magenta) to visualize F-actin and Hoechst (blue) to visualize DNA. Time lapse was recorded
every 54 sec immediately following the addition of DOC solution, pH 8.01, to live cells. The
addition of 2% DOC dissolves actin cytoskeleton and nuclei, and leaves Fn1 ECM fibrils

- (green). Fn1 fibrils collapse following the dissolution of the actin cytoskeleton due to the loss oftension.
- 749

Movie 4. 3D-rendering of STORM data. Fn1 fibril underlined in Fig. 4b1 is rotated around the
x-axis to show the arrangement of Fn1 localizations in 3D. The movie starts in x-y plane.
Yellow arrows point at Fn1 nanodomains in the fibrils. Red arrows point to the space between
the nanodomains.

754

Movie 5. Cells incubated with 11-IIIC, show robust fibrillogenesis. Fn1^{mEGFP} MEFs were plated on glass in 8-well Ibidi chambers for 4 hours, Medium containing 11-IIIC control peptide was then added and cells were filmed every 90 sec for about 15 hours, as described in Methods. The movie begins approximately 30 min after the 11-IIIC-containing medium was added, the time it takes to set up time-laps recording. Arrows point to cell periphery and examples of centripetally moving Fn1 fibrils.

761

762 **Movie 6. FUD interferes with linking centripetally moving Fn1+ nanodomains into fibrils.**

Fn1^{mEGFP} MEFs were plated on glass in 8-well Ibidi chambers for 4 hours. Medium containing
FUD peptide was then added and cells were filmed every 3 min for about 15 hours, as
described in Methods. The movie begins approximately 30 min after the FUD-containing
medium was added, the time it takes to set up time-laps recording. Note the dismantling of preexisting fibrils at the beginning of the movie. Yellow and red arrows point to cell periphery.
Note the presence of centripetally moving Fn1-mEGFP "beads" and the scarcity of Fn1 fibrils
for the majority of the duration of the movie.

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771 **References**

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Table 1

Distance between Fn1-containing nanodomains in fibrils and in non-fibrillar nanodomains

Cell type antibody	Median, nm	Interquartile range, nm	n
Distance be	etween nanodomains	in fibrils, nm	
Wild-type MEFs α III ₁₋₆ Fn1 (R184 ab)	148.5	108 – 187	202
Wild-type MEFs monoclonal α Fn1	140	105 – 166	134
Wild-type MEFs polyclonal 297.1	145	107 – 193	159
$\frac{\text{DOC-treated}}{\text{Wild-type MEFs}}$ monoclonal α Fn1	123	83 – 160	148
${\sf Fn1}^{{\sf mEGFP}/{\scriptsize +}}$ monoclonal $lpha$ Fn1	124	99 – 166	120
Fn1 ^{mEGFP/+} α GFP	115	89 – 155	142
Fn1 ^{mEGFP/+} 4-antibody cocktail	114	82 - 156	106
Distance between non-fibrillar nanodomains, nm			
<u>Non-fibrillar nanodomains</u> Fn1 ^{mEGFP/+} 4-antibody cocktail	329	260 – 410	132

Table 2

Area of Fn1 nanodomains

Cell type antibody	Median 10 ⁻³ μm²	Interquartile range, 10 ⁻³ μm ²	n
Area of	nanodomains in fibri	ils, 10⁻³ μm²	
Wild-type MEFs α III ₁₋₆ Fn1 (R184 ab)	5	3 – 7	60
Wild-type MEFs monoclonal α Fn1	5	3 – 6	66
Wild-type MEFs polyclonal 297.1	4	3 – 6	62
Fn1 ^{mEGFP/+} monoclonal α Fn1	3	2 – 5	65
Fn1 ^{mEGFP/+} α GFP	4	2 – 6	63
Fn1 ^{mEGFP/+} 4-antibody cocktail	4	3 – 6	62
Area of non-fibrillar nanodomains, 10 ⁻³ μm ²			
Untreated Fn1 ^{mEGFP/+} monoclonal α Fn1 <mark>16 hr plating</mark>	4.3	3– 6	127
11-IIIC treated Fn1 ^{mEGFP/+} monoclonal α Fn1 16 hr plating	3	2.4 - 4	93
FUD-treated Fn1 ^{mEGFP/+} monoclonal α Fn1 16 hr plating	4	3 – 6	666

Table 3

Number of localizations per Fn1 nanodomain

Cell type antibody	Median	Interquartile range	n	
Number of Fr	Number of Fn1 localizations per nanodomain within fibrils			
Wild-type MEFs α III ₁₋₆ Fn1 (R184 ab)	72	46.5 - 112	345	
Wild-type MEFs monoclonal α Fn1	60	43 – 81	271	
Wild-type MEFs polyclonal 297.1	66	42 – 106.5	381	
DOC-treated Wild-type MEFs monoclonal α Fn1	72	46 – 113	112	
${\sf Fn1}^{{\sf mEGFP/+}}$ monoclonal $lpha$ Fn1	63	45.25 – 114	216	
Fn1 ^{mEGFP/+} α GFP	61	39 – 92	351	
Fn1 ^{mEGFP/+} 4-antibody cocktail	73	50 – 103	314	
Number of Fn1 localizations per non-fibrillar nanodomain				
Fn1 ^{mEGFP/+} 4-antibody cocktail	58.5	39 – 95	62	
Untreated Fn1 ^{mEGFP/+} monoclonal α Fn1 Unpermeabilized cells	45	33 – 68	127	
11-IIIC treated Fn1 ^{mEGFP/+} monoclonal α Fn1	46	29 – 68.5	93	
FUD-treated Fn1 ^{mEGFP/+} monoclonal α Fn1 Unpermeabilized cells	54.5	37 – 92	102	



Figure 1: Beaded architecture of Fn1 fibrils in embryonic ECM wild spie EIS mouse embryos were fibre and statmet with the monotenial antibody to Fn1 (which) and CMPI, and marging integed and fibro and objective. At 1.46, periode al. And sampling rate of 40 pharyogen at 1. and 8. the backtop lay, and EdMeth region between the mycoardial and endoardaul layers of the outdow toat of the hant. Large arrowhards in A – A1 park to the ECM at the coldown reservicement boundary of the "thytharyogel act. The box in A1 is expanded in A1 to draw Fn1 microarchitecture. Arrow in A parties 16⁻¹ first within the act of the hant. EdM at the box in A1 is expanded in A1 to draw Fn1 microarchitecture. Arrow in A parties 16⁻¹ first within the act microarchitecture action profiles pid of a Fn1 fibri allows a regularly-undulating profiles, with trouch well above the roles au, authorizy units.



Figure 2. Integrin 45 and Frit co-localize in beaded adhesions. With-type MEFs were plated for 16 hours on glass coversitios, them fixed, statement with antibudies to Fin 1 and integrin or 6 (tigads), and imaged at the critical angle of incidence using 100x oil objective, NA 1.49. A - A 2 cell periphery. Arrows in A-A2 point at examples of non-fibrillar Frit adhesions. B - B2 medial portion of a cell cortaining beaded fibrillar adhesions (arrows). Mignifications in all panels are the same.

Figure 3



Figure 3. Beaded architecture of Fn1 fibrila is retained in the absence of cell contact. A -B. Fn1 scored by wild type MEFs and deposited A) on glass or B) between cells. C. Fn1 fibrils between endothelial cells. Boxes in A - C are magnified in A1-C1. Arxws point to fibrils site behalind on subtrastic A(1) or between cells (B1-C1). E. Fn1 fibril following DOC treatment was imaged using 100x oil objective, NA1.40, pinhole size 0.8.4 multis, and sampling resolution of 40 nmbjoxel in xy. Note beaded architecture (arrow). Solubilization of cell components is shown in Mové 4.



Figure 4. Fn1 fibrils are composed linear arrays of Fn1 nanodomains. Ent^{reactive}, MEFs were plated on glass for 18 hrs; then fixed and stained with antibodies to GFP followed by accordary antibodies conjugated to Alexa-647. A. Native GFP fluorescence imaged by TIRF using 100x objective NA 1.49. Arrow in a points to a thin, beaded Fn1 fibril resolved by STORM in B1 and B2. B. The entire field of view in a is imaged by STORM detecting Alexa-Fluor 647-conjugated antibody (see Methods). The box in B is expanded in B1. The box in B1 is expanded in B2. The arrow in B1 points to the same region as the arrow in A. Arrows in B2 point to Fn1 nanodomains; arrowheads point to Fn1 localizations between the nanodomains, notched arrowheads point to Fn1 nanodomains that are not in fibrils, and wide arrows point to Fn1-free areas between the nanodomains the fibril marked by the arrow in A. A, B, B1 and B2 are in x-y planes; B2' is in x-z plane, Images in B1' and B2' are depth-coded according to the x-ate in B1'. See Movie 4 (or 30 zotation of the fibril underlined in B1' and B2 are



Model 2 Fn1 fibrils form from nanodomains containing multiple Fn1 dimers



Figure 5. Wild-type or Fn1mEGFP/+ MEFs were plated on glass for 16 hrs, fixed and stained with different antibodies to Fn1 followed by Alexa 647conjugated secondary antibodies. Columns showing cells treated with DOC prior to fixation are marked. Cells were imaged using STORM. A. zoomout views to show the overall appearance of Fn1 fibrils. B-E. Successive magnifications of fibrils shown in (A). Arrows in D point to nanodomains magnified in E; arrowheads in D-E point to Fn1 molecules between nanodomains, wide open arrows point to Fn1-free zones between nanodomains in a fibril. F. distances between nanodomains within fibrils or non-fibrillar (NF) nanodomains,

**** p<10-4. Kruskal-Wallis test. with Dunn's correction for multiple testing. Inset in F is a plot profile of the fibril marked by the box in the column 6D showing regularly-spaced peaks of intensity. DOC-treated samples are marked. 4-antibody cocktail contains four antibodies as in Model 1. G. Diameter of nanodomains in fibrils. H. number of Fn1 localizations per nanodomain in fibrils. Model 1. Two Fn1 dimers in a fibril. Depiction of antibody coverage of Fn1 fibrils if dimers were oriented in end-to-end fashion with alternating N- and Ctermini. Model 2. Our data show that Fn1 fibrils consist of a linear array of nanodomains (black arrows) containing multiple Fn1 dimers (Fn1 dimers are depicted as balls and color-coded according to the antibody scheme in Model 1. Arrowheads point to Fn1 and open arrows point to Fn1-free areas between nanodomains.

Model 1 Two Fn1 dimers aligned in a periodical end-to-end fashion. If fibrils were composed of periodically aligned dimers, they would be uniformly stained by the antibody cocktail:





Figure 6. The N-terminal Fn1 assembly domain regulates the organization of Fn1 nanodomains into linear fibrillar arrays. Fn1mEGFP/+ MEFs were plated on glass and were either left untreated, or were incubated with the control 11-IIIC peptide, or the FUD peptide for 16 hrs. Cell were then fixed and stained with the monoclonal antibody to Fn1 followed by Alexa 647-conjugated secondary antibodies. Cells were imaged at the critical angle of incidence by STORM. A -A2. Untreated, unpermeabilized cells. B - B2. Cells incubated with controls 11-IIIC peptide. C - C1, C1-1 FUD-treated, unpermeabilized cells. Boxes marked 1 in A-B were expanded in A1-B1. Boxes marked 2 in A-B, were expanded in A2 - B2. The box in C is expanded in C1 and C1-1. Arrows in A1-B1 point to Fn1 nanodomains (NDs) in fibrils. Arrows in C1 points to non-fibrillar nanodomains expanded in C1-1; D. Quantification of the number of Fn1 localizations in NDs in fibrils and in non-fibrillar NDs after various conditions. Red lines mark medians. Differences are not statistically significant, Kruskal-Wallis test with Dunn's correction for multiple testing. E. Model of fibril formation: Fn1 dimers assemble into small nanodomains containing integrin a5ß1 at cell periphery, move rearward with actin flow, and become organized into linear arrays of nanodomains. Joining of the additional Fn1 nanodomains to these arrays leads to the generation of longer fibrils as the assembly moves toward the cell center. FUD does not interfere with the formation of Fn1 nanodomains. But instead, it blocks the organization of Fn1 nanodomains into linear arrays.

Èun

. Fn1 dimen's)