FAK promotes recruitment of talin to nascent adhesions to control cell motility

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Cell migration is a dynamic process that involves the continuous formation, maturation, and turnover of matrix-cell adhesion sites. New (nascent) adhesions form at the protruding cell edge in a tension-independent manner and are comprised of integrin receptors, signaling, and cytoskeletal-associated proteins. Integrins recruit focal adhesion kinase (FAK) and the cytoskeletal protein talin to nascent adhesions. Canonical models support a role for talin in mediating FAK localization and activation at adhesions. Here, alternatively, we show that FAK promotes talin recruitment to nascent adhesions occurring independently of talin binding to β1 integrins. The direct binding site for talin on FAK was identified, and a point mutation in FAK (E1015A) prevented talin association and talin localization to nascent adhesions but did not alter integrin-mediated FAK recruitment and activation at adhesions. Moreover, FAK E1015A inhibited cell motility and proteolytic talin cleavage needed for efficient adhesion dynamics. These results support an alternative linkage for FAK–talin interactions within nascent adhesions essential for the control of cell migration.

Introduction

Integrin-based adhesion assembly and turnover are highly dynamic events underlying cell movement (Parsons et al., 2010). Integrin signals control cell cycle progression, cell survival, and cell polarity, and mediate a linkage to the actin cytoskeleton in the control of cell shape. In spreading cells, edge protrusions are connected to the underlying extracellular matrix via integrin and cytoskeletal structures, termed nascent adhesions, that form concurrently with lamellipodium protrusion (Gardel et al., 2010). Nascent adhesions either disassemble or mature into focal adhesions in part by myosin II-dependent tension generation and actomyosin contractility (Choi et al., 2008). Signaling scaffolding proteins such as FAK, paxillin, RACK1, and talin are rapidly recruited to nascent adhesions (Serrels et al., 2010; Choi et al., 2011). Despite knowledge on protein composition within adhesions, how these sites are formed and remodeled during cell movement remains under investigation.

Talin is a large cytoskeletal protein comprised of an N-terminal head or a band 4.1, ezrin, radixin, moesin homology (FERM) domain that binds to β1 and β3 integrin cytoplasmic tails, type 1y phosphatidylinositol phosphate (PIP) kinase, and FAK (Calderwood et al., 1999; Di Paolo et al., 2002). The C-terminal talin rod domain binds vinculin and actin, and contains a second integrin-binding site (Critchley and Gingras, 2008). FAK is a cytoplasmic tyrosine kinase that phosphorylates targets such as paxillin in the regulation of adhesion dynamics (Schaller, 2010). FAK is one of the first cytoplasmic proteins recruited to clustered integrins in a tyrosine phosphorylation-independent manner (Miyamoto et al., 1995), and FAK-paxillin binding occurs within nascent adhesions (Choi et al., 2011). Although canonical models also support the importance of talin in the recruitment and activation of FAK at adhesions, FAK connections to talin remain uncharacterized (Chen et al., 1995; Zhang et al., 2008; Frame et al., 2010). As recent three-dimensional nanoscale fluorescent microscopy has co-localized FAK, paxillin, integrin tails, and talin FERM (head domain) to a proximal signaling layer at adhesions followed by the C-terminal part of talin (rod domain) and vinculin localized to a more distal layer (Kanchanawong et al., 2010), we set out to test the importance of FAK–talin binding at nascent adhesions.

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Results and discussion

FAK controls talin association with nascent adhesions

FAK knockout (FAK−/−) results in an early embryonic lethal phenotype, and FAK−/− mouse embryonic fibroblasts (MEFs) form an abundance of adhesions limiting cell movement (Sieg et al., 1999). To study new or “nascent” adhesion formation, FAK−/− and normal (FAK+/+) MEFs were trypsinized, held in suspension, and then replated onto fibronectin (FN)-coated glass slides in the absence of serum to synchronize adhesion site formation (Figs. 1 and S1). MEF binding to FN occurs rapidly, and by 15 min, MEFs were spread, and possessed cortical actin rings with low levels of vinculin staining within peripheral paxillin-containing nascent adhesions (Fig. S1 A). Talin strongly colocalized with FAK within nascent FAK+/+ MEF adhesions at 15 min (Fig. 1, A and B), and a talin–FAK complex was detected in cell lysates (Fig. 1 C). Strikingly, talin was not localized to nascent FAK+/− MEF adhesions formed at 15 min even though talin expression was equivalent in FAK−/− and FAK+/− MEFs (Fig. 1, A–C). Similar results were obtained upon transient FAK knockdown in human ovarian SKOV3.ip1 carcinoma cells, resulting in the significant reduction of talin–paxillin colocalization within nascent adhesions (Fig. S1, C and D). However, by 60 min on FN, more mature vinculin-containing adhesion sites with integrated actin stress fibers are formed within MEFs (Fig. S1 B) and SKOV3.ip1 cells, and the absence of FAK did not prevent talin localization to these sites (not depicted). Together, these results show that nascent adhesions can form without FAK and that talin recruitment to these sites is enhanced by FAK expression.

FAK localizes to nascent adhesions independent of talin and adhesion maturation

Previous studies showed that knockdown of talin-1 and talin-2 within MEFs did not prevent initial adhesion formation or spreading on FN, but resulted in reduced integrin activation and mature adhesion formation (Zhang et al., 2008). Interestingly, β1 integrin was localized (not depicted) and activated within nascent adhesions formed at 15 min within FAK−/− and FAK+/− MEFs (Fig. 1 D). Thus, β1 integrin is activated and localized to nascent adhesions without FAK and with limited talin involvement. Talin-1 is the primary isoform expressed in human umbilical vein endothelial cells (HUVECs), and talin-1 knockdown does not alter the kinetics of initial HUVEC spreading at 15 min, but results in reduced cell area and adhesion formation after 1 h (Kopp et al., 2010). Transfection of HUVECs with anti–talin-1 siRNA results in ~90% knockdown of talin without effects on FAK or paxillin expression (Fig. 1 E). HUVEC spreading on FN at 15 min was equivalent between control- and talin siRNA–expressing cells (Fig. 1 F), and FAK equally colocalized with paxillin at nascent adhesions in talin siRNA–expressing HUVECs (Fig. 1 G). Moreover, the addition of blebbistatin, a pharmacological inhibitor of myosin II that prevents adhesion maturation (Choi et al., 2008), did not prevent FAK colocalization with paxillin or FAK activation upon MEF adhesion to FN (Fig. S2, A and B). FAK transit to nascent adhesions is mediated in part by paxillin binding (Scheswolh et al., 2008), but this is not dependent on talin expression or adhesion maturation.

Integrin-talin binding is not required for talin recruitment to adhesions

Our results with FAK−/− MEFs and FAK knockdown in SKOV3.ip1 cells support an alternative model whereby FAK facilitates talin recruitment to nascent adhesions. Because talin can bind directly to the cytoplasmic domain of β integrins (Calderwood et al., 1999) and this could link talin to nascent adhesions, FAK–talin recruitment analyses were performed in cells expressing chimeric integrins that support direct talin binding (β1A wild type [WT]) or contain a point mutation in the cytoplasmic domain (β1A Y783A) disrupting talin binding (Fig. 2 A; Nieves et al., 2010). Chimeric integrin-mediated cell adhesion is initiated by plating cells on fibrinogen (FB), and equal numbers of adhesion sites are formed at 15 min in WT and β1A Y783A-expressing cells (unpublished data). At 60 min, β1A Y783A adhesion sites are few in number because of decreased stability (Nieves et al., 2010). Therefore, analyses were performed at 15 min on FB whereby a talin, FAK, and chimeric WT integrin complex was detected by coimmunoprecipitation and was colocalized to nascent adhesions (Fig. 2, B–D). Importantly, in β1A Y783A-expressing cells on FB for 15 min, talin and FAK associate (Fig. 2 B) and are colocalized to adhesions (Fig. 2, C and D) even though β1A Y783A mutation prevents direct talin binding. This result shows that direct talin binding to β1 integrin is not essential for talin recruitment to nascent adhesions.

Identification of the talin-binding site on FAK

Previous deletion studies identified the FAK C-terminal region (965–1012) and the talin FERM F3 lobe as required for binding (Chen et al., 1995; Borowsky and Hynes, 1998). Pull-down experiments confirmed that FAK 947–1,052 could bind full-length talin independently of paxillin (Fig. 3 A) and that FAK 1,011–1,042 was a minimal region that bound to the talin F2-F3 FERM domain (Fig. 3 B). A direct binding assay using recombinant full-length or C-terminal FAK (853–1,052) and a GST fusion protein of the talin FERM F2-F3 region revealed that FAK–talin interactions were disrupted by talin FERM F3 lobe W359A and A360E mutations (Fig. S3, A and D). As K357A or R358A talin FERM F3 lobe mutations had no effect on FAK–talin interactions (Fig. S2, E and F), these results support FAK binding to talin FERM F3 lobe at a site that may overlap with β integrins (Bouaouina et al., 2008).

Alanine scanning mutagenesis was used to determine residues within FAK 1,011–1,042 required for talin FERM F2-F3 binding (Fig. 3 C). FAK E1015A mutation prevented talin binding. FAK E1015 is a conserved and surface-exposed residue within the FAK focal adhesion targeting (FAT) domain (Fig. 3 D) that does not directly participate in paxillin binding (Hayashi et al., 2002; Gao et al., 2004). In a direct binding assay, in vitro translated full-length WT but not FAK E1015A formed a complex with the GST-talin F2-F3 FERM domain.
FAK is required for talin recruitment to nascent adhesions. (A) FAK+/+ and FAK−/− MEFs were serum starved, plated onto FN-coated coverslips for 15 min, and costained with antibodies to talin (green) and paxillin (red). The merged image shows colocalization (yellow). Inset, enlarged area of peripheral adhesion staining (circled). Bars, 10 µm. (B) Cells were analyzed for talin and paxillin colocalization at 15 min on FN (**, P < 0.01). (C) Co-immunoprecipitation of FAK with antibodies to talin after cell replating for 15 min on FN. FAK+/+ and FAK−/− MEF lysates were analyzed for talin, FAK, and actin expression. (D) Analyses of active β1 integrin (9EG7; red) and paxillin (green) staining of FAK+/+ and FAK−/− MEFs plated on FN for 15 min. Bars, 5 µm. (E) Control or anti-talin-1 siRNA transfection of HUVECs and immunoblotting for talin (antibody detects talin-1/-2), FAK, paxillin, and actin expression. Molecular mass is indicated in kilodaltons. (F) Control or anti-talin siRNA transfected HUVECs were replated on FN for 15 min and costained with antibodies to paxillin (green) and FAK (red). Merged image shows colocalization (yellow) and siGlo (blue) transfection marker. Bars, 10 µm. (G) Cells were analyzed for paxillin and FAK colocalization at 15 min on FN. Co-localization was measured on a pixel-by-pixel basis within all adhesions in at least 20 cells per experimental group encompassing at least three independent experiments. Box and whisker plots show the distribution of the data: black square, mean; bottom line, 25th percentile; middle line, median; top line, 75th percentile; whiskers, fifth and 95th percentiles.
talin but not paxillin colocalization with GFP-FAK E1015A at adhesions compared with MEFs expressing GFP-FAK WT (Fig. S3, B–D). Partial talin localization to mature adhesions in FAK E1015A MEFs is consistent with alternative localization mechanisms (Franco et al., 2006; Wang et al., 2011). However, as FAK E1015A expression prevents talin recruitment to nascent adhesions, this supports the importance of FAK–talin binding in this process.

FAK E1015A is activated by FN replating but does not rescue adhesion or motility defects of FAK−/− MEFs

FAK is activated in a cell adhesion–dependent manner as detected by phospho-specific antibodies to the FAK Y397 auto-phosphorylation site upon FN replating (Schaller, 2010). Although FAK WT and E1015A were not phosphorylated under suspended cell conditions, both were rapidly phosphorylated at Y397 within 15 min on FN and remained activated at
Figure 3. Talin FERM binds to FAK residues 1,011–1,042 within the FAT domain. [A] Transient transfection of the indicated GST-FAK C-terminal domain constructs in 293T cells and GST pull-down shows construct size and expression by Coomassie staining. Binding of endogenous paxillin or talin to GST-FAK constructs was determined by immunoblotting. [B] Transient cotransfection of the indicated GST-FAK C-terminal domain constructs in 293T cells and HA-tagged talin FERM F2-F3 was followed by GST pull-down and HA immunoblotting. Equal GST-FAK construct binding to beads and expression of HA-tagged talin FERM was verified by immunoblotting. [C] The indicated alanine point mutations were generated within GST-FAK (1,011–1,042) and cotransfected into 293T cells with HA-talin FERM F2-F3. GST pull-down was followed by HA immunoblotting. Equal GST-FAK construct binding to beads and expression of HA-tagged talin FERM was verified by immunoblotting. [D] Schematic representation of the crystal structure of the FAK FAT domain. E1015 (magenta) is located in the fourth helix of the FAK FAT domain. The paxillin-binding region is indicated (green). [E] In vitro translated full-length FAK WT or FAK E1015A in the presence of biotin-lysine were incubated with GST or GST-talin FERM F2-F3 beads. Direct binding was evaluated by streptavidin immunoblotting with 10% of FAK-WT or FAK E1015A input as a positive control. [F] 293T cells were transfected with GFP, GFP-FAK-WT, or GFP-FAK-E1015A; replated onto FN-coated dishes for 15 min before lysis; and evaluated for endogenous talin and paxillin association by blotting and anti-GFP immunoprecipitation. Molecular mass is indicated next to the gel blots in kilodaltons.
Figure 4. **FAK E1015A expression inhibits talin binding and recruitment to nascent adhesions.** (A) Cell lysates were prepared from FAK−/− MEFs stably re-expressing GFP, GFP-FAK WT, or GFP-FAK E1015A after replating on FN for 15 min. Anti-GFP immunoprecipitates were analyzed for endogenous talin and paxillin association. Lysates were also analyzed for talin, GFP-FAK, paxillin, and actin expression. (B) GFP-FAK WT and GFP-FAK E1015A MEFs were plated on FN for 15 min and analyzed for GFP-FAK (green) and paxillin (red) staining (Fig. S3 A). (C and D) In parallel, cells were analyzed for GFP-FAK (green) and talin (red) staining. Co-localization analyses and box and whisker plots are described in the legend for Fig. 1. Representative images of FN-replated GFP-FAK WT and GFP-FAK E1015A MEFs are shown. The merged images show colocalization (yellow). Inset, enlarged area of peripheral adhesion staining (circled). Bars, 10 µm. (E) FAK−/− MEFs expressing GFP, GFP-FAK WT, or GFP-FAK E1015A were serum starved, suspended (S), and FN replated for the indicated times. Anti-pY397 FAK immunoblotting shows FN-stimulated FAK activation with respect to total FAK and actin levels. Molecular mass is indicated next to the gel blots in kilodaltons. (F and G) Adhesion number (F) and size (G) were determined in the indicated cells at 60 min on FN by anti-paxillin staining in at least 10 cells per experimental group. Box and whisker plots are described in the legend to Fig. 1 (**, P < 0.01; *, P < 0.05).
60 min (Fig. 4 E). Thus, talin binding is not essential for integrin-stimulated FAK activation. FAK−/− MEFs exhibit excessive formation of adhesions that accumulate in size and exhibit slow turnover kinetics (Webb et al., 2004). FAK reexpression reverses these defects (Tomar and Schlaepfer, 2009), and adhesion number and size is significantly reduced by GFP-FAK WT compared with GFP-reconstituted FAK−/− MEFs (Fig. 4, F and G). However, GFP-FAK E1015A expression did not reduce adhesion number or size upon FN replating (Fig. 4, F and G). Increased adhesion formation was associated with a threefold reduction of haptotaxis cell motility (Fig. 5 A) and the inhibition of scratch wound closure (Fig. 5 B) by FAK E1015A compared with FAK WT expression. FAK knockdown and re-expression of FAK E1015A in SKOV3.ip1 cells did not promote cell motility initiated by β1 integrin–mediated cell attachment as did FAK WT re-expression (Fig. S3, E–H). Together, these results support the importance of FAK binding to talin in mediating β1 integrin–stimulated cell motility.

**FAK-talin interactions facilitate adhesion dynamics**

It is the coordination of adhesion formation and turnover that regulates cell movement, and FAK is intimately involved in these processes (Parsons et al., 2010). Live cell imaging of mCherry-paxillin at wound edge–associated adhesions revealed a mean fluorescence lifetime of 60 min within FAK−/− MEFs, and this was reduced to 20 min upon GFP-FAK WT reexpression (Fig. 5 C). Even though GFP-FAK E1015A was localized to adhesions, these sites exhibited a significantly extended lifetime at wound edges compared with GFP-FAK WT–expressing MEFs (Fig. 5 C). This result is consistent with the inability of FAK E1015A to decrease adhesion number or size. Thus, FAK binding to talin is associated with increased adhesion turnover.

A potential mechanism that may account for increased adhesion lifetime in MEFs lacking FAK or expressing FAK E1015A is that both FAK and talin are substrates of the calpain II protease involved in adhesion turnover (Franco et al., 2004; Chan et al., 2010). Talin cleavage and head domain fragment generation is associated with increased adhesion turnover and cell motility associated with talin cleavage. [A] FAK−/− MEFs were transfected with the indicated constructs and analyzed for FN haptotaxis cell migration over 3 h. Migratory cells identified by β-gal staining were counted and results normalized to control transfected FAK−/− MEFs. Data are the mean ± SEM of three independent experiments (**, P < 0.001). (B) Wound closure motility stimulated by GFP-FAK WT but not GFP-FAK E1015A expression in FAK−/− MEFs. Data are the mean ± SEM of three independent experiments (**, P < 0.001). [C] FAK WT but not FAK E1015A reexpression in FAK−/− MEFs decreased leading edge–associated adhesion lifetime. Real-time spinning disk confocal microscopy was used to image mCherry paxillin fluorescence (2 min intervals over 2 h) in wound closure assays. Pixel intensity analyses and adhesion tracking were performed using Image J (version 1.38). Box and whisker plots represent analyses of all newly formed adhesions at the leading edge in at least six cells per experimental group (**, P < 0.001). [D] Increased talin cleavage by FAK. Scrambled (Scr) or anti-FAK shRNA SKOV3.ip1 cells were transfected with HA-tagged FAK WT or FAK E1015A and replated onto FN for 60 min. An anti-talin head domain antibody was used for immunoprecipitation, followed by talin, FAK, and actin immunoblotting. Molecular mass is indicated in kilodaltons. (E) Graph showing the ratio of talin head domain to full-length talin by immunoprecipitation and immunoblotting. Data are the mean from two independent experiments (error bars indicate ± SEM; *, P < 0.03).
binding to β1 integrins. FAK recruitment and activation is associated with integrin clustering, and is independent of talin binding or adhesion maturation. A talin binding mutant of FAK (E1015A) is activated by integrins and is efficiently localized to nascent adhesions, but does not promote adhesion dynamics or cell motility. The presence of a FAK–talin complex within adhesions regulates a cycle of talin proteolysis and focal adhesion turnover enabling cell movement. These studies highlight a new linkage between integrins, FAK, talin, and the control of cell motility.

Materials and methods

Cells and constructs

FAK−/−/MEFs, FAK+/−/MEFs, and human 293T cells were grown and maintained in DME containing 10% FBS, 100 U/ml penicillin, 100 µg/ml streptomycin, MEM nonessential amino acids, and 1 mM sodium pyruvate at 37°C in a 5% CO2 incubator as described previously (Lim et al., 2008). Ovarian SKOV3.ip1 carcinoma cells were from J. Chien (Mayo Clinic, Rochester, MN) and were maintained as described earlier. CHO cells expressing αβ5/β3-1A chimeric integrin receptors were from S. LaFlamme (Albany Medical College, Albany, NY) and were cultured in F12 medium with supplements as already described. HUVECs were purchased from Lonza and grown according to the manufacturer’s instructions in Endothelial Cell Growth Medium 2 (Lonza) at 37°C in 10% CO2, and cultured on 0.2% gelatin and 10 µg/ml collagen type I coated plastic dishes. HA-tagged talin FERM F2-F3 WT, K357A, R358A, W359A, and A360E constructs were used as described previously (Garçia-Alvarez et al., 2003). mCherry paxillin was from C. Waterman (National Institutes of Health, Bethesda, MD). Primers used in the generation of pEeG FAK mammalian GST fusion proteins, GFP-FAK fusions in pEGFP-C1, GSTTalin F2-F3 FERM in pGEX-4T1, or shRNA knockdown constructs in plentilox 3.7 are detailed in Table S1. Constructs were verified by sequencing.

Antibodies and reagents

mAb to paxillin (clone M107) was obtained from BD. Rabbit mAb to paxillin (clone Y113) was from Abcam. mAbs to FAK (clone 4.47), to the talin head domain (clone 1676), integrin β1 (clone PAC10), integrin αvβ3 (clone LM609), integrin αvβ5 (clone P1f6), integrin β3 (clone SAP), integrin αβ (clone C32C), integrin α4 (clone HP2/1), and rabbit polyclonal antibodies to FAK (J6–543) were from Millipore. Cytochalasin D, latrunculin-A, β-actin (clone AC-17), pan-talin (clone B4d4), mAbs, purified bovine plasma FN, and mitomycin-C were obtained from Sigma-Aldrich. Integrin αv (clone 13C2), integrin α5 (clone P4C10), integrin αv (clone 9EG7), and integrin α5 (clone IIA1) mAbs were from BD. HA-tag (clone 16B12) mAb was from Covance. GST (clone Stag05) mAb was from Thermo Fisher Scientific. Phospho-specific rabbit mAb to FAK pY397 (44-625G) was from Invitrogen. Affinity-purified rabbit polyclonal antibodies to FAK 533–1052 were used as described previously (Lim et al., 2008). Integrin α5 (β5SS) antibody was from D. Stupack (University of California, San Diego, La Jolla, CA). Human FB (depleted of von Willebrand Factor and FN) was from Enzo Life Sciences.

RNA interference and lentivirus

Human PTK2 (FAK) siGENOME SMART pool siRNA (M-003164-02, siGENOME nontargeting siRNA pool (D-001206-13-05), and siGLO transfection indicator were used (Thermo Fisher Scientific). For siRNA knockdown of human talin1, Stealth Select RNAi (catalog no. 1299003; Invitrogen) oligo 804 (sequence: 5′-CCAAAGAGGAGAACCAGGAGCCAGGAGAAAGGC-3′) was used (Kopp et al., 2010). HUVECs or SKOV3.ip1 cells were transfected with 100 pmol siRNA + 50 pmol siGLO using JetPrime DNA and siRNA transfection reagent (Polyplus Transfection Inc). After 48 h, FAK knockdown was confirmed by immunoblotting. Lentiviral shRNA to human FAK and a scrambled control in plentilox 3.7 were created as described previously (Lim et al., 2008) with the primers listed in Table S1. Site-directed mutagenesis of FAK E1015A was performed using QuickChange (Agilent Technologies) and subcloned into pEGFP-C1. FAK as a Cλ–XbaI fragment. GFP and GFP–FAK were subcloned from pEGFP-C1 into the lentiviral vector pCDH1-MCS1-EGF-Puro (System Biosciences) via NheI sites. Lentiviral production was performed as described previously (Lim et al., 2008); target cells were selected with puromycin, sorted by flow cytometry for GFP expression, and maintained as pooled populations.

Cell replating, protein binding, and immunoblotting

For GST pull-down assays, 293T cells were transfected with plasmids for GST-FAK, HA-talin F2-F3, or empty vector control using JetPrime, and analyzed after 48 h. For replating or imaging experiments, cells were starved (0.5% serum) for 16 h at subconfluent densities, then treated with 0.06% trypsin and 2 mM EDTA in PBS (2.5 min at 37°C). Trypsin was inactivated by the addition of 0.5 mg/ml soybean trypsin inhibitor with 0.25% BSA in DME, collected by centrifugation, resuspended in migration medium (DME with 0.5% BSA), and held at 37°C (2 × 105 cells/ml) for 1 h. Acid-washed gelatin-coated slides or plastic culture dishes were coated with FN (10 µg/ml in PBS) or FB (15 µg/ml in PBS) overnight, blocked with 1% BSA in PBS for 30 min, and preheated to 37°C before use in cell experiments. Total protein lysates were prepared after the indicated times in extraction buffer containing 50 mM Heps, pH 7.4, 150 mM NaCl, 1% Triton X-100, 1% sodium deoxycholate, 0.1% SDS, and 1% glycerol. For immunoprecipitation and GST binding analyses, lysates were diluted twofold in HNTG buffer (50 mM Heps, pH 7.4, 150 mM NaCl, 0.1% Triton X-100, and 1% glycerol), then incubated with antibodies (1 µg) or glutathione-agarose beads (Sigma-Aldrich) for 3 h at 4°C. Antibodies were collected with either protein A or G Plus (Millipore) agarose beads, and beads washed at 4°C in 1% Triton X-100–only extraction buffer, followed by washes with HNTG buffer, or beads resolved by SDS-PAGE and transferred to PVDF membranes whereby the bait protein was detected by Coomassie staining. The bound prey was detected by streptavidin-horseradish peroxidase immunoblotting. Where indicated, FAK 533–1052 was expressed as a GST fusion protein, incubated with 50 U/ml thrombin, purified using anion exchange chromatography (BioRad Laboratories), and detected by anti-FAK immunoblotting.

Cell motility

For transient transfection studies, cells were cotransfected with a pCDNA3.1 or pCDNA3.1-LocZ, HA-tagged FAK-WT, or FAK-E1015A, and evaluated 36 h after transfection. MilliCell chambers (8 µm pores; Millipore) were coated on the membrane underside with 10 µg/ml FN in migration medium for 2 h, washed with PBS, and air dried (30 min) before use. Starved cells (0.5% serum, overnight) were suspended by limited trypsin-EDTA treatment. Soybean trypsin inhibitor (0.25 mg/ml in DME) was added, and cells were pelleted and washed in migration medium (DME with 0.5% BSA) and enumerated (ViCell XR; Beckman Coulter). Cells were held in suspension as pooled populations.

In vitro translation and direct binding assay

Prey proteins were in vitro translated using the TNT transcription-translation system (Promega). Expression constructs in 1 µg pCDNA3.1 were translated in a mixture containing biotin-labeled lysine and diluted 50-fold into binding buffer (50 mM Heps, pH 7.4, 150 mM NaCl, and 1% Triton X-100). Bait proteins were expressed as GST fusion proteins in bacteria, prebound to glutathione-agarose beads, incubated with diluted prey protein for 2 h at 4°C, washed three times in binding buffer, resolved by SDS-PAGE, and transferred to PVDF membranes whereby the bait protein was detected by Coomassie staining. The bound prey was detected by streptavidin-horseradish peroxidase immunoblotting. Where indicated, FAK 533–1052 was expressed as a GST fusion protein, incubated with 50 U/ml thrombin, purified using anion exchange chromatography (BioRad Laboratories), and detected by anti-FAK immunoblotting.

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Immunofluorescent staining

Cells replated for the indicated time on FN or FB glass coverslips (as described earlier) were fixed in 3.7% paraformaldehyde (15 min), permeabilized with 0.1% Triton X-100 in PBS (10 min), and incubated in blocking buffer (2% BSA in PBS) for 1 h. Talin (1:500), paxillin (1:250), activated β1 (9EG7, 1:50), or FAK (1:50) antibodies were diluted in blocking buffer and incubated overnight at 4°C. Coverslips were washed in PBS,
incubated with Alexa Fluor 488 goat anti-mouse and Alexa Fluor 594 goat anti-rabbit or Alexa Fluor 647 goat anti-rabbit (Invitrogen) diluted in blocking buffer (45 min). Images were acquired sequentially using a mercury lamp source, multiband dichroic, single-band exciter, and single-band emitter filter sets (Chroma Technology Corp.) on dual filter wheels. A spinning disc confocal microscope (IX81; Olympus) at 60x (Plan-Apochromat, NA 1.42 objective lens) and a camera (OrcaER; Hamamatsu) controlled by Slidebook (v5.0) software were used. Files were cropped, pseudo-colored, and contrast-adjusted using Photoshop (Adobe). The degree of association exhibited by patterns of fluorescence was measured on a pixel-by-pixel basis and calculated as a Pearson’s correlation coefficient using the “measure correlations” module (Cell Profiler, v2.0; Broad Institute). A value of 0 indicates no overlap and a value of 1 corresponds to 100% colocalization. Adhesion size (pixels) and the number remote within a cell were determined using Cell Profiler using a pipeline to threshold old images and reduce background fluorescent staining.

Flow cytometry
Cells were trypsinized, enumerated, and incubated with primary antibodies to integrins (10° cells/µg antibody) for 20 min on ice in 100 µl of PBS, followed by pelleting and washing using cold PBS. Allophycocyanin (APC)-conjugated goat anti-mouse IgG (BD) was used as a secondary antibody, and analyses were performed using a FACS Calibur (BD) with FlowJo software. Mouse IgG was used as a negative control.

Cell adhesion
Serum-starved cells were trypsinized, enumerated, and held in suspension in migration medium at 37°C for 10 min in the presence of 10 µg/ml of control mouse IgGs or antibodies specific to integrins (β1, αvβ3, αvβ5, α4, and α5). Cells (in the presence of antibodies) were evaluated for adhesion to FN-coated (2 µg/ml) dishes at 15 min by paraformaldehyde fixation, crystal violet staining, and counting.

Scratch-wound, time-lapse imaging, and quantification of FA dynamics
Glass-bottom dishes (MatTek) were coated with 2 µg/ml FN, and cells were plated at a subconfluent density (75%). After 24 h, cells were serum starved overnight, wounded with a pipette tip, washed with PBS, and incubated in growth media containing 0.5 µg/ml of mitomycin-C before imaging. For time-lapse wound healing experiments, images of cells in phase were acquired every 15 min in a humidified 5% CO2 environment at 37°C using a microscope (IX51; Olympus), xy-controlled stage and a camera (Hamamatsu) controlled by Slidebook (v5.0) software. Wound closure percentage was calculated by the change in area between 0 and 8 h. For confocal time-lapse microscopy of cells at the leading wound edge, GFP, GFP-FAK-WT, or GFP-FAK-E1015A reconstituted MEFs were transfected with mCherry paxillin as an adhesion marker. Sequential GFP and mCherry images were acquired every 2 min for 2 h in a humidified 5% CO2 environment at 37°C using a control spinning confocal microscope with zero drift compensation (IX81; Olympus) at 60x (Plan-Apochromat, NA 1.42 objective lens), as well as a mercury lamp source, multiband dichroic, single-band exciter, and single-band emitter filter sets (Chroma Technology Corp.) on dual filter wheels, and a camera (OrcaER) controlled by Slidebook (v5.0) software. Files were cropped, pseudo-colored, and contrast-adjusted using Photoshop. Quantification of adhesion dynamics at the leading edge was performed by background subtraction and pixel intensity analyses using ImageJ (version 1.38) software. Adhesion lifetime was determined by thresholding images to select for individual adhesions followed by tracking using an ImageJ (manual tracking) plug-in module.

Statistical analysis
A two-tailed unpaired Student’s t test was used to evaluate two groups. Significance between multiple groups was determined by one-way analysis of variance (ANOVA) followed by either Dunnett’s or Tukey’s multiple comparison test.

Online supplemental material
Fig. S1 shows nascent and mature adhesion formation upon FN repletion and the role of FAK in promoting talin recruitment to nascent adhesions in human ovarian carcinoma cells. Fig. S2 shows that nascent adhesion localization and activation of FAK occurs in a tension-independent manner, and shows experiments with direct tail FERM binding to FAK. Fig. S3 shows that FAK-E1015A localizes to nascent adhesions but does not promote β1 integrin-mediated cell motility. Table S1 summarizes the primers used for PCR cloning, PCR mutagenesis, FAK shRNA, and recombinant FAK protein production. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201108078/DC1.

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