Nuclear FAK and Runx1 cooperate to regulate IGFBP3, cell cycle progression and tumor growth

Marta Canel1,2,*; Adam Byron1,*; Andrew H. Sims1; Jessy Cartier3; Hitesh Patel1; Margaret C. Frame1; Valerie G. Brunton1; Bryan Serrels1,†‡; and Alan Serrels1,2,†‡

1Cancer Research UK Edinburgh Centre, Institute of Genetics and Molecular Medicine, University of Edinburgh, Edinburgh EH4 2XR, UK. 2MRC Centre for Inflammation Research, The Queen’s Medical Research Institute, University of Edinburgh, Edinburgh EH16 4TJ, UK. 3British Heart Foundation Centre for Cardiovascular Science, University of Edinburgh, The Queen’s Medical Research Institute, 47 Little France Crescent, Edinburgh EH16 4TJ, UK.

*These authors contributed equally to this work
†These authors contributed equally to this work
‡Correspondence to A.S. or B.S., email: a.serrels@ed.ac.uk or b.serrels@ed.ac.uk

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Running Title: Cooperation of nuclear FAK and Runx1 in IGFBP3 regulation

Key words: FAK, Runx1, IGFBP3, cell cycle, tumor growth

Abbreviations: FAK, focal adhesion kinase; FERM, four-point-one, ezrin, radixin, moesin; FUCCI, fluorescent ubiquitination-based cell-cycle indicator; GFP, green fluorescent protein; IGFBP3, insulin-like growth factor binding protein 3; Runx1, Runt-related transcription factor 1; SCC, squamous cell carcinoma
Abstract

Nuclear focal adhesion kinase (FAK) is a potentially important regulator of gene expression in cancer, impacting both cellular function and the composition of the surrounding tumor microenvironment. Here we report in a murine model of skin squamous cell carcinoma (SCC) that nuclear FAK regulates Runx1-dependent transcription of insulin-like growth factor binding protein 3 (IGFBP3), and that this regulates SCC cell cycle progression and tumor growth in vivo. Furthermore, we identified a novel molecular complex between FAK and Runx1 in the nucleus of SCC cells and showed that FAK interacted with a number of Runx1 regulatory proteins, including Sin3a and other epigenetic modifiers known to alter Runx1 transcriptional function through post-translational modification. These findings provide important new insights into the role of FAK as a scaffolding protein in molecular complexes that regulate gene transcription.
Introduction

Focal adhesion kinase (FAK) is a non-receptor protein tyrosine kinase that controls diverse cellular functions including cell adhesion, migration, invasion, polarity, proliferation, and survival (1,2). FAK is therefore involved in a number of processes that can impact on the malignant phenotype. Deletion of fak in mouse models of cancer has shown a requirement for FAK in tumor formation and progression (3-9). Over-expression of FAK has also been reported in a number of human epithelial tumors (10-12), and it has therefore emerged as a potential target for cancer therapy, with a number of FAK kinase inhibitors currently being developed (13).

FAK was identified as a protein highly phosphorylated in response to integrin activation and primarily located at cell–extracellular matrix adhesion sites termed focal adhesions (1). Recent reports have also identified that FAK contains nuclear localization signals within the F2 lobe of the four-point-one, ezrin, radixin, moesin (FERM) domain (14) and a nuclear export signal within the kinase domain (15). Therefore, FAK can translocate to the nucleus, where its function remains poorly characterized. Within the nucleus, the FAK FERM domain can bind to the transcription factors p53 and GATA4, resulting in their turnover and inactivation, thereby controlling cell survival and inflammatory signals (14,16). Recently, we reported that nuclear FAK was tethered to chromatin and regulated the expression of chemokines and cytokines, including Ccl5 and TGFβ2, that contribute to establishment of an immuno-suppressive tumor environment through driving elevated intra-tumoral regulatory T-cell numbers (17). Therefore, nuclear FAK protein complexes can act to regulate transcriptional programs important in controlling cellular responses and the composition of the tumor immune environment.

Runt-related transcription factor 1 (Runx1; also known as AML1) is one of a
family of three transcription factors (Runx1 - 3) that can either activate or repress transcription depending on the target gene, cell type, and associated co-factors (18,19). It has been shown to have a critical role in hematopoiesis and hematopoietic function (20), and is essential for mammalian development (21). In the context of cancer, Runx1 is best known for its role in acute myeloid leukemia (AML) where it is frequently found mutated (18). In recent years it has also become clear that Runx1 plays an important role in solid epithelial malignancies. For example, Runx1 deficiency impairs mouse skin tumorigenesis (22), while in contrast it acts as a tumor suppressor in the ApcMin mouse model of colorectal carcinogenesis (23). Therefore, it has an important but context dependent function in cancer. Here, we identify a novel molecular complex between FAK and Runx1 in the nucleus of skin squamous cell carcinoma (SCC) cells. We show that nuclear FAK and Runx1 cooperate to regulate expression of IGFBP3, and that IGFBP3 regulates cell cycle progression and tumor growth in vivo. Using proteomics and network biology approaches, we identify that nuclear FAK interacts with a number of Runx1 regulatory proteins that can alter Runx1 transcriptional function through post-translational modification. Further, we identify a novel interaction between nuclear FAK and Sin3a, a core component of the Sin3a/HDAC co-repressor complex, and show that Runx1/Sin3a interaction is enhanced in SCC FAK-wt cells. This study provides new insights into the potential mechanisms through which nuclear FAK regulates transcription factor function to control cell behavior.
Materials and Methods

Materials. All antibodies used are listed in Supplementary Materials and Methods. All siRNA was purchased from Dharmacon and shRNA from Open Biosystems.

Cell lines. Cell lines used in this study were generated, authenticated and characterized as previously described (17, 24). Cells were pathogen tested in September 2016 using the ImpactIII test (Idex Bioresearch) and were negative for all pathogens. Cell lines are routinely tested for mycoplasma every 2-3 months in-house and have never been found to be mycoplasma positive. Cell lines are cultured for no more than 3 months following freeze thawing. Runx1 or scrambled siRNA was transiently transfected into SCC cells using HiPerFect (Qiagen). Cells were left for 3 days and then immunoblotting or (q)RT-PCR analysis was carried out. Cell lines stably expressing FUCCI, Runx1, or empty-vector shRNA were generated by lentiviral infection and selected in 2 µg/ml puromycin.

Generation of nuclear-targeting FAK mutants. Point mutations were introduced into wild-type FAK in three different combinations, (1) R177A and R178A, (2) K190A and K191A, (3) K216A and K218A, using site-directed mutagenesis as previously described (17). Primer sequences are listed in Supplementary Materials and Methods.

Quantitative RT-PCR. RNA extracts were obtained using an RNeasy kit (Qiagen), following manufacturer’s instructions. cDNA was made using a first-strand cDNA synthesis kit (Invitrogen). (q)RT-PCR was performed as previously described (17). Analysis was performed using Rotor-Gene software, and expression relative to B2M was calculated using Excel (Microsoft). Standard PCR was performed using the above
conditions but substituting SensiFAST for a 2× PCR master mix. Primers used are listed in Supplementary Materials and Methods.

**Nuclear fractionation and total cell lysates.** Cells were collected by scraping and low-speed centrifugation (1,000 g at 4 °C for 5 min), followed by two washes with ice-cold PBS. Cell pellets were resuspended in buffer A (10 mM KCl, 10 mM HEPES, pH 7.9, 0.5 mM DTT, 1.5 mM MgCl₂) and incubated on ice for 15 min. Cells were lysed using a 25G needle and nuclei isolated by centrifugation at 12,000 g at 4 °C for 30 sec. The nuclear pellet was washed twice in buffer A and incubated in buffer C (25% (v/v) glycerol, 0.2 mM EDTA, pH 8.0, 0.42 M NaCl, 0.5 mM PMSF, 20 mM HEPES, pH 7.9, 0.5 mM DTT, 1.5 mM MgCl₂) at 4 °C for 1 h. Clarification of the nuclear extracts was by high-speed centrifugation (16,000 g at 4 °C for 5 min). Alternatively, cells were washed with PBS and lysed in RIPA lysis buffer (50 mM Tris–HCl, pH 7.4, 150 mM sodium chloride, 0.1% SDS and 1% sodium deoxycholate) with inhibitors (Complete Protease Inhibitor Cocktail and PhosSTOP; Roche). Clarification was by high-speed centrifugation (16,000 g at 4 °C for 15 min).

**Immunoblotting and immunoprecipitation.** Cell lysates or cell fractions (10–20 μg protein, as measured by Micro BCA Protein Assay kit (Pierce)) were supplemented with SDS sample buffer (Tris, pH 6.8, 20% glycerol, 5% SDS, β-mercaptoethanol, and bromophenol blue), separated by SDS–PAGE, transferred to nitrocellulose and immunoblotted with specific antibodies at 1:1000 dilution. For immunoprecipitation experiments, 0.25–1 mg of cell lysate or cell fraction was immunoprecipitated with either 5 μl of mouse Runx1 antibody, 10 μl of agarose-conjugated mouse FAK antibody, 10 μl of agarose-conjugated Myc-tag, or 10 μl of agarose-conjugated control IgG, and immune complexes collected. Beads were washed three times with lysis buffer, once with 0.6 M lithium chloride, and then added to SDS sample buffer.
2D-Gel Electrophoresis. Two-dimensional SDS–polyacrylamide gel electrophoresis was performed using the ZOOM IPGRunner System (ThermoFisher Scientific) according to the manufacturers protocol. Gels were transferred to nitrocellulose membranes, blocked (5 % BSA in TBST) and probed with anti-RUNX1 antibody.

Protein capture arrays. 2 × 10⁶ SCC cells were plated onto a 90-mm tissue culture dish and allowed to adhere overnight. Growth medium was removed and replaced with 5 ml of normal growth medium. Cells were cultured for a further 24 hours before conditioned medium was removed and centrifuged at 1000 rpm for 5 minutes. Supernatant was collected and used for analysis. To control for cell number, the 90-mm dish containing SCC cells was lysed in RIPA buffer and protein quantified using a BCA protein assay as described above. This was used to normalize loading of conditioned media onto the protein capture arrays. Secreted protein analysis was performed using mouse angiogenesis protein capture arrays (R & D Systems) according to the manufacturer’s protocol. The only modification was the use of a streptavidin DyLight-800 antibody for detection (1:10,000 dilution, Rockland Biochemicals). Image acquisition and analysis was performed using a Licor Imager. Mean spot intensities were calculated from duplicate arrayed spots per cell line. Data were median centered and subjected to unsupervised agglomerative hierarchical clustering on the basis of Pearson correlation computed with a complete-linkage matrix using Cluster 3.0 (C Clustering Library, version 1.50) (25). Clustering results were visualized using Java TreeView (version 1.1.6) (26).

Blood vessel immunostaining. Subcutaneous tumors were surgically excised, placed in an aluminum foil boat, submerged in OCT compound, and snap frozen using a dry ice and methanol bath. OCT-embedded samples were stored long-term at
−80°C. For analysis, sections were cut at −20°C onto siliconized microscope slides. For staining, sections were removed from the −20°C freezer, and 50 μl of fixative was applied directly onto the tissue. Sections were incubated for 8 minutes at 4°C and then the fixative removed and the slides allowed to equilibrate to room temperature for approximately 10–20 minutes. Sections were then rehydrated in PBS for 10 minutes. The area of tissue was surrounded with a hydrophobic barrier pen and blocked using 1% horse serum in PBS for 30 minutes at room temperature. Tissue was then incubated with anti-CD31 antibody (BD Biosciences, 1:100 dilution in incubation buffer (1% bovine serum albumin, 1% normal donkey serum, 0.3% Triton X-100, 0.01% sodium azide in PBS)) overnight at 4°C. Slides were washed three times for 15 minutes with PBS, and then incubated with anti-rat Alexa Fluor 594-conjugated secondary antibody (Molecular Probes, ThermoFisher Scientific) diluted at 1:200 in incubation buffer for 1 hour at room temperature. Slides were washed three times for 15 minutes in PBS and then counter-stained with DAPI for 5 minutes. Slides were washed once for 15 minutes in PBS and mounted in VectaShield anti-fade mounting media (Vectorlabs). Images were acquired using an Olympus FV1000 confocal microscope.

Transcription factor network analysis. Transcription factors were extracted from the DECODE database (Qiagen), selecting the most relevant transcription factors predicted by text mining to bind between 20 kb upstream and 10 kb downstream of the human transcription starting site. The transcription factor Runx1 was used to seed a network of 1000 related proteins using the GeneMANIA plugin (version 3.4.1; human interactions) in Cytoscape (version 3.3.0) (27), onto which proteins specifically isolated in nuclear FAK protein complexes (17) were mapped. The resulting interactome was extracted, and proteins with physical or predicted direct or
indirect interactions with Runx1 were analyzed and topological network parameters computed using the NetworkAnalyzer plugin (version 2.7). Networks were clustered using the yFiles Organic algorithm implemented in Cytoscape. Pathway enrichment analysis was performed using the Database for Annotation, Visualization and Integrated Discovery (version 6.8) (28). Pathway terms with Benjamini–Hochberg-corrected $P$-value < 0.01 were considered significantly overrepresented.

**Subcutaneous tumor growth.** $2.5 \times 10^5$ cells were injected subcutaneously into each flank of immune-compromised CD-1 nude mice, and tumor growth measured twice weekly using calipers. Measurements were taken from three mice (each bearing two tumors) for each cell line, and the volume of the tumor ($v$) was calculated in Excel using the formula $v = \frac{4}{3}\pi r^3$. Data were graphed and statistics calculated using Prism (GraphPad).

**In vivo cell cycle analyses.** Optical window chambers were implanted onto CD-1 nude mice as described previously (29). All animal work was carried out in compliance with UK Home Office guidelines. $1 \times 10^6$ FUCCI-expressing SCC cells were injected into the dermis and at the time of window implantation. 24 hours later, mice were anaesthetized using an isoflurane–oxygen mix and three-dimensional image stacks acquired using an Olympus FV1000 confocal microscope. Image analysis was performed using the spot detection tool in Imaris (Bitplane). The number of green, red, and double-positive nuclei were counted and calculated as a percentage of the total number of cells within the image stack.

**Longitudinal in vivo imaging of tumor angiogenesis.** Optical window chambers were implanted onto CD-1 nude mice as described above. Prior to sealing the window with a coverslip, a tagRFP-expressing tumor fragment (~1 mm in diameter) was placed
into the window. To obtain tumor fragments, a donor animal was injected with $2.5 \times 10^5$ tagRFP-expressing SCC cells 10 days prior to optical window implantation. Tumors were removed, cut into small pieces using a scalpel, and fluorescence checked using an Olympus OV110 whole-animal imaging system. For longitudinal imaging of tumor angiogenesis, mice bearing optical windows were anaesthetized using an isoflurane–oxygen mix. Images were acquired 2, 4, 7, and 9 days post-implantation using an Olympus OV110 whole-animal imaging system set to acquire both GFP and RFP signals using the zoom lens set to 1.6× magnification. For image analysis and identification of blood vessels, autofluorescence images acquired using the GFP channel were inverted in ImageJ. The tubness plugin was used to detect blood vessel structures and apply an image mask over these structures. The accuracy of this was checked manually. The vascular density was determined by calculating the percentage of the field of view covered by the vascular image mask. Data were graphed and statistics calculated using Prism.

**Results**

*FAK is required for cell cycle progression and angiogenesis in SCC tumors*

We have previously shown using a murine model of skin SCC that depletion of FAK expression can result in immune-mediated tumor regression in syngeneic mice, and a growth delay in immune-deficient mice (17,24). We therefore sought to further dissect the complex role of FAK in regulating SCC tumor growth. We measured tumor growth in CD-1 nude mice following injection of $0.25 \times 10^6$ FAK-deficient cells (SCC FAK-/-) and FAK-deficient cells that re-expressed wild type FAK (SCC FAK-wt) at close to endogenous levels. As previously reported, SCC FAK-/- tumors
exhibited a growth delay when compared to SCC FAK-wt tumors (24) (Fig. 1a) that was associated with an increased tumor doubling time (Fig. 1b). FAK is a protein that plays a pleiotropic role in regulating cancer development and progression, and we hypothesized that a number of its reported functions, including regulation of proliferation/cell cycle and tumor angiogenesis (1,2,30), may contribute to the observed tumor growth delay in SCC FAK-/ tumors when established on an immune deficient host background. Thus, we set out to use intravital imaging to analyze, in real-time, the cell cycle distribution and kinetics of tumor neo-angiogenesis in SCC FAK-wt and SCC FAK-/ tumors when grown on CD-1 nude mice.

To measure cell cycle distribution in real-time, SCC FAK-wt and SCC FAK-/ cells expressing the fluorescent ubiquitination-based cell-cycle indicator (FUCCI) reporter (31) were implanted via intra-dermal injection under dorsal-skinfold imaging windows. The FUCCI probe is based on the differential proteolysis of geminin and cdt1 at specific phases of the cell cycle, allowing differential profiling of cells at G1 (red), G1/S (yellow), and S/G2/M (green) phases. Images of FUCCI-expressing SCC FAK-wt and SCC FAK-/ tumors revealed an increased number of red fluorescent cells in SCC FAK-/ tumors (Fig. 1c). Analysis of the proportion of red, yellow, and green nuclei from three-dimensional image stacks acquired 24 hours after tumor cell implantation identified a specific delay in the G1 phase of the cell cycle in SCC FAK-/ tumor cells (Fig. 1d).

FAK, expressed in cancer cells, has been reported to influence tumor angiogenesis through regulating the expression of vascular endothelial growth factor (VEGF) (30). To ascertain whether defective tumor neo-angiogenesis may also be occurring in SCC FAK-/ tumors, we performed longitudinal imaging of tagRFP-labeled SCC FAK-wt and SCC FAK-/ tumors under dorsal-skinfold windows. Using acquisition
parameters similar to those required to image green fluorescent protein (GFP), we acquired images of tissue autofluorescence, in which blood vessels appear as dark non-fluorescent structures that can be readily visualized. Longitudinal imaging of neo-angiogenesis revealed a significant decrease in vascular density in SCC FAK-/- tumors when compared to SCC FAK-wt tumors (Figs. 1e and 1f). Thus, SCC FAK-/- tumors exhibit defective cell cycle progression and neo-angiogenesis that may contribute to defective tumor growth in CD-1 nude mice.

**FAK negatively regulates transcription of IGFBP3**

We and others have previously identified that FAK can regulate the expression of secreted proteins that have the potential to act in both an autocrine and paracrine manner to influence cancer cell behavior and the surrounding tumor microenvironment (17,32,33). To investigate whether FAK-dependent regulation of secreted factors in SCC cancer cells could influence cell cycle progression and tumor angiogenesis, we screened for secreted factors implicated in both processes using protein capture arrays. Analysis of 53 secreted factors revealed both positive and negative regulation as a consequence of FAK expression (Fig. 2a and Supplementary Fig. 1). Applying a four-fold cutoff for differential regulation, we identified IGFBP3 as the most highly upregulated protein in SCC FAK-/- conditioned medium, while PDGF-AA, PDGF-AB/BB, FGF-1, and MMP-3 were the most highly upregulated proteins identified in SCC FAK-wt conditioned medium. Using these findings, we focused on IGFBP3, a protein belonging to the family of insulin-like growth factor binding proteins (IGFBPs), as this factor has previously been reported to negatively regulate both cell cycle progression (34,35) and angiogenesis (36-38). Western blot analysis of conditioned media confirmed an upregulation of IGFBP3 in SCC FAK-/- conditioned samples (Fig. 2b), and q(RT)-PCR identified similar
regulation of Igfbp3 gene transcript levels (Fig. 2c). Thus, we conclude that FAK negatively regulates the transcription of Igfbp3, leading to reduced levels of extracellular secreted IGFBP3 protein.

IGFBP3 is a member of a family of six proteins, IGFBP1–6 (39). However, only antibody capture pairs specific for IGFBP1, 2, and 3 were present on the protein capture arrays used. Therefore, we sought to determine whether FAK regulated the expression of other IGFBP family members. Employing sequence-specific primers designed for each of the six IGFBPs, we used PCR to screen for expression in cDNA libraries prepared from SCC FAK-wt and SCC FAK-/ cells (Fig. 2d). Aside from Igfbp3, only expression of Igfbp4 and Igfbp6 could be detected following 20 cycles of PCR. All primer sets were confirmed to produce single products of the correct size using cDNA prepared from mouse liver (Supplementary Fig. 2). (q)RT-PCR analysis of Igfbp4 expression revealed no differential regulation (Fig. 2e), whereas Igfbp6 expression was observed to exhibit a small but significant decrease in SCC FAK-/- cells when compared to SCC FAK-wt (Fig. 2f). Therefore, IGFBP3 was the only IGFBP family member to be negatively regulated in response to FAK expression.

FAK kinase activity is not required for regulation of IGFBP3

There has been considerable interest in targeting FAK function as a cancer therapy, and a number of small molecule FAK kinase inhibitors are now in early-phase clinical development (13). To determine the role of FAKs kinase activity in regulation of IGFBP3, we tested IGFBP3 protein expression using an SCC FAK-/- cell line in which a FAK kinase-deficient mutant (FAK-kd) had been re-expressed to levels comparable with SCC FAK-wt cells. Anti-IGFBP3 western blotting from conditioned
media revealed that IGFBP3 expression in SCC FAK-kd cells was similar to that observed in SCC FAK-wt cells (Fig. 2g), implying that FAK-dependent regulation of IGFBP3 expression was independent of FAK kinase activity.

**IGFBP3 regulates cell cycle progression but not angiogenesis to influence tumor growth**

To determine whether increased IGFBP3 expression in SCC FAK-/- cells contributed to the defective growth of SCC FAK-/- tumors, we generated SCC FAK-/- cells with a stable knockdown of IGFBP3 using shRNA (Fig. 3a). Analysis of subcutaneous tumor growth showed that knock-down of IGFBP3 in SCC FAK-/- cells partially restored tumor growth *in vivo* (Fig. 3b), resulting in an average tumor volume of approximately double the size when compared to SCC FAK-/- tumors on day 12. Using intravital imaging together with the FUCCI cell cycle reporter, we identified that the G1 arrest observed in SCC FAK-/- cells *in vivo* was overcome following IGFBP3 depletion (Figs. 3c and 3d). However, the defect in blood vessel formation was still evident in the SCC FAK-/- shRNA-IGFBP3 tumors as measured by the presence of CD31+ vessels (Figs. 3e and 3f). Therefore, IGFBP3 likely contributes to regulation of SCC tumor growth through controlling cell cycle progression, but not angiogenesis. Furthermore, depletion of IGFBP3 in SCC FAK-/- cells was not sufficient to fully restore tumor growth, implying that regulation of angiogenesis is likely also important.

**FAK nuclear localization is required for regulation of IGFBP3 transcription**

We have recently reported in SCC cells that nuclear FAK can bind to transcription factors and transcriptional regulators with the potential to influence gene expression (17). To investigate the requirement for FAK nuclear localization in regulation of
Igfbp3, we made a series of FAK nuclear targeting mutants according to previously reported findings that identified putative nuclear localization sequences within the F2 lobe of the FERM domain (40), and expressed these in SCC FAK-/- cells. A total of three nuclear targeting mutants were constructed as follows: (1) by replacing arginine residues at positions 177 and 178 with alanines (FAK-177/178), (2) by replacing lysine residues at positions 190 and 191 with alanines (FAK-190/191), and (3) by replacing lysine residues at positions 216 and 218 with alanines (FAK-216/218). We have previously reported a fourth mutant deficient in nuclear targeting, in which all six of these amino acid residues have been replaced with alanines (17). Biochemical fractionation followed by western blotting was used to assess nuclear/cytoplasmic distribution of the resulting mutants when re-expressed in SCC FAK-/- cells, revealing that all four mutant FAK proteins were deficient in their ability to localize to the nucleus (Fig. 4a). Given that all of the double mutants appeared as defective in nuclear translocation as the mutant harboring all six mutations, we chose to move forward with these for further analysis. (q)RT-PCR analysis of Igfbp3 expression in cell lines expressing these mutant FAK proteins revealed significantly increased levels of Igfbp3 transcript in all three cell lines (Fig. 4b), implying a crucial role for FAK nuclear targeting in the transcriptional regulation of Igfbp3. The inability of these mutants to completely restore Igfbp3 transcript levels to that of SCC FAK-/- cells is likely the consequence of low levels of residual nuclear FAK (Fig. 4a).

Runx1 is required for FAK-dependent transcriptional regulation of IGFBP3

To better define the link between nuclear FAK and regulation of Igfbp3 transcription, we next sought to identify transcription factors with predicted binding sites in the promoter of the Igfbp3 gene. Using the DECODE database (Qiagen), we generated a list of transcription factors predicted by text mining to bind between 20 kb upstream
and 10 kb downstream of the human transcription start site (Fig. 4c). Of these transcription factors, three – Sp1, Runx1, and components of the transcription factor IID complex (TFIID) – were detected by mass spectrometry in a nuclear FAK interactome in SCC cells (Fig. 4c, dark grey bars) (17). We noted that the transcription factor Runx1, which has been reported to regulate Igfbp3 gene expression (41), had the most Igfbp3 promoter binding sites predicted by text mining (Fig. 4c). Moreover, Runx1 was not among the top transcription factors predicted to bind to the gene promoters of the other secreted proteins in the cluster highly upregulated in SCC FAK-/- conditioned medium (Supplementary Fig. 1, top cluster, and Supplementary Fig. 3), suggesting that these other angiogenesis-related proteins may be transcriptionally regulated by a distinct mechanism to Igfbp3.

Using shRNA-mediated stable knockdown of Runx1 expression, we next determined whether Runx1 was required for IGFBP3 expression in SCC FAK-wt and SCC FAK-/- cells. Biochemical fractionation followed by western blotting confirmed a substantial knockdown of Runx1 expression in both SCC FAK-wt shRNA-Runx1 and SCC FAK-/- shRNA-Runx1 cells (Fig. 4d) when compared to controls. Analysis of Igfbp3 expression in these cell lines using (q)RT-PCR showed a loss of Igfbp3 gene expression in SCC FAK-/- shRNA-Runx1 cells, reverting expression levels down to those observed in SCC FAK-wt vector-only control cells (Fig. 4e). Thus, Runx1 is required for increased Igfbp3 transcription following FAK loss in SCC cells. Using an independent method of expression knockdown, similar results were observed following depletion of Runx1 using siRNAs (Supplementary Fig. 4).

**FAK is in a complex with Runx1 in the nucleus**
Analysis of Runx1 expression in SCC FAK-wt and FAK-/- cells revealed that Runx1 is exclusively expressed in the nucleus, and that its expression level is not regulated by FAK (Fig. 4d). Hence, regulation of Runx1 expression / degradation is unlikely the mechanism through which FAK controls Runx1 transcriptional activity in SCC cells. Runx1 can act as either a transcriptional activator or repressor, and the composition of the proteins interacting with Runx1 at a given gene can modulate this function (18). Using a mass spectrometry dataset of the nuclear FAK interactome in SCC cells (17) we identified a potential FAK / Runx1 interaction. Co-immunoprecipitation of Runx1 and FAK in SCC FAK-wt but not in SCC FAK-/- cells confirmed a novel association between FAK and Runx1 (Figs. 5a and 5b). Thus, nuclear FAK exists in complex with Runx1 under steady-state conditions in SCC cancer cells.

A key mechanism of Runx1 regulation is post-translational modification. Runx1 interacts with a number of proteins including kinases, histone acetyltransferases, arginine methyltransferases, and histone deacetylases that can post-translationally modify Runx1, switching it between transcriptional activation and repression (18,19). To further explore how FAK influences Runx1 function, we examined the Runx1 interaction landscape in the context of FAK. To do this, we constructed a Runx1 protein interaction network in silico, onto which we mapped the experimentally derived nuclear FAK interactome. This integrated interaction network identified a subnetwork of Runx1 regulators and associated proteins that interact with FAK in the nuclei of SCC cells (Fig. 5c and Supplementary Fig. 5). Interestingly, a number of these proteins, including DNA (cytosine-5)-methyltransferase 1 (Dnmt1), Sin3a, Histone deacetylases 1, 2, and 3 (HDAC1, 2, and 3), and Nuclear receptor corepressor 1 and 2 (NCoR1 and 2), are linked to repression of Runx1 transcriptional...
activity (18,19). Therefore, nuclear FAK interacts with proteins that can regulate the post-translational modification and transcriptional function of Runx1. KEGG analysis also identified an enrichment for proteins with roles in the cell cycle (Fig. 5d and Supplementary Table 1), consistent with a complex that may play a wider role in the regulation of cell cycle progression.

To determine whether FAK regulated Runx1 tyrosine phosphorylation, we used immunoprecipitation of tyrosine phosphorylated proteins from SCC FAK-wt and FAK-/− cell lysates, followed by western blotting and detection with an anti-Runx1 antibody. No regulation of Runx1 tyrosine phosphorylation was observed (Fig. 5e). To assess Runx1 post-translational modification on a wider scale we performed 2D gel electrophoresis using nuclear extracts prepared from SCC FAK-wt and FAK-/− cells, and probed these gels with an anti-Runx1 antibody. This identified a number of potential differences in both the migration and intensity of spots detected (Fig. 5f, red arrows highlight changes), consistent with an altered state of post-translational modification.

We next investigated whether FAK regulated proteins associated with the post-translational modification of Runx1. Based on data presented in Fig. 5e, we focused on Sin3a, a transcriptional co-repressor known to interact with a number of HDACs (42) and suppress Runx1 transcriptional activity (43). Western blotting using an anti-Sin3a antibody identified reduced Sin3a levels in the nucleus of SCC FAK-/− cells when compared to SCC FAK-wt cells (Fig. 5g). Co-immunoprecipitation from SCC FAK-wt and FAK-/− nuclear extracts confirmed a novel association between Sin3a and FAK in SCC FAK-wt cells (Fig. 5h). Thus, FAK is in complex with and regulates the nuclear levels of a Sin3a. Immunoprecipitation of Runx1 from SCC FAK-wt and FAK-/− nuclear extracts
followed by western blotting and detection with an anti-Sin3a antibody confirmed an association between Runx1 and Sin3a in SCC FAK-wt cells (Fig. 5i). No interaction was observed in SCC FAK-/- cells. Therefore, we propose that one potential mechanism through which FAK may influence the post-translational modification and transcriptional activity of Runx1, is through regulating the expression and subsequent recruitment of Sin3a to Runx1 transcriptional complexes.

**Discussion**

Numerous studies have demonstrated the importance of FAK in regulating tumor cell behavior, which has largely been linked to its role in integrating signals from adhesion sites and growth factor receptors at the cell periphery to control cell adhesion, migration and survival (1,2). In addition, it is now becoming clear that FAK can also play a role within the nucleus to control gene expression (14,17,44). However, the mechanisms underpinning this function remain to be fully characterized. A number of nuclear FAK binding proteins have now been identified including the transcription factors p53 and GATA4 (14,16). Recently, we have shown that nuclear FAK is associated with chromatin and interacts with transcription factors, including the TBP-associated factor TAF9, and transcriptional regulators reported or predicted to regulate expression of the chemokine Ccl5. In doing so, it controls the transcription of chemokines that regulate the composition of the immuno-suppressive tumor environment required to evade the CD8+ T-cell anti-tumor immune response (17). Here, we identify a novel interaction between nuclear FAK and the transcription factor Runx1. We show that nuclear FAK controls the Runx1 dependent expression of IGFBP3, which in turn regulates cell cycle progression and SCC tumor growth in
Further, we identify that FAK interacts with and regulates the nuclear levels of Sin3a, and that FAK is required for recruitment of Sin3a into complex with Runx1. Therefore, nuclear FAK can regulate transcription factor activity, potentially via recruitment of proteins to transcriptional complexes that can alter transcription factor post-translational modification, thereby controlling expression of specific genes that can play an important role in regulating both tumor cell behavior, and how tumor cells influence the composition of the tumor immune environment.

Runx1 is a transcription factor with an important role in both normal development and disease (20,21). De-regulation of Runx1 function contributes to the development of hematological malignances, and in solid epithelial cancers, it has been identified as both a tumor promoter and a tumor suppressor (19). Runx1 can either activate or repress transcription depending on the composition of the protein complexes with which it is associated at a given gene (18), and this likely contributes to the complexity of its role in regulating tumorigenesis. It is known to interact with an array of proteins including kinases, histone acetyltransferases, arginine methyltransferases, histone deacetylases, and ubiquitin ligases, all of which can post-translationally modify Runx1 to regulate its transcriptional function (18,19). Using mass spectrometry, we show that FAK interacts with a number of these proteins including Dnmt1, Sin3a, HDACs 1, 2, and 3, and NCoR1 and 2, all of which are known to repress Runx1 function (18,19). Co-immunoprecipitation experiments confirmed a novel interaction between Sin3a and FAK, and further identified a requirement for FAK to promote interaction of Runx1 with Sin3a. Supporting this as a potential mechanism of FAK-dependent Runx1 regulation, 2D gel analysis of nuclear extracts from SCC FAK-wt and FAK/- cells using an anti-Runx1 antibody identified several protein species that show differential migration, consistent with altered post-
translational modification. Notably, post-translational modification is an important mechanism in the general regulation of transcription factor function (45). Thus, we conclude that this may represent a previously unknown mode of FAK-dependent regulation of gene expression.

FAK is known to regulate a number of cellular processes important for the malignant phenotype (1,2). Here, we identify a new role for nuclear FAK in regulation of SCC cell cycle progression in vivo via controlling Runx1-dependent expression of IGFBP3. IGFBP3 has been linked to cell cycle arrest in tumor cells previously where a G1 arrest was accompanied by reduction in a number of cyclins, including cyclin D1, CDKs, and increased p21 expression (34,35). IGFBP3 has also been linked to regulation of angiogenesis (36-38). However, knockdown of IGFBP3 in the SCC cells had no effect on tumor angiogenesis. Analysis of secreted proteins present in SCC conditioned media identified a number of changes including multiple factors that can influence tumor angiogenesis, implying that other factors may be more important in the regulation of angiogenesis in our SCC model. Interestingly, analysis of the predicted transcription factor binding sites in the promoters of other angiogenesis-related genes regulated by FAK identified that their mechanism of regulation is likely distinct to that of IGFBP3, highlighting the potential mechanistic diversity underpinning FAK-dependent transcriptional regulation.

The data presented here sheds new light on the possible mechanisms underpinning nuclear FAK-dependent regulation of gene expression, highlighting the importance of FAK as a scaffold for protein interactions in the nucleus. Future work should focus on FAK associated transcription factors to define how their interactome may change in the absence of FAK, or its kinase activity, and what the biological and potential clinical relevance of these changes may be.
Acknowledgments

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References


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Figure Legends

Figure 1 | FAK regulates SCC tumor growth, cell cycle and angiogenesis in vivo.
(a) Growth of SCC FAK-wt and SCC FAK-/− tumor xenografts in CD-1 nude mice. n = 5 - 6 tumors per group. (b) SCC FAK-wt and SCC FAK-/− tumor doubling time. Unpaired T-test, ****p < 0.0001. (c) Intra-vital imaging of FUCCI expressing SCC FAK-wt and SCC FAK-/− cells 24 hours post-implantation under dorsal skinfold windows. (d) Quantitation of FUCCI cell cycle distribution from 3-dimentional image stacks shown in panel c. Sidak’s corrected 2way ANOVA, ***p < 0.001. n = 4 tumors per group. (e) Longitudinal imaging of tumour angiogenesis following implantation of tumour fragments under dorsal skinfold windows. Red – tagRFP labelled SCC tumor, Green – tissue autofluorescence. (f) Quantitation of blood vessel density at day 9. Unpaired T-test, *p = 0.0306. Data in all graphs represented as mean +/- s.e.m. n = 3 tumors per group.

Figure 2 | FAK negatively regulates the expression of IGFBP3 but not other IGFBP family members. (a) Relative secreted levels of 53 angiogenesis-related proteins measured by antibody capture array from media conditioned by SCC FAK-wt and FAK-/− cells. Proteins are ordered by fold change. Dotted gray lines indicate four-fold enrichment; proteins changed by at least four fold are indicated. Box-and-whisker plot summarizes the median (line), 25th and 75th percentiles (box) and 5th and 95th percentiles (whiskers). (b) Representative anti-IGFBP3 western blot from concentrated conditioned media. (c) (q)RT-PCR analysis of IGFBP3 transcript levels in SCC FAK-wt and FAK-/− cells. n = 3. (d) Representative PCR analysis of IGFBP
family transcript levels in SCC FAK-wt and FAK-/− cells. (e) (q)RT-PCR analysis of IGFBP4 transcript levels in SCC FAK-wt and FAK-/− cells. n = 3. (f) (q)RT-PCR analysis of IGFBP6 transcript levels in SCC FAK-wt and FAK-/− cells. n = 3. (g) Representative anti-IGFBP3 western blot showing secreted IGFBP3 protein levels from SCC FAK-wt, SCC FAK-/−, and SCC FAK-kd cells. Data in all graphs represented as mean +/- s.e.m. Unpaired T-test, **p < 0.01.

**Figure 3** | IGFBP3 regulates cell cycle progression but not tumor angiogenesis.

(a) Representative anti-IGFBP3 western blot from concentrated media conditioned by either SCC FAK-wt, SCC FAK-/−, or SCC FAK-/− IGFBP3 shRNA cells. Anti-FAK western blot shows FAK expression status and Anti-tubulin western blot was used a loading control. (b) Left - Growth of SCC FAK-wt, SCC FAK-/−, and SCC FAK-/− IGFBP3 shRNA tumor xenografts in CD-1 nude mice. Right – Average volume of SCC FAK-/− and SCC FAK-/− IGFBP3 shRNA tumors at day 12. Unpaired T-test, ***p < 0.001. n = 6 tumors per group. (c) Intra-vital imaging of FUCCI expressing SCC FAK-wt, SCC FAK-/−, and SCC FAK-/− IGFBP3 shRNA cells 24 hours post-implantation under dorsal skinfold windows. (d) Quantitation of FUCCI cell cycle distribution from 3-dimentional image stacks shown in panel c. Values shown for SCC FAK-wt and SCC FAK-/− cells are repeated from Fig. 1d. n = 4 tumors per group. (e) Fluorescent staining of frozen tissue sections using anti-CD31 antibody (red). Nuclei labelled using DAPI (blue). (f) Quantitation of the % area occupied by CD31+ cells. Tukey’s corrected one-way ANOVA, ****p < 0.0001. Data in all graphs represented as mean +/- s.e.m. n = 3 tumors per group with an average of 3 fields measured per tumor.
Figure 4 | Nuclear FAK and RUNX1 regulate IGFBP3 expression.  (a) Representative anti-FAK western blot of cytoplasmic and nuclear fractions prepared from a series of SCC cells expressing FAK nuclear localization signal (NLS) mutants.  (b) (q)RT-PCR analysis of IGFBP3 expression in SCC cells expressing FAK NLS mutants. Tukey’s corrected 1way ANOVA, ****p < 0.0001. Data in all graphs represented as mean +/- s.e.m. n = 3. (c) Predicted transcription factor binding sites in the promoter of Igfbp3. Transcription factors that interact with nuclear FAK in SCC cells are displayed in dark grey. (d) Representative western blot showing Runx1 depletion using shRNA. (e) (q)RT-PCR analysis of IGFBP3 expression in control and Runx1 depleted SCC FAK-wt and SCC FAK-/- cells. Sidak’s corrected 2way ANOVA, ****p < 0.0001. ns = not significant. n = 3.

Figure 5 | Nuclear FAK interacts with Runx1. (a) Representative western blot of anti-FAK immunoprecipitation probed with anti-Runx1 antibody. (b) Representative western blot of anti-Runx1 immunoprecipitation probed with anti-FAK antibody. IgG control (Ctrl). (c) Interaction network analysis of physical or predicted direct binders of Runx1 that interact with FAK in the nucleus of SCC cells (see Supplementary Table 2). Runx1 is shown as a square node. Protein node size is proportional to fold enrichment in nuclear FAK immunoprecipitations. Node color indicates significance of enrichment in nuclear FAK immunoprecipitations. (d) Pathway enrichment analysis of KEGG terms in the nuclear FAK interactome of Runx1 binders (Q < 0.01) (see Supplementary Fig. 5). (e) Representative western blot of anti-phospho-tyrosine
(pTyr) immunoprecipitation probed with anti-Runx1 antibody. (f) 2D gel electrophoresis probed with anti-Runx1 antibody. (g) Representative western blot of SCC FAK-wt and FAK-/- nuclear lysates probed with anti-Sin3a antibody. (h) Western blot of anti-Sin3a immunoprecipitation from SCC FAK-wt and FAK-/- nuclear lysates probed with anti-FAK antibody. IgG control (Ctrl) was done using SCC FAK-wt nuclear lysates. (i) Western blot of anti-Runx1 immunoprecipitation from SCC FAK-wt, SCC FAK-/, and SCC FAK-wt Runx1 shRNA nuclear lysates probed with anti-Sin3a antibody. IgG control (Ctrl) was done using SCC FAK-wt nuclear lysates.
Figure 1: Graphs showing tumor volume and doubling time for SCC FAK-wt and SCC FAK-/- mice. Panels (c) and (d) display immunofluorescent analyses of cell cycle phases. Panel (e) illustrates vascular density over days post implantation. Panel (f) presents quantification of vascular density for SCC FAK-wt and SCC FAK-/- mice.
Figure 2

(a) 

(b) 

(c) 

(d) 

(e) 

(f) 

(g) 

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Figure 3

(a) Western blot analysis of IGFBP3, FAK, and tubulin in conditioned media and whole cell lysates from SCC FA-K-wt, SCC FA-K-/-, and SCC FA-K-/- IGFBP3 shRNA cells.

(b) Tumor volume (mm$^3$) over time (days) for SCC FA-K-wt, SCC FA-K-/-, and SCC FA-K-/- IGFBP3 shRNA cells. The graph shows a significant decrease in tumor volume for the SCC FA-K-/- IGFBP3 shRNA group compared to the SCC FA-K-wt and SCC FA-K-/- groups.

(c) Immunofluorescence images of SCC FA-K-wt, SCC FA-K-/-, and SCC FA-K-/- IGFBP3 shRNA cells stained for markers in the x, y, and z planes.

(d) Bar chart showing the percentage of cell cycle phases (S/G2/M, G1/S, G1) in SCC FA-K-wt, SCC FA-K-/-, and SCC FA-K-/- IGFBP3 shRNA cells. The SCC FA-K-/- IGFBP3 shRNA group shows a significant increase in G1 phase cells.

(e) Confocal images of SCC FA-K-wt, SCC FA-K-/-, and SCC FA-K-/- IGFBP3 shRNA cells stained for nuclei and cytoskeleton.

(f) Bar chart showing the percentage area occupied by nuclei in SCC FA-K-wt, SCC FA-K-/-, and SCC FA-K-/- IGFBP3 shRNA cells. The SCC FA-K-/- IGFBP3 shRNA group shows a significant decrease in nuclear area compared to the SCC FA-K-wt and SCC FA-K-/- groups.
Figure 4.
Figure 5.
Nuclear FAK and Runx1 cooperate to regulate IGFBP3, cell cycle progression and tumor growth

Marta Canel, Adam Byron, Andrew H. Sims, et al.

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