Nuclear FAK Controls Chemokine Transcription, Tregs, and Evasion of Anti-tumor Immunity

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Nuclear FAK Controls Chemokine Transcription, Tregs, and Evasion of Anti-tumor Immunity

Graphical Abstract

Highlights

- Depletion or kinase inhibition of FAK can cause squamous cell carcinoma regression
- FAK promotes tumor evasion by inducing an immuno-suppressive microenvironment
- Nuclear FAK promotes transcription of chemokines that drive recruitment of Tregs
- FAK-induced Tregs inhibit cytotoxic CD8+ T cells, allowing tumor tolerance and growth

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In Brief

Nuclear focal adhesion kinase (FAK) regulates transcription of chemokines that drive recruitment of tumor-associated regulatory T cells (Tregs), thereby creating a tumor suppressive microenvironment by inhibiting cytotoxic CD8+ T cell activity.

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Nuclear FAK Controls Chemokine Transcription, Tregs, and Evasion of Anti-tumor Immunity

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SUMMARY

Focal adhesion kinase (FAK) promotes anti-tumor immune evasion. Specifically, the kinase activity of nuclear-targeted FAK in squamous cell carcinoma (SCC) cells drives exhaustion of CD8+ T cells and recruitment of regulatory T cells (Tregs) in the tumor microenvironment by regulating chemokine/cytokine and ligand-receptor networks, including via transcription of Ccl5, which is crucial. These changes inhibit antigen-primed cytotoxic CD8+ T cell activity, permitting growth of FAK-expressing tumors. Mechanistically, nuclear FAK is associated with chromatin and exists in complex with transcription factors and their upstream regulators that control Ccl5 expression. Furthermore, FAK’s immuno-modulatory nuclear activities may be specific to cancerous squamous epithelial cells, as normal keratinocytes do not have nuclear FAK. Finally, we show that a small-molecule FAK kinase inhibitor, VS-4718, which is currently in clinical development, also drives depletion of Tregs and promotes a CD8+ T cell-mediated anti-tumor response. Therefore, FAK inhibitors may trigger immune-mediated tumor regression, providing previously unrecognized therapeutic opportunities.

INTRODUCTION

First described more than a decade ago (Onizuka et al., 1999; Shimizu et al., 1999), regulatory T cells (Tregs) have become recognized as a core component of the immuno-suppressive armory utilized by many tumors to keep the anti-tumor activity of antigen-primed CD8+ T cells at bay. Increased Treg numbers has been associated with poorer survival in ovarian (Curiel et al., 2004), gastrointestinal (Sasada et al., 2003), and esophageal (Kono et al., 2006) cancer. Indeed, the ratio of CD8+ T cells/Tregs correlates with poor prognosis, shifting the balance from anti-tumor immunity toward tumor tolerance (Quezada et al., 2006; Sato et al., 2005; Shah et al., 2011). Through secreting a range of chemokines and cytokines, cancer cells can promote the recruitment of Tregs into tumors and can also facilitate their peripheral expansion and retention (Darrasse-Jéze and Podsypanina, 2013; Ondondo et al., 2013). Thus, Tregs can act as a barrier to effective immune-based therapy aimed at activation of a CD8+ T cell anti-tumor immune response. However, the specific signals within tumor cells that stimulate elevated intra-tumoral Tregs, giving rise to tumor tolerance, remain elusive.

FAK is a tyrosine kinase that regulates diverse cellular functions, including adhesion, migration, invasion, polarity, proliferation, and survival (Frame et al., 2010). Using targeted gene deletion in mouse skin, we have previously shown a requirement for fak in tumor initiation and progression to malignant disease (McLean et al., 2004). FAK is also required for mammary tumor progression, intestinal tumorigenesis, and the androgen-independent formation of neuroendocrine carcinoma in a mouse model of prostate cancer (Ashton et al., 2010; Lahlou et al., 2007; Luo et al., 2009a; Provenzano et al., 2008; Pylyyeva et al., 2009; Slack-Davis et al., 2009). Expression of FAK is elevated in a number of tumor types (reviewed in McLean et al., 2005), and FAK inhibitors are being developed as potential cancer therapeutics (Roberts et al., 2008; Shapiro et al., 2014). Many of FAK’s functions in cancer are via its role in signaling downstream of integrins and growth factor receptors at the plasma membrane. FAK also contains putative nuclear localization sequences (NLS) within the F2 lobe of its FERM domain and can localize to the nucleus upon receipt of cellular stress, where it binds to p33 (Lim et al., 2008). However, the extent of FAK’s nuclear functions remains largely unknown. Here, we report a function for nuclear FAK in regulating transcription of inflammatory cytokines and chemokines, in turn promoting an immuno-suppressive, pro-tumorigenic microenvironment. This is mediated...
by recruitment and expansion of Tregs via FAK-regulated chemokine/cytokine networks, and we have found an important role for Ccl5 and TGF\(\beta\)2. Therefore, FAK controls the tumor environment, and suppressing FAK activity, including via a clinically relevant FAK inhibitor, may be therapeutically beneficial by triggering immune-mediated tumor regression.

**RESULTS**

**FAK-Deficient SCC Tumors Undergo Regression in an Immune-Competent Host**

We used a syngeneic model of SCC in which the \(fak\) gene had been deleted by Cre-lox recombination (McLean et al., 2004; Serrels et al., 2012) and mutant tumor cell lines generated. We monitored tumor growth following injection of \(1 \times 10^6\) FAK-deficient cells (FAK\(^{-/-}\)) or FAK-deficient cells that re-expressed wild-type FAK (FAK-WT) at comparable levels to endogenous FAK in both CD-1 nude and FVB (syngeneic) host mouse strains. In CD-1 nude mice, SCC FAK\(^{-/-}\) tumor growth was characterized by a modest growth delay (Figure 1A) as reported previously (Serrels et al., 2012). By contrast, in FVB mice, SCC FAK\(^{-/-}\) tumor growth was characterized by an initial period of growth in the first 7 days followed by complete regression by day 21 (Figure 1B). Thus, FAK expression is required for the survival and growth of SCC tumors in FVB mice with a functional adaptive immune system.

**SCC FAK\(^{-/-}\) Tumor Regression Is Dependent on CD8\(^+\) T Cells**

To characterize the role of adaptive immunity in FAK\(^{-/-}\) SCC tumor regression, we used antibody-mediated T cell depletion in animals bearing SCC FAK\(^{-/-}\) tumors (Figures 1C and S1). Depletion of CD4\(^+\) T cells had no effect on tumor growth. In contrast, depletion of CD8\(^+\) T cells, either alone or in combination with CD4\(^+\) T cells, restored SCC FAK\(^{-/-}\) tumor growth. This implies that cytotoxic CD8\(^+\) T cells were responsible for regression of FAK\(^{-/-}\) tumors (Figure 1C) but does not exclude an accessory role for CD4\(^+\) T cells. T cell depletion in mice bearing SCC FAK-WT tumors (Figure 1D) revealed that: (1) depletion of CD8\(^+\) T cells, either alone or in combination with CD4\(^+\) T cells, caused a significant increase in tumor growth when compared to isotype-treated controls at day 14, and (2) depletion of CD4\(^+\) T cells alone caused regression of FAK-WT SCC tumors by day 21. This implied that FAK-expressing tumors were also under negative pressure from the immune system and that cells from the CD4\(^+\) T cell compartment play a role in protecting FAK-WT tumors from immune-mediated regression (reason discussed later; Figure 3).

Next, we re-challenged mice with \(1 \times 10^6\) SCC FAK-WT cells after regression of primary FAK\(^{-/-}\) SCC tumors, following 7 days of tumor-free survival after the tumors had regressed (Figure 1E, top and middle graphs). Neither FAK-deficient nor FAK-expressing SCC cells were able to grow after the mice had been pre-challenged with SCC FAK\(^{-/-}\) cells. As controls, SCC FAK-WT and FAK\(^{-/-}\) cells were injected at day 28 into mice with no pre-challenge, and these grew as expected (Figure 1E, bottom).

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Figure 1. Loss of FAK or FAK Kinase Activity Results in CD8\(^+\) T Cell-Dependent SCC Tumor Clearance

(A and B) SCC FAK-WT and SCC FAK\(^{-/-}\) subcutaneous tumor growth in immune-deficient CD-1 nude mice (A) and immune-competent FVB mice (B).

(C and D) SCC FAK\(^{-/-}\) (C) and SCC FAK-WT (D) tumor growth in FVB mice treated with T-cell-depleting antibodies.

(E) Secondary tumor re-challenge with SCC FAK\(^{-/-}\) (top) and SCC FAK-WT (middle) cells following a pre-challenge with SCC FAK\(^{-/-}\) cells and a 7-day tumor-free period. Subcutaneous growth of SCC FAK-WT and SCC FAK\(^{-/-}\) tumors injected at day 28 without pre-challenge (bottom).

(F) Tumor growth in FVB mice following subcutaneous injection of SCC FAK-WT, SCC FAK\(^{-/-}\), and SCC FAK-KD cells.

*p < 0.05, **p or ++p < 0.01, ****p < 0.0001; Sidak-corrected two-way ANOVA (A and B) or Tukey-corrected two-way ANOVA (C, versus SCC FAK\(^{-/-}\); D, versus SCC FAK-WT; F, *, versus SCC FAK\(^{-/-}\) and +, versus SCC FAK-KD). Data are represented as mean ± SEM; n = 5–6 tumors.
This implies that, following FAK−/− SCC tumor regression, host mice remain immunized against further tumor challenge because immunological memory had been established. It is possible that either broad immunization against SCCs may have occurred or, more likely, that the FAK−/− and FAK-WT SCCs shared common antigen(s) that are expressed irrespective of FAK status. We conclude that FAK enables SCC cancer cells to suppress an adaptive immune response rather than to circumvent it through evading recognition per se. SCC FAK−/− cells in which a FAK kinase-deficient mutant was re-expressed (SCC FAK-KD) initially grew and then regressed with kinetics that were only modestly delayed when compared to FAK−/− cells, indicating that immune suppression depends on FAK kinase activity (Figure 1F).

We next investigated the nature of the T cell response within tumors derived from all three SCC cell lines using FACS analysis on disaggregated tumor tissue taken at day 7. We did not observe a significant change in the percentage of total CD4+ T cells (Figures 2A and S2 and Table S2) or the percentage of CD4+ T cells that expressed the activation marker CD69 (Figure 2B). In contrast, we did observe a significant increase in the proportion of effector CD4+CD44hiCD62Llow T cells in SCC FAK−/− and FAK-KD tumors when compared to FAK-WT tumors (Figures 2C and S2 and Table S2). Analysis of tumor-infiltrating CD8+ T cells revealed a significant increase in SCC FAK−/− and SCC FAK-KD tumors when compared to SCC FAK-WT tumors (Figures 2D and S2 and Table S2), indicative of a heightened cytotoxic anti-tumor immune response. Staining with the activation marker CD69 identified the presence of CD8+CD69+ T cells in all tumors (Figure 2E). Further analysis revealed an increase in percentage of effector CD8+CD44hiCD62Llow T cells in SCC FAK−/− and SCC FAK-KD tumors when compared to SCC FAK-WT tumors (Figures 2F and S2 and Table S2), especially when effector CD8+ T cell numbers were normalized to account for the observed changes in total CD8+ T cells and presented as a “fold change” (Figure 2G). However, while SCC FAK−/− and SCC FAK-KD tumors had increased effector CD8+ T cells, there were activated CD8+ T cells present in all of the SCC tumors, raising the question of why SCC FAK-WT tumors do not succumb to the cytotoxic CD8+ T cell response.

It is now established that not only the quantity of tumor-infiltrating CD8+ T cells is important, but also their “quality.” Tumor-induced T cell exhaustion has been reported in a number of tumor types, including melanoma (Fourcade et al., 2010) and ovarian cancer (Matsuzaki et al., 2010), and is characterized by expression of co-inhibitory surface receptors, including programmed death receptor 1 (PD-1), lymphocyte-activation gene 3 (LAG-3), and T cell immunoglobulin mucin-3 (Tim-3), either alone or in combination (Fourcade et al., 2010; Sakushi et al., 2010; Wherry, 2011). Analysis of these markers on antigen-primed CD8+CD44hi T cells infiltrating SCC FAK-WT, FAK−/−, and FAK-KD tumors revealed increased surface expression of PD-1, LAG-3, and Tim-3 in CD8+CD44hi T cells present in SCC FAK-WT tumors (Figures 2H–2J). Together, our data imply that antigen-primed CD8+CD44hi T cells infiltrating SCC FAK-WT tumors exhibit a heightened state of exhaustion indicative of a dysfunctional T cell response. Linked to their exhausted state, there was also evidence of decreased proliferation of CD8+ T cells isolated from SCC FAK-WT tumors (judged by Ki-67 staining in Figure 2K).

Histological staining of tumor sections taken at day 7 revealed that: (1) CD8+ T cells are present throughout all tumors, and (2) while CD8+ T cells infiltrating SCC FAK-WT tumors appear predominantly as individual cells, CD8+ T cells infiltrating SCC FAK−/− and FAK-KD tumors are clustered (Figure 2L). Thus, the ability of SCC FAK-WT tumors to evade the anti-tumor immune response is not due to limited CD8+ T cell penetration into these tumors.

**FAK Expression Drives Establishment of an Immuno-Suppressive Environment**

Macrophages, myeloid-derived suppressor cells (MDSC), and Tregs with intrinsic immuno-suppressive capabilities can promote tumor development by inhibiting cytotoxic CD8+ T cell activity in mouse and humans (Beyer and Schultze, 2006; Biragyn and Longo, 2012; Marigo et al., 2008). Flow cytometric analysis revealed no differences in macrophage or MDSC populations that correlated with tumor regression (Figures 3A, 3B, S3, and S4 and Table S2), although this does not rule out an accessory role for these cells in eventual tumor clearance. However, we did find a significantly greater number of CD4+FoxP3+CD25+ Tregs in SCC FAK-WT tumors (Figures 3C and S4 and Table S2) when compared with FAK−/− and FAK-KD tumors (Figure 3C). Tregs have been associated with the development of CD8+ T cell exhaustion (Sakushi et al., 2013) and may therefore be linked to the CD8+ T cell exhaustion that we observed in SCC FAK-WT tumors (Figures 2H–2J). We next calculated the ratio of CD8+ T cells to Tregs (Figure 3D), as this has been reported to correlate with poor prognosis in a number of tumor types (Sato et al., 2005; Shah et al., 2011). We found a substantially lower CD8+ T cell to Treg ratio in SCC FAK-WT tumors when compared to SCC FAK−/− and SCC FAK-KD tumors, which correlated with outcome in terms of tumor tolerance versus immune-mediated tumor regression.

**Tregs Protect FAK-WT Tumors from Immune-Mediated Regression**

We next examined SCC FAK-WT tumor growth in animals treated with an anti-CD25 antibody to deplete Tregs (Figure 3E). Depletion of CD25+ cells led to regression of SCC FAK-WT tumors (Figure 3E). Further analysis revealed an increase in percentage of effector CD8+CD44hiCD62Llow T cells in SCC FAK−/− and SCC FAK-KD tumors when compared to SCC FAK-WT tumors (Figures 2F and S2 and Table S2), especially when effector CD8+ T cell numbers were normalized to account for the observed changes in total CD8+ T cells and presented as a “fold change” (Figure 2G). However, while SCC FAK−/− and SCC FAK-KD tumors had increased effector CD8+ T cells, there were activated CD8+ T cells present in all of the SCC tumors, raising the question of why SCC FAK-WT tumors do not succumb to the cytotoxic CD8+ T cell response.

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FAK Regulates the Transcription of Chemokines and Cytokines to Control Tregs

To address how FAK activity in SCC cancer cells promotes elevated intra-tumoral Tregs, we next analyzed global transcriptional profiles of SCC FAK-WT and SCC FAK−/− cells using Affymetrix GeneChip microarrays (Figure 4A). FAK expression resulted in the upregulation of 498 genes and the downregulation of 598 genes (p < 0.01). The upregulated transcript set in SCC FAK-WT cells included genes involved in immune cell activation and infiltration. Conversely, the downregulated transcript set in SCC FAK−/− cells included genes associated with immune cell suppression. These findings suggest that FAK activity modulates the transcription of genes that control the balance between pro-inflammatory and anti-inflammatory responses, thereby influencing the proportion of Tregs within the tumor microenvironment.

Figure 2. FAK-Depleted Tumors Exhibit a Heightened CD8+ T Cell Response

(A) FACS quantification of total intra-tumoral CD4+ T cells.
(B) FACS quantification of CD69+ cells as a percentage of CD4+ T cells.
(C) FACS quantification of CD4+CD44hiCD62Llow, CD4+CD44hiCD62Lhi, CD4+CD44lowCD62Llow T cell subpopulations.
(D) FACS quantification of total intra-tumoral CD8+ T cells.
(E) FACS quantification of CD69+ cells as a percentage of CD8+ T cells.
(F) Quantification of CD8+CD44hiCD62Llow, CD8+CD44hiCD62Lhi, CD8+CD44lowCD62Llow T cell subpopulations.
(G) Changes in effector (CD8+CD44hiCD62Llow) CD8+ T cells normalized to total CD8+ T cell proportions.
(H) FACS quantification of PD-1+LAG-3+ T cells as a percentage of CD8+CD44hi tumor-infiltrating T cells. n = 6 tumors.
(I) FACS quantification of PD-1+Tim-3+ T cells as a percentage of CD8+CD44hi tumor-infiltrating T cells. n = 3 tumors.
(J) FACS quantification of PD-1+Tim-3+LAG-3+ T cells as a percentage of CD8+CD44hi tumor-infiltrating T cells. n = 3 tumors.
(K) FACS quantification of Ki-67+ cells as a percentage of tumor-infiltrating CD8+ T cells. n = 3 tumors.
(L) Representative histological staining of CD8 in frozen sections from SCC FAK-WT, SCC FAK−/−, and SCC FAK-KD tumors. Dashed white lines demark tumor boundary. Scale bars, 500 μm. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001; ns, not significant; Tukey-corrected one-way ANOVA (C and F, CD44hiCD62Llow only). Data are represented as mean ± SEM; n = 5 tumors unless stated.
FAK-WT cells was associated with a number of processes, including cell migration, receptor binding, secretion, wounding, and ovulation (Figure 4B, top). Analysis of this gene set revealed the chemokine ligand group of genes to be significantly overrepresented (Figure 4B, bottom), which is interesting given that a number of these chemokines and cytokines mediate both Treg recruitment to tumors and induction of peripheral Tregs within tumors (Goldstein et al., 2013; Ondondo et al., 2013).

To establish which chemokines and cytokines were regulated by FAK and to address whether the FAK-dependent transcriptional profile was linked to chemokine receptor expression on tumor-infiltrating Tregs, we performed quantitative (q)RT-PCR analysis. Comparison of chemokine/cytokine transcript levels between SCC FAK-WT and SCC FAK−/− cells revealed a subset of ligands increased >2-fold in SCC FAK-WT cells (Figure 4C). Several of these (Ccl1, Ccl5, Ccl7, Cxcl10) have roles in Treg recruitment (Ondondo et al., 2013) (green arrowheads, Figure 4C), while one (Tgfβ2) has a reported role in peripheral inflammation (Figure 4B, top). Analysis of this gene set revealed that efficient knockdown of Ccl5 using two independent shRNA hairpins (P1 and P2, Figure 5A) resulted in a substantial reduction in both Ccl5-depleted tumors when compared with empty vector control SCC FAK-WT pLKO tumors at day 7 and found that there was a substantial reduction in both Ccl5-depleted tumors when compared with empty vector control SCC FAK-WT pLKO tumors (Figure 5C).

Expanding on these findings, shRNA-mediated knockdown of Tgfβ2 expression in SCC FAK-WT cells also influenced tumor growth (Figures 5B and SSC). Partial knockdown of TGFβ2 had complex effects, which resulted in one of two outcomes. One group (Figure S5C, dashed blue line), grew more rapidly and ulcerated, leading to removal from study at day 14.

**Figure 3. FAK Regulates the Immuno-Suppressive Microenvironment**

(A) FACS quantification of Ly6Ghi and Ly6Glow macrophage populations expressed as a percentage of tumor-infiltrating CD45+ leukocytes.

(B) FACS quantification of Ly6C+Gr1low (M-MDSC) and Ly6C−Gr1hi (G-MDSC) populations expressed as a percentage of tumor-infiltrating CD45+ leukocytes.

(C) Quantification of CD4+CD25+FoxP3+ Tregs expressed as a percentage of tumor-infiltrating CD4+ T cells.

(D) CD8+ T cell-to-Treg ratio calculated using mean values from Figures 2D and 3C.

(E) SCC FAK-WT tumor growth in FVB mice treated with anti-CD25 depleting antibody.

n = 6 tumors. * or p < 0.05, **p < 0.01, ***p < 0.001, **** or ++++p < 0.0001; Tukey-corrected one-way ANOVA (A, *, Ly6Ghi; +, Ly6Glow). Data are represented as mean ± SEM; n = 5 tumors unless stated.
Figure 4. FAK Regulates Transcription of Cytokines Implicated in Treg Recruitment and Expansion

(A) Transcriptomic profiling of SCC FAK-WT and SCC FAK-/- cells.

(B) Functional enrichment analysis of genes upregulated in SCC FAK-WT cells (bottom gray bar in A). Overrepresented biological processes are displayed as a heatmap (log_{10}-transformed color scale) (top); asterisks indicate presence of cytokine-related genes. Overrepresented gene families are displayed as a bar chart (bottom). p < 0.05; Benjamini–Hochberg-corrected hypergeometric tests.

(C) qRT-PCR array analysis of cytokine and chemokine expression in SCC FAK-WT and SCC FAK-/- cells. Gray bar indicates cluster of genes upregulated in SCC FAK-WT cells; cytokine and chemokine gene names are listed. Green arrowheads indicate reported roles in Treg recruitment; red arrowhead indicates reported role in peripheral Treg induction.

(D) qRT-PCR array analysis of chemokine and receptor expression in tumor- and thymus-derived Tregs. Gray bar indicates cluster of genes upregulated in tumor-derived Tregs; receptor gene names are listed.

(E) Interaction network analysis of chemokine ligand gene expression detected in SCC cells (circles, left) and corresponding receptor gene expression detected in Tregs (squares, right). Genes are ordered vertically by fold change. Light gray lines connect receptor-ligand pairs; green lines indicate pairs upregulated at least 2-fold in SCC FAK-WT cells and tumor-derived Tregs.

(F) qRT-PCR analysis of selected cytokine and chemokine gene expression in SCC cells. ***p < 0.001, ****p < 0.0001; Tukey-corrected one-way ANOVA. Data are represented as mean ± SEM.
other group that did not display such frank ulceration, we observed tumor regression by day 27 (Figure S5C, dashed red line). Analysis of Treg levels in SCC FAK-WT shRNA-TGFβ2 tumors at day 7 (regardless of initial growth characteristics) revealed that TGFβ2 knockdown was also associated with a reduction in CD4+ FoxP3+CD25+ Tregs (Figure S5D). Therefore, while the effects of reducing TGFβ2 expression are more complicated than for Ccl5, FAK-dependent TGFβ2 expression does contribute to elevated CD4+FoxP3+CD25+ Tregs in SCC FAK-WT tumors; and in the subset of mice bearing tumors that were able to complete the study, TGFβ2 knockdown also caused tumor regression.
Our findings that FAK regulated the transcription of cytokines and chemokines (including Ccl5 and TGFβ2) that were associated with elevated intra-tumoral Tregs and tumor tolerance led us to consider a possible role for nuclear FAK in regulating the transcription of these genes. Based on previous reports (Lim et al., 2008), which identified putative NLSs within the FERM domain of FAK, we constructed an optimally nuclear targeting-impaired mutant FAK by replacing two arginines (positions 177 and 178) and four lysines (positions 190, 191, 216, and 218) with alamines (termed FAK-NLS). Western blotting of cytoplasmic and nuclear fractions confirmed that the FAK-NLS mutant was indeed defective in nuclear localization (Figure 5D). Subsequent (q)RT-PCR analysis of Ccl5 and Tgfb2 expression in SCC cells expressing only FAK-NLS revealed that FAK nuclear localization was required for transcription of these genes (Figures 5E and S5E, respectively). Thus, nuclear FAK drives the transcription of Ccl5 and TGFβ2, which are required for recruitment and expansion of immuno-suppressive Tregs into SCC tumors, altering the balance between CD8+ T cells and Tregs in favor of tumor tolerance. In support of this, growth of SCC FAK-NLS tumor cells was similar to that of SCC FAK−/−, with ultimate tumor regression (Figure 5F). This confirmed that it was nuclear FAK that afforded protection from the anti-tumor immune response. Western blotting of cytoplasmic and nuclear fractions from SCC FAK-KD showed that the kinase-deficient tumor cells was able to localize to the nucleus, so we conclude that the immune modulatory effects of FAK are dependent on FAK kinase activity in the nucleus (Figure 5G).

We next examined nuclear FAK levels in primary skin keratinocytes, the normal cellular counterparts of the SCC cells used here, and did not find detectable nuclear FAK (Figure 5H). Thus, abundant nuclear localization, and therefore the capacity to exert regulatory control over chemokine and cytokine expression, is likely a feature of oncogenic transformation in skin keratinocytes. This suggests that the nuclear functions of FAK that we have identified—namely, regulating transcription of chemokine/ cytokine networks—may be associated with the cancerous state when FAK is highly expressed.

**Nuclear FAK Interacts with a Network of Ccl5 Transcriptional Regulators**

Having established an important role for the nuclear FAK-dependent transcription of Ccl5 in mediating recruitment and expansion of intra-tumoral Tregs, we wanted to determine how nuclear FAK could exert control over Ccl5 transcription. Using sucrose gradients, we fractionated the nuclei of SCC FAK-WT cells and demonstrated that nuclear FAK was present in the chromatin-containing fraction (Figure 6A). Transcriptional regulation of Ccl5 is mediated predominantly through six short regulatory elements contained within a region of the Ccl5 promoter spanning ~300 base pairs (Fessele et al., 2002). These regulatory elements contain binding sites for a number of TFs, including AP-1, C/EBP, IRF-1, NF-kB, and TATA box-binding protein (TBP), which is part of the transcription factor IID complex (TFIID). Using FAK immunoprecipitation and quantitative label-free mass spectrometry, we identified FAK binding partners in purified nuclear extracts and contextualized these by mapping onto a network of proteins associated with predicted Ccl5 TFs (constructed in silico; Figure 6B). This integrative approach identified a subset of Ccl5 TFs and regulators of these that interact with FAK in SCC cell nuclei (Figures 6C, S6 and Table S1). Interaction network analysis of this protein subset revealed nuclear FAK binding partners with roles in multiple transcriptional pathways, including regulators of AP-1, C/EBP, IRF-1/-7, NF-kB/Rel, and TFIID. Thus, we identified nuclear FAK binding partners that can interact, directly or indirectly, with five of the six main regulatory elements reported to control transcription of Ccl5 in multiple cell types (Fessele et al., 2002). Given that our interaction network was somewhat dominated by proteins associated with the TFIID pathway, including three TBP-associated factors (TAFs) (Figures 6C and S6), we used co-immunoprecipitation to confirm the interaction of nuclear FAK with one of these, TAF9, a core component of the TFIID complex (D’Alessio et al., 2009) (Figure 6D). Our data show that FAK binds to core components of the transcriptional machinery, many of which are known to be located on the promoter of genes undergoing active transcription and that are known or predicted to regulate Ccl5. Therefore, in SCC cells, nuclear FAK associates with chromatin and is physically linked to a network of TFs and their regulators known to modulate Ccl5 expression.

**Small-Molecule FAK Kinase Inhibitor Promotes Immune-Mediated Tumor Clearance**

Therapeutic targeting of FAK kinase activity using small-molecule inhibitors will inhibit FAK signaling not only in tumor cells, but also potentially in multiple host cell types. To complement expression of the FAK-KD mutant protein in the cancer cells and investigate whether a FAK inhibitor could induce immune-mediated regression of SCC tumors, we used the FAK/Pyk2 kinase inhibitor VS-4718 (Shapiro et al., 2014), which is currently in clinical development. Mice were treated with VS-4718 at 75 mg/kg for 24 hr prior to injection of 1 × 106 FAK-WT or FAK−/− SCC tumor cells and twice daily thereafter. This resulted in VS-4718-induced regression of SCC FAK-WT tumors by day 24 (Figure 7A). Following cessation of VS-4718 treatment, no tumor regrowth was observed (data not shown). SCC FAK−/− tumor growth and clearance was not greatly affected by VS-4718 treatment, suggesting that the anti-tumor effects of VS-4718 can be explained by FAK inhibition in tumor cells. Activity of VS-4718 was confirmed using an ELISA to measure FAK autophosphorylation on tyrosine-397 in tumor lysates from mice treated with 75 mg/kg VS-4718 (Figure 7B). Regression of VS-4718-treated SCC tumors was not accompanied by loss of cell viability at day 7, as measured by FACS using a viability stain following tumor disaggregation (Figure 7B). There was a significant but small increase in leukocytes in VS-4718-treated SCC FAK-WT tumors (Figure 7C) and a significant increase in total CD4+ T cells (Figures 7D and S2 and Table S2) and effector CD4+CD44hiCD62Llow T cells (Figures 7E and S2 and Table S2). A significant increase in CD8+ T cells was also evident in SCC FAK-WT VS-4718-treated tumors (Figures 7F and S2 and Table S2), although there was no change in effector CD8+CD44hiCD62Llow T cells (Figures 7G and S2 and Table S2). Crucially, there was a significant reduction in CD4+CD25+FoxP3+ Treg cells in VS-4718-treated SCC FAK-WT tumors, which was similar to that observed in vehicle and...
VS-4718-treated SCC FAK−/− tumors (Figures 7H and S4 and Table S2).

Thus, VS-4718 promoted robust anti-tumor activity, with similar immune cell changes to that observed upon FAK deletion or expression of a kinase-deficient form of FAK. Furthermore, anti-tumor efficacy of VS-4718 was also dependent on CD8+ T cells, and SCC FAK-WT tumors treated with VS-4718 on a CD8+ T cell-depleted background exhibited a growth delay but did not undergo tumor regression (Figure 7I). We conclude that the FAK kinase inhibitor targets mechanisms of immune suppression and may therefore represent a form of effective “immuno-modulatory” therapy that reduces Tregs in the tumor environment. Importantly, the FAK kinase inhibitor does not affect the cytotoxic function of antigen-primed CD8+ T cells. We also found that VS-4718 treatment that was initiated 5 days post-inoculation of 1 × 10^6 SCC FAK-WT cells, when these had
Figure 7. The FAK Kinase Inhibitor VS-4718 Leads to Immune-Mediated SCC Clearance

(A) SCC FAK-WT and SCC FAK−/− tumor growth in FVB mice treated with either vehicle or VS-4718. Treatment started 24 hr pre-tumor cell inoculation and continued for the duration of the experiment.

(B) FACS analysis of cell viability from disaggregated tumors treated with either vehicle or VS-4718.

(C) FACS analysis of vehicle- or VS-4718-treated tumor-infiltrating leukocytes expressed as a percentage of viable CD45+ cells relative to the total number of single cells.

(D) FACS analysis of tumor-infiltrating CD4+ T cells from vehicle- or VS-4718-treated tumors.

(E) FACS sub-categorization of tumor-infiltrating CD4+ T cells into CD45+CD3+CD4−CD8−CD44hiCD62Llow, CD45+CD3+CD4−CD8−CD44hiCD62Lhi, and CD45+CD3+CD4−CD8−CD44lowCD62Llow populations.

(F) FACS analysis of tumor-infiltrating CD8+ T cells from vehicle- or VS-4718-treated tumors.

(G) FACS sub-categorization of tumor-infiltrating CD8+ T cells into CD45+CD3+CD4−CD8+CD44hiCD62Llow, CD45+CD3+CD4−CD8+CD44hiCD62Lhi, and CD45+CD3+CD4−CD8+CD44lowCD62Llow populations.

(H) FACS analysis of tumor-infiltrating CD4+FoxP3+ Tregs expressed as a percentage of tumor-infiltrating CD4+ T cells.

(I) SCC FAK-WT tumor growth in FVB mice treated with either vehicle or VS-4718 and either isotype control or CD8-depleting antibodies.

(J) SCC FAK-WT and SCC FAK−/− tumor growth in FVB mice treated with either vehicle or VS-4718. Treatment started 5 days post-tumor cell inoculation (gray dashed line) and continued for the duration of the experiment.

*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001; ns, not significant; Tukey-corrected one-way ANOVA (E and G, CD44hiCD62Llow only). Data are represented as mean ± SEM; n = 6 tumors.
already formed palpable tumors of ~50 mm³, led to complete tumor regression (Figure 7J).

**DISCUSSION**

We show that nuclear FAK in SCC cancer cells drives the transcription of chemokines and cytokines, including Ccl5 and TGFβ2, which promote the formation of an immuno-suppressive, pro-tumorigenic microenvironment. This is dependent on FAK kinase activity, and expression of a catalytically inactive mutant FAK protein (FAK-KD) or treatment with a small-molecule inhibitor causes tumor regression. This is effective even when tumors are already established, raising the exciting possibility that targeting of FAK kinase activity may have immune-mediated anti-tumor efficacy in patients. We established that nuclear FAK is associated with chromatin and interacts with a number of TFs and transcriptional regulators, including components of the TFIID complex, that are linked to regulation of Ccl5 expression. Our data imply that FAK interacts with core transcriptional machinery to influence gene transcription and promote tumor immune escape.

Historically, FAK has been recognized as an adhesion-related non-receptor protein tyrosine kinase that clusters at focal adhesion (FA) structures and regulates cancer-associated processes, including adhesion, migration, invasion, survival, and proliferation (reviewed in Frame et al., 2010). FAK was also found to translocate to the nucleus (Lim et al., 2008; Luo et al., 2009b), leading to the idea of nuclear functions for FAK within the nucleus. Our data show that, at least in cancer cells, FAK regulates inflammatory transcriptional programs associated with generation and maintenance of a pro-tumorigenic and immuno-suppressive microenvironment. FAK associates with chromatin, and in the context of Ccl5 expression, it interacts with a number of TFs, and regulators of TFs, that bind regulatory elements in the Ccl5 promoter (Fessele et al., 2002). Our data imply that FAK exists in complexes with a number of TAF proteins, including TAF9 and TAF12, key components of the core promoter complex TFIID that serves to initiate transcription by driving recruitment of chromatin remodeling complexes, coactivators, and RNA polymerase II to the promoter (D’Alessio et al., 2009). Therefore, FAK interacts with components of the core transcriptional machinery in order to drive transcription of chemokines and cytokines that contribute to recruitment of Tregs into the tumor environment, promoting immunological tolerance and permitting tumor growth.

Recently, nuclear accumulation of active FAK (phosphorylated on Tyr-397) within tumor cells of patients with colorectal cancer was reported to correlate with poor prognosis (Albasri et al., 2014), highlighting the need to understand the nature of FAK’s role within the nucleus. Studies using endothelial cells, muscle cells, and fibroblasts have previously reported low steady-state levels of nuclear FAK that are substantially increased in response to cellular stress (Lim, 2013; Lim et al., 2008; Luo et al., 2009b). Our work implies that oncogenic stress is another route to inducing high levels of nuclear FAK and that this, in turn, can influence transcriptional programs, such as the chemokine and cytokine networks that control the tumor microenvironment.

A number of therapeutic strategies targeting components of the immuno-suppressive tumor microenvironment are currently being tested, with the aim of restoring anti-tumor immunity by releasing the break on CD8+ T cell cytotoxicity. In pre-clinical models of cancer, targeting Tregs (Ali et al., 2014; Bos et al., 2013) has shown anti-tumor efficacy, either alone or when used in combination with agents that enhance CD8+ T cell activation. A clinical study combining agents targeting cytokotic-T lymphocyte-associated antigen 4 (CTLA-4), which is thought to influence Treg function (Peggs et al., 2009; Quezada et al., 2006; Simpson et al., 2013; Wing et al., 2008), and PD-1, which blocks signals that inhibit T cell function, has reported impressive responses in patients with advanced melanoma (Wolchok et al., 2013). However, this combination of checkpoint blockade antibodies elicits substantial side effects in >50% of patients, highlighting the need to find alternative combinations with improved tolerability. We have shown that targeting FAK kinase activity has the potential to modulate intra-tumoral Treg levels, resulting in robust CD8+ T cell anti-tumor immunity, while others have reported previously that FAK kinase inhibitors block monocyte/macrophage and cancer-associated fibroblast recruitment into tumors by virtue of FAK’s role in regulating their migration (Stokes et al., 2011). Taken together, these findings suggest that targeting the pleiotropic cellular functions of FAK may have a broad impact on the immuno-suppressive tumor microenvironment, differentiating these agents from many therapeutic approaches that target single immune cell populations.

Targeting a molecular pathway that is upregulated in cancer cells may provide tumor specificity and help to overcome some of the potential issues with severe autoimmunity when modulating immune cell populations. FAK inhibitors, such as VS-4718, are in clinical development. VS-4718 is currently in a phase I dose escalation clinical trial in patients with solid tumors (www.clinicaltrials.gov NCT01849744). Our findings provide good rationale for pre-clinical and clinical testing of FAK kinase inhibitors alongside agents that stimulate CD8+ T cell activity, such as the checkpoint blockade therapies that target PD-1 and CTLA-4, which are both in clinical development (Pardoll, 2012).

**EXPERIMENTAL PROCEDURES**

Experiments involving animals were carried out in accordance with the UKCCCR guidelines by approved protocol (HO PL 60/4248). Brief experimental procedures are listed here. For details, please see the Supplemental Experimental Procedures.

**Generation of FAK Nuclear Localization Mutant**

Mutations were introduced into FAK-WT at R177A, R178A, K190A, K191A, K216A, and K218A using PCR-based site-directed mutagenesis.

**Cell Lines**

Isolation and generation of the FAK SCC cell model is described in Serrels et al. (2012). Keratinocyte cultures were prepared as detailed in McLean et al. (2004).

**Western Blot Analysis**

To prepare whole-cell lysates, cells were washed in cold PBS and lysed in RIPA buffer. Cytoplasmic and nuclear extracts were prepared as described in Lim et al. (2008). Lysates were resolved by gel electrophoresis, transferred to nitrocellulose, and probed with respective antibodies.
Subcutaneous Tumor Growth
Cells were injected into both flanks of either CD-1 nude mice or FVB mice and tumor growth measured twice-weekly. Animals were sacrificed when tumors reached maximum allowed size or when signs of ulceration were evident. For treatment with VS-4718, drug was prepared in 0.5% carboxymethyl cellulose + 0.1% Tween 80 and mice treated at 75 mg/kg BID by gavage. No signs of toxicity were observed.

Tumor Growth following Re-Challenge
SCC FAK WT or FAK-/- cells were injected into the right flank of FVB mice. Following tumor regression, mice were left for 7 days before being challenged with SCC FAK-WT or FAK-/- cells injected into the left flank. Tumor growth was measured twice-weekly. Control groups were injected into both flanks at day 28 using mice that had not been pre-challenged with SCC FAK-/- cells.

CD4+, CD8+, and CD25+ T Cell Depletion
Twice-weekly. Control groups were injected into both flanks at day 28 using (2

FACS Analysis of Immune Cell Populations
Tumors established following injection of SCC cells into both flanks of an FVB mouse were removed at day 7. Tumor tissue was processed to obtain single cell suspension for staining and subsequent FACS analysis (antibodies listed in Table S2).

Gene Expression Profiling
RNA was analyzed using the GeneChip Mouse Genome 430 2.0 Array. Normalized data for differentially expressed genes were median combined and clustered using Cluster 3.0 and Java TreeView. Functional enrichment analysis was performed using TopGene.

Quantitative RT2-PCR Array Analysis of Cytokine, Chemokine, and Chemokine Receptor Expression
RNA prepared from SCC cells was analyzed using the mouse cytokine and chemokine RT2 Profiler PCR Array and that from isolated Tregs was analyzed using the mouse cytokine and chemokine RT2 Profiler PCR Array. Relative gene expression (2

shRNA-Mediated TGFβ2 and Ccl5 Knockdown
Cells were subject to two rounds of lentiviral infection prior to selection with puromycin. shRNA constructs used were part of the pLKO lentiviral TRC library.

Preparation and Fractionation of Nuclei and Chromatin
Nuclei were prepared as described (Gilbert et al., 2003) but with a reduced concentration (0.05%) of NP-40 in nuclei buffer B. Soluble chromatin was prepared as described (Gilbert et al., 2004) and fractionated on a sucrose step gradient to separate soluble and chromatin-associated nuclear proteins. DNA was recovered from fractions and subjected to agarose gel electrophoresis. Protein was purified using TCA precipitation. Samples were analyzed by SDS-PAGE and blotted using anti FAK, HP1α, and histone H3 antibodies.

Proteomic Analysis of Nuclear FAK Protein Complexes
FAK nuclear protein complexes were subjected to on-bead proteolytic digestion, desalting, and liquid chromatography-tandem mass spectrometry, as described (Turriziani et al., 2014). For interaction network analysis, Ccl5 transcription factors were extracted from the DECODE database and used to seed a network of 1,000 transcription factor-related proteins using the GeneMANIA plugin in Cytoscape. Proteins specifically isolated in nuclear FAK protein complexes were mapped onto the interactome, and those with physical or predicted direct or indirect interactions with Ccl5 transcription factors were analyzed using the NetworkAnalyzer plugin in Cytoscape.

CD8 T Cell Fluorescent Immunohistochemistry
Tumors were removed 7 days post-implantation and frozen by submersing in liquid nitrogen. Tumor sections were cut, processed and stained. They were imaged using an Olympus FV1000 confocal microscope.

ACCESSION NUMBERS
The microarray data discussed in this manuscript has been deposited in NCBI’s Gene Expression Omnibus and is accessible through GEO series accession number GEO: GSE71662.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, seven figures, and two tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2015.09.001.

AUTHOR CONTRIBUTIONS

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Figure S1. T Cell FACS Analysis Post Antibody-Mediated T Cell Depletion, Related to Figure 1
(A) FACS analysis of spleen and thymus tissue from non-tumor-bearing animals 6 days after commencing antibody treatment. (B) FACS analysis of T cell populations from spleen and thymus tissue from tumor-bearing animals at the end of T cell depletion studies in Figures 1C and 1D.
Figure S2. T Cell FACS Gating Strategy, Related to Figures 2 and 7

(A) FACS gating strategy applied for identification of T cell sub-populations. E = effector, CM = central memory, and N = naive. (B) FMO (full antibody set minus one) control samples used to determine correct gating for T cell sub-population identification.
Figure S3. Macrophage FACS Gating Strategy, Related to Figure 3

(A) FACS gating strategy applied for identification of macrophage sub-populations. (B) FMO control samples used to determine correct gating for macrophage sub-population identification.
Figure S4. MDSC and Treg FACS Gating Strategy, Related to Figures 3 and 7

(A) FACS gating strategy applied for identification of MDSC sub-populations. M-MDSC – Monocytic Myeloid Derived Suppressor Cell; G-MDSC – Granulocytic Myeloid Derived Suppressor Cell. (B) FMO control samples used to determine correct gating for MDSC sub-population identification. (C) FACS gating strategy applied for identification of regulatory T cells (Treg).
Figure S5. Tregs Infiltrating SCC FAK-WT Tumors Express the Thymic Marker Helios; Nuclear FAK Regulates Transcription of TGFβ2, which Contributes to Treg Expansion and Tumor Growth, Related to Figures 3 and 5

(A) FACS analysis of Helios expression in CD4+CD25+FOXP3+ SCC FAK-WT tumor-infiltrating Tregs and Tregs isolated from the thymus of tumor-bearing mice. Control represents background signal from a sample stained with CD4, CD25, and FoxP3 conjugated antibodies but not Helios. Representative replicates are shown in different colors for thymus and SCC FAK-WT samples. (B) qRT-PCR analysis of Tgfb2 gene expression knockdown in SCC cells. ****p < 0.0001 (Tukey-corrected one-way ANOVA). (C) SCC FAK-WT shRNA-TGFβ2 tumor growth in FVB mice. Blue dashed line indicates growth of SCC FAK-WT shRNA-TGFβ2 tumors that had to be sacrificed due to ulceration at day 14. Red dashed line indicates growth of SCC FAK-WT shRNA-TGFβ2 tumors that showed no ulceration. Green solid line indicates the mean growth of all SCC FAK-WT shRNA-TGFβ2 tumors up until cohort numbers were reduced due to ulceration. n = 6 tumors / group. (D) FACS analysis of SCC FAK-WT shRNA-TGFβ2 tumor infiltrating Tregs. ****p < 0.0001 (Tukey-corrected one-way ANOVA) (E) qRT-PCR analysis of Tgfb2 gene expression in SCC FAK-NLS mutant cells. ***p < 0.001, ****p < 0.0001 (Tukey-corrected one-way ANOVA). Data are represented as mean ± SEM.
Figure S6. Nuclear FAK Interactome in the Context of Ccl5 Transcription Factors, Related to Figure 6 and Table S1

(A) Interaction network analysis of proteins that bind FAK in the nucleus of SCC cells. Predicted Ccl5 transcription factors (TFs) (squares; bottom) and respective TF binders (circles; top) enriched by at least two-fold in nuclear FAK immunoprecipitations (SCC FAK-WT over SCC FAK-/- controls; p < 0.05) are shown. Ccl5 TFs not detected (ND) are shown as gray squares. TF complexes or groups are indicated; proteins are labeled with gene names for clarity. TF binders are aligned above TF groups with which there are the greatest number of reported interactions. Overrepresented molecular functions determined by functional enrichment analysis are displayed as a heat map (log10-transformed color scale) (inset). Displayed terms satisfy p < 0.01 (Benjamini–Hochberg-corrected hypergeometric test) with >5 proteins assigned per term. (B) Topological analysis of Ccl5 TF–associated proteins identified in the nuclear FAK interactome. Ccl5 TF binders were clustered using the yFiles Organic algorithm implemented in Cytoscape. Topological parameters were computed using NetworkAnalyzer, excluding self-interactions. Protein node size is proportional to the number of interaction partners in the network (degree); node color indicates betweenness centrality (normalized number of shortest paths between proteins; a measure of the control a protein exerts over the interactions of other proteins in the network). Box-and-whisker plots (inset) show the distributions of degree and betweenness centrality for Ccl5 TF–associated proteins that bind nuclear FAK compared to those that were not enriched in nuclear FAK immunoprecipitations, indicating that FAK binders tend to have more interactions and be more central in the interaction network than undetected Ccl5 TF–associated proteins. Plots display the median (line), interquartile range (box) and 1.5 × interquartile range (whiskers) (n = 169 and 761 Ccl5 TF–associated proteins detected and not detected, respectively, with degree ≥ 1 based on physical or predicted interactions). ****p < 0.0001 (two-tailed Mann–Whitney test).

See also Table S1.
Figure S7. Analysis of FAK pY397 Phosphorylation in Tumors following Treatment with VS-4718, Related to Figure 7
Phosphorylation of FAK on Y397 was measured in protein lysates isolated from tumors following treatment with VS-4718 using ELISA. Tumors were removed within 30 min of treatment. n = 5. Data are represented as mean ± SEM.
Cell
Supplemental Information

Nuclear FAK Controls Chemokine Transcription, Tregs, and Evasion of Anti-tumor Immunity

SUPPLEMENTAL EXPERIMENTAL PROCEDURES

Animal Procedures
All experiments involving animals were carried out in accordance with the UK Coordinating Committee on Cancer Research guidelines by approved protocol (Home Office Project Licence no. 60/4248).

Generation of FAK Nuclear Localization Mutant (FAK-NLS)
Point mutations were introduced into wild-type FAK (FAK-wt) at R177A, R178A, K190A, K191A, K216A, and K218A using site directed mutagenesis. Specifically, 20 ng of pWZL (Hygro) FAK-wt DNA template was mixed with 0.5 µM of each primer in PfuUltra Hotstart master mix (Stratagene) and subjected to PCR with cycle conditions as follows: 95°C (30 s), 18 cycles; 95°C (30 s), 50°C (1 min), 68°C (12 min), final incubation 4°C (infinite). Following PCR, samples were incubated with 10 units of DpnI restriction enzyme at 37°C for 1 hour. The resulting product was transformed into TOP10 chemically competent bacteria (Life Technologies) and plated onto ampicillin agar plates. Colonies were selected and expanded in L broth containing ampicillin overnight, DNA was isolated using a QIAprep DNA miniprep kit (Qiagen), and mutational status was confirmed by sequencing. Three rounds of mutagenesis were required to generate FAK-NLS (R177A, R178A, K190A, K191A, K216A, K218A). Primer sequences were as follows:

FAK R177A/R178A forward, CTAGGTTGCCTTGAAATCGCGCATCCTACGGAGAGATGAG; FAK R177A/R178A reverse, CTCATCTCTCCGTAGGATGCGCGATTTCAAGGCAACCTAG; FAK K190A/K191A forward, GAGGCAATGCATTAGAGGCGGCATCCAACTATGAAGTGCTAG; FAK K190A/K191A reverse, CTAGCACTTCATAGTTGGATGCGCGATTTCAAGGCAACCTAG; FAK K216A/K218A forward, GAAGAGTTTGCTAGATTCAGTGGCGGCCGCAACACTACGAAAATTAATC; FAK K216A/K218A reverse, GATTAATTTTCGTAGTTGCGCGCCGCACTGAATCTAGCAAACCTCTTC.

Cell Lines
Isolation and generation of the FAK SCC cell model is described in detail in Serrels et al. (Serrels et al., 2012). Briefly, SCCs were induced in K14CreER FAK^{flox/flox} mice on the FVB background using the DMBA/TPA two-stage skin chemical carcinogenesis protocol and cells
isolated. Following treatment with 4-hydroxytamoxifen, a FAK-null (FAK-/-) cell clone was isolated, and retroviral transduction was used to stably re-express FAK-wt, FAK kinase-dead (FAK-kd), and FAK-NLS. Briefly, Phoenix Ecotropic cells were transfected with pWZL (Hygro) FAK constructs using Lipofectamine 2000 (Life Technologies) according to manufacturer's instructions. Twenty-four hours post-transfection, cell culture supernatant was removed, filtered through a 0.45-μm Millex-HA filter (Millipore), diluted at a 1:1 ratio in normal SCC cell culture medium, supplemented with 5 μg/ml polybrene, and added to SCC FAK-/- cells for 24 hours. A total of two rounds of infection were performed to generate each cell line. Cells were cultured at 37°C in Glasgow minimum essential medium (MEM) (Sigma-Aldrich) supplemented with 2 mM L-glutamine, MEM vitamins, 1 mM sodium pyruvate (all Sigma-Aldrich), MEM amino acids, and 10% fetal bovine serum (FBS) (both Life Technologies), and maintained under selection using 0.25 mg/ml hygromycin.

Western Blotting
To prepare whole cell lysates, cells were washed twice in ice-cold phosphate-buffered saline (PBS) and lysed in RIPA buffer [50 mM Tris-HCl (pH 7.6), 150 mM sodium chloride, 1% Triton X-100, 0.5% sodium deoxycholate, 0.1% SDS, cOmplete Ultra protease inhibitor cocktail (Roche), PhosSTOP phosphatase inhibitor cocktail (Roche)]. Cytoplasmic and nuclear extracts were prepared as described previously (Lim et al., 2008) and summarized below. Lysates were resolved by 4–15% Bis-Tris gel electrophoresis (Biorad), and proteins were transferred to nitrocellulose then blocked [5% bovine serum albumin in PBS–Tween 20 (BSA/PBS-T)] and probed with either anti-Bcl2 (Cell Signaling Technology), anti-FAK (Becton Dickinson), anti-PARP (Cell Signaling Technology), anti-GAPDH (Cell Signaling Technology), rabbit anti-TAF9 (Abcam), or anti-tubulin (Becton Dickinson) primary antibodies (all 1:1000 in 5% BSA/PBS-T). Bound antibody was detected by incubation with anti-rabbit or anti-mouse DyLight800-conjugated secondary antibody or HRP-conjugated secondary antibody (Cell Signaling Technology) and visualized using a Licor Odyssey imaging system or a BioRad ChemiDoc MP Imaging System.

Nuclear Fractionation
SCC FAK-wt and SCC FAK-/- cells were grown on tissue culture dishes for 48 hours at 37°C prior to fractionation. Keratinocyte cultures were prepared from two-day-old FVB pups as described previously (McLean et al., 2004). Keratinocytes pooled from seven pups were grown on two tissue culture dishes for 72 hours at 37°C prior to fractionation. For subcellular fractionation, cells were washed twice with ice-cold PBS, scraped into ice-cold cyto buffer [10 mM Tris-HCl (pH 7.5), 100 mM NaCl, 0.05% NP-40, 3 mM MgCl2, 1 mM EGTA, cOmplete Ultra protease inhibitor cocktail (Roche), PhosSTOP phosphatase inhibitor cocktail
(Roche)) and incubated for 5 min on ice. Cytosolic supernatants were collected following centrifugation at 800 \(\times\) g for 5 min at 4\(^\circ\)C. Nuclear pellets were washed twice in cyto buffer, re-suspended in RIPA buffer and incubated for 15 min at 4\(^\circ\)C. Nuclear supernatants were collected following centrifugation at 18,000 \(\times\) g for 15 min at 4\(^\circ\)C. For whole cell lysates, cells were washed twice with ice-cold PBS, scraped into RIPA buffer, and incubated for 15 min at 4\(^\circ\)C. Clarified supernatants were collected by centrifugation at 18,000 \(\times\) g for 15 min at 4\(^\circ\)C. Protein concentration was determined by BCA protein assay (Thermo Fisher Scientific), and Western blot analysis performed as described above.

**Subcutaneous Tumor Growth**

Cells were injected subcutaneously into both flanks of either CD-1 nude mice (0.25 \(\times\) 10\(^6\) cells) or FVB mice (1 \(\times\) 10\(^6\) cells) and tumor growth measured twice-weekly using calipers. Animals were sacrificed when tumors reached maximum allowed size, or more commonly when signs of ulceration were evident. For studies involving treatment with VS-4718, drug was prepared in 0.5% carboxymethyl cellulose containing 0.1% Tween 80 (Sigma-Aldrich), and mice were treated at 75 mg/kg BID by oral gavage. Animals were visually monitored for signs of toxicity and weighed prior to each dose of VS-4718. No signs of toxicity were observed. Group sizes ranged from 3–5 mice, each bearing two tumors, and tumor volume was calculated in Excel (Microsoft) using the formula \(v = \frac{4}{3}\pi r^3\). Statistics and graphs were calculated using Prism (GraphPad).

**Tumor Growth Following Re-challenge**

1 \(\times\) 10\(^6\) SCC FAK-/- cells were injected subcutaneously into the left flank of FVB mice and tumor growth measured twice-weekly as described above. Following tumor regression, mice were left for 7 days before being challenged with 1 \(\times\) 10\(^6\) SCC FAK-wt or FAK-/- cells injected subcutaneously into the right flank. Control groups were injected subcutaneously into both flanks at day 28 using mice that had not been pre-challenged with SCC FAK-/- cells. Tumor volume was calculated as described above.

**CD4\(^+\), CD8\(^+\), and CD25\(^+\) T-cell Depletion**

Anti-mouse CD4 (GK1.5, ATCC TIB-207) and CD8 (2.43, ATCC TIB-210) depleting antibodies were purified in-house from conditioned supernatant using a Melon Gel IgG purification kit or purchased from eBioscience (anti-CD4 clone GK1.5, anti-CD8 clone 53-6.7). Rat IgG isotype control and anti-mouse CD25 (clone PC61.5) antibodies were purchased from eBioscience. T-cell depletion was achieved following intra-peritoneal (IP) injection of 150 \(\mu\)g of depleting antibody (same for all antibodies) into female age-matched FVB mice for 3 consecutive days, and maintained by further IP injection at 3-day intervals.
until the study was terminated. $1 \times 10^6$ SCC FAK-wt or FAK-/- cells were injected subcutaneously into both flanks 6 days after initial antibody treatment, and tumor growth was measured twice-weekly as described above. The extent of T-cell depletion was determined at the end of the study using FACS analysis of disaggregated spleen and thymus tissue (Figure S1).

**FACS Analysis of Immune Cell Populations**

Tumors established following injection of $1 \times 10^6$ SCC cells into both flanks of FVB mice were removed at day 7 into RPMI (Sigma-Aldrich) supplemented with 10% FBS (Life Technologies). Tumor tissue was mashed into a pulp using a scalpel and re-suspended in DMEM (Sigma-Aldrich) supplemented with 2 mg/ml collagenase D (Roche). Samples were incubated for 1 hour at 37°C, pelleted by centrifugation at 1600 rpm for 5 min at 4°C, re-suspended in 5 ml of red blood cell lysis buffer (Pharm Lysis Buffer, Becton Dickinson) for 5 min at 37°C, pelleted by centrifugation at 1600 rpm for 5 min at 4°C, re-suspended in PBS and passed through a 70-μm cell strainer. The resulting single cell suspension was pelleted by centrifugation at 1600 rpm for 5 min at 4°C and re-suspended in FACS buffer (PBS containing 1% FBS and 0.1% sodium azide). This step was repeated a total of three times. A sample of the suspension was counted using trypan blue exclusion and the remaining cell suspension pelleted by centrifugation at 1600 rpm for 5 min at 4°C and re-suspended in FACS buffer at a concentration of $1 \times 10^6$ viable cells/ml. One hundred microlitres of each sample were pipetted into each well of a 96-well plate and the plate centrifuged at 1600 rpm for 5 min at 4°C. Cell pellets were re-suspended in 50 μl of Fc block [1:200 dilution of Fc antibody (eBioscience) in FACS buffer] and incubated for 15 min at 4°C. Fifty microlitres of antibody mixture [1:200 dilution of antibodies (listed in Table S2) in FACS buffer] were added to each well and the samples incubated for 30 min in the dark at 4°C. The plate was then centrifuged at 1600 rpm for 5 min at 4°C and the cells re-suspended in FACS buffer and analyzed using a BD FACS Aria II. Regulatory T-cell staining was performed using a Treg staining kit (eBioscience). Staining of spleen and thymus tissue was performed as above. Data analysis was performed using FlowJo software. All antibodies were from eBioscience. For absolute counting, the same protocol as above was used, except tumors were weighed following surgical removal, and prior to FACS analysis samples were resuspended in varying volumes of FACS buffer (dependent on weight) containing CountBright Absolute Counting Beads (Life Technologies). Statistics and graphs were calculated using Prism (GraphPad).

**Gene Expression Profiling**

RNA was prepared from SCC FAK-wt and SCC FAK-/- cells using an RNAeasy kit (Qiagen) according to the manufacturer’s instructions. RNA was analysed using the GeneChip Mouse
Genome 430 2.0 Array (Affymetrix). Gene expression data were analyzed using packages within Bioconductor (Gentleman et al., 2004) implemented in the R statistical programming language. The gene expression data were summarised from CEL files using Ensembl gene identifiers and normalized using the Robust Multi-array Average algorithm (Wu and Irizarry, 2004) within the ‘affy’ package. The data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus (Edgar et al., 2002) and are accessible through GEO Series accession number GSE71662. Data for differentially expressed genes ($p < 0.01$) were median centered and subjected to unsupervised agglomerative hierarchical clustering on the basis of Euclidean distance computed with a complete-linkage matrix using Cluster 3.0 (de Hoon et al., 2004). Clustering results were visualized using Java TreeView (version 1.1.1) (Saldanha, 2004). Functional enrichment analysis against the mouse genome background was performed using ToppGene (Chen et al., 2009).

**Quantitative RT²-PCR Array Analysis of Cytokine, Chemokine, and Chemokine Receptor Expression**

RNA prepared from SCC cells was analyzed using the mouse cytokine and chemokine RT² Profiler PCR Array (PAMM-150Z; Qiagen) and that from isolated Tregs (isolated using Treg isolation kit, 130-091-041; Miltenyi Biotech) was analyzed using the mouse chemokine and receptor array (PAMM-022Z; Qiagen) according to the manufacturer’s instructions. Relative gene expression ($2^{-\Delta\Delta C_t}$) values were log transformed, median centered and subjected to hierarchical clustering as for microarray analysis. For interaction network analysis, an interactome of chemokine ligands and receptors was constructed using the IUPHAR/BPS Guide to Pharmacology database (Alexander et al., 2013; Pawson et al., 2014) and curated from the literature (Bachelerie et al., 2014), onto which expression data for detected genes were mapped and visualized using Cytoscape (version 3.0.2) (Shannon et al., 2003). In addition, expression of selected cytokine and chemokine genes was assessed by standard quantitative RT-PCR. TGFβ2-specific primers were purchased from Qiagen (catalog no. PPM02992A). Ccl5 primers used were: forward, CCCTCACCATCATCCTCACT; reverse, CCTTCGAGTGACAAACACGA. Cxcl10 primers used were: forward, CCCACGTGTTGAGATCATTG; reverse, CACTGGGTAAAGGGGAGTGA. B2M primers used were: forward, GGGAAGCCGAACATACTGAA; reverse, TGCTTAACTCTGCAGGCGTAT. Briefly, a reaction mix consisting of 10 μl SensiFAST SYBR Hi-ROX reagent (Bioline), 0.4 μl of 10 μM stock forward primer, 0.4 μl of 10 μM stock reverse primer, 4.4 μl water, and 4 μl of 20 ng/μl stock random hexamer primed cDNA was run on a Rotor-Gene qRT-PCR machine (Qiagen) using the following cycling conditions: 94°C for 10 min; 40 cycles of 94°C for 10 s, 57°C for 20 s, 72°C for 20 s; and 72°C for 7 min.
Analysis was performed using Rotor-Gene software, and expression relative to B2M was calculated using Excel (Microsoft).

**shRNA-mediated TGFβ2 and Ccl5 Knockdown**

To generate lentiviral particles, $2 \times 10^6$ HEK293FT cells were transfected with a mixture of 10 μg shRNA (RMM4534-EG21808; GE Healthcare), 6.5 μg HIV, and 3 μg VSVG plasmid DNA using Lipofectamine 2000 as per manufacturer’s protocol. Twenty-four hours post-transfection, medium was removed and filtered through a 0.45-μm Millex-AC filter (Millipore) and mixed at a 1:1 ratio with normal SCC growth medium. This mixture was supplemented with polybrene to a final concentration of 5 μg/ml and added to SCC cells for 24 hours. Cells were subject to two rounds of lentiviral infection prior to selection with puromycin at a final concentration of 2 μg/ml. All shRNA constructs used were part of the pLKO lentiviral TRC library (GE Healthcare).

**Preparation and Fractionation of Nuclei and Chromatin**

Nuclei were prepared as previously described ([Gilbert et al., 2003](#)) but with a reduced concentration (0.05%) of NP-40 in nuclei buffer B. Soluble chromatin was prepared as described previously ([Gilbert et al., 2004](#)) and fractionated on a 5-ml sucrose step gradient to separate soluble and chromatin-associated nuclear proteins. Essentially, the soluble chromatin was layered over 10% and 50% sucrose in TEP80 buffer [10 mM Tris-HCl (pH 8.0), 1 mM EDTA, 80 mM NaCl, 100 μM PMSF] and centrifuged at 50,000 rpm for 1 h 50 min in an MLS-50 rotor (Beckman). Five hundred–microlitre fractions were collected by upward displacement in a gradient fractionator (Teledyne ISCO). DNA was recovered from fractions using a PCR purification kit (Qiagen) and subjected to agarose (1.5%) gel electrophoresis. Protein was purified using TCA precipitation. Specifically, one volume of TCA stock solution was mixed with four volumes of protein solution, incubated at 4°C for 10 min, and spun at 14,000 rpm for 5 min at 4°C. The supernatant was discarded, and the pellet was washed twice with ice-cold acetone and dried for 5 min at room temperature. Samples were analysed by SDS-PAGE and western blotted using anti-FAK (Cell Signalling Technologies), anti-HP1α (Millipore), and anti-histone H3 (Cell Signalling Technologies) antibodies.

**Proteomic Analysis of Nuclear FAK Protein Complexes**

Two milligrams of nuclear lysates were incubated with 10 μl agarose-conjugated anti-FAK antibody (clone 4.47; Merck Millipore) overnight at 4°C with rotation. Beads were washed twice with ice-cold RIPA buffer and twice with ice-cold PBS. Bead-bound complexes from biological triplicate experiments were subjected to on-bead proteolytic digestion, desalting
and liquid chromatography–tandem mass spectrometry in technical duplicate as described previously (Turriziani et al., 2014). Mean label-free MS intensities were calculated from technical duplicate data acquisitions for each biological replicate. Peptide and protein false discovery rates were set to 1%. Proteins enriched from SCC FAK-wt nuclei by at least two-fold over SCC FAK-/- control nuclei (p < 0.05) were considered specifically isolated by FAK immunoprecipitation. For interaction network analysis, Ccl5 transcription factors were extracted from the DECODE database (Qiagen) (most relevant transcription factors predicted to bind between 20 kb upstream and 10 kb downstream of the Ccl5 transcription starting site for human, mouse and rat) and used to seed a network of 1000 transcription factor–related proteins using the GeneMANIA plugin (version 2014-08-12-core; human interactions) in Cytoscape. Proteins specifically isolated in nuclear FAK protein complexes were mapped onto the interactome, and those with physical or predicted direct or indirect interactions with Ccl5 transcription factors were analyzed using the NetworkAnalyzer plugin (version 2.7) in Cytoscape.

**CD8 T-cell Fluorescent Immunohistochemistry**

SCC tumors were surgically removed 7 days post-implantation and snap-frozen in a cryovial by submersing in liquid nitrogen. Tumor sections were cut using a cryostat and stored at −80°C. Prior to staining, sections were moved to −20°C overnight. Slides were air-dried for 30 min, fixed with ice-cold acetone (Sigma-Aldrich) for 10 min at room temperature, and washed three times for 5 min each in PBS. Tumor sections were outlined using a hydrophobic barrier pen (DAKO), then incubated with mouse background block for 1 h at room temperature. Samples were incubated with primary antibody [anti-mouse CD8α (clone 53-6.7), rat IgG2a; R&D Systems; 1:100 dilution in 5% goat serum–2.5% BSA in PBS] for 1 h at room temperature, then washed three times with PBS-T, each for a period of 5 min, followed by three 5-min PBS washes. Samples were incubated with secondary antibody [Alexa Fluor 647 goat anti-rat IgG (H+L) (Life Technologies), 1:200 dilution] for 1 h at room temperature in the dark, then subjected to three 5-min washes with PBS-T, followed by one 5-min wash with PBS and one wash with sterile water. Samples were mounted under a coverslip (Thermo Scientific) using ProLong Gold Antifade Mount containing DAPI (Life Technologies) and imaged using an Olympus FV1000 confocal microscope. Images were acquired using an UPLSAPO 20× lens. DAPI and Alexa Fluor 647 were imaged using 405-nm and 635-nm excitation lasers. Three-dimensional image stitching was performed using the Olympus multi-site time-lapse module, with the final image comprising an 8 × 8 matrix with 20% overlap between adjacent images. Images were rendered using Imaris (Bitplane).
### Table S2. Markers and Antibody Combinations Used to Define Immune Cell Populations, Related to Experimental Procedures

(A) T-cell subsets for FACS analysis. (B) Macrophage subsets for FACS analysis. (C) MDSC subsets for FACS analysis. (D) Treg subsets for FACS analysis.

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SUPPLEMENTAL REFERENCES


