GFI1 downregulation promotes inflammation-linked metastasis of colorectal cancer

Wenjing Xing1,2, Yun Xiao3,7, Xiliang Lu1,2,4, Hongyan Zhu1,2, Xiangchuan He1,2, Wei Huang1,2, Elsa S Lopez5, Jerry Wong5, Huanyu Ju1,2, Linlu Tian1,2, Fengmin Zhang6, Hongwei Xu1,2, Sheng Dian Wang4, Xia Li*3, Michael Karin*5 and Huan Ren*1,2,6

Inflammation is frequently associated with initiation, progression, and metastasis of colorectal cancer (CRC). Here, we unveil a CRC-specific metastatic programme that is triggered via the transcriptional repressor, GFI1. Using data from a large cohort of clinical samples including inflammatory bowel disease and CRC, and a cellular model of CRC progression mediated by cross-talk between the cancer cell and the inflammatory microenvironment, we identified GFI1 as a gating regulator responsible for a constitutively activated signalling circuit that renders CRC cells competent for metastatic spread. Further analysis of mouse models with metastatic CRC and human clinical specimens reinforced the influence of GFI1 downregulation in promoting CRC metastatic spread. The novel role of GFI1 is uncovered for the first time in a human solid tumour such as CRC. Our results imply that GFI1 is a potential therapeutic target for interfering with inflammation-induced CRC progression and spread.

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Chronic inflammation is an important risk factor for colorectal cancer (CRC) development and progression.1–3 Colitis-associated CRC shows more rapid progression, lower sensitivity to treatment and higher mortality than sporadic CRC.4 The tumour microenvironment contains cytokines, chemokines, inflammatory mediators, and so on, which have critical roles in almost every stage of progression to malignancy and metastasis.5–11 Approximately 50% of CRC patients develop metastatic disease, and only a minority of patients who undergo resection of metastases attain long-term survival.12

CRC progression to metastasis is a multi-step process involving extensive tumour–stroma cross-talk. Potent metastasis-promoting factors, including cytokines and extra-cellular matrix (ECM) proteins, may trigger epithelial mesenchymal transition (EMT), which drives cancer cell dissemination. ECM-degrading proteases and the c-MET, Notch, and TGFβ signalling pathways regulate tumour–stroma interactions and metastasis.13–15 Among these, TGFβ signalling is essential for the metastasis of CRC cells. Mice treated with a TGFβR1 inhibitor were resilient to metastasis formation.15 Within the tumour microenvironment, amplified cyclooxygenase (COX)-converted prostaglandin E2 (PGE2) further induces the secretion of IL6, IL8, VEGF, iNOS, MMP-2, and MMP-9, leading to increased tumour invasion and metastasis.16 PGE2 exerts its effect by binding to EP receptors. Recent studies have shown that PGE2–EP2 signalling in tumour-infiltrating neutrophils promotes CRC cell growth via amplifying inflammation and shaping the tumour microenvironment.17,18 Moreover, tumour-associated macrophages (TAMs) are a major component promoting cancer progression in many aspects. The association between high TAMs and poor prognosis is observed in CRC.6 Nevertheless, tumour–stroma cross-talk during metastatic progression is highly complex, and the signalling events that control metastasis may be stage-specific. Eventually, potent microenvironmental factors may initiate a programme in which responsive cancer cells are selected as metastatic initiators that may travel to a secondary site and establish metastatic growth.19

Transcription factors (TF) have key roles in determining cell fate and behaviour. Given a significant link between inflammatory bowel disease (IBD) and CRC,20 we aimed to define key TF-regulated networks that may mediate transition programmes during CRC progression by analysing data sets from a large cohort of clinical specimens (Supplementary Figure S1).21 We thus defined a 24-core TF-regulated network comprising inflammation- and/or CRC-related genes.21 To validate the TF network in a cellular model, we applied an inflammation-promoting culture medium,22 LSMCM (LPS-stimulated Monocyte Conditioned Medium), derived from human THP1 cells, to a series of CRC cell lines (Supplementary Figure S2A). LSMCM exposure triggered changes in CRC cell behaviour, including the EMT, and greatly increased cell migration and invasion.21 The expression of two
TFs, including GFI1 (Growth factor independence 1) and STAT3, was significantly altered after LMSCM application (Supplementary Figure S1B and Table 1).

GFI1 is a six-zinc-finger transcription repressor belonging to the SNAG domain family.23 GFI1 has primarily been examined for its roles in different haematopoietic compartments,24,25 cell cycle regulation,23 and haematopoiesis.26 Outside of the haematopoietic system, GFI1 is involved in lineage decisions during intestinal epithelial cell (IEC) differentiation.20,21 Until recently, however, it is not known whether GFI1 contributes to CRC development and progression. Here, we described the establishment of a constitutively activated feed-forward inflammatory signalling circuit normally harnessed through GFI1 during CRC progression. This circuit has crucial roles in promoting CRC metastatic spread.

Results

Monocyte-derived TGFβ induces inflammation-linked CRC cell metastatic behaviours. We generated a previously established cellular model using LMSCM culture medium conditioned by LPS-stimulated monocytes (Supplementary Figure S2A). Four (HT29, LoVo, HCT116, and SW480) of the five CRC cell lines (including SW620) actively responded to LMSCM, morphologically changing into spindle-shaped fibroblast-like cells (Figure 1a). Further examination revealed that LSMCM greatly induced EMT and increased cell migration and invasion in response cells (Figure 1b–i and Supplementary Figure S2B); however, SW620 cells were the least responsive to LSMCM (Figure 1j and k), in which remnant LPS had a minor effect (Supplementary Figure S2C). Moreover, the conditioned medium derived from LPS-stimulated macrophages differentiated from human peripheral blood mononuclear cells (PBMCs), or the U-937 human monocyte cell line significantly induced EMT and increased cell migration and invasion in HT29 cells (Supplementary Figure S2D–F). These supplementary models provided proof of concept on the cellular model system to study the effect of TAMs and the tumour microenvironment in vitro.

To understand how LSMCM elicited these phenotypic changes, we analysed the supernatants of LMSCM-stimulated HT29 cells on 40 selected cytokines highly relevant to activated macrophages.32 The results indicated that the concentrations of TGFβ1, IL1α, IFNγ, IL309 (CCL1), and TNFβ were significantly decreased within the first 6 h of treatment and subsequently returned to levels equivalent to those present in the original LSMCM after 24 h of incubation, whereas GM-CSF, IL8, MCP1, PDGFβB, and sTNFR1 remained elevated for up to 24 h (Figure 2a). These data suggested that HT29 cells may consume or degrade certain cytokines in the medium before secreting these cytokines; these cells rapidly produced additional GM-CSF and IL8.

To identify the most important factors in LSMCM-affecting CRC cell behaviour, we referred to a previously identified core TF network linking IBD and CRC (Supplementary Figure S1).21 Among the 24-core TFs and their regulated network in LMSCM-stimulated HT29 cells, two genes, TGFBR2 and TLR8, were significantly upregulated (Table 1). Consistently, TGFBR2 mRNA expression increased upon LSMCM addition (Supplementary Figure S3A). Furthermore, the application of anti-TGFβ-neutralising antibodies prior to LMSCM significantly blunted EMT-related behavioural changes and migration/invasion in CRC cells (Figure 2b and c and Supplementary Figure S3B, S3C). Thus, TGFβ signalling may be an important contributor to the observed changes in CRC cell behaviour.

Inflammation-linked changes in CRC cell behaviour resulting from multiple factors in the cell model system. CRC cells exhibited maximal behavioural changes after treatment for 24 h with LMSCM. To investigate the factors driving these changes, we examined the expression of multiple effectors in HT29 cells for up to 96 h. The expressions of GFI1 and STAT3 were significantly altered upon LMSCM treatment, and the GFI1-associated factors PTGER2 (coding for EP2), TGFBR2, and STAT3-associated AKT1 were upregulated (Table 1). In addition, we examined ERK1/2, NF-κB (p65 subunit), EMT-related Snail, Zeb1, and Twist in the expression analysis, considering their effects on inflammation and metastasis.33,34 We observed sequential changes in the expression or activities of almost all of these effectors for up to 96 h at the protein level (Figure 2d and Supplementary Figure S3D). Two of the earliest changes were decreased GFI1 and increased ERK phosphorylation. Moreover, the expression of the GFI1-repressed gene PTGER2 was expectedly increased upon LMSCM addition (Figure 2e). Other changes subsequently occurred, including the activation of STAT3, EP2, AKT, and p65 (Figure 2d);

| Gene     | Fold changea | Gene     | Fold changea |
|----------|--------------|----------|--------------|
| GFI1b    | −2.58        | BUB1B    | 1.32         |
| TGFBR2   | 2.19         | TLR8     | 2.19         |
| TCF7L2   | 1.32         | RUNX3b   | −1.14        |
| REG4     | 1.09         | CD40     | −1.24        |
| HDAC5    | 1.15         | CXCR4    | −1.15        |
| SMAD2    | 1.21         | HDAC5    | −1.09        |
| MUC2     | 1.34         | CXCL13   | −1.32        |
| PTGER2   | 4.80         | CCR7     | −1.17        |
| TCF7L2b  | −1.00        | SMAD2b   | 1.44         |
| RPS9     | −1.02        | SOX10b   | 1.04         |
| APC      | −1.12        | TFF3b    | 1.37         |
| PTPRH    | −1.20        | SMAD7b   | 1.16         |
| ARHGEF4  | −1.06        | ETS1b    | 1.12         |
| NAT1     | −1.18        | POU5F1Bb | −1.61        |
| STAT3b   | 2.58         | FOX1b    | −1.41        |
| AKT1     | 3.58         | HOXA4b   | −1.30        |
| TP53I3   | 2.58         | NFYBb    | −1.05        |
| XBPIb    | −1.60        | FOXC1b   | −1.20        |
| C5       | −1.42        | ZNF589b  | 1.07         |
| CCL24    | −1.06        | TCF4b    | 1.12         |
| CHST2    | −1.20        | SREBF2b  | −1.13        |
| IL1B     | −1.16        | CEBPGb   | 1.16         |
| PBX1b    | 1.80         | NFATCb   | −1.07        |
| IL23R    | 1.14         | PROM1b   | 1.05         |
| CCX1     | 1.21         | BACH2b   | −1.26        |
| TYMS     | 1.64         | POU2F2b  | 1.02         |

*aCells were treated with LMSCM for 24 h. Gene expression showing at least twofold change was recognised in red as being significantly altered. Consistent results from independent real-time PCR assays were obtained. *Transcriptional factor, others are their related genes. The bold entries indicate the transcriptional factors
Figure 1  LSMCM increases CRC cell metastatic behaviour. (a) Culture with LSMCM induces morphology changes in HT29, LoVo, HCT116, SW480, and SW620 cells. (b) qRT-PCR of E-cadherin, Fibronectin, and Vimentin mRNA in HT29 with LSMCM for different times or prepared from supernatant of THP1 cells that were incubated with different LPS concentrations. (c) HT29 cells were treated with LSMCM for different times; or prepared from supernatant of THP1 cells that were incubated with different LPS concentrations. Cell migratory and invasive activities were measured and quantified. (d) LoVo cells were treated as in (c) and their migratory and invasive activity was measured and quantified. 
(e) qRT-PCR of E-cadherin, Fibronectin and Vimentin mRNA in LoVo cells treated as in (b). Results of similar experiments were exhibited for HCT116 cells, SW480 cells and SW620 cells, respectively, in (f–k). (b–k) show means ± S.D. of three independent experiments. Statistically significant differences are indicated. *P<0.05; **P<0.01
Figure 2  Monocyte-derived TGFβ downregulates GFI1 and induces EMT and cell metastatic behaviour. (a) Cytokines in culture medium from HT29 cells treated with LSMCM for 6 or 24 h were measured by Cytokine antibody arrays. Results shown are fold changes compared with freshly collected LSMCM. (b) qRT-PCR of E-cadherin, Fibronectin, and Vimentin mRNA in HT29 cells treated with LSMCM with/without anti-TGFβ-neutralising antibodies for 24 h. (c) Quantification of data shown in Supplementary Figure S3B, depicting the role of TGFβ in LSMCM-induced changes in cell behaviour. (d) HT29 cells were incubated with LSMCM for up to 96 h. Whole-cell lysates were prepared at the indicated time points and analysed for expression and phosphorylation of the indicated proteins. (e) Time course of GFI1 and PTGER2 mRNA expression in HT29 cells treated with LSMCM by qRT-PCR. (f) Effect of anti-TGFβ-neutralising antibodies on LSMCM-induced suppression of GFI1 expression. (g) qRT-PCR of GFI1 mRNA in HT29 cells treated with LSMCM plus indicated concentrations of anti-TGFβ-neutralising antibodies. (h) HT29 cells were treated with LSMCM for up to 96 h, and the expression or phosphorylation of SMAD2/3/4 were analysed. (i) qRT-PCR of SMAD2, SMAD3, SMAD4, SMAD7, and RPS6KB1 mRNA in HT29 cells treated with LSMCM or LSMCM plus 2.5 μg/μl anti-TGFβ-neutralising antibodies. (b), (c), (e), (g), (i) show means ± S.D. of three independent experiments. Statistically significant differences are indicated. *P < 0.05; **P < 0.01; ***P < 0.001.
increased expressions of Snail and Zeb1, but not Twist (Supplementary Figure S3D).

Significantly, the LSMCM-induced GFI1 decrease was reversed by TGFβ neutralisation in a dose-dependent manner in HT29 cells (Figure 2f and g). Further assays indicated that TGFβ might exert this effect via SMAD-dependent pathways. Notably, phospho-SMAD3 was significantly elevated as early as 6 h and remained activated for 96 h; phospho-SMAD2 and the expression of SMAD4 were also elevated (Figure 2h). Conversely, TGFβ neutralisation reversed the LSMCM-mediated SMAD3 elevation (Figure 2i). Notably, the expression of RPS9KB1, a factor in SMAD-independent pathways, was unaffected by either LSMCM or additional TGFβ neutralisation (Figure 2j). Furthermore, in the supernatant, three other cytokines, IL1α, TNFβ, and IFNγ, whose expression changes were similar to those of TGFβ under LSMCM treatment (Figure 2a), led to GFI1 downregulation and the activation of cellular downstream effectors to varied extents (Supplementary Figure S3E–G). Collectively, these results showed that effects of multiple factors contributed to the transformation of CRC cells under LSMCM treatment.

**GFI1 is a gating regulator in the cellular transformation of CRC cells.** We next examined whether decreased GFI1 expression affected the aforementioned changes in HT29 cells. Forced GFI1 expression decreased PTGER2 (gene name for EP2) mRNA (Figure 3a) but not vice versa (Figure 3b), although a potential GFI1 binding site was detected in the STAT3 promoter but not in PTGER2 (Figure 3c). Chromatin immunoprecipitation (ChiP) assays confirmed less and less GFI1 bound to the endogenous STAT3 promoter during LSMCM stimulation (Figure 3d). GFI1 expression significantly decreased phosphorylated and total STAT3 (Figure 3e). Luciferase reporter assays revealed specific binding between GFI1 and the indicated sequence (Figure 3c) in the STAT3 promoter (Figure 3f). Furthermore, STAT3 siRNA treatment moderately reduced EP2 protein expression (Figure 3g) but not its mRNA expression (Supplementary Figure S4A). Conversely, PTGER2 siRNA treatment did not affect STAT3 expression or activation by LSMCM (Figure 3h, Supplementary Figure S4B). We subsequently detected the physical interaction between STAT3 and EP2, which was further enhanced after LSMCM treatment (Figure 3i, Supplementary Figure S4C), but not transcriptional regulation of STAT3 on PTGER2 (Supplementary Figure S4D). Thus, the GFI1-STAT3-EP2 signalling cascade is activated in HT29 cells under LSMCM stimulation.

Moreover, although GFI1-expressing cells showed decreased ERK1/2 phosphorylation (Figure 3j), ERK1/2 siRNA, or U0126 (an MEK inhibitor) treatment reduced EP2 expression at the protein level (Figure 3k, Supplementary Figure S4E) but not mRNA (Supplementary Figure S4F). However, PTGER2 siRNA had no effect on ERK1/2 phosphorylation (Figure 3l). In addition, we observed physical interactions between ERK1/2 and EP2 (Figure 3m). These data suggest that, similar to STAT3, ERK1/2 activation upon GFI1 reduction contributed to increased EP2 expression. In response to LSMCM treatment, the activation of AKT and p65 was observed later than EP2 upregulation (Figure 2d). PTGER2 siRNA treatment reduced both AKT and p65 phosphorylation (Figure 3n). Furthermore, the inhibition of AKT phosphorylation using Ly294002 reduced p65 phosphorylation; RELA (coding for p65) siRNA, or PDTC (an inhibitor of NF-κB activation) had only a minor effect on AKT phosphorylation (Figure 3o, Supplementary Figure S4G). These data suggest the existence of an EP2-AKT-NF-κB cascade.

**GFI1 downregulation enhances the metastasis of human CRC cells.** To determine whether the activation of signalling effectors affects the consequent production of cytokines, we examined expressions of 10 cytokines in HT29 cells incubated in LSMCM for 24 h. The results showed that, except for IFNγ (data not shown), the mRNA expression of GM-CSF, IL8, MCP1, PDGFBB, sTNFR1, TGFβ1, IL1α, I309, and TNFβ significantly increased (Figure 4a). Moreover, ELISA showed a significant decrease in TGFβ production at 6 h and a return to basal amounts at 24 h after LSMCM addition (Figure 4b), consistent with previous data (Figure 2a). We next applied shRNA, siRNA or gene overexpression to GFI1, PTGER2, STAT3, or RELA in the cell model. The results showed that forced GFI1 expression suppressed all 9 cytokines at the mRNA level, but GFI1 knockdown induced the expression of these molecules; PTGER2 siRNA had only moderate inhibition effects. In comparison, gene knockdown of STAT3 or RELA had variable effects, inhibiting the expression of certain cytokines while increasing others (Figure 4c). Collectively, the most consistent effect of LSMCM incubation was the repression of GFI1, which acts as a suppressor of cytokine gene expression. The effect of decreased GFI1 may be mediated via STAT3 and ERK1/2, which stimulate EP2 expression (Figure 4d).

Moreover, GFI1 overexpression in LSMCM-stimulated HT29 cells reversed the changes in the expression of EMT markers and greatly inhibited cell migration and invasion (Figure 4e and f); conversely, GFI1 knockdown significantly increased cell migration & invasion, induced EMT (Figure 4g and h), and increased EP2 expression and STAT3, p65, AKT, and ERK phosphorylation (Supplementary Figure S4H). Keeping consistent, we further showed that although SW480 cells exhibited the weakest response and SW620 cells exhibited the strongest response to LSMCM stimulation, the expression of GFI1 was the highest in SW480 cells and lowest in SW620 cells among the tested CRC cells (Figure 5a and b). In parallel, forced GFI1 expression in SW620 cells significantly reduced cell migration and invasion and vice versa (Figure 5c–f). Whereas SW620 cells poorly responded to LSMCM stimulation (Figure 1a, j and k), forced GFI1 expression greatly enhanced SW620 sensitivity to LSMCM (Figure 5g). Collectively, these data indicated an essential role for GFI1 downregulation in coordinating the changes in the cell behaviour and metastatic potential of CRC cells.

**GFI1/Gfi1 inhibits CRC metastasis to distal sites.** We further examined GFI1 effect in vivo. GFI1 overexpression and control HT29 cells incubated with LSMCM for 24 h were subcutaneously (s.c.) transplanted into DKO (BALB/c- Rag2−/−; gc−/−) mice, and the s.c. tumour masses were harvested. The tumour growth rates, volume and mouse body weight were similar between the two groups of animals.
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(a) PTGER2 mRNA expression in NC and GF1 treated cells.

(b) PTGER2 siRNA knockdown in NC and GF1 treated cells.

(c) Table of sequence information.

(d) Western blot analysis of STAT3 expression.

(e) GF1 binding site on STAT3 promoter.

(f) Luciferase activity assay for WT and mutation.

(g) STAT3 siRNA knockdown in HT29 cells.

(h) PTGER2 siRNA knockdown in HT29 cells.

(i) Western blot analysis of EP2 and STAT3 expression.

(j) GF1 knockdown in HT29 cells.

(k) MAPK1/2 siRNA knockdown in HT29 cells.

(l) PTGER2 siRNA knockdown in HT29 cells.

(m) Western blot analysis of ERK1/2 expression.

(n) Western blot analysis of PTGER2 expression.

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before and after tumour cell transplantation (Supplementary Figure S5A-E). Immunoblot analysis confirmed GFI1 overexpression and the decreased expression of STAT3, Snail, and Zeb1 in GFI1-positive tumours in vivo (Supplementary Figure S5F).

Next, we injected cultured GFI1-expressing (GFI1-T) and NC-T s.c. tumour cells via the tail vein (t.v.) into DKO mice (Supplementary Figure S5G). LSMCM stimulation for 24 h prior to tumour cell injection greatly accelerated lung metastasis in the NC-T group, reducing latency from 90 to 30 d compared with cells not subjected to LSMCM stimulation (data not shown). Moreover, significantly more lung metastases were observed in the NC-T than GFI1-T group at 30 d (Supplementary Figure S5H, I). Furthermore, at 60 d, all eight NC-T mice and only one of the eight GFI1-T mice showed lung metastases, whereas three out of eight NC-T and no GFI1-T mice showed secondary metastasis to the liver, with significantly larger metastases in the lungs of NC-T mice (Supplementary Figure S6A-F). In addition, NC-T mice exhibited more weight loss compared with GFI1-T mice (Supplementary Figure S6G). H&E staining of the lung biopsies showed that the NC-T mice had more macroscopic tumours than GFI1-T mice; and NC-T liver metastases were also large (Supplementary Figure S6B, E).

To further investigate the role of murine Gfi1 in CRC cell metastasis in immune competent mice, we constructed MC38-Gfi1 and MC38-con cell lines using CRISPR/Cas9 technology. Forced Gfi1 activation in metastatic MC38 CRC cells reduced the rates of cell migration and invasion, diminished the expressions of EP2 and STAT3, and attenuated STAT3, AKT, and p65 phosphorylation (Figure 6a and b). We next intrasplenically (i.s.) injected these cells into syngeneic C57BL/6 mice and subsequently examined liver metastases and disease progression. Gfi1 overexpression greatly reduced tumour metastatic incidence (Supplementary Figure S6H), liver metastases sizes and body weight in age-matched groups (Figure 6c–f). In addition, MC38-Gfi1 mice survived much longer than the control mice (Figure 6g).

**GFI1 downregulation increased the metastatic ability of CRC cells in vivo.** HT29 cells harbouring GFI1 gene knockdown by shRNA or control plasmid were s.c. transplanted into nude mice and harvested at 20 d. The results showed no significant differences between the two groups of mice in animal body weight, tumour formation rate, and tumour weight and volume (Supplementary Figure S7A–E), although shGFI1 tumours seemed smaller than the controls (Supplementary Figure S7C, D). Next, we injected these two groups of cells via the t.v. into the mice. At 40 d after injection, all of the lungs from the shGFI1 mice and none of the lungs from control mice developed pronounced lung metastases (Supplementary Figure S7F, G). GFI1 knockdown had a minor effect on the body weight of the mice; H&E staining revealed that the lungs from shGFI1 mice showed aggregated CRC cells around the blood vessels to form larger metastases (Supplementary Figure S7H, I).

To determine whether GFI1 downregulation correlated with CRC metastatic progression in humans, we analysed a TCGA data set including 155 patient biopsies from primary tumour sites. The results indicated that the primary tumours from patients with metastatic CRC (M1) expressed significantly less GFI1 than those from patients who did not progress to metastatic disease (M0) (Figure 6h). Collectively, these data strongly suggest that GFI1 downregulation promotes CRC metastasis to distant organs.

**Loss of GFI1 in advanced human CRC samples.** To further relate GFI1 expression with CRC metastasis, we examined a series of human biopsies, including normal colon tissues (n = 10), cancer adjacent normal colonic tissue (n = 10), precancerous colon lesions (n = 5 adenomatous and n = 5 polyp tissues), CRC (n = 23) and CRC metastatic lesions (n = 23) using IHC (Figure 7a). The loss of GFI1 expression was primarily observed in advanced CRC (stage IIb and higher) and metastases compared with normal and lower stage tissues, respectively (Figure 7b and c). Decreased GFI1 expression was initially observed at stage IIb and further decreased at stage III (Figure 7a), suggesting that GFI1 downregulation likely precedes metastatic spread. In addition, in the same set of human samples, the markedly increased expression of STAT3 was observed in CRC compared with normal biopsies (Supplementary Figure S8A, B), although the expression was similar between early- and late-stage tumour samples (Supplementary Figure S8C). Moreover, a negative correlation between GFI1 and STAT3 expression at the protein level was observed across all categorised sample sets, and significantly correlated expression was only observed in normal colon specimens and metastases (Supplementary Figure S8D). These data...
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a) mRNA expression

b) TGFβ (pg/ml)

(c) mRNA expression

d) Tumor-associated monocyte

e) HT29

f) HT29

g) HT29

h) HT29

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Figure 4  GFI1 controls an inflammatory signalling circuit that enhances invasive and migratory behaviour of human CRC Cells. (a) Invasive and migratory activity of different CRC cell lines (SW480, HT29, LoVo, SW620). (b) qRT-PCR of GFI1 mRNA in three other human CRC cells shown relative to GFI1 mRNA content of HT29 cells. (c) GFI1 protein expression in SW620 cells transfected with GFI1-encoding plasmid or control. (d) GFI1 protein expression in SW480 cells transfected with GFI1 shRNA plasmid or control. (e) Migratory and invasive behaviour of SW620 cells transfected with negative control or GFI1-encoding plasmids or control. (f) Migratory and invasive behaviour of SW480 cells transfected with negative control or GFI1 shRNA plasmids or control. (g) Migratory and invasive behaviour of SW620 cells transfected with negative control or GFI1-encoding plasmids upon LSMCM stimuli. (b), (e–g) are shown as means ± S.D. of three independent experiments. Statistically significant differences are indicated. *P<0.05; **P<0.01

Figure 5  GFI1 controls invasive and metastatic behaviour of CRC cells. (a) Invasive and migratory activity of different CRC cell lines (SW480, HT29, LoVo, SW620). (b) qRT-PCR of GFI1 mRNA in three other human CRC cells shown relative to GFI1 mRNA content of HT29 cells. (c) GFI1 protein expression in SW620 cells transfected with GFI1-encoding plasmid or control. (d) GFI1 protein expression in SW480 cells transfected with GFI1 shRNA plasmid or control. (e) Regulation of HT29 invasive and migratory behaviour by ectopically expressed GFI1. (f) Regulation of HT29 invasive and migratory behaviour by knocking down of GFI1 expression. (h) qRT-PCR of E-cadherin, Fibronectin and Vimentin mRNA in HT29 cells transfected with GFI1 shRNA or control. (a–c), (e–h) are shown as means ± S.D. of three independent experiments. Statistically significant differences are indicated. *P<0.05; **P<0.01
**Figure 6** Gfi1 inhibits MC38 metastasis to liver. (a) Migratory and invasive activity of MC38 cells transfected with control or Gfi1-activation plasmids. (b) Immunoblot analysis of MC38-con and MC38-Gfi1 cells demonstrating effects of Gfi1 on indicated signalling proteins. (c) Liver metastases in mice injected intrasplenicly (i.s.) with MC38-con or MC38-Gfi1 cells. Arrows pointed to metastases in the liver. (d) Representative H&E-stained sections of the livers of mice injected with MC38-Gfi1 or MC38-con cells. (e) Weights of livers in mice injected with MC38-Gfi1 or MC38-con cells 16 days after i.s. injection. (f) Weights of mice injected with MC38-Gfi1 or MC38-con cells 16 days after i.s. injection. (g) Kaplan-Meier survival curve in respective MC38-Gfi1 and -con mice, log-rank test, $P < 0.0001$. (h) GFI1 expression levels in the primary tumour of human CRC in patients who did (M1) or did not (M0) develop distal metastases in a TCGA data set.

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suggest that GF11 downregulation may be highly correlated with increased STAT3 expression in CRC human samples. To confirm the role of GF11 in an even larger number of clinical samples, we examined the public databases of colon cancer data sets, including CRC data from TCGA, GSE17538, GSE41258, and GSE14333, comprising a total of 918 clinical samples. Notably, a significant decrease in GF11 expression was observed in advanced CRC relative to early-stage tumours (Figure 7d). To evaluate the clinical prognosis value of GF11 expression, we used another set of CRC data containing 177 samples harbouring relapse-free survival time (GSE39582). By stratifying cancer patients according to GF11 mRNA expression (median), we observed that lower GF11 mRNA expression was associated with decreased patient survival (Figure 7e). Interestingly, a significant GF11 loss was also observed in adenomas and polyps compared with normal tissues (Figure 7f), suggesting that this loss might be an early event in colon carcinogenesis that needs to exceed a certain threshold to enable cancer cells to acquire metastatic properties.

Discussion
Cancer metastasis is a multi-step process relying on tumour–stroma interactions. Inflammation greatly contributes to CRC development and progression. In the present study, we identified an inflammatory environment-mediated transcriptional programme that controls the metastatic potential of CRC through GF11, a transcriptional repressor. Curiously, a key cytokine responsible for GF11 downregulation is TGFβ. Although in early CRC cells and premalignant lesions, the TGFβ signalling pathway has tumour-suppressive activities, the pro-metastatic activity of this signalling pathway is observed in more advanced cancer. These results suggest that tumour stromal cytokines may accomplish pro-metastatic activity through GF11. Importantly, the analysis of human clinical specimens confirmed that GF11 downregulation correlated with increased CRC metastasis.

Previously, GF11 demonstrated importance in the lineage decision process during IEC differentiation. but the results of the present study are the first to implicate GF11 as a negative regulator of CRC metastasis. GF11 downregulation during the metastatic progression of CRC may reflect increased TGFβ1 expression in TGFβR- and SMAD2/3-expressing CRC cells. In addition, using a cell model of inflammation-driven metastasis, we verified that other cytokines, such as IL1α, TNFβ or IFNγ in the tumour microenvironment also contributed to GF11 decrease, thereby activating CRC cell metastatic behaviours; yet the effect of the cytokine combination or other factors needs further verification. Consistently, some of these cytokines, or conditioned medium from TAMs, were implicated in EMT-like processes and promoting metastatic behaviours in CRC cells. Moreover, NF-κB and/or STAT3 signalling and the feed-forward activation circuit in CRC cells may support metastasis to remote organs.

STAT3 has been demonstrated as an important player in inflammation-associated CRC progression. Our data highlight STAT3 as a core TF in the inflammation-mediated TF network that controls CRC metastasis and showed that GF11 expression inhibits STAT3 activation. Previous studies showed that GF11 enhanced STAT3 activation through interactions with the STAT3 inhibitor PIAS3. In contrast, the results of the present study showed that although GF11 controlled STAT3 transcription through binding to its promoter, GF11 downregulation triggered via LMSCM abrogated the effect, resulting in STAT3 activation. This variation suggests the operation of a context-dependent regulatory circuit. Consistent with other data, we verified that EP2 signalling activation was highly relevant to inflammation-associated CRC progression.

It has been suggested that disseminated cancer cells are selected via stroma–tumour interactions before adapting to new microenvironmental conditions and giving rise to a metastatic clone. There is a debate as to whether this selection occurs within or outside the primary tumour. We observed that once GF11 downregulation within CRC cells reaches a certain threshold, CRC cells become competent for metastatic spread as a result of the long-term activation of a regulatory circuit that is normally repressed by GF11. The ectopic overexpression of GF11/Gf11 impaired the function of this signalling circuit and inhibited the metastatic activity of CRC cells, and vice versa. Although we observed that GF11 expression is lost in human metastatic CRC and confirmed the ability of GF11 to suppress metastatic spread, the CRC metastasis process is complex and likely to involve additional regulation. For example, despite a strongly reverse relationship between GF11 and STAT3 expression in normal human colorectal tissue samples, such a relationship was less pronounced in CRC samples (Supplementary Fig S8D and data not shown). As a gating regulator of CRC metastasis, GF11 downregulation may elicit further feedback and/or network regulations on its downstream effectors, such as STAT3, besides other layers of disease complexity (Supplementary Figure S6I). Thus far, the dynamics of cellular adaptation resulting from genetic/epigenetic changes or phenotypic plasticity remain poorly understood in patients with advanced cancer.

In summary, these findings elucidate a new mechanism contributing to the acquisition of metastatic potential during CRC progression. Therapeutic interventions that restore GF11 expression may be suitable for the treatment of metastatic CRC, a condition that conveys high rates of mortality.

Materials and Methods
Cell culture, transfection, and treatment. THP1, U-937, HT29, LoVo, SW480, and SW620 cells from ATCC and MC38 cells granted from Dr. Shoshana Yakar were cultured at 37 °C and 5% CO2 in Dulbecco’s modified Eagle’s medium or RPMI 1640 medium (HyClone, Foster City, CA, USA) supplemented with 10% fetal bovine serum (FBS). Human cells were transfected using lipofectamine (Invitrogen, Grand Island, NY, USA). LMSCM were prepared from the supernatants of human THP1 cells mock stimulated or stimulated with Salmonella enteridis-derived LPS (Sigma-Aldrich, St Louis, MO, USA) for 6 h. THP1 cells subsequently were centrifuged, and the supernatant was filtrated to eliminate any remaining cells. PBMC were isolated from human blood using lymphocyte separation solution, and then were induced into macrophage via IL4 and IL10 stimulation. U-937 cells also were pretreated as PBMC. LSMC-M were prepared like LMSCM using macrophages derived from PBMC or U-937 cells. CRC cells then were cultivated in the presence of regular medium or half LMSCM or LSMC-M and half regular medium. Where indicated, cells were treated with anti-TGFβ-neutralising antibody (Raybiotech, Norcross, GA, USA), 10 μM of U0126, 20 μM of Ly294002 or 10 μM of PDTC (Sigma-Aldrich) for the indicated times.
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GFI1 mRNA expression. In normal colon and different tumour stages from TCGA, GSE17538, GSE41258, and GSE14333 data sets. (c) Number of GFI1 expression positive cells in human early-stage (I and II) CRC and late-stage (III and IV) CRC (four sections evaluated per sample). (d) GFI1 expression levels in normal colon and different tumour stages from TCGA, GSE17538, GSE41258, and GSE14333 data sets. (e) Relapse-free survival of CRC patients with relatively high and low GFI1 mRNA expression. (f) Number of GFI1-expressing cells in human normal, and pre-cancer tissues including adenoma, and polyp tissues (four sections evaluated per sample).

Construction of MC38-Gfi1 cells with CRISPR activation plasmids. To generate MC38-Gfi1 and -con cells, we used CRISPR Gfi1-activation plasmid (sc-437272) and control CRISPR/Cas9 plasmid (sc-418922, Santa Cruz Ltd., Dallas, CA, USA). Gfi1 CRISPR Activation Plasmid is a synergistic activation mediator transcription activation system designed to specifically upregulate gene expression according to manufacturer’s instructions (Santa Cruz Ltd.). In brief, we infected MC38 cells with three plasmids at a 1:1:1 mass ratio: a plasmid encoding the deactivated Cas9 (dCas9) nuclease (D10A and N683A) fused to the transactivation domain VP64, and a blastidcin-resistant gene; a plasmid encoding the MS2-p65-HSF1 fusion protein, and a hygromycin-resistant gene; a plasmid encoding a target-specific 20 nt guide RNA (gRNA), and a puromycin-resistant gene. The positive cells were selected and cultured. Similar procedures were applied to generate MC38-con cells using control CRISPR activation plasmid.

Real-time PCR analysis. Total RNA was extracted using TRIzol (Invitrogen). RNA was reverse transcribed using a Reverse Transcription kit (Applied Biosystems, Foster city, CA, USA). Real-time PCR was performed using SYBR green (Roche, Mannheim, Germany) on a Step One Real-time PCR System (Applied Biosystems). Expression data were normalised to GAPDH mRNA expression. Data were calculated as 2 ΔΔ Ct. Primer sequences are listed in Supplementary information.

ELISA and human antibody cytokine arrays. Culture medium was collected and subjected to ELISA analysis. Human TGF-β1 and TGF-β2 concentration were measured by ELISA (Raybiotech) according to the manufacturer’s protocols. Measurement and analysis of Antibody Cytokine Arrays for 40 kinds of cytokines associated with macrophage were provided by Raybiotech.

Luciferase reporter assays. The pGL3-STAT3 or pGL3-STAT3-mut (BOSHI Ltd., Harbin, China) was transfected into HT29, HT29-NC, and HT29-GFI1 cells for 40 kinds of cytokines associated with macrophage were provided by Raybiotech.

Migration and invasion assays. About 4000 cells were seeded into the upper chambers in 100 μl of the culture medium containing 1% FBS. Transmigration of cells into the lower compartment containing 600 μl medium (9% FBS) occurred at 37°C. After 24 h, transwells were removed and cleaned with cotton swabs to remove non-migrated cells. Migrated cells adherent to the bottom of the membrane were stained with crystal violet (Sigma-Aldrich). Each determination was done at least in duplicates. Data are expressed as the number of migrating cells per field. For invasion assays, the upper chambers were coated with matrigel before using and the procedures were as above.

For monolayer wound-healing assays, a total of 3×105 cells were collected and plated in a six-well plate. At 100% confluence, two parallel wounds of 1 mm were made using a pipette tip. The number of cells crawling into the wound was measured after 24 h, in three independent experiments.

Statistical analysis. Statistical analyses were performed using the GraphPad Prism Software. D’Agostino & Pearson normality test was performed prior to statistical analysis. For comparisons, Student’s t-test (two-tailed) was used when normality test was achieved. Non-parametric test Mann-Whitney U-test was performed when normality test failed. Log-rank (Mantel-Cox) test was used for survival analysis. Pearson’s correlation test was used to evaluate the expression relationships. A P<0.05 was considered significant.

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Conflict of Interest
The authors declare no conflict of interest.
imunohistochemistry. EL and JW helped with the CRC liver metastasis models and reviewed the manuscript. ZH and JH performed the cell culture and CRC cellular metastatic behaviour experiments. HK and HW performed the immunohistochemistry analysis and animal models. MK, LX, WX, ZF assisted with data analysis and manuscript preparation. RH led the study, supervised the overall project and reviewed the manuscript. All authors reviewed and approved of the submitted manuscript.

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