

Original research

Novel TCF21^{high} pericyte subpopulation promotes colorectal cancer metastasis by remodelling perivascular matrix

Additional supplemental material is published online only. To view, please visit the journal online (http://dx.doi.org/10.1136/gutjnl-2022-327913).

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Received 22 May 2022 Accepted 20 August 2022 Published Online First 7 September 2022



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To cite: Li X, Pan J, Liu T, et al. Gut 2023;**72**:710–721.

ABSTRACT

Objective Haematogenous dissemination is a prevalent route of colorectal cancer (CRC) metastasis. However, as the gatekeeper of vessels, the role of tumour pericytes (TPCs) in haematogenous metastasis remains largely unknown. Here, we aimed to investigate the heterogeneity of TPCs and their effects on CRC metastasis.

Design TPCs were isolated from patients with CRC with or without liver metastases and analysed by single-cell RNA sequencing (scRNA-seq). Clinical CRC specimens were collected to analyse the association between the molecular profiling of TPCs and CRC metastasis. RNAsequencing, chromatin immunoprecipitation-sequencing and bisulfite-sequencing were performed to investigate the TCF21-regulated genes and mechanisms underlying integrin α5 on TCF21 DNA hypermethylation. Pericyteconditional *Tcf21*-knockout mice were constructed to investigate the effects of TCF21 in TPCs on CRC metastasis. Masson staining, atomic force microscopy, second-harmonic generation and two-photon fluorescence microscopy were employed to observe perivascular extracellular matrix (ECM) remodelling. **Results** Thirteen TPC subpopulations were identified by scRNA-seg. A novel subset of TCF21^{high} TPCs, termed 'matrix-pericytes', was associated with liver metastasis in patients with CRC. TCF21 in TPCs increased perivascular ECM stiffness, collagen rearrangement and basement membrane degradation, establishing a perivascular metastatic microenvironment to instigate colorectal cancer liver metastasis (CRCLM). Tcf21 depletion in TPCs mitigated perivascular ECM remodelling and CRCLM, whereas the coinjection of TCF21^{high} TPCs and CRC cells markedly promoted CRCLM. Mechanistically, loss of integrin $\alpha 5$ inhibited the FAK/PI3K/AKT/DNMT1 axis to impair TCF21 DNA hypermethylation in TCF21high TPCs.

Conclusion This study uncovers a previously unidentified role of TPCs in haematogenous metastasis and provides a potential diagnostic marker and therapeutic target for CRC metastasis.

INTRODUCTION

Colorectal cancer (CRC) is the third most common malignant tumour worldwide, and colorectal cancer

SIGNIFICANCE OF THIS STUDY

WHAT IS ALREADY KNOWN ON THIS SUBJECT?

- ⇒ Colorectal cancer (CRC) metastasises mainly to the liver, which results from haematogenous dissemination.
- ⇒ Tumour pericytes (TPCs) are the major components of tumour vessels; their heterogeneity and functions in CRC metastasis remain largely unknown.

WHAT ARE THE NEW FINDINGS?

- ⇒ The heterogeneity of TPCs derived from patients with CRC was dissected by single-cell RNA sequencing.
- ⇒ This study elucidated an undefined role of TPCs in promoting CRC metastasis.
- ⇒ We identified a novel TPC-related prometastatic signalling pathway for CRC metastasis.

HOW MIGHT IT IMPACT ON CLINICAL PRACTICE IN THE FORESEEABLE FUTURE?

- ⇒ This study provides new insights into the heterogeneity of TPCs and their prometastatic effects on CRC.
- ⇒ Therapeutic targeting of the prometastatic TCF21^{high} TPCs may represent a new paradigm for cancer therapy.

liver metastasis (CRCLM) counts as the leading cause of disease mortality. CRCLM results from haematogenous dissemination,² which refers to tumour cell intravasation into circulation and extravasation from blood vessels, followed by the formation of metastatic foci in liver.³ Tumour intravasation is a critical and rate-limiting step of haematogenous metastasis, during which tumour cells invade the perivascular extracellular matrix (ECM), breach the endothelial barrier and enter blood circulation.^{4 5} During intravasation, the invading tumour cells contact endothelial and immune cells to form a tumour microenvironment of metastasis (TMEM).⁶ Pericytes are contractile cells embedded within the capillary walls and wrapped around endothelial cells, where they act as the supervisors of endothelial cells to regulate vessel stabilisation



and vascular permeability. However, the role of tumour pericytes (TPCs) in haematogenous metastasis remains controversial.

TPCs attached to endothelial cells act as physiological barriers to inhibit tumour cell intravasation. Genetic depletion of NG2+ or PDGFRβ⁺ pericytes or pharmaceutical inhibition of pericyte recruitment has been shown to enhance vascular permeability and intratumoral hypoxia, which promotes tumour cell epithelial-tomesenchymal transition (EMT) and tumour metastasis.8-10 Tumorderived PDGF-BB induces pericyte-fibroblast transition (PFT), leading to cell detachment from blood vessels or secretion of interleukin-33 to recruit tumor-associated macrophages, enhancing haematogenous metastasis. 11 12 Tumor-derived exosomes activate KLF4 in pericytes, resulting in cell detachment and abundant deposition of fibronectin in secondary organs, thereby establishing a premetastatic niche to facilitate haematogenous metastasis. However, TPCs also play a prometastatic role. CD45 VLA-1 bri or endosialin-expressing pericytes in primary tumours facilitate haematogenous metastasis by promoting tumour cell intravasation in a cell contact-dependent manner without altering the tumour vasculature structure or permeability. 14 15 These contradictory effects of TPCs on haematogenous metastasis may be associated with their heterogeneity, ¹⁶ which has not been determined.

TCF21, a member of the basic helix-loop-helix family of transcription factors, is critical for embryogenesis. Deficiency of TCF21 leads to abnormalities in multiple organs and neonatal lethality. High expression of TCF21 in coronary artery smooth muscle cells inhibits their differentiation and promotes their migration and proliferation, contributing to stable atherosclerotic plaques and reducing the risk of coronary artery disease. TCF21 also drives the expression of inflammatory genes and deposition of collagen IV in visceral adipose stem cells. TCF21 acts as a tumour suppressor in various types of tumours, which is low expressed in tumour cells owing to its promoter hypermethylation. However, the expression profile and function of TCF21 in TPCs remain unknown.

Here, we employed single-cell RNA sequencing (scRNA-seq) to dissect the heterogeneity of TPCs derived from patients with CRC with or without liver metastases, and 13 distinct TPC subpopulations were identified. Among them, a novel subset of TCF21^{high} TPCs, termed matrix–pericytes, was correlated with CRCLM. TCF21 in TPCs increased perivascular collagen deposition and rearrangement and basement membrane degradation, facilitating the establishment of the perivascular metastatic microenvironment (PMM) to enhance tumour cell intravasation. Pericyte-specific knockout of *Tcf21* inhibited CRCLM by reducing perivascular ECM remodelling, maintaining the integrity of the basement membrane, and decreasing circulating tumour cells (CTCs). Moreover, the expression of TCF21 in matrix–pericytes was regulated by the loss of integrin α*S*, which conferred to the DNA hypermethylation of *TCF21* via the FAK/PI3K/AKT-DNMT1 axis.

MATERIALS AND METHODS

The detailed materials and methods can be found in the online supplemental materials.

RESULTS

Identification of a distinct subpopulation of TPCs associated with CRC metastasis

To dissect the heterogeneity of TPCs and assess their contribution to CRC metastasis, TPCs were isolated from primary tumour tissues of patients with CRC with or without liver metastases and were analysed by scRNA-seq using the 10× chromium platform. Single-cell transcriptomes were generated from 50000 cells, which were juxtaposed in t-stochastic neighbour embedding (t-SNE) space to

identify distinct clusters (figure 1A). The gene expression profiles of TPCs were classified into 13 distinct clusters (figure 1B), and their origins were determined using known pericyte markers (online supplemental figures 1A, B). The top 10 cluster-specific genes are shown in figure 1C. The proportion of cells in cluster 2 showed the greatest difference among all subsets between patients with CRC with (18.5%) and without (3.4%) liver metastases (figure 1D), indicating that this subset was associated with CRCLM. Gene Ontology (GO) enrichment analysis indicated that the upregulated genes in samples from patients with CRC with liver metastases were associated with several GO terms related to the ECM (figure 1E), including MATN2, CHI3L1, COL3A1, COL1A2, CILP, MMP2, MFAP4, FBLN1 and FBLN2 (figure 1F and online supplemental figure 2). Therefore, cells in cluster 2 were defined as matrix-pericytes. One of the genes exclusively enriched in cluster 2, MATN2, which encodes the protein matrilin-2 (MATN2), was selected as a biomarker for matrix-pericytes. The ratio of MATN2⁺ TPCs was higher in tumour sections derived from patients with CRC with liver metastases than in those without liver metastases (figure 1G and online supplemental table 1). In addition, receiver operating characteristic (ROC) curve analysis showed that the optimal cut-off percentage of MATN2⁺ TPCs to predict CRCLM with high sensitivity and specificity was 30% (figure 1H). Kaplan-Meier survival analysis showed that overall survival (OS) (figure 1I) and diseasefree survival (DFS) (figure 1]) were shorter in patients with a high (>30%) ratio of MATN2+ TPCs. Furthermore, the correlation analysis between the MATN2+ TPC ratio and clinicopathological parameters showed that the MATN2+ TPC ratio had a positive correlation with the TNM stage and liver metastasis of patients with CRC (online supplemental table 2). Taken together, these data indicate that matrix-pericytes are notably associated with CRC metastasis.

TCF21 in TPCs is associated with CRC metastasis

Key regulators of matrix-pericytes were identified using the singlecell regulatory network inference and clustering (SCENIC) pipeline, which connects cis-regulatory sequence information with scRNA-seq data.²¹ SCENIC analysis showed that the regulon activity of TCF21 was the highest in matrix-pericytes (figure 2A), which was confirmed in the t-SNE space of all cells (figure 2B). In addition, the number of TCF21high TPCs was markedly increased in samples from patients with CRC with liver metastases compared with those without liver metastases (figure 2C). In addition, the ratio of TCF21⁺ TPCs was increased in tumour sections from patients with CRC with liver metastases (figure 2D and online supplemental table 1). However, TCF21 was undetectable in TPCs in hepatic metastatic tumours derived from patients with CRC (online supplemental figure 3). Pearson's correlation coefficient analysis showed a positive correlation between the ratio of TCF21⁺ TPCs and MATN2⁺ matrix-pericytes (r=0.805, P<0.001; figure 2E). Additionally, flow cytometry analysis showed that the proportion of TCF21⁺MATN2⁺ TPCs was significantly increased in TPCs derived from patients with CRC without metastases (TPC $_{NM}$) infected with lentivirus harbouring TCF21-overexpressing plasmid (TPC_{NM} TCF21) compared with those harbouring vector (TPC_{NM}) (online supplemental figure 4A). In contrast, knockdown of TCF21 in TPCs derived from patients with CRC with liver metastases (TPC_{1M}) showed opposite effects (online supplemental figure 4B). These data indicate that TCF21⁺ TPCs are closely associated with matrix-pericytes. ROC curve analysis revealed that the optimal percentage of TCF21⁺ TPCs for predicting liver metastases in patients with CRC was 44% (figure 2F). Kaplan-Meier survival analysis suggested that OS (figure 2G) and DFS (figure 2H) were significantly shorter in

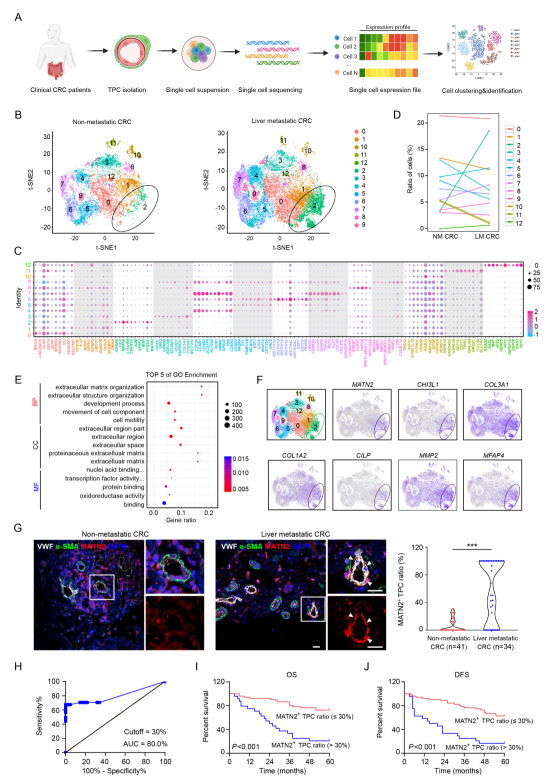


Figure 1 Transcriptomic characterisation of matrix-pericytes in TPCs derived from CRC patients. (A) Schematic diagram of scRNA-seq for TPCs. (B) t-SNE visualisation of TPC subsets derived from primary tumour tissues of CRC patients with (n=2) or without (n=2) liver metastasis. (C) Dot plots of the top ten marker genes in each TPC subset. (D) Percentage of each subset of TPCs. (E) GO analysis of the upregulated genes in cluster 2. (F) t-SNE visualisation of TPCs at all stages is combined, overlaid with the expression of indicated genes. (G) Immunofluorescence analysis and quantification of MATN2⁺ matrix-pericyte ratio in primary tumour tissues from CRC patients with (n=34) or without (n=41) liver metastasis. white arrows indicate the staining of MATN2 in TPCs. scale bar, 20 µm. each sample on the violin plots represents individual patient data. ***P < 0.001 by two-tailed Mann-Whitney test. (H) ROC curve analysis of MATN2⁺ TPC ratio in CRC patients (n=75). (I, J) Kaplan-Meier analysis of the OS (I) or DFS (J) in CRC patients with high or low MATN2⁺ matrix-pericyte ratio (based on 30% cut-off, n=75). P<0.001 by Log-rank (Mantel-Cox) test. BP, biological process; CC, cellular component; CRC, colorectal cancer; DAPI, 4',6-diamidino-2-phenylindole; DFS, disease-free survival; GO, Gene Ontology; LM CRC, liver metastatic colorectal cancer; MF, molecular function; NM CRC, non-metastatic colorectal cancer; OS, overall survival; ROC, receiver operating characteristic; TPC, tumour pericyte; t-SNE, t-stochastic neighbour embedding; VWF, von willebrand factor.

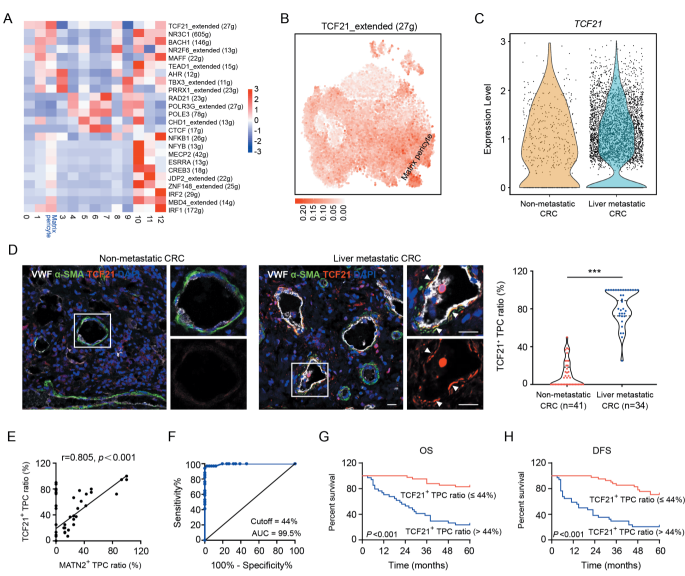


Figure 2 TCF21 in matrix—pericytes is positively associated with liver metastasis of CRC. (A) Heat map analysis of the regulon activities in distinct subsets of TPCs. (B) t-SNE visualisation of TCF21 regulon modulus in all TPC subsets. (C) Expression of *TCF21* in matrix—pericytes derived from CRC patients with or without liver metastasis. (D) Immunofluorescence staining and quantification of TCF21⁺ TPC ratio in tumour sections from CRC patients with (n=34) or without (n=41) liver metastases. White arrows indicate TCF21 staining in TPCs. Scale bar, 20 μm. Each sample on the violin plots represents individual patient data. ***P<0.001 by two-tailed Mann-Whitney test. (E) Pearson's correlation analysis of TCF21⁺ TPC ratio and MATN2⁺ matrix—pericyte ratio (n=75). (F) ROC curve analysis of TCF21⁺ TPC ratio in CRC patients (n=75). (G, H) Kaplan-Meier analysis of OS (G) or DFS (H) in CRC patients with high or low TCF21⁺ TPC ratio (based on the 44% cut-off, n=75). P<0.001 by log-rank (Mantel-Cox) test. CRC, colorectal cancer; DAPI, 4',6-diamidino-2-phenylindole; DFS, disease-free survival; OS, overall survival; ROC, receiver operating characteristic; TPC, tumour pericyte; t-SNE, t-stochastic neighbour embedding; VWF, von willebrand factor.

patients with higher (>44%) ratio of TCF21⁺ TPCs. Furthermore, the TCF21⁺ TPC ratio was notably correlated with TNM stage and liver metastasis in patients with CRC (online supplemental table 3). These findings indicate that TCF21 in TPCs can serve as a predictive biomarker for CRC metastasis.

TCF21 in TPCs contributes to the phenotype transition of matrix-pericytes and the remodelling of perivascular ECM

The positive correlation between TFC21 upregulation in TPCs and CRCLM suggested that TCF21 may be involved in the activation of TPCs and may promote the phenotype transition of matrix-pericytes. Depletion or overexpression of TCF21 in TPCs had negligible effects on cell proliferation, adhesion and migration (online supplemental figure 5). The

levels of nine genes in matrix–pericytes, including *IGFBP5*, *CILP*, *MFAP4*, *c11orf96*, *A2M*, *SFRP2*, *MAF*, *PTGDS* and *MATN2*, were significantly higher in $\text{TPC}_{\text{NM}}^{\text{TCF21}}$ than in $\text{TPC}_{\text{NM}}^{\text{Vector}}$ (figure 3A). Conversely, the levels of these genes were significantly lower in TPC_{LM} transfected with siTCF21 ($\text{TPC}_{\text{LM}}^{\text{siTCF21}}$) than in those transfected with negative control siRNA ($\text{TPC}_{\text{LM}}^{\text{siNC}}$) (online supplemental figure 6A). In addition, the percentage of MATN2⁺ matrix–pericytes was increased by overexpression of TCF21 in TPC_{NM} and decreased by the knockdown of TCF21 in TPC_{LM} (figure 3B and online supplemental figure 6B).

Genes acting downstream of TCF21 were investigated in TPCs using whole-transcriptome RNA-seq and ChIP-seq assays (online supplemental figure 6C, D). RNA-seq assays showed that 1276

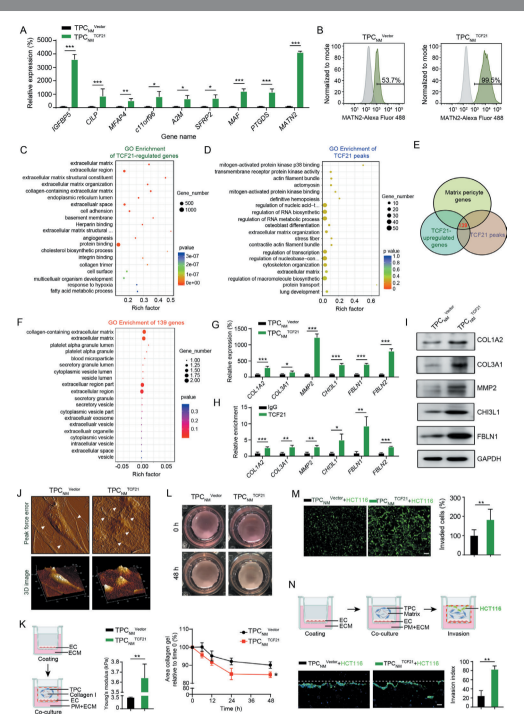


Figure 3 TCF21 is essential for the formation of matrix—pericytes and facilitates CRC cell invasion by ECM remodelling. (A) RT-qPCR analysis for matrix—pericyte-related genes in TPC_{NM} ^{Vector} or TPC_{NM} (n=3). (B) FCM analysis for MATN2 expression in TPC_{NM} (D) GO terms of differentially expressed genes derived from RNA-seq in TPC_{NM} (TCF21) compared with those in TPC_{NM} (D) GO terms of TCF21 peaks derived from ChIP-Seq in TPC_{NM} (E) Venn diagram showing the number of overlapped genes derived from TCF21-upregulated genes, TCF21 peaks and matrix—pericyte genes. (F) GO analysis for the intersected genes derived from E. (G) RT-qPCR analysis for the indicated genes in TPC_{NM} (n=3). (H) ChIP-qPCR analysis for the binding of TCF21 with the promoter of indicated genes in TPC_{NM} (n=3). (I) Western blotting analysis for the expression of indicated proteins in TPC_{NM} (n=3). (J) AFM imaging of topography and roughness of the collagen fibres incubated with TPC_{NM} (n=3). (N) Schematics of the 3D coculture model (left) and Young's modulus detection (right) are shown. Red dotted line indicates the removed part. (L) Representative images and quantification of collagen contraction caused by TPC_{NM} (N) Schematics of organotypical culture system for studying the invasion of tumour cells (upper). Representative images and quantification of HCT116 cells that invaded into the TPC-containing matrix (bottom) (n=3). Scale bar, 20 μm. Red dotted line indicates the removed part; white lines indicate the initiation. Data are presented as mean±SEM. *P<0.05, **P<0.01, ***P<0.001 by two-tailed unpaired t-test. AFM, atomic force microscopy; ChIP-qPCR, chromatin immunoprecipitation-quantitative PCR; CRC, colorectal cancer; EC, endothelial cell; ECM, endothelial culture medium; GO, Gene Ontology; RT-qPCR, real-time quantitative PCR; TPC, tumour pericyte.

genes were upregulated, and 1397 genes were downregulated in $\text{TPC}_{\text{NM}}^{\text{TCF21}}$ compared with $\text{TPC}_{\text{NM}}^{\text{Vector}}$ (online supplemental figure 6E). A heatmap of the RNA-seq results showed that the expression of matrix-pericyte-specific genes, including MFAP4, CILP, MATN2, MMP2, COL3A1, COL1A2, CHI3L1 and FBLN1, was significantly higher in TPC_{NM}^{TCF21} (online supplemental figure 6F). GO enrichment analysis of the RNA-seq and ChIP-seq results showed that genes regulated by TCF21 were associated with the categories 'ECM organisation' and 'ECM' (figure 3C,D), which were similar to the expression profiles in matrix-pericytes. Moreover, comparison of the TCF21 peaks detected by ChIP-seq analysis, TCF21-induced genes detected by RNA-seq and matrix-pericyte genes detected by scRNA-seq identified 139 overlapping genes (figure 3E). GO analysis of these overlapping genes showed their associations with the categories 'collagen-containing ECM' and ECM (figure 3F). These results were confirmed by RT-qPCR and ChIP-qPCR, and the binding of TCF21 to the promoter of matrix-pericyte-specific genes was significantly upregulated in TCF21-overexpressing TPC_{NM} (figure 3G,H, and online supplemental figure 6G and H), which were confirmed by western blotting (figure 3I and online supplemental figure 6I). These results indicate that TCF21 is critical for the generation of matrix-pericytes.

ECM remodelling is crucial for tumour metastasis, ²² which is represented by aberrant collagen production and cross-linking leading to tissue stiffness, thus promoting tumour metastasis.²³ ²⁴ Moreover, degradation of the vascular basement membrane by proteases such as matrix metalloproteinases facilitates the escape of tumour cells from primary tumour sites.²⁵ We proposed that TCF21 may play a role in perivascular ECM remodelling and CRC metastasis. Neither overexpression nor knockdown of TCF21 in TPCs altered tumour cell proliferation, migration, EMT or endothelial cell tube formation in vitro (online supplemental figure 7A-H). Atomic force microscopy (AFM) revealed that TPC_{NM}^{TCF21} possessed greater ability than TPC_{NM}^{Vector} in inducing local alignment of collagen, reorganisation of curly and sheet-like collagen fibres into radial and bundling structures, and increased roughness (figure 3J). Moreover, TPC_{NM} TCF21 was superior to TPC_{NM} Vector in stiffening the mechanical properties of collagen (figure 3K) and increasing collagen contraction (figure 3L), thereby enhancing ECM stiffness. To further assess the effects of TCF21-mediated perivascular ECM remodelling on the transmigration of CRC cells, PKH-67-labelled HCT116 or DLD-1 cells mixed with TPC_{NM} Vector, TPC_{NM} TCF21, TPC_{SINC} or TPC_{LM} were seeded into Matrigel. TPC_{NM} TCF21 was better than TPC_{NM} Vector in facilitating CRC cell invasion through the Matrigal costed transportal manufacture (forms 2) for the last control to the matricel costed transportal manufacture (forms 2) for the last control transportation of the Matrigal costed transpor the Matrigel-coated transwell membranes (figure 3M and online supplemental figure 8A), whereas $\mathrm{TPC}_{\mathrm{LM}}^{\mathrm{siTCF21}}$ reduced the invasion of CRC cells compared with $\mathrm{TPC}_{\mathrm{LM}}$ (online supplemental figure 8B, C). The experiments performed on an organotypical culture system showed that the number of invaded HCT116 cells was significantly increased when matrix, consisting of collagen I and Matrigel, was premixed with TPC_{NM} TCF21 compared with TPC_{NM} (figure 3N). Collectively, these results indicate that TCF21 in matrix-pericytes induces perivascular ECM remodelling and facilitates CRC cell invasion through the perivascular

We further investigated whether MATN2 exerted prometastatic effects similar to those of TCF21. Given that TCF21^{high} TPCs facilitated tumour metastasis through ECM remodelling, the levels of genes encoding ECM proteins, such as *COL1A2*, *COL3A1*, *MMP2*, *CHI3L1* and *FBLN1*, were examined by RT-qPCR in TPCs with overexpression or knockdown of MATN2 (online supplemental figure 9A, B). Our results showed that MATN2 in TPCs had negligible effects on the levels of these ECM-related genes (online supplemental figure 9C, D). Additionally, MATN2 in TPCs had negligible effects on tumour cell invasion (online supplemental figure 9E, F). These data indicate that MATN2⁺ TPCs could not give metastatic phenotypes to tumour cells similar to TCF21^{high} TPCs, which may merely serve as a characteristic marker for matrix–pericytes.

Knockout of Tcf21 in TPCs inhibits CRC metastasis

To determine whether TCF21 in TPCs is essential for CRC metastasis in vivo, pericyte lineage-tracing mice (PClin) and tamoxifen-inducible Cspg4-driven pericyte-specific Tcf21 knockout mice (PClin-KO) were generated (figure 4A and online supplemental figure 10A, B). Tamoxifen administration to both PC^{lin} and PC^{lin-KO} mice resulted in the permanent labelling of pericytes with tdT fluorescence, and the knockout of Tcf21 in PClin-KO mice. To examine the effects of TCF21 in TPCs on CRC metastasis, PClin and PClin-KO mice were orthotopically injected with luciferase-labelled tumour cells (MC38-luc-LM3), followed by treatment with tamoxifen for 1 week (online supplemental figure 11A). Tamoxifen treatment had negligible effects on the non-tumour tissues (online supplemental figure 11B) but induced Cre activity in TPCs, as indicated by tdT fluorescence in both PClin and PClin-KO mice and loss of TCF21 in PClin-KO mice (online supplemental figure 11C, D). In vivo, the TPC-specific deletion of Tcf21 significantly inhibited the formation of liver metastases (figure 4B). The area and number of liver metastatic foci were significantly decreased in PClin-KO mice compared with those in PClin mice (figure 4C). Furthermore, the number of EpCAM+CD45- CTCs was significantly lower in PClin-KO mice than that in PC lin mice (figure 4D and online supplemental figure 12A, B). Knockout of Tcf21 in TPCs had negligible effects on primary tumour growth and EMT, as indicated by equal levels of E-cadherin, vimentin, and Ki67 in tumour sections from PClin and PClin-KO mice (online supplemental figure 13A, B). These data indicate that TCF21 in TPCs promotes CRC metastasis, independent on tumour growth and EMT.

The effects of TCF21 in TPCs on vascular structure and function were further evaluated by assessing pericyte coverage and the percentage of MATN2⁺ matrix-pericytes in tumour vessels. Immunofluorescence staining showed that deletion of Tcf21 in TPCs had negligible effects on total pericyte coverage (online supplemental figure 13C), whereas the number of MATN2⁺ matrix-pericytes in primary tumour tissues of MC38-luc-LM3 CRCLM xenografts was lower in PC^{lin-KO} mice than in PC^{lin} mice (figure 4E and online supplemental figure 12C). In addition, the expression of ECM remodelling-related proteins, including MMP2, COL1A2, COL3A1 and CHI3L1, in TPCs from PClin mice was significantly higher than those from PClin-KO mice (figure 4F and online supplemental figure 12D). Masson staining showed that the density of fibrillar collagen components was significantly decreased in tumour sections from PClin-KO mice compared with that in PClin mice (figure 4G and online supplemental figure 12E). Transmission electron microscopy analysis indicated that the collagen bundles at the perivascular region were thinner in tumour sections from PClin-KO than in those from PClin mice (figure 4H). Collagen deposition is required for tissue stiffness and the local invasiveness of tumour cells, and changes in collagen orientation also contribute to tumour metastasis. ²⁶ The structure of perivascular collagen was further examined by second-harmonic generation and two-photon excited fluorescence. Perivascular collagen fibres in PC in-KO mice were arranged irregularly around the tumour vessels with curly and

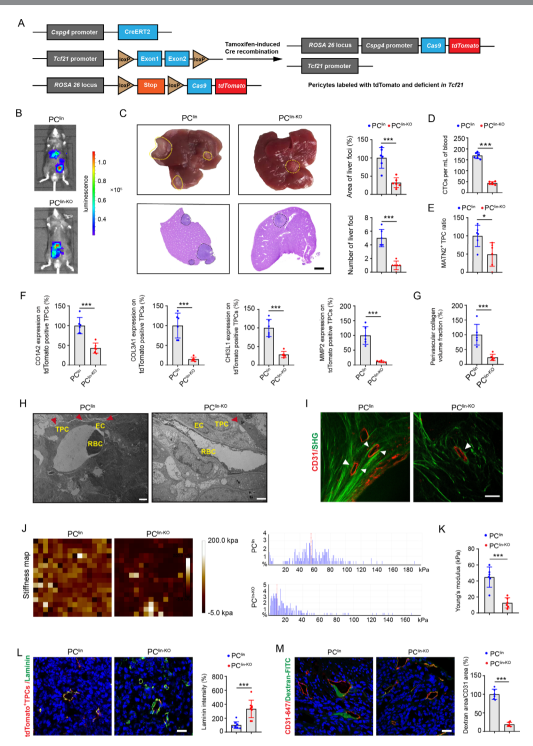


Figure 4 Pericyte-specific knockout of *Tcf21* inhibits the remodelling of perivascular ECM and CRC metastasis. (A) schematic diagram of the construction of *Cspg4* (NG2) lineage-tracing *Tcf21* inducible knockout mice. (B) Representative images and quantification of bioluminescence signals of mice orthotopically injected with MC38-luc-LM3 cells (n=6). (C) Representative images and H&E analysis of livers derived from MC38-luc-LM3 xenograft-bearing mice (n=6). Yellow and black dotted lines indicate the liver metastatic loci. Scale bar, 2 mm. (D) Quantification of CTCs (n=6). (E) Quantification of MATN2⁺ TPC ratio in primary tumour sections (n=6). (F) Quantification of the expression of COL3A1, MMP2, COL1A1 and CHI3L1 on TPCs (tdTomato) in primary tumour sections (n=6). (G) Quantification of perivascular collagen in primary tumour sections (n=6). (H) Representative images of collagen fibres surrounding TPCs (n=6). Red arrowheads indicate the perivascular collagen fibres. Scale bar, 2 μm. (I) Representative images of perivascular collagen organisation in tumours imaged by SHG (green) and CD31 (red) (n=6). White arrowheads indicate the perivascular collagen fibres. Scale bar, 50 μm. (I,K) Representative AFM images and quantification of the stiffness of the perivascular area in primary tumour sections (n=6). (L) Immunofluorescence staining and quantification of laminin (green) surrounding TPCs (tdTomato) in primary tumour sections (n=6). Scale bar, 20 μm. Data are presented as mean±SEM. *P< 0.05, *** P< 0.001 by two-tailed unpaired t-test. AFM, atomic force microscopy; CRC, colorectal cancer; CTC, circulating tumour cell; EC, endothelial cell; ECM, extracellular matrix; RBC, red blood cell; SHG, second-harmonic generation; TEM, transmission electron microscopy; TPC, tumour pericyte.

unfixed outlines, whereas perivascular collagen in PClin mice had an almost parallel orientation and were tightly embraced surrounded the tumour vessels (figure 4I). The stiffness of perivascular areas was determined by AFM and analysed with Young's modulus patterns, and we found that the perivascular areas in PClin-KO mice were significantly softer than those in PClin mice (figure 4[,K). As the basement membrane serves as a physical barrier for tumour cell intravasation, degradation of the basement membrane could weaken its barrier function and increase vessel permeability.²⁷ Our results showed that the expression of laminin, the most abundant component in the vascular basement membrane, was significantly higher in PClin-KO mice than that in PClin mice (figure 4L), and deletion of Tcf21 in TPCs resulted in decreased vessel permeability in PClin-KO mice compared with PClin mice (figure 4M). Taken together, TCF21 in matrix-pericytes might be a key regulator of perivascular ECM remodelling, which establishes a PMM to facilitate CRC cell intravasation.

Loss of integrin α5 attenuates DNA methylation of *TCF21* promoter and increases TCF21 expression in TPCs

Integrins, the key receptors of ECM components, function as mechanical transducers to facilitate tumour metastasis. Our results showed that ITGA2, ITGA5 and ITGB1 were significantly decreased in matrix—pericytes among the 13 clusters of TPCs (figure 5A). Integrin $\alpha 2$ and integrin $\beta 1$ were unrelated to TCF21, whereas integrin $\alpha 5$ expression was inversely correlated with TCF21 expression in TPCs (online supplemental figure 14A, B). Gain-of-function and loss-of-function experiments showed that TCF21 was negatively regulated by integrin $\alpha 5$ at both the mRNA and protein levels (figure 5B and online supplemental figure 14C-E), whereas integrin $\alpha 2$ and integrin $\beta 1$ had negligible effects on TCF21 expression (online supplemental figure 14F).

Hypermethylation of *TCF21* DNA suppresses TCF21 expression in various types of tumour cells, including non-small cell lung cancer, head and neck squamous cell carcinoma, and CRC. ^{29–31} Evaluation of the methylation of CpG islands within the promoter of *TCF21* in TPCs showed that TCF21 expression in TPCs was inversely correlated with its methylation status, as indicated by the decreased 5-mC level of *TCF21* promoter of TPC_{LM} compared with TPC_{NM} (online supplemental figure 14G). In addition, integrin α 5 positively regulated *TCF21* DNA hypermethylation, and loss of integrin α 5 in TPCs attenuated DNA hypermethylation of *TCF21* promoter in TPCs (figure 5C and online supplemental figure 14H).

DNMT1 has been reported to regulate the DNA hypermethylation of TCF21 in lung cancers. ³² Our results showed that depletion of integrin $\alpha 5$ reduced the expression of DNMT1 in TPC_{NM} by inhibiting the FAK/PI3K/AKT axis, whereas overexpression of integrin $\alpha 5$ in TPC_{LM} exerted opposite effects, which could be reversed by the FAK inhibitor Y15 (figure 5D and online supplemental figure 14I). Furthermore, either Y15 or DNMT inhibitor SGI1027 decreased the expression of DNMT1 and suppressed the DNA hypermethylation of TCF21 (figure 5E), followed by an increase in the expression of TCF21 in TPC_L. ITGAS (figure 5D). These results indicate that the loss of integrin $\alpha 5$ upregulates TCF21 in TPCs by suppressing the DNA hypermethylation of TCF21.

The effects of integrin $\alpha 5$ in TPCs on CRC metastasis in vivo were investigated in an orthotopic xenograft model generated by the co-injection of HCT116-luc-LM3 or DLD1-luc-LM3 cells and integrin $\alpha 5$ -overexpressing or -knockdown TPCs (figure 5F). Compared with TPC $_{NM}^{shIVGA5}$, TPC $_{NM}^{shIVGA5}$

TCF21^{high} matrix-pericytes correlate with perivascular ECM remodelling and liver metastasis in patients with CRC

To determine whether the above findings were applicable to patients with CRC, perivascular collagen deposition and alignment were assessed in patients with CRC with and without liver metastases. Compared with patients with CRC without liver metastases, the perivascular collagen abundance, and the reorientation of perivascular collagen fibres into a radial alignment were more prominent in patients with CRC with liver metastases (figure 6A,B, and online supplemental figure 16A, B). In addition, the stiffness of perivascular areas was significantly enhanced in tumour tissues from patients with CRC with liver metastases compared with those without liver metastases (figure 6C,D). Moreover, the expression of COL1A2, COL3A1 and CHI3L1 (figure 6E), as well as MMP2, a key protease involved in the degradation of basement membrane (figure 6F), was higher in TPCs from patients with CRC with liver metastases than those without liver metastases. Correspondently, the expression of integrin as was lower in TPCs from patients with CRC with liver metastases than in those without liver metastases (figure 6G). Pearson correlation analysis indicated that the levels of MMP2, COL1A2, COL3A1 and CHI3L1 in TPCs were positively correlated, whereas the level of integrin $\alpha 5$ in TPCs was negatively correlated with the ratio of TCF21^{high} TPCs (figure 6E–G). These clinical data indicate that integrin α5 loss-induced upregulation of TCF21 is an important regulator of perivascular ECM remodelling and the establishment of the PMM, thus facilitating CRC metastasis.

DISCUSSION

Cellular heterogeneity in the tumour microenvironment (TME) may result in distinct pathological phenotypes and different responses to cancer therapy.³³ The heterogeneity of tumour cells,³⁴ tumor-infiltrating immune cells,³⁵ cancer-associated fibroblasts (CAFs),³⁶ and endothelial cells³⁷ has been extensively investigated by scRNA-seq, which has also been employed to analyse pericytes derived from tumours, enteritis, and normal GI tract.^{38–41} However, the specific phenotype and function of pericytes in tumour progression are largely unknown. Therefore, evaluating the heterogeneity of TPCs may provide new insights into the mechanisms underlying haematogenous metastasis. Here, the heterogeneity of TPCs was first revealed by scRNA-seq. By comparing the subpopulations of TPCs in our study with those in previous studies,³⁹ ⁴² four novel subpopulations of TPCs were specifically identified in our work (online supplemental figure 17A–C).

Loss of NG2 is an important hallmark of PFT, during which pericytes shed from the blood vessels where they were originally attached. We found that the expression of NG2 in clusters 10, 11 and 12 was extremely low, suggesting that PFT may have occurred in these subpopulations. However, the expression of NG2 in cluster 2 was not significantly decreased, indicating that

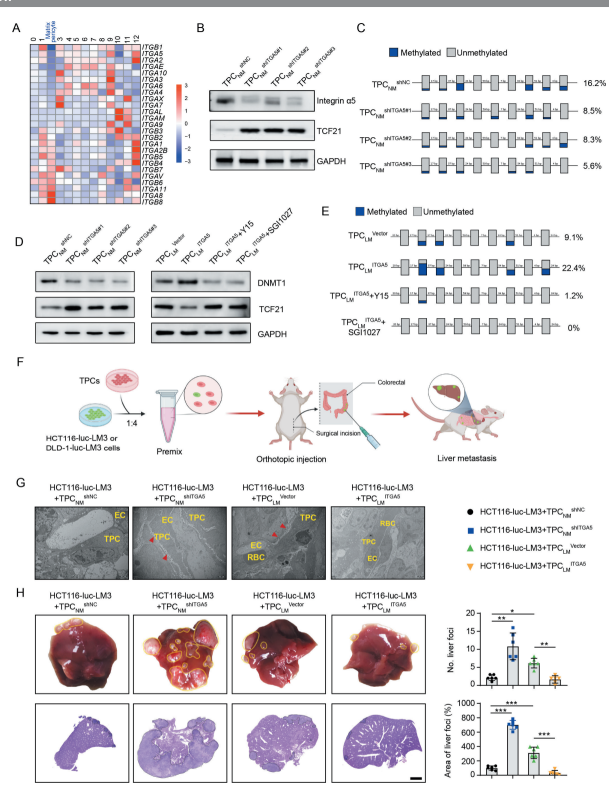


Figure 5 Loss of integrin α5 inhibits hypermethylation of *TCF21* promoter and upregulates the expression of TCF21 in TPCs. (A) Heat map of the expression of integrins in all subsets of TPCs. (B) Western blotting analysis of integrin α5 and TCF21 in integrin α5-knockdown TPC $_{NM}$ (n=3). (C) BSP analysis for *TCF21* promotor region in integrin α5-knockdown TPC $_{NM}$ (n=3). Blue and grey circles represent the methylated and unmethylated CpGs, respectively. The percentage of total methylated CpGs is given on the right. (D) Western blotting analysis of DNMT1 and TCF21 in integrin α5-knockdown (left) or integrin α5-overexpressing (right) TPCs treated with or without FAK inhibitor (Y15) or DNMT1 inhibitor (SGI1027) (n=3). (E) BSP analysis for the methylation and unmethylation levels of *TCF21* in TPC $_{LM}$ treated with or without Y15 and SGI1027 (n=3). (F) Schematic diagram of the in vivo experiments. (G) Representative TEM images of collagen fibres surrounding TPCs in primary HCT116 orthotopic xenografts (n=6). Red arrowheads indicate perivascular collagen fibres. Scale bar, 2 μm. (H) Representative images of the whole liver and H&E analysis of liver metastatic foci (n=6). Yellow and black dotted lines indicate the metastatic loci. Scale bar, 2 mm. Data are presented as mean±SEM. *P<0.05, **P<0.01, ***P<0.001 by one-way analysis of variance followed by Tukey's post hoc test. BSP, bisulfite sequencing PCR; CRC, colorectal cancer; EC, endothelial cell; RBC, red blood cell; TEM, transmission electron microscopy; TPC, tumour pericyte.

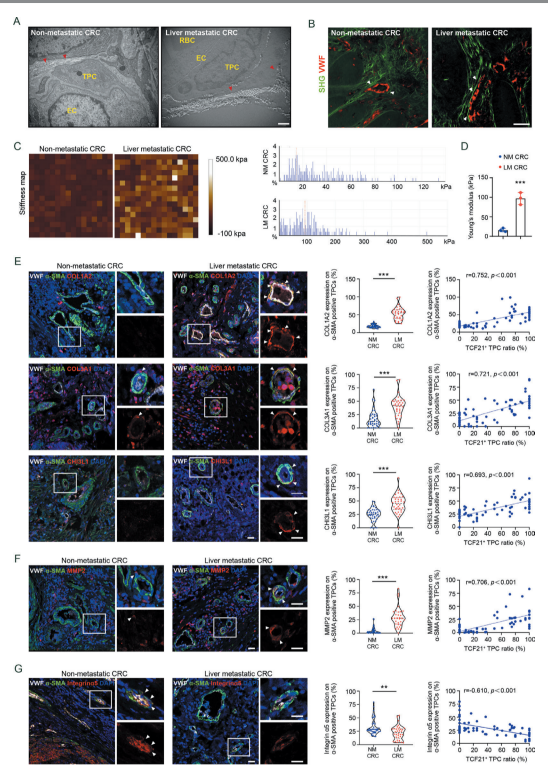


Figure 6 TCF21 in matrix–pericytes is associated with the remodelling of perivascular ECM in primary tumours derived from patients with CRC. (A) Representative TEM images of collagen fibres surrounding TPCs in primary tumour tissues from patients with CRC (n=3). Red arrowheads indicate the perivascular collagen fibres. Scale bar, 1 μm. (B) Representative images of perivascular collagen structure and density in patients with CRC imaged by SHG (green) and VWF (red) (n=3). White arrowheads indicate perivascular collagen fibres. Scale bar, 50 μm. (C,D) Representative AFM images and quantification of stiffness in perivascular area in primary tumour tissues from patients with CRC. Data are presented as mean±SEM (n=3).***P<0.001 by two-tailed unpaired t-test. (E–G) Immunofluorescence staining and quantification of COL1A2, COL3A1, CHI3L1 (E), MMP2 (F) and integrin α5 expression (G) in αSMA⁺ TPCs in primary tumour sections from patients with CRC (n=75), and Pearson's correlation analysis for COL1A2, COL3A1, CHI3L1, MMP2 or integrin α5 expression in αSMA⁺ TPCs with TCF21⁺ TPC ratio (%) are shown (n=75). White arrowheads indicate the staining of indicated proteins in TPCs. Scale bar, 20 μm. Each sample on the violin plots represents individual patient data. **P<0.01, ***P<0.001 by two-tailed Mann-Whitney test. AFM, atomic force microscopy; DAPI, 4',6-diamidino-2-phenylindole; EC, endothelial cell; ECM, extracellular matrix; LM CRC, liver metastatic colorectal cancer; NM CRC, non-metastatic colorectal cancer; RBC, red blood cell; SHG, second-harmonic generation; TEM, transmission electron microscopy; TPC, tumour pericyte; VWF, von willebrand factor.

these TPCs retained perivascular characteristics despite being activated. Furthermore, trajectory analysis indicated that cluster 2 originated from cluster 9 and later evolved into cluster 12 (online supplemental figure 18A, B), further demonstrating that cluster 2 was a newly identified subset of TPCs. Therefore, our findings improve understanding of the heterogeneity of TPCs.

To initiate metastasis, tumour cells must break through the collagen-enriched ECM by matrix deposition and degradation, which generate 'tracks' or 'tunnels' that help tumour cells pass. ⁴³ On reaching blood vessels, these tumour cells interact with proangiogenic TIE2^{high}/VEGF^{high} macrophages and luminal endothelial cells to construct a TMEM, which supports the transendothelial migration of tumour cells into blood circulation. ⁶ During intravasation, tumour cells physically contact with tumour endothelial cells via juxtracrine and paracrine signalling. ⁵ Although TPCs attach to endothelial cells, their roles in tumour cell intravasation and the underlying mechanisms remain unclear. Our study demonstrates that matrix–pericytes are one of the TPC subpopulations promote CRC metastasis by establishment of PMM, providing an unidentified function of TPCs in haematogenous metastasis.

TCF21 in tumour cells is considered to inhibit tumour growth and metastasis, ¹⁷ whereas the expression and function of TPCs in haematogenous metastasis remain largely unknown. In contrast to its suppressive role in tumour cells and CAFs, 44 TCF21 promoted phenotypical transition of TPCs into matrix-pericytes, which facilitated the haematogenous metastasis of CRC. The distinct functions of TCF21 in TPCs in haematogenous metastasis may be caused by the different methylation levels of the TCF21. DNA methylation of TCF21 was lower in matrixpericytes than in tumour cells, which increased the expression of TCF21 in TPCs from patients with CRC with liver metastases. DNA methylation of TCF21 was regulated by the FAK/ AKT/DNMT1 signalling pathway, whereas the expression and activation of FAK in TPCs are negatively correlated with tumour angiogenesis, tumour growth, and metastasis. 45 However, the mechanism by which cancer cell regulates integrin α5 and TCF21 expression in TPCs has not been revealed. Metastatic CRC cells could rely on gene mutations⁴⁶ including TP53, BRAF and KRAS or extracellular vehicles (EVs)⁴⁷ to promote tumour metastasis. We found that the TP53, BRAF and KRAS mutations were not independent predictors of CRCLM (online supplemental table 4), and the above gene mutations were not associated with the TCF21⁺ TPC ratio or the expression of integrin α5 in TPCs (online supplemental figure 19A and B). Interestingly, we found that metastatic CRC cells could reduce the expression of integrin α5 while increase the expression of TCF21 in an EV-dependent manner (online supplemental figure 19C, D). Nevertheless, the mechanisms underlying tumor-derived EVs and other factors involved in the regulation of TPCs require further investigation.

In conclusion, this study used scRNA-seq to reveal the heterogeneity of TPCs in patients with CRC and to identify a novel subpopulation of TCF21^{high} TPCs associated with CRCLM. These findings revealed the effects and mechanisms of TCF21^{high} TPCs in the construction of the PMM to facilitate CRC metastasis by remodelling the perivascular ECM and provided a potential diagnostic marker for haematogenous metastasis.

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Correction notice This article has been corrected since it published Online First. Figure one and the supplementary files have been corrected.

Acknowledgements We are grateful to Guangzhou Genedenovo Biotechnology Co. for assistance with sequencing and/or bioinformatics analysis of single-cell RNA sequencing. We thank Guangzhou LC-Bio Technology Co. for their assistance with RNA-seq and CHIP-seq. We thank GL for assistance with atomic force microscopy and data analysis.

Contributors DZ, WY, MC and YC designed and supervised the experiments, revised the manuscript and were responsible for the drafts of the manuscript, as well as all requested revisions. QQ, MH, LD and PM-KT critically revised the manuscript. MC, XL, JP and TL wrote the manuscript and analysed the data. XL, MC, WY, QM, ZZ and SQ performed experiments. JP, DH, RD and YZ collected human colorectal cancer tissues, reviewed the pathological sections and assessed preclinical and clinical samples. YG and WZ directed the second-harmonic generation and two-photon-excited fluorescence experiments. HL directed the Young modulus measurements.

Funding This work was supported by the National Natural Science Foundation of China (81973340, 81803566, 81973341, 81773758 and U1801287), Local Innovative and Research Teams Project of Guangdong Pearl River Talents Program (2017BT01Y036), Natural Science Foundation of Guangdong Province (2019A1515010144, 2019A1515110543, A1515110543, 2021A1515110242, 2019A1515011934 and 2020A1515010071), Ministry of Science and Technology of China (2018ZX09711001-008-008), National High-Level Personnel of Special Support Program (Zhang Dongmei), National Key R&D Program of China (2017YFC1703800), Key-Area Research and Development Program of Guangdong Province (2020B1111110004, 2021B1111110004), Science and Technology Program of Guangzhou (202002030010), Guangdong Province Key Laboratory of Pharmacodynamic Constituents of TCM and New Drugs Research, College of Pharmacy (2020B1212060076), Science and Technology Projects in Guangzhou (202102070001), Young S&T Talent Training ProgramProgramme of Guangdong Provincial Association for S&T, China (SKXRC202216).

Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, conduct, reporting or dissemination plans of this research.

Patient consent for publication Not applicable.

Ethics approval Not applicable.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available in a public, open access repository. scRNA-seq data have been deposited in NCBI's Gene Expression Omnibus and are accessible through the GEO accession number GSE199726. RNA-seq and ChIP-seq data were deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession numbers GSE200064 and GSE200065, respectively.

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Supplemental materials for 1 Novel TCF21^{high} pericyte subpopulation promotes colorectal cancer metastasis 2 3 by remodeling perivascular matrix 4 5 Supplemental methods 6 Cell lines and cell culture 7 Human CRC cell lines HCT116, DLD-1, RKO, SW480, SW620, Caco-2 and human 8 microvascular endothelial cell-1 (HMEC-1) were purchased from the American Type 9 Culture Collection (Manassas, VA). Mouse colon cancer cell line MC38 (Cat. 10 BNCC337716) was from BeNa Culture Collection (Beijing, China). HCT116, DLD-1, 11 RKO, SW480, SW620 and Caco-2 cells were cultured with DMEM. MC38 cells were 12 maintained in RPMI-1640. DMEM and RPMI-1640 medium were supplemented with 13 14 10% FBS (Cat. FSP500, ExCell Bio, Shanghai, China) and 1% penicillin-streptomycin (PS). HMEC-1 cells were cultured in endothelial cell medium (ECM, Cat. 1001, 15 Sciencell research laboratories, Corte Del Cedro Carlsbad, CA) supplemented with 5% 16 FBS, 1% endothelial cell growth supplement (ECGS), and 1% PS. All cell lines were 17 cultured at 37 °C in incubator with 5% CO2. MC38, HCT116 and DLD-1 cells were 18 19 infected with lentivirus harboring luciferase (Genechem, Shanghai, China) to generate the MC38-luc, HCT116-luc and DLD-1-luc cells, which were then selected with 20 21 puromycin (2 μg/mL) for 2 days. All cell lines were authenticated to have no crosscontamination using a STR Multi-amplification Kit (MicroreaderTM21 ID System) and 22 tested negative for mycoplasma by the TransDetect® PCR Mycoplasma Detection Kit 23 (Cat. FM311-01 Transgen, Beijing, China). 24 25

Human samples and specimens

- 27 Human CRC surgical samples (12 cases, patients' information was listed in
- 28 Supplemental Table 5 and Supplemental Table 6) and specimens (75 cases, patients'
- 29 information was listed in Supplemental Table 1) were obtained from the First
- 30 Affiliated Hospital of Jinan University (Guangzhou, China).
- 32 Mice

- 33 Male C57BL/6JGpt mice, male BALB/c nude mice (4-6 weeks, 20-22 g), Rosa26-
- 34 CAG-LSL-Cas9-tdTomato mice (B6/JGpt-Rosa26^{tm1(CAG-LSL-Cas9-tdTomato)}/ Gpt; Cat.
- 35 T002249), Cspg4-CreERT2 mice (B6/JGpt-Cspg4^{em1Cin(CreERT2-P2A)/Gpt}; T006187), and
- 36 Tcf21-flox mice (B6/JGpt-Tcf21^{em1Cflox}/Gpt; T013083) were obtained from
- 37 GemPharmatech Co., Ltd (Nanjing, Jiangsu, China). Pericyte lineage tracing mice
- 38 (PClin) were generated by crossing mice carrying a tamoxifen-inducible Cre
- 39 recombinase driven by the pericyte-specific *Cspg4* promoter (Tg^{Cspg4-CreERT2}) with mice
- 40 carrying a Cre-responsive reporter gene (tandem dimer Tomato (tdT)) inserted at the
- 41 ROSA26 locus (ROSA^{tdT/+}). The PC^{lin} mice were further crossed with mice harboring
- both Tcf21 alleles flanked by LoxP sites (Tcf21^{flox/flox}) to generate tamoxifen-inducible
- 43 Cspg4-driven pericyte-specific Tcf21 knockout mice (PClin-KO). All mice were
- 44 maintained in a specific pathogen-free (SPF) facility. Mouse genotyping was detected
- by PCR (The primer sequences were listed in **Supplemental Table 7**). Cre activity was
- induced in tumor-bearing mice (6-7 weeks, weight 22-25 g) via oral gavage every other
- 47 day for 3 times (10 mg/kg of tamoxifen in peanut oil). The animal experiments were
- 48 complied with the ARRIVE Guidelines 2.0: updated guidelines for reporting animal
- 49 research¹.

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Isolation and culture of TPCs

TPCs were isolated from CRC patients though a microdissection combined with pericyte medium-based approach that developed by our lab. Briefly, fresh surgical tumor specimens were obtained from CRC patients with or without liver metastasis. Information of the CRC patients was listed in **Supplemental Table 5 and Supplemental Table 6.** Tumor tissues were kept in serum-free DMEM containing PS and placed on the ice, and then washed with pre-chilled PBS in a sterile hood to remove the blood, adipose tissues. Tumor vessels were separated from perivascular adipose tissues under a stereomicroscope (Olympus, SZX7). The acquired tumor vessels were cultured in Pericyte Medium (PM, Cat. 1201, Corte Del Cedro Carlsbad, CA, USA, Sciencell research laboratories) with 5% FBS, 1% PGS and 1% PS at 37 °C with 5% CO₂. TPCs were migrated from the tumor vessels within 14 days, which were then disassociated by trypsin once the confluence reaches 80%. The purity of the isolated TPCs were authenticated by STR Multi-amplification Kit (**Supplemental Table 8**).

Construction of single cell cDNA libraries

For single cell cDNA libraries construction, passage 1 TPCs derived from four CRC patients (patient information was listed in **Supplemental Table 9**) were prepared and analyzed by a 10×Genomics GemCode Single-cell instrument, generating single-cell Gel Bead-In-EMlusion (GEMs). The libraries were generated and sequenced by Chromium Next GEM Single Cell 3' Reagent Kits v3.1 and Illumina HiSeq 4000 by

Genedenovo Biotechnology Co., Ltd (Guangzhou, China) with a custom paired-end

73 sequencing mode 26 bp (read 1) \times 98 bp (read 2).

Bioinformatic analysis of scRNA-seq

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Reads uniquely mapped to the transcriptome and intersecting an exon at least 50% were considered for UMI counting. Before quantification, the UMI sequences would be corrected for sequencing errors, and valid barcodes were identified based on the EmptyDrops method. The cells by gene matrices were produced via UMI counting and cell barcodes calling. The cells by gene matrices for each sample were individually imported to Seurat version 3.1.1 for downstream analysis. After removing the unwanted cells from the dataset, data normalization and batch effect correction, the integrated expression of matrix was then scaled and performed on principal component analysis (PCA) for dimensional reduction, those had a strong enrichment of low P-value genes for downstream clustering were identified as significant PCs. Seurat implemented a graph-based clustering approach. Distances between the cells were calculated based on previously identified PCs. For visualization of clusters, t-distributed Stochastic Neighbor Embedding (t-SNE) were generated using the same PCs. Expression value of each gene in given clusters were compared against the rest of cells using Wilcoxon rank sum test. Significantly upregulated genes were identified using several criteria. First, genes had to be at least 1.28-fold overexpressed in the target cluster. Second, genes had to be expressed in more than 25% of the cells belonging to the target cluster. Third, P value is less than 0.05. The Gene ontology (GO) enrichment analysis was performed with the GO database (http://www.geneontology.org/). GO has three ontologies: molecular function, cellular component, and biological process. The calculated P-values were false discovery rate (FDR)-corrected, taking FDR ≤ 0.05 as a threshold. GO terms meeting this criterion were defined as significantly enriched GO terms in differentially expressed genes.

Analysis of transcription factor network inference was performed with the SCENIC R package². In brief, log-normalized expression matrix generated using Seurat was used as input, and the pipeline was implanted in three steps. First, gene coexpression network was established via GENIE3³. Second, each module was pruned based on a regulatory motif near a transcription start site via RcisTarget. Precisely, the networks were retained if the transcription factor (TF)-binding motif was enriched among its targets, while target genes without direct TF-binding motifs were removed. The retained networks were called regulons. Third, the activity of each regulon in each single cell was scored (AUC score) using AUCell R package. Gene regulatory network (GRN) plots of all regulons were done using the cytoscape software⁴.

Analysis of the public datasets

scRNA-seq data of colon (GEO accession GSM3140596, GSM3140595, GSM3140594, and GSM3140593)⁵ and intestine (GEO accession GSM4159165 and GSM4159164)⁶ acquired using 10× Chromium protocol were download, and the sequencing reads were realigned, and cell clustering was performed as described above.

Construction of MC38-luc-LM3, HCT116-luc-LM3 cells and DLD-1-luc-LM3

cells

To establish highly metastatic MC38-luc-LM3 cells, MC38-luc cells (1×10^5) suspended in 100 μ L of Matrigel (Cat. 354248, Corning, NY) were injected into the spleen of male C57BL/6JGpt mice anesthetized with isoflurane inhalation. Liver metastasis was detected by bioluminescence imaging. MC38-luc-LM1 cells from the metastatic foci were isolated by mouse tumor dissociation kit (Cat. 130-096-730, Miltenyi Biotec, Bergisch Gladbach, Germany) and cultured in complete RPMI-1640 medium and

selected by puromycin (2 µg/mL). MC38-luc-LM1 cells were inoculated into the spleen of male C57BL/6JGpt mice, and MC38-luc-LM2 cells were obtained from the liver metastatic foci. Tumor cells isolated from the third round of liver metastatic foci were termed MC38-luc-LM3 cells, which were employed for the subsequent experiments. The HCT116-luc-LM3 and DLD-1-luc-LM3 cells were isolated by human tumor dissociation kit (Cat. 130-095-929, Miltenyi Biotec) and acquired with BALB/C nude mice by the same pattern of MC38-luc-LM3. The cellular morphology, nucleus size, cell size, cell migration, proliferation, EpCAM expression, stemness and EMT were assessed to evaluate the phenotypical/biological differences between the parental cells and the LM3 cells (Supplemental Figure 20). The origin of all LM3 cells was further validated by short tandem repeat (STR) (Supplemental table 10-12) and luciferase activity (Supplemental Figure 21).

Flow cytometry

Cells were collected, re-suspended in flow cytometry staining buffer, and distributed into 1.5 mL EP tubes. Following fixation with 4% paraformaldehyde on ice and permeabilization with 0.1% Triton X-100 in PBS for 5 min, the cells were incubated with anti-TCF21 antibody (Cat. AB_182134, Abcam) or anti-MATN2 antibody (Cat. AF3044, R&D system,) for 1 h on ice. Then, cells were washed with PBS twice and incubated with Donkey anti-Rabbit IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 546 (Cat. AB_2534016, Invitrogen, Carlsbad, CA, USA) or Donkey anti-Goat IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 488 (Cat. AB_2534102, Invitrogen) for 1 h and analyzed by flow cytometry (BD Biosciences, San Jose, CA). The size of HCT116-luc, DLD-1-luc, MC38-luc, HCT116-luc-LM3, DLD-1-luc-LM3 and MC38-luc-LM3 cells was directly evaluated by forward

scatter using flow cytometry

Animal studies

MC38-luc-LM3 cells (1×10^5) suspended in 100 μ L of Matrigel were orthotopically injected into the cecum wall of PC^{lin} mice and PC^{lin-KO} mice anesthetized with isoflurane inhalation. At the end of the experiment, tumors were collected and subjected to immunohistochemistry and immunofluorescence analysis. The livers were harvested, photographed, and prepared for H&E staining. For co-injection assays, HCT116-luc-LM3, DLD-1-luc-LM3 cells, TPC_{NM} transfected with lentivirus harboring negative control shNC (TPC_{NM}^{shNC}) or shITGA5 (TPC_{NM}^{shITGA5}), TPC_{LM} transfected with Vector (TPC_{LM}^{Vector}) or lentivirus expressing ITGA5 (TPC_{LM}^{ITGA5}) were collected. HCT116-luc-LM3 or DLD-1-luc-LM3 cells (4×10^5) were premixed with TPCs (1.6×10^6) in 100 μ L of Matrigel, which was then injected into the cecum wall of BALB/C nude mice. At the end of the experiment, the mice were sacrificed with CO₂ and the metastatic foci in mouse liver were analyzed by H&E staining. Orthotopic tumor tissues were obtained for Masson staining, immunohistochemical staining, immunofluorescence, and transmission electron microscope analysis.

In vivo cell tracking

For the whole animal imaging *in vivo*, mice were intraperitoneal (i.p.) injected with 3

mg of D-luciferin (Cat. 40901ES01, Yeason Biotechnology, Shanghai, China)

dissolved in 200 µL saline and were anesthetized by isoflurane after injection for 5 min.

Luminescence signals were collected with Xenogen IVIS 200 (Alameda, CA, USA)

and analyzed by the Xenogen Living Image software (Alameda, CA, USA).

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Vessel permeability assay

176 PClin and PClin-KO mice bearing MC38 orthotopic xenografts were intravenously (i.v.)

injected with 1 mg of FITC-labeled Dextran-40 kDa (Cat. F6434, Thermo Scientific)

for 10 min. Then, the mice were perfused with 4% PFA and tumors were obtained and

then frozen. Tumor tissues were sectioned, and tumor vessels were stained for CD31

180 (RRID: AB_2161028, RD, Minneapolis, MN, USA) followed by Donkey anti-Goat

IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor™ 647 (RRID:

AB_2534102, Invitrogen, Carlsbad, CA, USA), and the double staining of FITC-

labeled Dextran over CD31-positive vessels indicates vessel permeability.

Isolation and identification of circulating tumor cells (CTCs)

CTC isolation was performed according to a previous study 7 . Briefly, blood (500 µL) was collected from each of PC^{lin} and PC^{lin-KO} mice bearing MC38-luc-LM3 allografts by cardiac puncture and immediately released into heparin-coated tube to avoid coagulation. The red blood cells were removed by blood red cell lysing reagent before cells were seeded in a 12-well plate and cultured with DMEM supplemented with 20% FBS. The adherent tumor cells were identified and counted within 12 h prior to no tumor cell growth. Adherent cells were stained with cancer cell-associated surface marker EpCAM and leukocyte marker CD45 and identified by confocal microscopy as described previously 8,9 . Cells positive for EpCAM but not CD45 were scored as CTCs and subsequently subjected to manual counting, and the CTC counts were presented as CTCs per milliliter of whole blood.

Isolation efficiency of CTCs

To determine the isolation efficiency of CTCs¹⁰⁻¹², 500 MC38-luc-LM3 cells were

spiked into 500 µL of blood collected from the healthy C57BL/6JGpt mice by cardiac puncture. The spiked blood was then treated with blood red cell lysing reagent and the remaining cells were seeded on a 12-well plate and cultured with DMEM containing 20% FBS. The adherent tumor cells were stained with EpCAM and CD45 and the EpCAM+CD45- CTCs were identified by confocal microscopy. The number of EpCAM⁺CD45⁻ CTCs was counted within 12 h at a time of no tumor cell growth. The efficiency of CTC recovery was calculated using the following equation: Cell recovery (%) = counts of isolated MC38-luc-LM3 cells/ 500×100 %.

Chromatin immunoprecipitation (ChIP) and ChIP-Seq

ChIP assay was performed according to the manufacture manual of SimpleChIP® Enzymatic Chromatin IP Kit (Cat. 9003, Cell Signaling Technology, MA, USA). Briefly, TCF21-overexpressing TPCs (TPC $_{NM}^{TCF21}$) were washed twice in clod PBS buffer and cross-linked with 1% formaldehyde for 10 min at room temperature and then stopped by addition of glycine (125 mM). Afterwards, samples were lysed, and chromatins were obtained on ice. Chromatins were then sonicated to get soluble sheared chromatin (average DNA length of 150-900 bp). Then, 20 μ L of chromatin was saved as input and 100 μ L of chromatin was harvested for immunoprecipitation by anti-TCF21 antibody (RRID: AB_10601215, Sigma, Shanghai, China), and anti-IgG was served as the negative control. 10 μ g of anti-TCF21 was used in the immunoprecipitation reactions at 4 °C overnight. Then 30 μ L of protein A beads was added and the samples were further incubated for 2 h. After reverse cross-linking and DNA purification, immunoprecipitated DNA was quantified by real-time PCR. Immunoprecipitated DNA was used to construct sequencing libraries following the protocol provided by the NEXTflex® ChIP-Seq kit (Cat. NOVA-5143-02, BioScientific, TX , USA) and

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sequenced on Illumina Xten with PE 150 method (LC-Bio Technology CO., Ltd., Hangzhou, China). For data analysis, Trimmomatic (version 0.38) was used to filter out low-quality read. MACS2 software (version 2.1.1.20160309) was used to call peaks by default parameters (bandwidth, 300 bp; model fold, 5, 50; q value, 0.05). If the summit of a peak located closest to the TSS of one gene, the peak will be assigned to that gene. GO enrichment analysis was performed using the EasyGO gene ontology enrichment analysis tool (http://bioinformatics.cau.edu.cn/easygo). The GO term enrichment was calculated using hypergeometric distribution with a *P* value cutoff of 0.01. *P* values obtained by Fisher's exact test were adjusted with FDR for multiple comparisons to detect overrepresented GO terms.

RT-qPCR assay

- Total RNA was collected by E.Z.N.A.® Total RNA Kit I (Cat. R6834-02, Omega Bio-
- 238 Tek, Norcross, GA, USA). The purity and concentration of RNA was examined by
- Nanodrop Lite micro spectrophotometer. RNA (2 µg) was reversely transcribed to
- 240 cDNA with All-in-One cDNA Synthesis SuperMix (Cat. B24408-1000, Bimake,
- 241 Houston, TX, USA). Reverse transcription quantitative PCR (RT-qPCR) was
- 242 performed in triplicate using 2× SYBR Green qPCR Master Mix (Cat. B21202,
- Bimake). Samples were loaded into a Roche LightCycler 480 II real-time polymerase
- chain reaction detection system (Roche, Basel, Switzerland) and the data is analyzed
- by $2^{-\Delta \triangle^{Ct}}$ method. The primer sequences were listed in **Supplemental Table 13.**

Cell infection and transfection

- 248 TPCs derived from CRC patients with non-metastasis (TPC_{NM}) were infected with
- 249 lentivirus harboring TCF21 for 48 h and selected by puromycin (2 mg/mL). Detailed

250 information of TCF21 lentivirus and vector was listed as follows: TCF21 (NM 004460) 251 Human Untagged Clone (Cat. SC117372, Origene. Rockwell, MD, USA), Cloning 252 vector PCMV6-XL5 (Cat. PCMV6XL5, Origene). For TCF21- or MATN2-knockdown 253 experiments, TPCs derived from CRC patients with liver metastasis (TPC_{LM}) were 254 transfected with siRNA for 48 h followed by subsequent analysis. For ITGA2- and 255 ITGB1-knockdown experiments, TPC_{NM} were transfected with siRNA for 48 h 256 followed by subsequent analysis. Transfection was performed with LipofectamineTM 257 3000 (Cat. L3000015, Invitrogen, Carlsbad, CA, USA) according to the manufacturer 258 instructions, and the siRNA sequences were listed in Supplemental Table 14. For 259 MATN2-overexpression experiments, TPC_{NM} were transfected with pCMV6-MATN2-260 overexpressing plasmid (Cat. RC203833, Origene) or pCMV6-Entry as empty vector 261 for 48 h (Cat. PS100001, Origene). For ITGA5 overexpression experiments, TPC_{LM} 262 were infected with lentivirus harboring ITGA5 or its corresponding Vector (pGC-FU-263 3FLAG-CBh-gcGFP-IRES-puromycin) (Genechem, Shanghai, China). For ITGA5-264 knockdown experiments, TPC_{NM} were infected with lentivirus harboring shITGA5 or 265 pFU-GW-016 as Vector (Genechem, Shanghai, China).

Y15 and SGI1027 treatment

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Y15 and SGI1027 were purchased from Selleck (Shanghai, China) and dissolved in DMSO. Integrin α 5-overexpressing TPCs (2×10^5) were seeded into 6-well plates and cultured overnight. The next day, cells were treated with Y15 (5 μ M) or SGI1027 (2.5 μ M) for 24 h, and then cells were applied for Western blotting assay and bisulfite

Western blotting assay

sequencing.

Cells were lysed in RIPA lysis buffer on ice for 30 min. Total protein concentration was measured with PierceTM BCA Protein Assay Kit (Cat. 23225, Thermo Scientific, Waltham, MA, USA). Equal amounts of protein (20 μg) were separated in SDS-PAGE gel (Cat. G2004, Solarbio, Beijing, China) and transferred onto polyvinylidene fluoride (PVDF) membranes (Cat. IPVH00010, Millipore, Boston, MA, USA). Following blocking with 5% BSA, the membranes were incubated with indicated antibodies. The blots were detected by Amersham Imager 600 (GE, Boston, MA, USA). The antibodies were listed in **Supplemental Table 15**.

Immunofluorescence analysis

Tissue slices were deparaffinized and incubated with 1×Tris-EDTA (pH 9.0) and 0.05% Tween for 3 min for antigen retrieval. After that, tumor sections were permeabilized in 0.1% TritonTM X-100, blocked with QuickBlockTM immunostaining blocking solution (Cat. ST797, Beyotime, Shanghai, China) and incubated with the corresponding primary antibody overnight at 4 °C. Then, the sections were incubated with the corresponding secondary antibody for 1 h at room temperature. For nucleus staining, sections were incubated with 1 μg/mL DAPI (Cat. MBD0015, Sigma) for 15 min. The slides were photographed with a Zeiss LSM 800 confocal microscope and analyzed with Image J software (RRID: SCR_003070, Rawak Software Inc., Stuttgart, Germany). The primary and secondary antibodies were listed in Supplemental Table 16. For phalloidin immunofluorescence assay, HCT116-luc, DLD-1-luc, MC38-luc, HCT116-luc-LM3, DLD-1-luc-LM3 or MC38-luc-LM3 cells were plated on the glass bottom cell culture dish and incubated with DMEM for 24 h. The next day, cells were fixed, permeabilized with 0.1% TritonTM X-100 and then incubated with Alexa FluorTM 594-phalloidin (Cat. A12381, Thermo) for 1 h. Cell nucleus were stained with 1 μg/mL

DAPI for 15 min. The cytoskeleton elements were photographed with a Zeiss LSM 800 confocal microscope.

H&E staining, immunohistochemistry, and Masson staining

Fixed tissues were embedded in paraffin and sectioned (5 μm). Following deparaffinized, the sections were subjected to antigen retrieval procedures with an EDTA antigen retrieval solution (Cat. P0086, Beyotime, Shanghai, China). Then, the slides were incubated with hematoxylin followed by counterstaining with eosin. For immunohistochemistry assay, tumor sections were incubated with primary antibodies overnight at 4 °C followed by incubation with HRP-conjugated secondary antibodies. The primary and secondary antibodies were listed in **Supplemental Table 17.** Protein expression in tumor sections was detected using a DAB kit (Cat. G1212, Servicebio, Wuhan, Hubei, China), followed by counterstaining with hematoxylin (Cat. G1004, Servicebio, Wuhan, Hubei, China). Images were acquired with an Olympus BX 53 microscope and analyzed with Image J software. For Masson staining, tissue sections were prepared with Masson Tricolor Staining Solution (Fast Green Method) kit (Cat. G1343, Solarbio, Beijing, China). Images were acquired with Olympus BX 53 microscope and analyzed with Image J software.

RNA sequencing analysis

Total RNA was isolated and purified by TRIzol reagent (Cat. 15596018, Invitrogen, Carlsbad, CA, USA) following the manufacturer manual. The RNA concentration and integrity were evaluated by NanoDrop ND-1000 (NanoDrop, Wilmington, DE, USA) and Bioanalyzer 2100 (Agilent, CA, USA). Then, poly (A) RNA was purified from 1µg total RNA by Dynabeads Oligo (dT) 25-61005 (Thermo Fisher, CA, USA) using two

rounds of purification. Then the poly(A) RNA was cut into pieces using Magnesium RNA Fragmentation Module (Cat. e6150, NEB, NY, USA) under 94 °C for 5-7 min. Then the fragmented RNA pieces were reversely transcribed into cDNA by SuperScriptTM II Reverse Transcriptase (Cat.1896649, Invitrogen, USA) and sequenced with illumina NovaseqTM 6000 (LC-Bio Technology CO., Ltd., Hangzhou, China). Then, StringTie and edgeR were used to evaluate the expression levels of all transcripts. The differentially expressed mRNAs and genes were picked with log2 (fold change) >1 or log2 (fold change) <-1 and with statistical significance (P value < 0.05) by R package-edgeR. The volcano plot revealed the distributions of log2 fold change and P values for the differentially expressed genes. The GO terms (http://www.geneontology.org) of these differentially expressed genes were annotated.

Migration and invasion assay

Migration assay was performed with 24-well Boyden chambers (Corning, NY, USA) containing inserts of polycarbonate membranes with 8 μ m-pores. Cells suspended with 100 μ L of serum-free medium were seeded in the upper compartment (3×10⁴ HCT116 cells or 2×10⁴ TPCs). The bottom chamber was filled with different chemoattractants. For invasion assay, the upper chamber was pre-coated with 30 μ L of Matrigel (diluted at 3:1 using PBS) and incubated for 30 min. Then, PKH67-labeled HCT116 cells or DLD-1 cells (5×10³) mixed with TPCs (2.5×10⁴) were seeded into the upper chamber. The bottom chamber was filled with PM and DMEM (5:1). Following incubation for 48 h, the upper chamber was fixed with 4% paraformaldehyde for 30 min and then the cells were stained with 0.1% crystal violet. The non-migrated cells on the upper side of the membrane were removed with a cotton swab. The cells remaining on the lower

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surface were photographed under an inverted microscope and analyzed with Image Pro Plus 6 software. Collagen gel contraction assay This experiment was performed with Cell Contraction Assay kit (Cat. CBA-5020, CELL BIOLABS, San Diego, CA, USA). TCF21-overexpressing or -knockdown TPCs were harvested and resuspended in PM at 5×10^6 cells/mL. Cold collagen gel working solution was prepared according to the instructions and mixed with the cell suspension at a ratio of 4: 1. 0.5 mL of the cell-collagen mixture per well was added in a 24-well plate. After incubating 1 h at 37 °C, 1.0 mL culture medium was added into the collagen gel. Cultures were incubated for two days, and the collagen gels were gently released from the sides of the culture dishes with a sterile spatula. The collagen gel size (contraction index) was measured at 0, 6, 12, 24 and 48 h and quantified with Image J. Cell proliferation assay Cells (5×10^3) were cultured overnight in 96-well plates. The next day, cells were treated with or without the culture supernatant of TPCs (48-h culture medium) and cell proliferation was determined by BeyoClick™ EdU Cell Proliferation kit (Cat. C0071S, Beyotime, Shanghai, China) and analyzed with Image Pro Plus 6 software. Adhesion assay TPCs with the overexpression or knockdown of TCF21 were collected, washed, and stained with PKH67 (Cat. MINI67, Sigma). PKH67-labeled TPCs (2×10⁴) were seeded in a 96-well plate for 2 h. Then, the media was removed, and cells were washed with

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PBS twice to remove the non-adherent cells. Images were acquired with a fluorescent microscope (ZEISS) and analyzed with Image Pro Plus 6 software. **Tube formation assay** Tube formation was performed with a 96-well plate. Matrigel was first coated in the plates at 37 °C for 30 min. Then, HMEC-1 cells (2×10⁴) supplemented with 100 μL ECM were seeded in the Matrigel-coated plates. After 2-h incubation, ECM was replaced, and HMEC-1 cells were incubated with the conditioned medium of TPCs. The capillary tubes were photographed under an inverted light microscopy, and the number of tubes was analyzed by Image Pro Plus 6 software. Transmission electron microscopy analysis Tumor tissues were acquired and fixed in 2.5% glutaraldehyde (Cat. PH9003, Maya Reagent, Zhejiang, China). All samples were post-fixed in 1% osmium tetroxide (Cat. 23311-10, Polysciences, USA), dehydrated in graded concentration of alcohols, and then embedded in low-viscosity resin. The embedded tissues were sectioned and stained with saturated uranyl-acetate and Sato's lead-citrate. Sections were imaged using JEM1200EX transmission electron microscope equipped with BioScan600W digital camera (JEOL, Tokyo, Japan). Second harmonic generation and two-photon excited fluorescence (SHG/TPEF) Tumor tissues were acquired and fixed in 4% paraformaldehyde overnight followed by washing with PBS twice and sectioned (5 μm) using Vibration slice (Leica, VT1000S). Following blocking with 5% BSA solution for 1 h, the sections were incubated with anti-CD31 antibody at 4 °C overnight. The slides were then incubated with Donkey

anti-Goat IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor™ 488 (RRID:

399 AB_2534102, Invitrogen) and immersed in PBS for SHG/TPEF microscopy (Nikon,

Tokyo, Japan). TPEF was utilized for visualization of tumor vessels stained by CD31

(red) and SHG was used to visualize collagen structure (green) at 790 nm excitation

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Atomic Force Microscopy (AFM) measurement

For collagen organization, coatings of isotropic collagen I and AFM detection were performed according to previous report¹³. In vitro fibrillogenesis of collagen-I (Cat. 08-115, Merck; 4.4 mg/mL) was initiated by diluting the collagen I solution tenfold in fibrillogenic buffer (50 mM glycine, 200 mM KCl, pH 9.0). After mixture for 10 min at room temperature, 60 µL of the diluted solution were added to the 22 mm-silicide coverslips and incubated overnight at 37 °C. The next day, coverslips were washed with PBS twice and plated in 12-well plate. Then, TPCs (1×10^4) were seeded on the Collagen I-coated coverslip and cultured for 5 days. AFM Imaging was performed with a NanoWizard II AFM (JPK-Instruments, Berlin, Germany) mounted on an inverted microscope (Axiovert 200, Zeiss, Jena, Germany). Scanning of samples was performed at a scan rate of 0.25 Hz and five fields were recorded for each sample. For perivascular stiffness measurements, AFM was performed according to modified published procedures¹⁴. Tissues were acquired and embedded within OCT. Then, the frozen tissue blocks were sectioned at a thickness of 20 µm and immersed in proteinase inhibitorcontained PBS at room temperature. The next day, tumor sections were applied for AFM quantification of Young's modulus (Bruker, USA). Briefly, silicon nitride cantilevers with a spring constant of 0.15 N/m were attached by a borosilicate glass spherical tip with a diameter of 5 µm. Cantilevers were tapping on the perivascular

region of tumor sections and five 15 μ m \times 15 μ m AFM stiffness map (16 \times 16 raster series) were acquired for each sample. The Young's modulus of the perivascular region in each section were determined by Hertz model. Tissue samples were assumed to be incompressible and a Poisson's ratio of 0.5 was used in the calculation of the Young's modulus.

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DNA extraction and bisulfite sequencing

- 430 DNA was extracted using the Genomic DNA Purification Kit (Cat. A1120, Promega,
- 431 WI, USA), which was followed by treatment with sodium bisulfite (Zymo Research,
- 432 CA, USA). The converted DNA was purified and amplified for sequencing by Biossci
- Biotechnology Co. Ltd (Wuhan, Hubei, China). Primers of TCF21 promoter bisulfite-
- 434 modified regions were: Forward primer (5'-3'): TTTTTGATGTTTTGAAAATGATT -
- 435 AGG; Reverse primer (5'-3'): CAACCACCTTC TCCCAACTATAA.

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Organotypic culture system

- 438 Organotypic culture system was constructed with 12-well Boyden chambers (Corning,
- NY, USA) containing inserts of polycarbonate membranes with 0.4 µm-pores. HMEC-
- 1 cells (2×10^4) supplemented with 100 µL ECM were seeded in the upper chamber and
- 441 incubated overnight. The next day, ECM was removed, and TPCs (5 \times 10⁵) were
- embedded in 1.0 mL collagen I and plated in the chamber. The lower chamber was
- 443 filled with complete PM and ECM (PM: ECM, 5:1). Following incubation at $37\,^{\circ}\text{C}$, 5%
- 444 CO₂ for 5 days, the mechanical properties of the matrix, complex modulus (G*), were
- determined with a rheometer (Malvern Kinexus pro⁺, USA) according to previous
- report¹⁵. The elastic modulus (E) was determined from G* by assuming a poison's ratio
- 447 (v) of 0.5 with the expression $E=2G^*(1+v)$ to allow comparison to other published

work. For invasion assay, TCF21-overexpressing TPCs (5 × 10⁵) were embedded in a matrix mixture of 0.25 mL collagen I and 0.75 mL Matrigel and plated in the chamber. Following incubation for 5 days, PKH67-labeled HCT116 cells (3 × 10⁴) supplemented with 100 μL complete DMEM were plated on the top of matrix and further cultured for 1 day. At the end of experiment, the whole matrix was fixed in 4% overnight and subjected to immunofluorescence analysis. The migrated HCT116 cells were detected by staining of EpCAM. The invaded cells were observed under a Zeiss LSM 800 confocal microscope and analyzed with Image Pro Plus 6 software.

Statistical analysis

The statistical values were calculated with GraphPad Prism 8.0 (GraphPad Software, Inc., San Diego, CA). Differences between two groups were evaluated with two-tailed unpaired t-test or Mann Whitney test. Differences among three groups or more were evaluated using one-way ANOVA followed by Tukey's post hoc test. Survival curves were plotted using the Kaplan Meier method and compared using the log-rank test. The receiver operating characteristic (ROC) curves were performed and the area under ROC curve (AUC) was calculated by logistic regression model to evaluate the diagnostic accuracy. Comparisons of variables were performed using Fisher's exact test or chi-squared test based on their categorical data. Multivariable logistic regression was used to analyze the predictors of CRC metastasis. P < 0.05 was considered as significant

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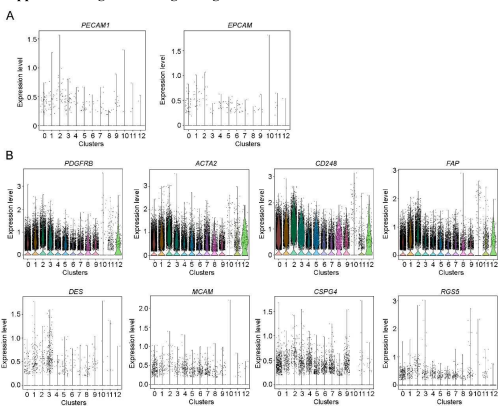
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Supplemental figures and figure legends



Supplemental Figure 1. Transcriptomic characterization of 13 subsets of TPCs. (A)

Gene expression profiles of *PECAM1* and *EPCAM* in distinct subsets of TPCs. **(B)** Gene expression profiles of *PDGFRB*, *ACTA2*, *CD248*, *FAP*, *DES*, *MCAM*, *CSPG4*, and *RGS5* in distinct subsets of TPCs.

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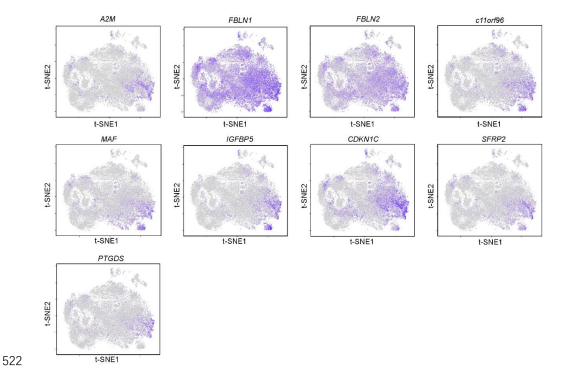
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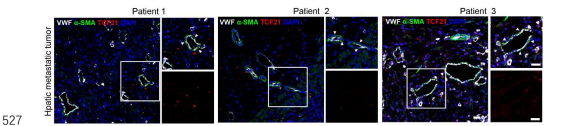
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Supplemental Figure 2. Distribution of the matrix-pericyte-related genes in all subsets of TPCs. t-SNE visualization of gene distribution of A2M, FBLN1, FBLN2, c11orf96, MAF, IGFBP5, CDKN1C, SFRP2 and PTGDS in all subsets of TPCs.



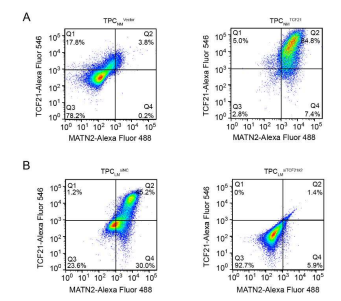
Supplemental Figure 3. Determination of TCF21 in TPCs in the hepatic metastatic tumors from CRC patients. Representative images of TCF21 staining (red) in TPCs (αSMA^+ , green) in the hepatic metastatic tumors from CRC patients (n = 20). Scale bar, 20 μm .

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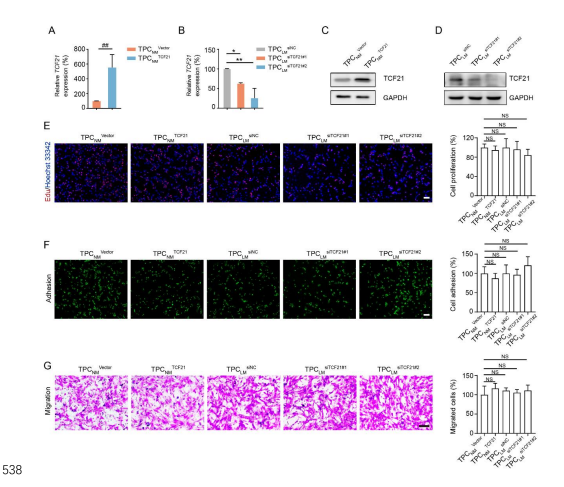
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Supplemental Figure 4. The expression of TCF21 is positively correlated with MATN2 in TPCs. (A) FCM analysis of the TCF21⁺MATN2⁺ TPCs in TPC_{NM} infected with TCF21 lentivirus (TPC_{NM}^{TCF21}) or Vector (TPC_{NM}^{Vector}) (n = 3). (B) FCM analysis of the TCF21⁺MATN2⁺ TPCs in TPC_{LM} transfected with siRNA targeting TCF21 (TPC_{LM}^{siTCF21}) or negative control (TPC_{LM}^{siNC}) (n = 3).



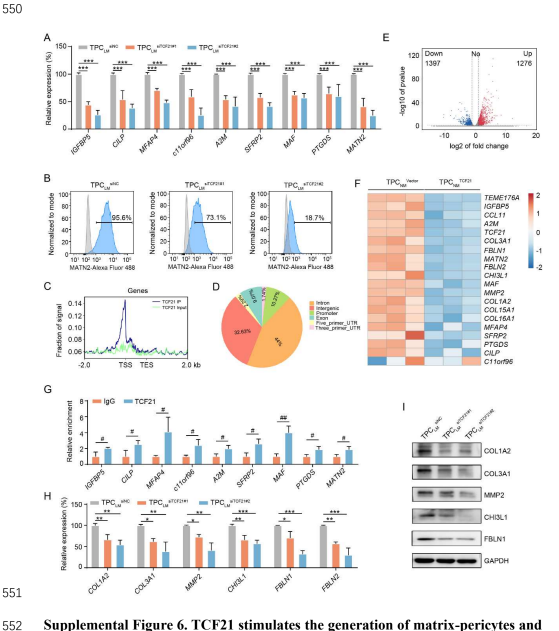
Supplemental Figure 5. Effects of TCF21 on the proliferation, adhesion, and migration of TPCs. (A, B) RT-qPCR analysis of TCF21 mRNA levels in TCF21-overexpressing (A) or -knockdown (B) TPCs (n = 3). (C, D) Western blotting analysis of TCF21 in TCF21-overexpressing (C) or -knockdown (D) TPCs (n = 3). (E) Representative images and quantification of cell proliferation in TCF21-overexpressing and -knockdown TPCs (n = 3). Scale bar, 100 μ m. (F) Representative images and quantification of TPCs adhesive to fibronectin (n = 3). Scale bar, 100 μ m. (G) Transwell assay for the migration of TPCs. Quantification of the migrated TPCs is shown (n = 3). Scale bar, 200 μ m. Data are presented as mean \pm SEM. NS, not significant. *#*P < 0.01 by two-tailed unpaired t-test; NS, *P < 0.05, **P < 0.01 by one-way ANOVA followed by Tukey's post hoc test.

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Supplemental Figure 6. TCF21 stimulates the generation of matrix-pericytes and induces ECM remodeling. (A) RT-qPCR analysis of matrix-pericyte-specific genes in TPC_{LM} transfected with siRNA targeting TCF21 (TPC_{LM}^{siTCF21}) or negative control (TPC_{LM}^{siNC}) (n = 3). (B) FCM analysis of MATN2 expression in TCF21-knockdown TPCs (n = 3). (C) ChIP-seq summary plot of TCF21 enrichment across the indicated genomic distance in TCF21-overexpressing TPCs (n = 3). (D) The distribution of TCF21 peaks on gene elements (n = 3). (E) Volcano Plot of TCF21 regulated genes (n

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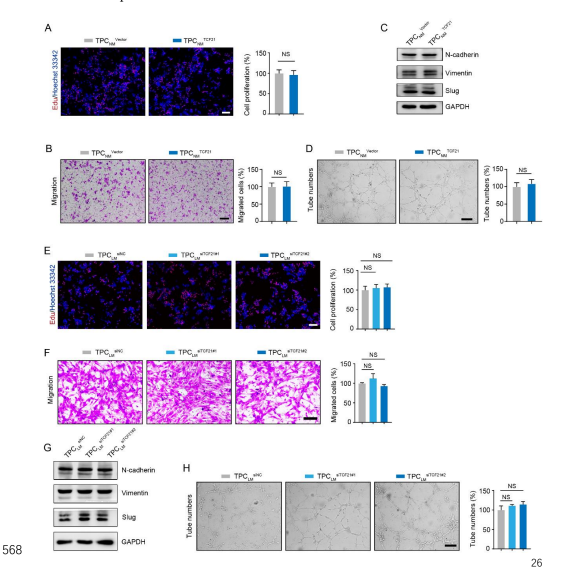
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= 3); Red dots represent the up-regulated genes and blue dots represent the down-regulated genes. **(F)** Heat maps of the differentially expressed genes between TPC_{NM}^{Vector} and TPC_{NM}^{TCF21} (n = 3). **(G)** ChIP-qPCR analysis of TCF21 binding at the promoter of indicated genes in TPC_{NM}^{TCF21} (n = 3). **(H)** RT-qPCR analysis of differentially expressed genes in TPC_{LM}^{siNC} and TPC_{LM}^{siTCF21}. **(I)** Western blotting analysis of COL1A2, COL3A1, MMP2, CHI3L1, and FBLN1 in TPC_{LM}^{siNC} and TPC_{LM}^{siTCF21} (n = 3). Data are presented as mean \pm SEM, $^*P < 0.05$, $^{**}P < 0.01$, $^{***}P < 0.001$ by one-way ANOVA followed by Tukey's post hoc test, $^*P < 0.05$, $^{**}P < 0.01$ by two-tailed unpaired t -test.



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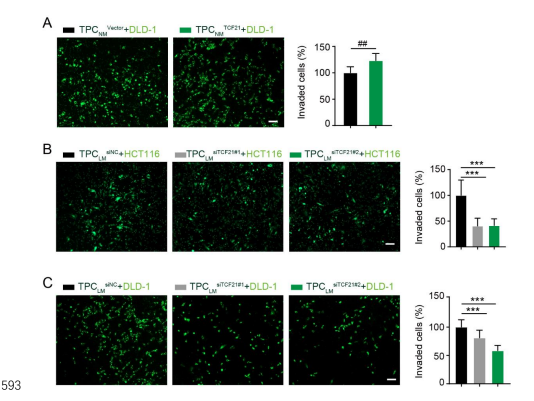
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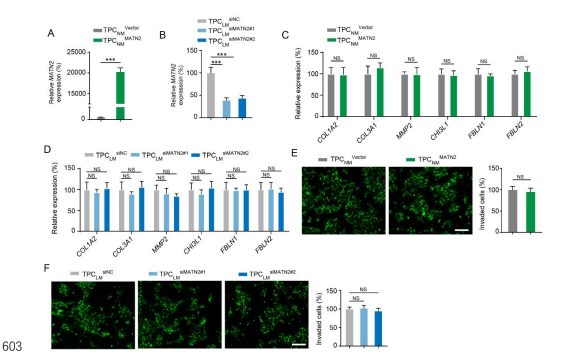
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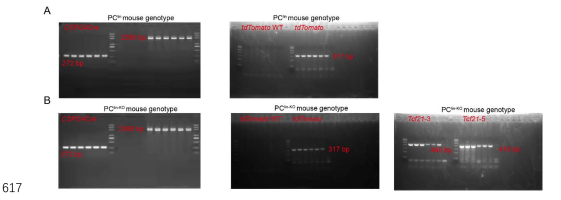
Supplemental Figure 7. TCF21 in TPCs has negligible effects on cell migration and angiogenesis. (A) EdU assay for the proliferation of HCT116 cells primed with conditioned medium from TPC_{NM}^{Vector} and TPC_{NM}^{TCF21} for 48 h (n = 3). Scale bar, 100 μm. (B) Transwell assay for cell migration of HCT116 cells. HCT116 cells were seeded on the upper chamber of the transwell and the bottom compartment was filled with conditioned medium of TPC_{NM}^{Vector} and TPC_{NM}^{TCF21}. After 48 h, the migrated cells were imaged and counted (n = 3). Scale bar, 100 μ m. (C) Western blotting analysis of EMT markers in HCT116 cells primed with conditioned medium of TPC_{NM}^{Vector} and TPC_{NM}^{TCF21} (n = 3). **(D)** Representative images and quantification of tube numbers formed by HMEC-1 cells. HMEC-1 cells suspended with ECM were seeded on the Matrigel coated 96-well plated. After 2 h, ECM were replaced with the conditioned medium of TPC_{NM}^{Vector} and TPC_{NM}^{TCF21}, and the number of formed tubes was calculated 2 hours later (n = 3). Scale bar, 100 μ m. (E) EdU assay for the proliferation of HCT116 cells primed with conditioned medium of TPC_{LM}siNC and TPC_{LM}siTCF21 (n = 3). Scale bar, 100 µm. (F) Representative images and quantification of migrated HCT116 cells. HCT116 cells were seeded on the upper chamber of the transwell and the bottom chamber was filled with the conditioned medium of TPC_{LM}siNC and $TPC_{LM}^{siTCF21}$. After 48 h, the migrated cells were imaged and counted (n = 3). Scale bar, 100 μm. (G) Western blotting analysis of EMT markers in HCT116 cells primed with conditioned medium of TPC_{LM}^{siNC} and $TPC_{LM}^{siTCF21}$ (n = 3). (H) Tube formation assay for HMEC-1 cells treated with conditioned medium of TPC_{LM}siNC and TPC_{LM}siTCF21 as indicated in (D) (n = 3). Scale bar, 100 μ m. Data are presented as mean \pm SEM, NS, not significant. Two-tailed unpaired t-test (A, B, D), one-way ANOVA followed by Tukey's post hoc test (E, F, H).



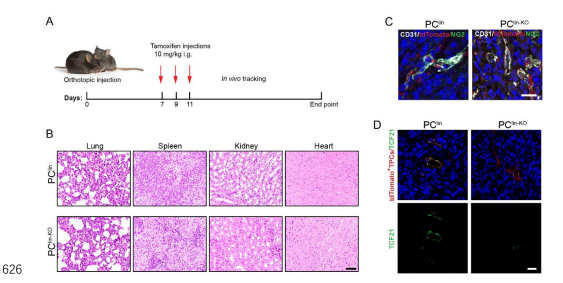
Supplemental Figure 8. TCF21 in TPCs promotes invasion of CRC cells. (A) Representative images and quantification of invaded DLD-1 cells (green). DLD-1 cells mixed with TPC_{NM}^{Vector} or TPC_{NM}^{TCF21} were seeded into the Matrigel-coated transwell. The invaded DLD-1 cells were photographed and counted after 48 h (n = 3). Scale bar, 100 μ m. (B, C) Transwell assay for invasion of HCT116 cells and DLD-1 cells. HCT116 cells (B) or DLD-1 cells (C) were pre-mixed with TPC_{LM}^{SiNC} or TPC_{LM}^{SiTCF21} and subjected to invasion assay as indicated in (A) (n = 3). Scale bar, 100 μ m. Data are presented as mean \pm SEM. ***P < 0.01 by two-tailed unpaired t-test, ***P < 0.001 oneway ANOVA followed by Tukey's post hoc test



Supplemental Figure 9. MATN2 has negligible effects on ECM remodeling and CRC metastasis. (A, B) RT-qPCR analysis of MATN2 in MATN2-overexpressing (A) or -knockdown (B) TPCs (n = 3). (C) RT-qPCR analysis of the ECM-related genes in TPC_{NM}^{Vector} and TPC_{NM}^{MATN2} (n = 3). (D) RT-qPCR analysis of the indicated genes in TPC_{LM}^{siNC} and TPC_{LM}^{siMATN2} (n = 3). (E, F) Representative images and quantification of the invaded DLD-1 cells (green). DLD-1 cells pre-mixed with MATN2-overexpressing (E) or -knockdown (F) TPCs were seeded into the Matrigel-coated transwell. The invaded DLD-1 cells were photographed and counted after 48 h (n = 3). Scale bar, 100 μ m. Data are presented as mean \pm SEM, NS, not significant. ***P<0.001. Two-tailed unpaired *t*-test (A, C, E), one-way ANOVA followed by Tukey's post hoc test (B, D, F).

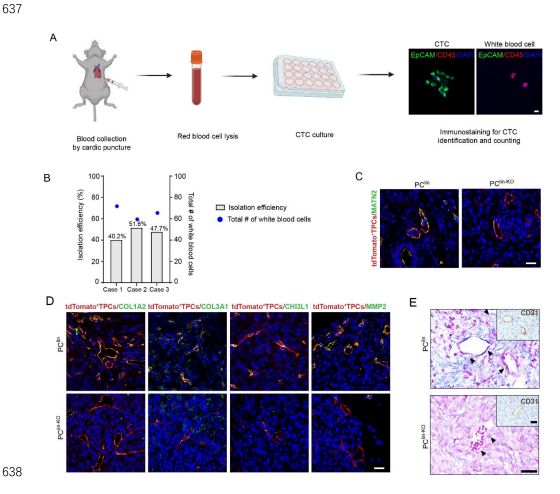


Supplemental Figure 10. Mouse genotyping. (A) PCR analysis of the genotype of PC^{lin} mouse (n = 6). **(B)** PCR analysis of the genotype of PC^{lin-KO} mice (n = 6). All mice were analyzed by PCR genotyping. PCR analysis of Cspg4Cre showed two bands identifying homozygous Cspg4Cre at 2386 bp (knock in) and 272 bp (wild type); PCR analysis of tdTomato showed homozygous tdTomato with a band at 317 bp (knock in) and no signal at 479 bp (wild type); PCR analysis of Tcf21 indicated homozygous Tcf21 flox/flox at 418 bp and 440bp.



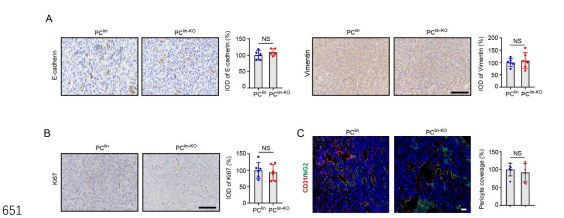
Supplemental Figure 11. Characterization of PClin and PClin-KO mice. (A) Schematic diagram describing the experimental design of *in vivo* experiments. PClin mice and PClin-KO mice were orthotopically injected with MC38-luc-LM3 cells. After 7

days, all mice were treated with tamoxifen (10 mg/kg) through intragastric administration every other day for three times. *In vivo* tracking was performed to detect tumor liver metastasis. **(B)** Representative images of H&E staining of lung, spleen, kidney, heart, and liver derived from PC^{lin} and PC^{lin-KO} mice (n = 6). Scale bar, 50 µm. **(C)** Immunofluorescence analysis of TPCs (tdTomato) in tumor sections by NG2 (green) staining (n = 6). Scale bar, 20 µm. **(D)** Immunofluorescence analysis of TCF21 (green) in TPCs (tdTomato) from PC^{lin} mice and PC^{lin-KO} mice (n = 6). Scale bar, 20 µm.



ECM remodeling and CRCLM. (A) Schematic diagram of the isolation and identification of CTCs. Scale bar, 20 μm. **(B)** Isolation efficiency of CTCs in the blood

spiked with MC38-luc-LM3 cells (n = 3). **(C)** Immunofluorescence staining for MATN2 (green) in TPCs (tdTomato) in primary tumor sections from MC38 allografts (n = 6). Scale bar, 20 μ m. **(D)** Immunofluorescence staining for COL3A1, MMP2, COL1A1 and CHI3L1 (green) in TPCs (tdTomato) from primary tumor sections of MC38 allografts (n = 6). Scale bar, 20 μ m. **(E)** Masson staining for perivascular collagen in in primary tumor sections from MC38 allografts. Tumor vessels were labeled with CD31 (n = 6). Black arrows indicate the perivascular collagen fibers. Scale bar, 50 μ m.



Supplemental Figure 13. Pericyte-specific knockout of *Tcf21* has negligible effects on EMT and proliferation of CRC cells. (A) Immunohistochemical staining and quantification of E-cadherin and vimentin in orthotopic MC38 tumor tissues (n = 6). Scale bar, 50 μ m. (B) Representative images and quantification of Ki67 staining in orthotopic MC38 tumor sections (n = 6). Scale bar, 50 μ m. (C) Immunofluorescence staining and quantification of pericyte coverage as indicated by CD31 (red) and NG2 (green) in tumor sections (n = 6). Scale bar, 20 μ m. Data are presented as mean \pm SEM. NS, not significant. NS by two-tailed unpaired *t*-test.

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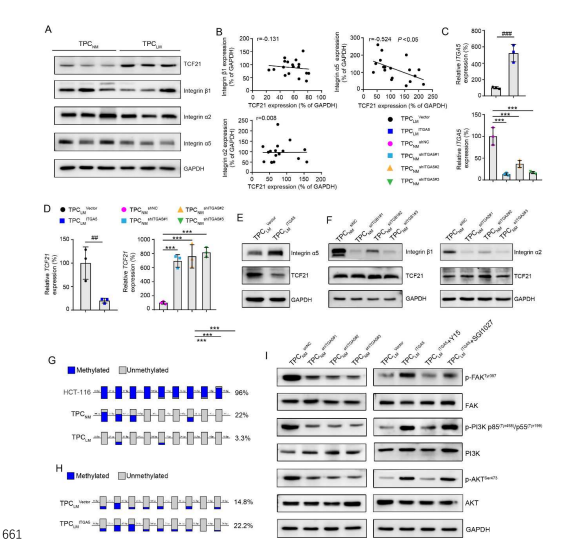
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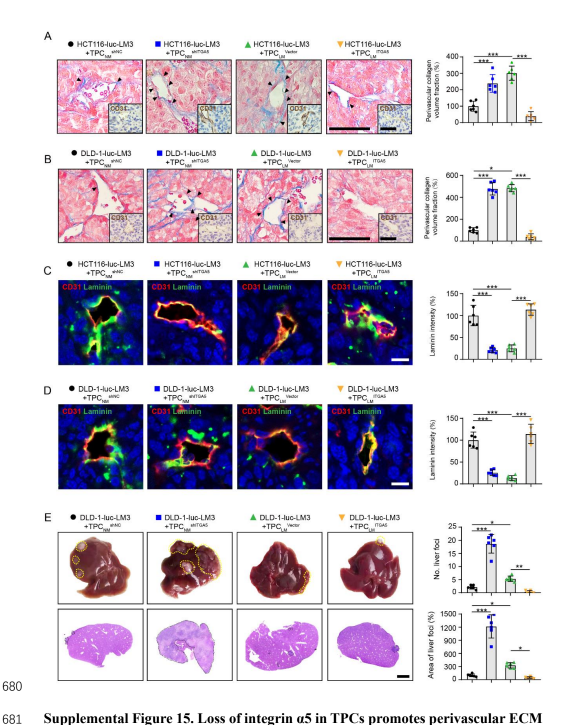
Supplemental Figure 14. Effects of integrin $\alpha 2$ and integrin $\beta 1$ on TCF21 in TPCs.

(A) Western blotting analysis of TCF21, integrin β 1, integrin α 2, and integrin α 5 in TPC_{NM} and TPC_{LM} (n = 3). (B) Pearson's correlation analysis of the expression of integrin β 1, integrin α 2 and integrin α 5 with TCF21 in TPCs (n = 3). (C) RT-qPCR analysis of the levels of *ITGA5* in the integrin α 5-knockdown or -overexpressing TPCs (n = 3). (D) RT-qPCR analysis of the levels of *TCF21* in integrin α 5-knockdown or -overexpressing TPCs (n = 3). (E) Western blotting analysis of integrin α 5 and TCF21 in integrin α 5-overexpressing TPC_{LM} (n = 3). (F) Western blotting analysis of integrin α 2, integrin β 1 and TCF21 in integrin α 2 or integrin β 1-knockdown TPC_{NM} (n = 3). (G)

671 Bisulfite DNA sequencing analysis of TCF21 promoter region in TPCs or HCT116 cells. 672 Blue and gray circles represent methylated and unmethylated CpGs, respectively. The percentage of total methylated CpGs is given on right of each dataset (n = 3). (H) 673 674 Bisulfite DNA sequencing analysis of TCF21 promoter region in integrin α5-675 overexpressing TPC_{LM} (n = 3). (I) Western blotting analysis of FAK/PI3K/AKT 676 signaling axis in integrin α5-knockdown or -overexpressing TPCs with or without FAK inhibitor (Y15) or DNMT1 inhibitor (SGI1027) treatment (n = 3). Data are presented 677 as mean \pm SEM, ##P < 0.01, ###P < 0.001 by two-tailed unpaired t-test, ***P < 0.001 by 678 679 one-way ANOVA followed by Tukey's post hoc test.

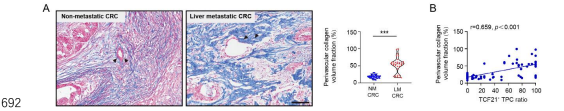
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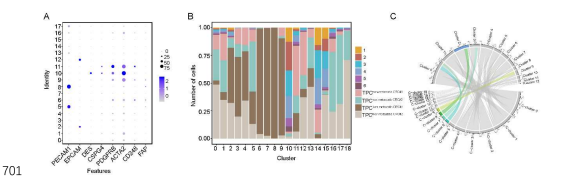


Supplemental Figure 15. Loss of integrin α 5 in TPCs promotes perivascular ECM remodeling and CRCLM. (A, B) Representative images of Masson and CD31 staining in primary tumor sections (n = 6). Black arrows indicate the perivascular collagen fibers. Scale bar, 50 μ m. The quantification of perivascular collagen volume fraction was shown in the right. (C, D) Immunofluorescence staining and quantification of laminin

(green) around the CD31⁺ tumor vessels (red) in HCT116-luc-LM3 xenografts (**C**) and DLD-1-luc-LM3 xenografts (**D**) (n = 6). Scale bar, 20 μ m. (**E**) Representative images and H&E staining of liver metastatic foci (n = 6). Yellow and black dotted lines indicate the metastatic loci. Scale bar, 2 mm. Data are presented as mean \pm SEM. *P < 0.05, **P < 0.01, ***P < 0.001 by one-way ANOVA followed by Tukey's post hoc test.



Supplemental Figure 16. TCF21 in TPCs is associated with perivascular ECM deposition. (A) Masson staining and quantification of perivascular collagen in tumors derived from CRC patients with non-metastasis or liver metastasis (n = 75). Scale bar, 50 μ m. ***P < 0.001 by two-tailed Mann-Whitney test. (B) Pearson's correlation analysis of perivascular collagen volume fraction and TCF21⁺ TPC ratio (n = 75). NM CRC, non-metastatic colorectal cancer, LM CRC, liver metastatic colorectal cancer.



Supplemental Figure 17. Comparison of the scRNA-seq data derived from TPCs and the previous published data. (A) Dot plots for gene expressions in pericytes

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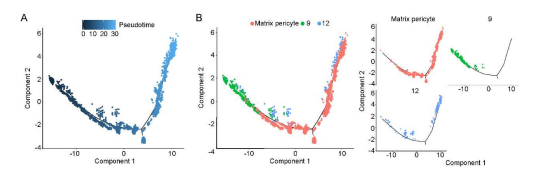
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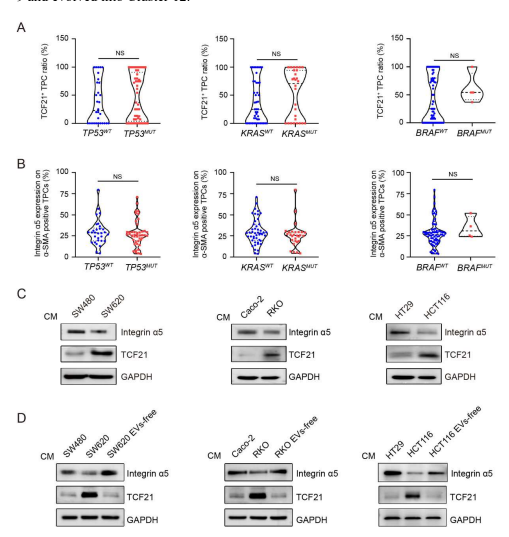
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acquired from previous studies. Raw data of pericyte scRNA-seq were collected from six samples derived from previous studies (GEO accession GSM3140596, GSM3140595, GSM3140594, and GSM3140593) and (GEO accession GSM4159165 and GSM4159164). Total cells derived from previous studies were classified into 18 clusters named L-Cluster 0 to L-Cluster 17. Among them, L-Cluster 3, L-Cluster 9, L-Cluster 10, and L-Cluster 11 were subjected for further analysis as these four populations were positive for DES, CSPG4, PDGFRB, ACTA2, CD248 and FAP (pericyte markers), but negative for PECAM1 (endothelial cell marker) and EPCAM (epithelial cell marker). (B) Analysis of the data of the four populations (L-Cluster 3, L-Cluster 9, L-Cluster 10, and L-Cluster 11) derived from six samples (Sample 1-6) in (A) and our scRNA-seq data (Cluster 0-12). The combined pericytes were categorized into 19 subpopulations, termed C-Cluster 0 to C-Cluster 18. Among them, C-cluster 0-5, C-cluster 10-18 were presented both in our data and the extended data (Sample1-6); however, C-cluster 6-9 were specifically revealed in our data, indicating that the existing pericyte clusters originated from the previous research^{5, 6} were included in our scRNA-seq data and we discovered four new subsets. (C) Comparative analysis of pericyte cluster derived from (B) (C-cluster 0-18) with our data (Cluster 0-12). Among them, C-Cluster 8 was included in Cluster 2 (matrix pericytes).



Supplemental Figure 18. Pseudo-time trajectory for dynamic changes in matrix-

pericytes. (A) Trajectory analysis plot for matrix-pericytes. Cells are ordered in pseudo-time colored in a gradient from dark blue to light blue (B) The trajectory of the differentiation state of matrix-pericytes. Matrix-pericytes were originated from Cluster 9 and evolved into Cluster 12.



Supplemental Figure 19. Effects of metastatic CRC cells on the expressions of integrin $\alpha 5$ and TCF21 in TPCs. (A) Quantification of TCF21⁺ TPC ratio in CRC patients with or without *TP53*, *BRAF* and *KRAS* mutation (n = 75). (B) Quantification of integrin $\alpha 5$ expression in TPCs derived from CRC patients with or without *TP53*, *BRAF* and *KRAS* mutation (n = 75). (C) Western blotting analysis of integrin $\alpha 5$ and

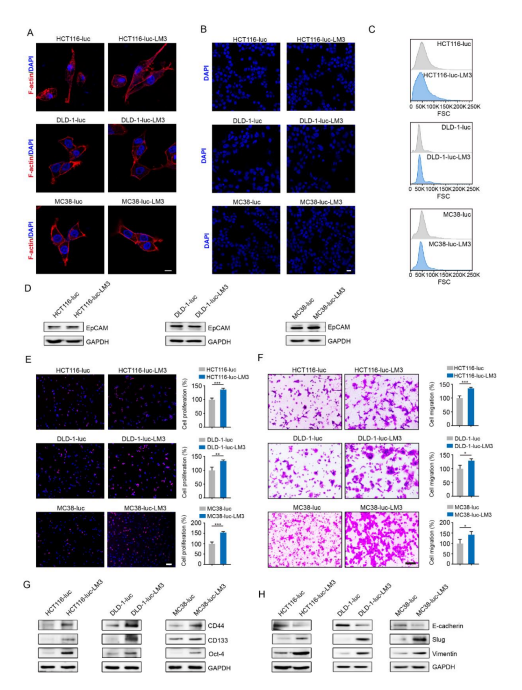
TCF21 in TPCs primed with CM from weakly-metastatic (SW480, Caco-2, HT29) and 735 highly-metastatic (SW620, HCT116, RKO) CRC cells (n = 3). (D) Western blotting 736 analysis of integrin $\alpha 5$ and TCF21 in TPCs primed with or without the EVs-free CM of 737 738 weakly- or highly-metastatic CRC cells (n = 3). The EVs-free CM of highly-metastatic CRC cells were generated by centrifugation at 1×10⁵ g to remove the EVs. EVs, 739 740 extracellular vesicles; CM, conditioned medium. Each sample on the violin plots represents individual patient data. NS, not significant. NS by two-tailed Mann-Whitney 741 742 test. WT, wildtype; MUT, mutant.

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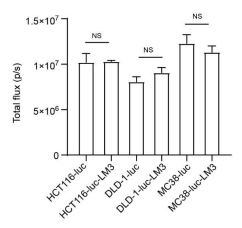
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Supplemental Figure 20. Comparison of parental cells with LM3 cells. (A) Immunofluorescence analysis of cellular morphology (n = 3). Phalloidin-rhodamine was used to identify F-actin. Scale bar, 10 μ m. (B) Immunofluorescence analysis of nucleus size (n = 3). Cell nucleus was measured after DAPI staining of fixed cells. Scale bar, 20 μ m. (C) FCM analysis of the cell size of parental cells and LM3 cells by Forward

scatter (n = 3). **(D)** Western blotting analysis of EpCAM in parental cells and LM3 cells (n = 3). **(E)** EdU assay for the proliferation of parental cells and LM3 cells (n = 3). Scale bar, 100 μ m. **(F)** Transwell assay for the migration of parental cells and LM3 cells (n = 3). Scale bar, 100 μ m. **(G)** Western blotting analysis of CD44, CD133 and Oct-4 in parental cells and LM3 cells (n = 3). **(H)** Western blotting analysis of E-cadherin, slug and vimentin in parental cells and LM3 cells (n = 3). Data are presented as mean \pm SEM. *P < 0.05, **P < 0.01, ***P < 0.001 by two-tailed unpaired t-test.



Supplemental Figure 21. Comparison of luciferase activities between parental cells and LM3 cells. Bioluminescence detection of parental cells and LM3 cells. Data are presented as mean \pm SEM. NS, not significant. NS by two-tailed unpaired t-test.

768 **Supplemental Table 1.** Clinical characteristics of CRC specimens.

Character	Overall population
Gender, number of patients (%)	
Male	33 (44.0)
Female	42 (56.0)
Age, median (range)	69 (34 – 91)
Site of primary tumor, number of patients (%)	
Right hemicolon	17 (22.7)
Left hemicolon	58 (77.3)
Histological grade, number of patients (%)	
High/Moderate	47 (62.7)
Low	28 (37.3)
Size (cm), median (range)	4.3(2-8.6)
TNM stage, number of patients (%)	
I-II	41 (54.7)
IV	34 (45.3)
Liver metastasis (%)	
No	41 (54.7)
Yes	34 (45.3)
TP53 mutation (%)	
No	30 (40)
Yes	45 (60)
KRAS mutation (%)	
No	47 (62.7)
Yes	28 (37.3)
BRAF mutation (%)	
No	71 (94.7)
Yes	4 (5.3)

Supplemental Table 2. Correlation analysis between the MATN2⁺ TPC ratio and the clinicopathologic data.

Character	MATN2 ⁺ TPC	ratio MATN2+ TPC	ratio
Character	(≤30%)	(>30%)	P value
Gender			
Female	22 (66.7%)	11 (33.3%)	0.826
Male	29 (69.0%)	13 (31.0%)	0.820
Age			
< 60	15 (75.0%)	5 (15.0%)	0.422
≥60	36 (65.5%)	19 (34.5%)	0.433
Location			
Right hemicolon	13 (76.5%)	4 (23.5%)	0.395
Left hemicolon	38 (65.5%)	20 (34.5%)	
Differentiatio			
n			
High/Moderate	33 (70.2%)	14 (29.8%)	0.595
Low	18 (64.3%)	10 (35.7%)	0.373
Size			
< 5 cm	35 (79.5%)	9 (20.5%)	0.011
≥5 cm	16 (51.6%)	15 (48.4%)	0.011
TNM stage			
I-II	40 (97.6%)	1 (2.4%)	< 0.001
IV	11 (32.4%)	23 (67.6%)	₹ 0.001
Liver			
metastasis			
No	40 (97.6%)	1 (2.4%)	< 0.001
Yes	11 (32.4%)	23 (67.6%)	` 0.001
TP53 mutation			
No	23 (76.7%)	7 (23.3%)	0.189

Yes	28 (62.2%)	17 (37.8%)	
KRAS			
mutation			
No	35 (74.5%)	12 (25.5%)	0.120
Yes	16 (57.1%)	12(42.9%)	0.120
BRAF			
mutation			
No	49 (69%)	22 (31%)	0.000
Yes	2 (50%)	2 (50%)	0.808

Supplemental Table 3. Correlation analysis between the TCF21⁺ TPC ratio and the clinicopathologic data.

Character	TCF21 ⁺	TPC	ratio	TCF21 ⁺	TPC	ratio	P
Character	(:	≤44%)		((>44%)		value
Gender							
Female	16	(48.5%)		17	(51.5%)		0.340
Male	25	(59.5%)		17	(40.5%)		0.340
Age							
< 60	11	(55.0%)		9 ((45.0%)		0.972
≥60	30	(54.5%)		25 (45.5%)			0.972
Location							
Right hemicolon	6	(35.3%)		11	(64.7%)		0.068
Left hemicolon	35	(60.3%)		23 (39.7%)		0.008	
Differentiation							
High/Moderate	29	(61.7%)		18	(38.3%)		0.113
Low	12	(42.9%)		16	(57.1%)		0.113
Size							
< 5 cm	27	(61.4%)		17	(38.6%)		0.165
≥5 cm	14	(45.2%)		17 (54.8%)			0.105
TNM stage							

I-II	40 (97.6%)	1 (2.4%)	< 0.001
IV	1 (2.9%)	33 (97.1%)	₹ 0.001
Liver			
metastasis			
No	40 (97.6%)	1 (2.4%)	< 0.001
Yes	1 (2.9%)	33 (97.1%)	₹ 0.001
TP53 mutation			
No	18 (60.0%)	12 (40.0%)	0.449
Yes	23 (51.1%)	22 (48.9%)	0.449
KRAS mutation			
No	29 (61.7%)	18 (38.3%)	0.112
Yes	12 (42.9%)	16 (57.1%)	0.113
BRAF mutation			
No	40 (56.3%)	31 (43.7%)	0.479
Yes	1 (25.0%)	3 (75.0%)	0.478

Supplemental Table 4. Multivariable logistic regression for clinical and demographic factors between CRC patients with or without liver metastasis.

	β	S.E.	Wald	P	OR	95% CI
TCF21 ⁺ TPC ratio (%)	7.112	1.435	24.558	<0.001	1226.464	73.636-20427.781
TP53 mutation	0.074	1.536	0.002	0.962	1.077	0.053-21.851
KRAS mutation	0.448	1.561	0.082	0.774	1.565	0.073-33.387
BRAF mutation	0.875	3.392	0.067	0.796	2.399	0.003-1850.29

Abbreviations: S.E., standard error; OR, odds ratio; CI, confidence interval.

Supplemental Table 5. Clinical characteristics of non-metastatic CRC specimens

Characteristics	Case 1	Case 2	Case 3	Case 4	Case 5	Case 6
Gender/Age (yr)	Male (58)	Male (78)	Male (59)	Female (53)	Male (73)	Female (75)
Date of diagnosis	20190429	20191031	20200331	20210323	20210402	20210420
Tumor type	Colorectal	Colorectal	Colon	Colon	Colon	Colon
Tumor type	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma
Location	Rectum	Sigmoid colon	Sigmoid colon	Sigmoid colon	Sigmoid colon	transverse colon
Tumor size	5 cm	4 cm				
(maximum diameter)	3 CIII	4 CIII	2 cm	4 cm	5 cm	5 cm
Differentiation	Moderate	Moderate	Moderate	Moderate	Moderate	Moderate
TNM stage	T2N0M0	T4aN0M0	T3N1bM0	T4aN1bM0	T3N0M0	T3N1aM0
Clinical stage	I	IIB	IIIB	IIIB	IIA	IIIB
Clinical metastasis	No metastasis	No metastasis	No metastasis	No metastasis	No metastasis	No metastasis
Treatment status	Chemotherapy,	No treatment				

radiotherapy	before surgery				

Supplemental Table 6. Clinical characteristics of liver-metastatic CRC specimens

Characteristics	Case 1	Case 2	Case 3	Case 4	Case 5	Case 6
Gender/Age (yr)	Female (61)	Male (51)	Male (78)	Male (64)	Female (51)	Female (59)
Date of diagnosis	20210324	20190516	20191010	20190103	20190626	20200320
T.,,,,,,,,	Rectal	Colorectal	Colorectal	Colorectal	Colorectal	Colorectal
Tumor type	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma
Landin	D	Descending colon	<u>C</u>	G: :1 1	Descending colon	Descending
Location	Rectum	and ileum		Sigmoid colon	and ileum	colon and ileum
Tumor size	C	4	(5	<i>(</i>	7
(maximum diameter)	6 cm	4 cm 6 cm		5 cm	6 cm	7 cm
Differentiation	Moderate	Moderate	Low	Low	Moderate	Low

TNM stage	T3N2M1a	T4bN1M1a	T3N2bM1a	T3N2M1a	T4bN1M1a	T4bN1aM1a
Clinical stage	IVa	IVa	IVa	IVa	IVa	IVa
Clinical metastasis	Liver metastasis					
T	Chemotherapy,	No treatment				
Treatment status	radiotherapy	before surgery				

Supplemental Table 7. Primers for mouse genotyping

Mice	Primer name	Primer Sequence	Product size	Gene type	
1	Rosa26-tF1	CCCAAAGTCGCTCTGAGTTGTTA	VV. 4701		
I	Rosa26-tR1	TCGGGTGAGCATGTCTTTAATCT	Wt=479bp	T002249 <i>Rosa26-CAG-</i>	
2	tdTomato-tF1	CGGCATGGACGAGCTGTACAAG		LSL-Cas9- tdTomato	
2	WPRE-tR2	TCAGCAAACACAGTGCACACCAC	KI=317bp		
3	JS04431-Tcf21-5wt-tF1	GATCCTTCAAATGACTCCAGGCC	WT: 314bp Fl: 418bp	T013083 <i>Tcf21</i> -flox	

	JS04431-Tcf21-5wt-tR1	GTTTGCTAACTTGCTGCCACACAC		
4	XM003792- Cspg4-TF1	AAATCTAAGCGCGGGTCTGGC	WT:0bp	
4	XM003792- Cspg4-TR1	TGCGAACCTCATCACTCGTTGC	KI:272bp	T006187 <i>Cspg4</i> - CreERT2
5	XM003792- Cspg4-TF2	AAATCTAAGCGCGGGTCTGGC	WT:352bp	
3	XM003792- Cspg4-TR2	GGACCATGAGTGCAGTCCCCATA	(KI=2386bp)	

Supplemental Table 8. STR profiles of TPCs.

Marker	Allele 1	Allele 2	Allele 3	Allele 4
D19S433	13	14		
D5S818	11	13		
D21S11	29	31.2		
D18S51	13	14		
D6S1043	12	13		
AMEL	X	Y		
D3S1358	16	17		
D13S317	10			
D7S820	8	12		
D16S539	9	11		
CSF1PO	12	14		
Penta D	9	12		
D2S441	10	11		
vWA	14	15		
D8S1179	10	14		
TPOX	8	10		
Penta E	11	12		
TH01	6	8		
D12S391	19	23		
D2S1338	22	23		
FGA	20	23		

Conclusion of cell identification: The results of STR typing showed that there were no multiple alleles at each locus. No cross contamination of human cells was found in the cells.

Supplemental Table 9. Clinical characteristics of the resected CRC patients

Characteristics	Case 1	Case 2	Case 3	Case 4
Gender/Age (yr)	Male (58)	Male (78)	Male (51)	Male (78)
Date of diagnosis	20190429	20191031	20190516	20191010
Tumor type	Colorectal Adenocarcinoma	Colorectal Adenocarcinoma	Colorectal Adenocarcinoma	Colorectal Adenocarcinoma
Location	Rectum	Sigmoid colon	Descending colon and ileum	Sigmoid colon
Tumor size (maximum diameter)	5 cm	4 cm	4 cm	6 cm
Differentiation	Moderate	Moderate	Moderate	Low
TNM stage	T2N0M0	T4aN0M0	T4bN1M1a	T3N2bM1a
Clinical stage	I	IIB	IVa	IVa
Clinical metastasis	No metastasis	No metastasis	Liver metastasis	Liver metastasis
Treatment status	No treatment before surgery			

Supplemental Table 10. STR profiles of HCT116-luc cells and HCT116-luc-LM3 cells.

Locus		HCT116-luc	HCT116-luc-LM3	
Amelogenin	X	Y	X Y	
D5S818	10	11	10 11	
D13S317	10	12	10 12	
D7S820	11	12	11 12	
D16S539	11		11	
vWA	17		17	
TH01	8	9	8 9	
TPOX	8		8	
CSF1PO	7	10	7 10	

The number of matched peaks

15

Percent match between the query and the database profile:

100%

Conclusion of cell identification: ①The results of STR typing of the cell DNA of this strain show that there is no multi-allelic phenomenon at each locus. No human cell cross-contamination is found in the cells. ②The submitted HCT116-luc-LM3 cells are 100% match for HCT116-luc cells.

Supplemental Table 11. STR profiles of DLD-1-luc cells and DLD-1-luc-LM3 cells.

Locus		DLD-1-luc	DLD-1-luc-LM3	3
Amelogenin	X	Y	X Y	
D5S818	13		13	
D13S317	8	11	8 11	
D7S820	10	12	10 12	

D16S539	12	13	12	13	
vWA	18	19	18	19	
TH01	7	9.3	7	9.3	
TPOX	8	11	8	11	
CSF1PO	12		12		
	and a	1 6 1	,		1.6

The number of matched peaks

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Percent match between DLD-1-luc and DLD-1-luc-LM3:

100%

Conclusion of cell identification: ①The results of STR typing of the cell DNA of this strain show that there is no multi-allelic phenomenon at each locus. No human cell cross-contamination is found in the cells. ②The submitted DLD1-luc-LM3 cells are 100% match for DLD-1-luc cells.

Supplemental Table 12. STR profiles of MC38-luc cells and MC38-luc-LM3 cells.

Locus		MC38-luc	MC38-luc-LM3
18-3	16		16
4-2	19.3	20.3	19.3 20.3
6-7	14	15	14 15
19-2	13		13
1-2	19		19
7-1	26.2		26.2
8-1	16		16
1-1	16		16
3-2	13	14	13 14
2-1	16		16
15-3	22.3		22.3
6-4	18		18
13-1	17.1		17.1
11-2	16		16
17-2	15		15
12-1	17		17

5-5	17	17	
X-1	27	27	
The number of matched peaks			21
Percent match between the query and the database profile			100%

Conclusion of cell identification: ①The results of STR typing of the cell DNA of this strain show that there is no multi-allelic phenomenon at each locus. No human cell cross-contamination is found in the cells. ②The cells of MC38-luc and MC-38-luc-LM3 are mouse cells from a single source, and there is no human-derived cell contamination. ③The submitted MC38-luc-LM3 cells are 100% match for MC38-luc cells.

Supplemental Table 13. Primers for qPCR.

Genes	Forward (5'-3')	Reverse (5'-3')
ACTB	GTTGCTATCCAGGCTGTGCTATCC	GGTGGCAGTGATGGCATGGAC
MFAP4	TGAAGGCACAAGGAGTTCTCT	GGGTAGATGAGGTACACGCC
MMP2	CGACCACAGCCAACTACGATGATG	GTGCCAAGGTCAATGTCAGGAGAG
FBLNI	TGCTCCATCAACGAGACCTG	AGCACTCCCGATTCTCATGG
COL1A2	CCGTGGCAGTGATGGAAGTGTG	CCTTGTTACCGCTCTCTCTTTGG
CHI3L1	GGCTTCTTCTGAGACTGGTGTTGG	CGCTTTCCTGGTCGTCGTATCC
COL3A1	TGTACCAGCCAGACCAGGAAGAC	TGTACCAGCCAGACCAGGAAGAC
FBLN2	GACCGAGGACAGTGAGGAGGAAG	CAGGCAGTGATGTGGACAGGATG
IGFBP5	GTACCTGCCCAATTGTGACC	AAGTCCCCGTCAACGTACTC
MATN2	AGAGGTGTGGGCTGTGGACTAC	GAGCACTGGCAGACGAAGGAATC
ITGA5	GTCGGGGGCTTCAACTTAGAC	ACAGAGGTAGACAGCACCAC
CILP	CTTTGAGAACCTCCGGGCAT	TCGATCCCCCTCAATCTGGT

C11orf96	TCCAGTTACCAGGCGGTGAT	TGCGTCTTGAAGCGAGACTG
A2M	GAGGCAGAAGGACAATGGCT	ATAGGCGGAGAGGGTCACTT
SFRP2	GCCCGACTTCTCCTACAAGC	CTCCTTCATGGTCTCGTGGC
MAF	CGTCCTCTCCCGAGTTTTTCA	GGCTTCCAAAATGTGGCGTA
PTGDS	CCATGTGCAAGTCTGTGGTG	CATGGTTCGGGTCTCACACT
TCF21 TCCTGGCTAACGACAAATACGA		TTTCCCGGCCACCATAAAGG

Supplemental Table 14. Sequences of siRNAs.

SiRNA	Sense (5'-3')	Antisense (5'-3')
siTCF21-1	GGAUUCGAACAAGGAAUUUTT	AAAUUCCUUGUUCGAAUCCTT
siTCF21-2	GCUAACGACAAAUACGAGATT	UCUCGUAUUUGUCGUUAGCTT
siITGB1-1	GAACAGAUCUGAUGAAUGATT	UCAUUCAUCAGAUCUGUUCTT
siITGB1-2	GUGGUUUCGAUGCCAUCAUTT	AUGAUGGCAUCGAAACCACTT

siITGB1-3	GAUCAUUGAUGCAUACAAUTT	AUUGUAUGCAUCAAUGAUCTT
siITGA2-1	CCCGAGCACAUCAUUUAUATT	UAUAAAUGAUGUGCUCGGGTT
siITGA2-2	GCUGGUGACAUCAGUUGUATT	UACAACUGAUGUCACCAGCTT
siITGA2-3	GUGGUUGUGUGAUGAAUTT	AUUCAUCACACACAACCACTT
siMATN2-1	GCAUCCUAAUCUUUGCCAUTT	AUGGCAAAGAUUAGGAUGCTT
siMATN2-2	GCAGUUUGUCACUGGAAUUTT	AAUUCCAGUGACAAACUGCTT

Supplemental table 15. Antibodies for Western blotting

Antibody	RRID	Company
TCF21	AB_10601215	SIGMA
COL1A2	AB_10679394	Abcam
N-cadherin	AB_1310479	Abcam
Vimentin	AB_10562134	Abcam
Slug	AB_777968	Abcam
FBLN2	AB_11153545	Invitrogen
FBLN1	AB_2553938	Invitrogen
Integrin α5	AB_2233962	Cell Signaling Technology
GAPDH	AB_10622025	Cell Signaling Technology
p-FAK ^{Tyr397}	AB_10891442	Cell Signaling Technology
FAK	AB_2799801	Cell Signaling Technology
$p\text{-PI3K }p85^{(Tyr458)}/p55^{(Tyr199)}$	AB_2895293	Cell Signaling Technology
PI3K	AB_2165248	Cell Signaling Technology
p-AKT ^{Ser473}	AB_2315049	Cell Signaling Technology
AKT	AB_2225340	Cell Signaling Technology
EpCAM	Cat. GB12274	Servicebio
E-cadherin	AB_2728770	Cell Signaling Technology
CHI3L1	Cat. AF8379	Beyotime
COL3A1	Cat. AF6531	Beyotime
Integrin alpha 2	Cat. AF2332	Beyotime
Integrin β1/CD29	Cat. AF0207	Beyotime
MMP2	Cat. GB11130	Servicebio
CD44	AB_2750879	Cell Signaling Technology

CD133	AB_2721172	Cell Signaling Technology	
Oct-4	AB_823583	Cell Signaling Technology	
HRP conjugated Goat Anti-	Cat. GB23303	Servicebio	
Rabbit IgG (H+L)	Cat. GB25305	Serviceolo	
HRP conjugated Goat Anti-	Cat. GB23301	Servicebio	
Mouse IgG (H+L)	Cat. GB25301		
HRP conjugated Donkey Anti-	Cat. GB23404	Servicebio	
Goat IgG (H+L)	Cat. GB23404	Serviceoro	

Supplemental table 16. Antibody list for immunofluorescence.

Antibody	RRID	Company
MATN2	AB_2811126	Abcam
VWF	AB_298501	Abcam
COL1A2	AB_10679394	Abcam
Laminin	AB_298179	Abcam
α-SMA	AB_2799045	Cell Signaling Technology
Integrin α5	AB_2233962	Cell Signaling Technology
TCF21	AB_10601215	SIGMA
CD31	AB_2161028	R&D Systems
CD45	AB_306361	Abcam
NG2	AB_11213678	MERK
MMP2	Cat. GB11130	Servicebio
CHI3L1	Cat. AF8379	Beyotime
COL3A1	Cat. AF6531	Beyotime

EpCAM	Cat. GB12274	Servicebio
Alexa Fluor 647-Donkey	AB 2535865	Invitrogen
anti-Sheep IgG (H+L)	AB_2333003	mvuogen
Alexa Fluor 546-Donkey	AB 2534016	Invitrogen
anti-Rabbit IgG (H+L)	AB_2334010	mvinogen
Alexa Fluor 488-Donkey	AB 2535792	Invitrogen
anti-Rabbit IgG (H+L)	NB_2333172	mvinogen
Alexa Fluor 488-Donkey	AB 2534102	Invitrogen
anti-Goat IgG (H+L)	122201102	m, mogen

Supplemental table 17. Antibody list for immunohistochemistry.

Antibody	RRID	Company
CD31	AB_2161028	R&D Systems
E-Cadherin	AB_2728770	CST
Vimentin	AB_10562134	Abcam
Ki67	Cat. GB11030	Servicebio
HRP- Goat Anti-Rabbit IgG (H+L)	Cat. GB23303	Servicebio
HRP -Goat Anti-Mouse IgG (H+L)	Cat. GB23301	Servicebio
HRP-Donkey Anti-Goat IgG (H+L)	Cat. GB23404	Servicebio

Supplemental materials for 1 Novel TCF21^{high} pericyte subpopulation promotes colorectal cancer metastasis 2 3 by remodelling perivascular matrix 4 5 Supplemental methods 6 Cell lines and cell culture 7 8 Human CRC cell lines HCT116, DLD-1, RKO, SW480, SW620, Caco-2, HT29 and human microvascular endothelial cell-1 (HMEC-1) were purchased from the American 9 Type Culture Collection (Manassas, VA). Mouse colon cancer cell line MC38 (Cat. 10 BNCC337716) was from BeNa Culture Collection (Beijing, China). HCT116, DLD-1, 11 RKO, SW480, SW620 and Caco-2 cells were cultured with DMEM. MC38 cells were 12 maintained in RPMI-1640. DMEM and RPMI-1640 medium were supplemented with 13 14 10% FBS (Cat. FSP500, ExCell Bio, Shanghai, China) and 1% penicillin-streptomycin (PS). HMEC-1 cells were cultured in endothelial cell medium (ECM, Cat. 1001, 15 Sciencell research laboratories, Corte Del Cedro Carlsbad, CA) supplemented with 5% 16 FBS, 1% endothelial cell growth supplement (ECGS), and 1% PS. All cell lines were 17 cultured at 37 °C in incubator with 5% CO2. MC38, HCT116 and DLD-1 cells were 18 19 infected with lentivirus harboring luciferase (Genechem, Shanghai, China) to generate the MC38-luc, HCT116-luc and DLD-1-luc cells, which were then selected with 20 21 puromycin (2 μg/mL) for 2 days. All cell lines were authenticated to have no crosscontamination using a STR Multi-amplification Kit (MicroreaderTM21 ID System) and 22 tested negative for mycoplasma by the TransDetect® PCR Mycoplasma Detection Kit 23 (Cat. FM311-01 Transgen, Beijing, China). 24 25

Human samples and specimens

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- 27 Human CRC surgical samples (12 cases, patients' information was listed in
- 28 Supplemental Table 5 and Supplemental Table 6) and specimens (75 cases, patients'
- 29 information was listed in Supplemental Table 1) were obtained from the First
- 30 Affiliated Hospital of Jinan University (Guangzhou, China).
- 32 Mice

- 33 Male C57BL/6JGpt mice, male BALB/c nude mice (4-6 weeks, 20-22 g), Rosa26-
- 34 CAG-LSL-Cas9-tdTomato mice (B6/JGpt-Rosa26^{tm1(CAG-LSL-Cas9-tdTomato)}/ Gpt; Cat.
- 35 T002249), Cspg4-CreERT2 mice (B6/JGpt-Cspg4^{em1Cin(CreERT2-P2A)/Gpt}; T006187), and
- 36 Tcf21-flox mice (B6/JGpt-Tcf21em1Cflox/Gpt; T013083) were obtained from
- 37 GemPharmatech Co., Ltd (Nanjing, Jiangsu, China). Pericyte lineage tracing mice
- 38 (PClin) were generated by crossing mice carrying a tamoxifen-inducible Cre
- 39 recombinase driven by the pericyte-specific *Cspg4* promoter (Tg^{Cspg4-CreERT2}) with mice
- 40 carrying a Cre-responsive reporter gene (tandem dimer Tomato (tdT)) inserted at the
- 41 ROSA26 locus (ROSA^{tdT/+}). The PC^{lin} mice were further crossed with mice harboring
- both *Tcf21* alleles flanked by LoxP sites (*Tcf21* flox/flox) to generate tamoxifen-inducible
- 43 Cspg4-driven pericyte-specific Tcf21 knockout mice (PClin-KO). All mice were
- 44 maintained in a specific pathogen-free (SPF) facility. Mouse genotyping was detected
- by PCR (The primer sequences were listed in **Supplemental Table 7**). Cre activity was
- 46 induced in tumor-bearing mice (6-7 weeks, weight 22-25 g) via oral gavage every other
- day for 3 times (10 mg/kg of tamoxifen in peanut oil). The animal experiments were
- 48 complied with the ARRIVE Guidelines 2.0: updated guidelines for reporting animal
- 49 research¹.

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Isolation and culture of TPCs

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TPCs were isolated from CRC patients though a microdissection combined with pericyte medium-based approach that developed by our lab. Briefly, fresh surgical tumor specimens were obtained from CRC patients with or without liver metastasis. Information of the CRC patients was listed in **Supplemental Table 5 and Supplemental Table 6.** Tumor tissues were kept in serum-free DMEM containing PS and placed on the ice, and then washed with pre-chilled PBS in a sterile hood to remove the blood, adipose tissues. Tumor vessels were separated from perivascular adipose tissues under a stereomicroscope (Olympus, SZX7). The acquired tumor vessels were cultured in Pericyte Medium (PM, Cat. 1201, Corte Del Cedro Carlsbad, CA, USA, Sciencell research laboratories) with 5% FBS, 1% PGS and 1% PS at 37 °C with 5% CO₂. TPCs were migrated from the tumor vessels within 14 days, which were then disassociated by trypsin once the confluence reaches 80%. The purity of the isolated TPCs were authenticated by STR Multi-amplification Kit (**Supplemental Table 8**).

Construction of single cell cDNA libraries

- For single cell cDNA libraries construction, passage 1 TPCs derived from four CRC patients (patient information was listed in **Supplemental Table 9**) were prepared and
- analyzed by a 10×Genomics GemCode Single-cell instrument, generating single-cell
- 70 Gel Bead-In-EMlusion (GEMs). The libraries were generated and sequenced by
- 71 Chromium Next GEM Single Cell 3' Reagent Kits v3.1 and Illumina HiSeq 4000 by
- 72 Genedenovo Biotechnology Co., Ltd (Guangzhou, China) with a custom paired-end
- 73 sequencing mode 26 bp (read 1) \times 98 bp (read 2).

Bioinformatic analysis of scRNA-seq

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Reads uniquely mapped to the transcriptome and intersecting an exon at least 50% were considered for UMI counting. Before quantification, the UMI sequences would be corrected for sequencing errors, and valid barcodes were identified based on the EmptyDrops method. The cells by gene matrices were produced via UMI counting and cell barcodes calling. The cells by gene matrices for each sample were individually imported to Seurat version 3.1.1 for downstream analysis. After removing the unwanted cells from the dataset, data normalization and batch effect correction, the integrated expression of matrix was then scaled and performed on principal component analysis (PCA) for dimensional reduction, those had a strong enrichment of low P-value genes for downstream clustering were identified as significant PCs. Seurat implemented a graph-based clustering approach. Distances between the cells were calculated based on previously identified PCs. For visualization of clusters, t-distributed Stochastic Neighbor Embedding (t-SNE) were generated using the same PCs. Expression value of each gene in given clusters were compared against the rest of cells using Wilcoxon rank sum test. Significantly upregulated genes were identified using several criteria. First, genes had to be at least 1.28-fold overexpressed in the target cluster. Second, genes had to be expressed in more than 25% of the cells belonging to the target cluster. Third, P value is less than 0.05. The Gene ontology (GO) enrichment analysis was performed with the GO database (http://www.geneontology.org/). GO has three ontologies: molecular function, cellular component, and biological process. The calculated P-values were false discovery rate (FDR)-corrected, taking FDR ≤ 0.05 as a threshold. GO terms meeting this criterion were defined as significantly enriched GO terms in differentially expressed genes.

Analysis of transcription factor network inference was performed with the SCENIC R package². In brief, log-normalized expression matrix generated using Seurat was used as input, and the pipeline was implanted in three steps. First, gene coexpression network was established via GENIE3³. Second, each module was pruned based on a regulatory motif near a transcription start site via RcisTarget. Precisely, the networks were retained if the transcription factor (TF)-binding motif was enriched among its targets, while target genes without direct TF-binding motifs were removed. The retained networks were called regulons. Third, the activity of each regulon in each single cell was scored (AUC score) using AUCell R package. Gene regulatory network (GRN) plots of all regulons were done using the cytoscape software⁴.

Analysis of the public datasets

scRNA-seq data of colon (GEO accession GSM3140596, GSM3140595, GSM3140594, and GSM3140593)⁵ and intestine (GEO accession GSM4159165 and GSM4159164)⁶ acquired using 10× Chromium protocol were download, and the sequencing reads were realigned, and cell clustering was performed as described above.

Construction of MC38-luc-LM3, HCT116-luc-LM3 cells and DLD-1-luc-LM3

cells

To establish highly metastatic MC38-luc-LM3 cells, MC38-luc cells (1×10^5) suspended in 100 μ L of Matrigel (Cat. 354248, Corning, NY) were injected into the spleen of male C57BL/6JGpt mice anesthetized with isoflurane inhalation. Liver metastasis was detected by bioluminescence imaging. MC38-luc-LM1 cells from the metastatic foci were isolated by mouse tumor dissociation kit (Cat. 130-096-730, Miltenyi Biotec, Bergisch Gladbach, Germany) and cultured in complete RPMI-1640 medium and

selected by puromycin (2 µg/mL). MC38-luc-LM1 cells were inoculated into the spleen of male C57BL/6JGpt mice, and MC38-luc-LM2 cells were obtained from the liver metastatic foci. Tumor cells isolated from the third round of liver metastatic foci were termed MC38-luc-LM3 cells, which were employed for the subsequent experiments. The HCT116-luc-LM3 and DLD-1-luc-LM3 cells were isolated by human tumor dissociation kit (Cat. 130-095-929, Miltenyi Biotec) and acquired with BALB/C nude mice by the same pattern of MC38-luc-LM3. The cellular morphology, nucleus size, cell size, cell migration, proliferation, EpCAM expression, stemness and EMT were assessed to evaluate the phenotypical/biological differences between the parental cells and the LM3 cells (Supplemental Figure 20). The origin of all LM3 cells was further validated by short tandem repeat (STR) (Supplemental table 10-12) and luciferase activity (Supplemental Figure 21).

Flow cytometry

Cells were collected, re-suspended in flow cytometry staining buffer, and distributed into 1.5 mL EP tubes. Following fixation with 4% paraformaldehyde on ice and permeabilization with 0.1% Triton X-100 in PBS for 5 min, the cells were incubated with anti-TCF21 antibody (Cat. AB_182134, Abcam) or anti-MATN2 antibody (Cat. AF3044, R&D system,) for 1 h on ice. Then, cells were washed with PBS twice and incubated with Donkey anti-Rabbit IgG (H+L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor 546 (Cat. AB_2534016, Invitrogen, Carlsbad, CA, USA) or Donkey anti-Goat IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor 488 (Cat. AB_2534102, Invitrogen) for 1 h and analyzed by flow cytometry (BD Biosciences, San Jose, CA). The size of HCT116-luc, DLD-1-luc, MC38-luc, HCT116-luc-LM3, DLD-1-luc-LM3 and MC38-luc-LM3 cells was directly evaluated by forward

scatter using flow cytometry

Animal studies

MC38-luc-LM3 cells (1×10^5) suspended in 100 μ L of Matrigel were orthotopically injected into the cecum wall of PC^{lin} mice and PC^{lin-KO} mice anesthetized with isoflurane inhalation. At the end of the experiment, tumors were collected and subjected to immunohistochemistry and immunofluorescence analysis. The livers were harvested, photographed, and prepared for H&E staining. For co-injection assays, HCT116-luc-LM3, DLD-1-luc-LM3 cells, TPC_{NM} transfected with lentivirus harboring negative control shNC (TPC_{NM} shIC) or shITGA5 (TPC_{NM} shITGA5), TPC_{LM} transfected with Vector (TPC_{LM} Vector) or lentivirus expressing ITGA5 (TPC_{LM} ITGA5) were collected. HCT116-luc-LM3 or DLD-1-luc-LM3 cells (4×10^5) were premixed with TPCs (1.6×10^6) in 100 μ L of Matrigel, which was then injected into the cecum wall of BALB/C nude mice. At the end of the experiment, the mice were sacrificed with CO₂ and the metastatic foci in mouse liver were analyzed by H&E staining. Orthotopic tumor tissues were obtained for Masson staining, immunohistochemical staining, immunofluorescence, and transmission electron microscope analysis.

In vivo cell tracking

169 For the whole animal imaging *in vivo*, mice were intraperitoneal (i.p.) injected with 3

mg of D-luciferin (Cat. 40901ES01, Yeason Biotechnology, Shanghai, China)

dissolved in 200 µL saline and were anesthetized by isoflurane after injection for 5 min.

Luminescence signals were collected with Xenogen IVIS 200 (Alameda, CA, USA)

and analyzed by the Xenogen Living Image software (Alameda, CA, USA).

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Vessel permeability assay

- 176 PClin and PClin-KO mice bearing MC38 orthotopic xenografts were intravenously (i.v.)
- injected with 1 mg of FITC-labeled Dextran-40 kDa (Cat. D1845, Thermo Scientific)
- for 10 min. Then, the mice were perfused with 4% PFA and tumors were obtained and
- then frozen. Tumor tissues were sectioned, and tumor vessels were stained for CD31
- 180 (RRID: AB_2161028, RD, Minneapolis, MN, USA) followed by Donkey anti-Goat
- 181 IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa FluorTM 647 (RRID:
- 182 AB_2534102, Invitrogen, Carlsbad, CA, USA), and the double staining of FITC-
- labeled Dextran over CD31-positive vessels indicates vessel permeability.

Isolation and identification of circulating tumor cells (CTCs)

- 186 CTC isolation was performed according to a previous study⁷. Briefly, blood (500 μL)
- was collected from each of PClin and PClin-KO mice bearing MC38-luc-LM3 allografts
- 188 by cardiac puncture and immediately released into heparin-coated tube to avoid
- coagulation. The red blood cells were removed by blood red cell lysing reagent before
- cells were seeded in a 12-well plate and cultured with DMEM supplemented with 20%
- FBS. The adherent tumor cells were identified and counted within 12 h prior to no
- 192 tumor cell growth. Adherent cells were stained with cancer cell-associated surface
- marker EpCAM and leukocyte marker CD45 and identified by confocal microscopy as
- described previously^{8, 9}. Cells positive for EpCAM but not CD45 were scored as CTCs
- and subsequently subjected to manual counting, and the CTC counts were presented as
- 196 CTCs per milliliter of whole blood.

Isolation efficiency of CTCs

To determine the isolation efficiency of CTCs¹⁰⁻¹², 500 MC38-luc-LM3 cells were

spiked into 500 μL of blood collected from the healthy C57BL/6JGpt mice by cardiac puncture. The spiked blood was then treated with blood red cell lysing reagent and the remaining cells were seeded on a 12-well plate and cultured with DMEM containing 20% FBS. The adherent tumor cells were stained with EpCAM and CD45 and the EpCAM⁺CD45⁻ CTCs were identified by confocal microscopy. The number of EpCAM⁺CD45⁻ CTCs was counted within 12 h at a time of no tumor cell growth. The efficiency of CTC recovery was calculated using the following equation: Cell recovery (%) = counts of isolated MC38-luc-LM3 cells/500 × 100%.

Chromatin immunoprecipitation (ChIP) and ChIP-Seq

ChIP assay was performed according to the manufacture manual of SimpleChIP® Enzymatic Chromatin IP Kit (Cat. 9003, Cell Signaling Technology, MA, USA). Briefly, TCF21-overexpressing TPCs (TPC_{NM}^{TCF21}) were washed twice in clod PBS buffer and cross-linked with 1% formaldehyde for 10 min at room temperature and then stopped by addition of glycine (125 mM). Afterwards, samples were lysed, and chromatins were obtained on ice. Chromatins were then sonicated to get soluble sheared chromatin (average DNA length of 150-900 bp). Then, 20 μ L of chromatin was saved as input and 100 μ L of chromatin was harvested for immunoprecipitation by anti-TCF21 antibody (RRID: AB_10601215, Sigma, Shanghai, China), and anti-IgG was served as the negative control. 10 μ g of anti-TCF21 was used in the immunoprecipitation reactions at 4 °C overnight. Then 30 μ L of protein A beads was added and the samples were further incubated for 2 h. After reverse cross-linking and DNA purification, immunoprecipitated DNA was quantified by real-time PCR. Immunoprecipitated DNA was used to construct sequencing libraries following the protocol provided by the NEXTflex® ChIP-Seq kit (Cat. NOVA-5143-02, BioScientific, TX , USA) and

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sequenced on Illumina Xten with PE 150 method (LC-Bio Technology CO., Ltd., Hangzhou, China). For data analysis, Trimmomatic (version 0.38) was used to filter out low-quality read. MACS2 software (version 2.1.1.20160309) was used to call peaks by default parameters (bandwidth, 300 bp; model fold, 5, 50; q value, 0.05). If the summit of a peak located closest to the TSS of one gene, the peak will be assigned to that gene. GO enrichment analysis was performed using the EasyGO gene ontology enrichment analysis tool (http://bioinformatics.cau.edu.cn/easygo). The GO term enrichment was calculated using hypergeometric distribution with a *P* value cutoff of 0.01. *P* values obtained by Fisher's exact test were adjusted with FDR for multiple comparisons to detect overrepresented GO terms.

236 RT-qPCR assay

- Total RNA was collected by E.Z.N.A.® Total RNA Kit I (Cat. R6834-02, Omega Bio-
- 238 Tek, Norcross, GA, USA). The purity and concentration of RNA was examined by
- Nanodrop Lite micro spectrophotometer. RNA (2 µg) was reversely transcribed to
- 240 cDNA with All-in-One cDNA Synthesis SuperMix (Cat. B24408-1000, Bimake,
- 241 Houston, TX, USA). Reverse transcription quantitative PCR (RT-qPCR) was
- 242 performed in triplicate using 2× SYBR Green qPCR Master Mix (Cat. B21202,
- Bimake). Samples were loaded into a Roche LightCycler 480 II real-time polymerase
- chain reaction detection system (Roche, Basel, Switzerland) and the data is analyzed
- by $2^{-\Delta \triangle^{Ct}}$ method. The primer sequences were listed in **Supplemental Table 13.**

Cell infection and transfection

- 248 TPCs derived from CRC patients with non-metastasis (TPC_{NM}) were infected with
- lentivirus harboring TCF21 overexpression plasmid for 48 h and selected by puromycin

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listed as follows: TCF21 (NM_198392) Human Untagged Clone (Cat. RC220002, Origene. Rockwell, MD, USA), Cloning vector pCMV6-Entry (Cat. PS100001, Origene). For TCF21- or MATN2-knockdown experiments, TPCs derived from CRC patients with liver metastasis (TPC_{LM}) were transfected with siRNA for 48 h followed by subsequent analysis. For ITGA2- and ITGB1-knockdown experiments, TPC_{NM} were transfected with siRNA for 48 h followed by subsequent analysis. Transfection was performed with Lipofectamine™ 3000 (Cat. L3000015, Invitrogen, Carlsbad, CA, USA) according to the manufacturer instructions, and the siRNA sequences were listed in Supplemental Table 14. For MATN2-overexpression experiments, TPC_{NM} were transfected with pCMV6-MATN2-overexpressing plasmid (Cat. RC203833, Origene) or pCMV6-Entry as empty vector for 48 h (Cat. PS100001, Origene). For ITGA5 overexpression experiments, TPC_{LM} were infected with lentivirus harboring ITGA5 or its corresponding Vector (pGC-FU-3FLAG-CBh-gcGFP-IRES-puromycin) (Genechem, Shanghai, China). For ITGA5-knockdown experiments, TPC_{NM} were infected with lentivirus harboring shITGA5 or pFU-GW-016 as Vector (Genechem, Shanghai, China).

(2 mg/mL). Detailed information of TCF21 overexpression plasmid and vector was

Y15 and SGI1027 treatment

Y15 and SGI1027 were purchased from Selleck (Shanghai, China) and dissolved in DMSO. Integrin $\alpha 5$ -overexpressing TPCs (2×10^5) were seeded into 6-well plates and cultured overnight. The next day, cells were treated with Y15 (5 μ M) or SGI1027 (2.5 μ M) for 24 h, and then cells were applied for Western blotting assay and bisulfite sequencing.

Western blotting assay

Cells were lysed in RIPA lysis buffer on ice for 30 min. Total protein concentration was measured with PierceTM BCA Protein Assay Kit (Cat. 23225, Thermo Scientific, Waltham, MA, USA). Equal amounts of protein (20 μg) were separated in SDS-PAGE gel (Cat. G2004, Solarbio, Beijing, China) and transferred onto polyvinylidene fluoride (PVDF) membranes (Cat. IPVH00010, Millipore, Boston, MA, USA). Following blocking with 5% BSA, the membranes were incubated with indicated antibodies. The blots were detected by Amersham Imager 600 (GE, Boston, MA, USA). The antibodies were listed in **Supplemental Table 15**.

Immunofluorescence analysis

Tissue slices were deparaffinized and incubated with 1×Tris-EDTA (pH 9.0) and 0.05% Tween for 3 min for antigen retrieval. After that, tumor sections were permeabilized in 0.1% TritonTM X-100, blocked with QuickBlockTM immunostaining blocking solution (Cat. ST797, Beyotime, Shanghai, China) and incubated with the corresponding primary antibody overnight at 4 °C. Then, the sections were incubated with the corresponding secondary antibody for 1 h at room temperature. For nucleus staining, sections were incubated with 1 μg/mL DAPI (Cat. MBD0015, Sigma) for 15 min. The slides were photographed with a Zeiss LSM 800 confocal microscope and analyzed with Image J software (RRID: SCR_003070, Rawak Software Inc., Stuttgart, Germany). The primary and secondary antibodies were listed in **Supplemental Table 16.** For phalloidin immunofluorescence assay, HCT116-luc, DLD-1-luc, MC38-luc, HCT116-luc-LM3, DLD-1-luc-LM3 or MC38-luc-LM3 cells were plated on the glass bottom cell culture dish and incubated with DMEM for 24 h. The next day, cells were fixed, permeabilized with 0.1% TritonTM X-100 and then incubated with Alexa FluorTM 594-

phalloidin (Cat. A12381, Thermo) for 1 h. Cell nucleus were stained with 1 μ g/mL DAPI for 15 min. The cytoskeleton elements were photographed with a Zeiss LSM 800 confocal microscope.

H&E staining, immunohistochemistry, and Masson staining

Fixed tissues were embedded in paraffin and sectioned (5 μm). Following deparaffinized, the sections were subjected to antigen retrieval procedures with an EDTA antigen retrieval solution (Cat. P0086, Beyotime, Shanghai, China). Then, the slides were incubated with hematoxylin followed by counterstaining with eosin. For immunohistochemistry assay, tumor sections were incubated with primary antibodies overnight at 4 °C followed by incubation with HRP-conjugated secondary antibodies. The primary and secondary antibodies were listed in **Supplemental Table 17.** Protein expression in tumor sections was detected using a DAB kit (Cat. G1212, Servicebio, Wuhan, Hubei, China), followed by counterstaining with hematoxylin (Cat. G1004, Servicebio, Wuhan, Hubei, China). Images were acquired with an Olympus BX 53 microscope and analyzed with Image J software. For Masson staining, tissue sections were prepared with Masson Tricolor Staining Solution (Fast Green Method) kit (Cat. G1343, Solarbio, Beijing, China). Images were acquired with Olympus BX 53 microscope and analyzed with Image J software.

RNA sequencing analysis

Total RNA was isolated and purified by TRIzol reagent (Cat. 15596018, Invitrogen, Carlsbad, CA, USA) following the manufacturer manual. The RNA concentration and integrity were evaluated by NanoDrop ND-1000 (NanoDrop, Wilmington, DE, USA) and Bioanalyzer 2100 (Agilent, CA, USA). Then, poly (A) RNA was purified from 1µg

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total RNA by Dynabeads Oligo (dT) 25-61005 (Thermo Fisher, CA, USA) using two rounds of purification. Then the poly(A) RNA was cut into pieces using Magnesium RNA Fragmentation Module (Cat. e6150, NEB, NY, USA) under 94 °C for 5-7 min. 328 Then the fragmented RNA pieces were reversely transcribed into cDNA by 329 SuperScriptTM II Reverse Transcriptase (Cat.18064022, Invitrogen, USA) and sequenced with illumina Novaseq[™] 6000 (LC-Bio Technology CO., Ltd., Hangzhou, 330 China). Then, StringTie and edgeR were used to evaluate the expression levels of all transcripts. The differentially expressed mRNAs and genes were picked with log2 (fold 333 change) >1 or log2 (fold change) <-1 and with statistical significance (P value < 0.05) by R package-edgeR. The volcano plot revealed the distributions of log2 fold change and P values for the differentially expressed genes. The GO terms (http://www.geneontology.org) of these differentially expressed genes were annotated.

Migration and invasion assay

Migration assay was performed with 24-well Boyden chambers (Corning, NY, USA) containing inserts of polycarbonate membranes with 8 µm-pores. Cells suspended with 100 μL of serum-free medium were seeded in the upper compartment $(3\times10^4 \text{ HCT}116$ cells or 2×10^4 TPCs). The bottom chamber was filled with different chemoattractants. For invasion assay, the upper chamber was pre-coated with 30 µL of Matrigel (diluted at 3:1 using PBS) and incubated for 30 min. Then, PKH67-labeled HCT116 cells or DLD-1 cells (5×10^3) mixed with TPCs (2.5×10^4) were seeded into the upper chamber. The bottom chamber was filled with PM and DMEM (5:1). Following incubation for 48 h, the upper chamber was fixed with 4% paraformaldehyde for 30 min and then the cells were stained with 0.1% crystal violet. The non-migrated cells on the upper side of the membrane were removed with a cotton swab. The cells remaining on the lower

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surface were photographed under an inverted microscope and analyzed with Image Pro Plus 6 software. Collagen gel contraction assay This experiment was performed with Cell Contraction Assay kit (Cat. CBA-5020, CELL BIOLABS, San Diego, CA, USA). TCF21-overexpressing or -knockdown TPCs were harvested and resuspended in PM at 5×10^6 cells/mL. Cold collagen gel working solution was prepared according to the instructions and mixed with the cell suspension at a ratio of 4: 1. 0.5 mL of the cell-collagen mixture per well was added in a 24-well plate. After incubating 1 h at 37 °C, 1.0 mL culture medium was added into the collagen gel. Cultures were incubated for two days, and the collagen gels were gently released from the sides of the culture dishes with a sterile spatula. The collagen gel size (contraction index) was measured at 0, 6, 12, 24 and 48 h and quantified with Image J. Cell proliferation assay Cells (5×10^3) were cultured overnight in 96-well plates. The next day, cells were treated with or without the culture supernatant of TPCs (48-h culture medium) and cell proliferation was determined by BeyoClick™ EdU Cell Proliferation kit (Cat. C0071S, Beyotime, Shanghai, China) and analyzed with Image Pro Plus 6 software. Adhesion assay TPCs with the overexpression or knockdown of TCF21 were collected, washed, and stained with PKH67 (Cat. MINI67, Sigma). PKH67-labeled TPCs (2×10⁴) were seeded in a 96-well plate for 2 h. Then, the media was removed, and cells were washed with

374 PBS twice to remove the non-adherent cells. Images were acquired with a fluorescent 375 microscope (ZEISS) and analyzed with Image Pro Plus 6 software. 376 377 **Tube formation assay** 378 Tube formation was performed with a 96-well plate. Matrigel was first coated in the plates at 37 °C for 30 min. Then, HMEC-1 cells (2×10⁴) supplemented with 100 μL 379 380 ECM were seeded in the Matrigel-coated plates. After 2-h incubation, ECM was 381 replaced, and HMEC-1 cells were incubated with the conditioned medium of TPCs. 382 The capillary tubes were photographed under an inverted light microscopy, and the 383 number of tubes was analyzed by Image Pro Plus 6 software. 384 385 Transmission electron microscopy analysis 386 Tumor tissues were acquired and fixed in 2.5% glutaraldehyde (Cat. PH9003, Maya 387 Reagent, Zhejiang, China). All samples were post-fixed in 1% osmium tetroxide (Cat. 388 23311-10, Polysciences, USA), dehydrated in graded concentration of alcohols, and 389 then embedded in low-viscosity resin. The embedded tissues were sectioned and stained 390 with saturated uranyl-acetate and Sato's lead-citrate. Sections were imaged using 391 JEM1200EX transmission electron microscope equipped with BioScan600W digital 392 camera (JEOL, Tokyo, Japan). 393 394 Second harmonic generation and two-photon excited fluorescence (SHG/TPEF) 395 Tumor tissues were acquired and fixed in 4% paraformaldehyde overnight followed by 396 washing with PBS twice and sectioned (250 µm) using Vibration slice (Leica, 397 VT1000S). Following blocking with 5% BSA solution for 1 h, the sections were incubated with anti-CD31 antibody at 4 °C overnight. The slides were then incubated 398

with Donkey anti-Goat IgG (H+L) Cross-Adsorbed Secondary Antibody, Alexa Fluor™ 488 (RRID: AB_2534102, Invitrogen) and immersed in PBS for SHG/TPEF microscopy (Nikon, Tokyo, Japan). TPEF was utilized for visualization of tumor vessels stained by CD31 (red) and SHG was used to visualize collagen structure (green) at 790 nm excitation light.

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Atomic Force Microscopy (AFM) measurement

For collagen organization, coatings of isotropic collagen I and AFM detection were performed according to previous report¹³. In vitro fibrillogenesis of collagen-I (Cat. 08-115, Merck; 4.4 mg/mL) was initiated by diluting the collagen I solution tenfold in fibrillogenic buffer (50 mM glycine, 200 mM KCl, pH 9.0). After mixture for 10 min at room temperature, 60 µL of the diluted solution were added to the 22 mm-silicide coverslips and incubated overnight at 37 °C. The next day, coverslips were washed with PBS twice and plated in 12-well plate. Then, TPCs (1×10^4) were seeded on the Collagen I-coated coverslip and cultured for 5 days. AFM Imaging was performed with a NanoWizard II AFM (JPK-Instruments, Berlin, Germany) mounted on an inverted microscope (Axiovert 200, Zeiss, Jena, Germany). Scanning of samples was performed at a scan rate of 0.25 Hz and five fields were recorded for each sample. For perivascular stiffness measurements, AFM was performed according to modified published procedures¹⁴. Tissues were acquired and embedded within OCT. Then, the frozen tissue blocks were sectioned at a thickness of 20 µm and immersed in proteinase inhibitorcontained PBS at room temperature. The next day, tumor sections were applied for AFM quantification of Young's modulus (Bruker, USA). Briefly, silicon nitride cantilevers with a spring constant of 0.15 N/m were attached by a borosilicate glass spherical tip with a diameter of 5 µm. Cantilevers were tapping on the perivascular

region of tumor sections and five 15 μ m \times 15 μ m AFM stiffness map (16 \times 16 raster series) were acquired for each sample. The Young's modulus of the perivascular region in each section were determined by Hertz model. Tissue samples were assumed to be incompressible and a Poisson's ratio of 0.5 was used in the calculation of the Young's modulus.

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DNA extraction and bisulfite sequencing

- 431 DNA was extracted using the Genomic DNA Purification Kit (Cat. A1120, Promega,
- 432 WI, USA), which was followed by treatment with sodium bisulfite (Zymo Research,
- 433 CA, USA). The converted DNA was purified and amplified for sequencing by Biossci
- Biotechnology Co. Ltd (Wuhan, Hubei, China). Primers of TCF21 promoter bisulfite-
- 435 modified regions were: Forward primer (5'-3'): TTTTTGATGTTTTGAAAATGATT -
- 436 AGG; Reverse primer (5'-3'): CAACCACCTTC TCCCAACTATAA.

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Organotypic culture system

- 439 Organotypic culture system was constructed with 12-well Boyden chambers (Corning,
- NY, USA) containing inserts of polycarbonate membranes with 0.4 μm-pores. HMEC-
- 1 cells (2×10^4) supplemented with 100 µL ECM were seeded in the upper chamber and
- 442 incubated overnight. The next day, ECM was removed, and TPCs (5 \times 10⁵) were
- 443 embedded in 1.0 mL collagen I and plated in the chamber. The lower chamber was
- 444 filled with complete PM and ECM (PM: ECM, 5:1). Following incubation at 37 °C, 5%
- 445 CO₂ for 5 days, the mechanical properties of the matrix, complex modulus (G*), were
- determined with a rheometer (Malvern Kinexus pro+, USA) according to previous
- report¹⁵. The elastic modulus (E) was determined from G* by assuming a poison's ratio
- (v) of 0.5 with the expression E=2G*(1+v) to allow comparison to other published

work. For invasion assay, TCF21-overexpressing TPCs (5×10^5) were embedded in a matrix mixture of 0.25 mL collagen I and 0.75 mL Matrigel and plated in the chamber. Following incubation for 5 days, PKH67-labeled HCT116 cells (3×10^4) supplemented with 100 μ L complete DMEM were plated on the top of matrix and further cultured for 1 day. At the end of experiment, the whole matrix was fixed in 4% overnight and subjected to immunofluorescence analysis. The migrated HCT116 cells were detected by staining of EpCAM. The invaded cells were observed under a Zeiss LSM 800 confocal microscope and analyzed with Image Pro Plus 6 software.

Statistical analysis

The statistical values were calculated with GraphPad Prism 8.0 (GraphPad Software, Inc., San Diego, CA). Differences between two groups were evaluated with two-tailed unpaired t-test or Mann Whitney test. Differences among three groups or more were evaluated using one-way ANOVA followed by Tukey's post hoc test. Survival curves were plotted using the Kaplan Meier method and compared using the log-rank test. The receiver operating characteristic (ROC) curves were performed and the area under ROC curve (AUC) was calculated by logistic regression model to evaluate the diagnostic accuracy. Comparisons of variables were performed using Fisher's exact test or chisquared test based on their categorical data. Multivariable logistic regression was used to analyze the predictors of CRC metastasis. P < 0.05 was considered as significant difference.

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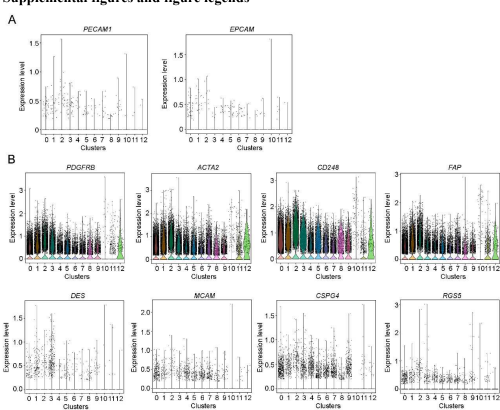
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Supplemental figures and figure legends



Supplemental Figure 1. Transcriptomic characterization of 13 subsets of TPCs. (A)

Gene expression profiles of *PECAM1* and *EPCAM* in distinct subsets of TPCs. **(B)** Gene expression profiles of *PDGFRB*, *ACTA2*, *CD248*, *FAP*, *DES*, *MCAM*, *CSPG4*, and *RGS5* in distinct subsets of TPCs.

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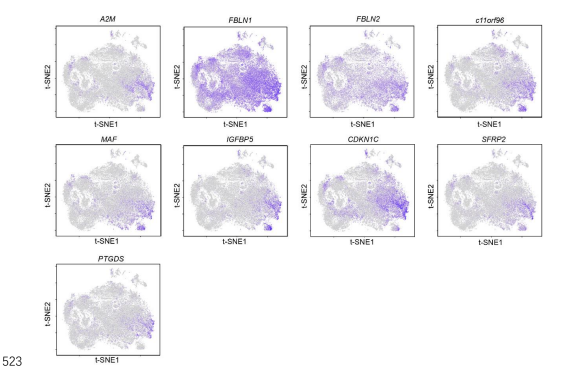
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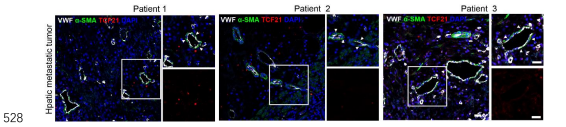
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Supplemental Figure 2. Distribution of the matrix-pericyte-related genes in all subsets of TPCs. t-SNE visualization of gene distribution of A2M, FBLN1, FBLN2, c11orf96, MAF, IGFBP5, CDKN1C, SFRP2 and PTGDS in all subsets of TPCs.



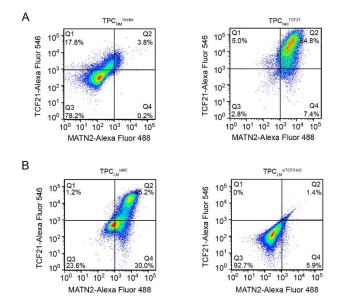
Supplemental Figure 3. Determination of TCF21 in TPCs in the hepatic metastatic tumors from CRC patients. Representative images of TCF21 staining (red) in TPCs (αSMA^+ , green) in the hepatic metastatic tumors from CRC patients (n = 20). Scale bar, 20 μm .

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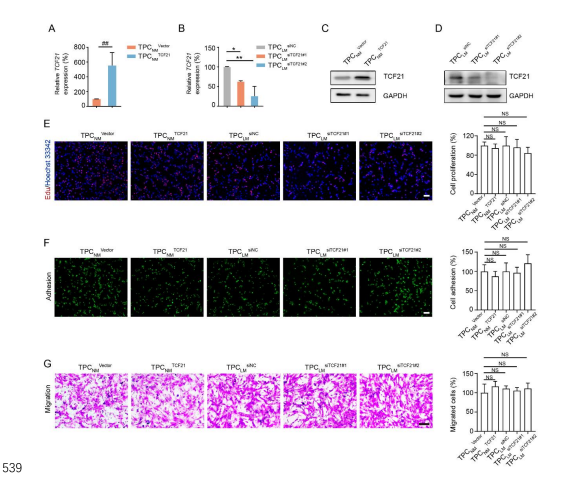
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Supplemental Figure 4. The expression of TCF21 is positively correlated with MATN2 in TPCs. (A) FCM analysis of the TCF21⁺MATN2⁺ TPCs in TPC_{NM} infected with TCF21 lentivirus (TPC_{NM}^{TCF21}) or Vector (TPC_{NM}^{Vector}) (n = 3). (B) FCM analysis of the TCF21⁺MATN2⁺ TPCs in TPC_{LM} transfected with siRNA targeting TCF21 (TPC_{LM}^{siTCF21}) or negative control (TPC_{LM}^{siNC}) (n = 3).



Supplemental Figure 5. Effects of TCF21 on the proliferation, adhesion, and migration of TPCs. (A, B) RT-qPCR analysis of TCF21 mRNA levels in TCF21-overexpressing (A) or -knockdown (B) TPCs (n = 3). (C, D) Western blotting analysis of TCF21 in TCF21-overexpressing (C) or -knockdown (D) TPCs (n = 3). (E) Representative images and quantification of cell proliferation in TCF21-overexpressing and -knockdown TPCs (n = 3). Scale bar, 100 μ m. (F) Representative images and quantification of TPC adhesion (n = 3). Scale bar, 100 μ m. (G) Transwell assay for the migration of TPCs. Quantification of the migrated TPCs is shown (n = 3). Scale bar, 200 μ m. Data are presented as mean \pm SEM. NS, not significant. *#P < 0.01 by two-tailed unpaired *t*-test; NS, *P < 0.05, **P < 0.01 by one-way ANOVA followed by Tukey's post hoc test.

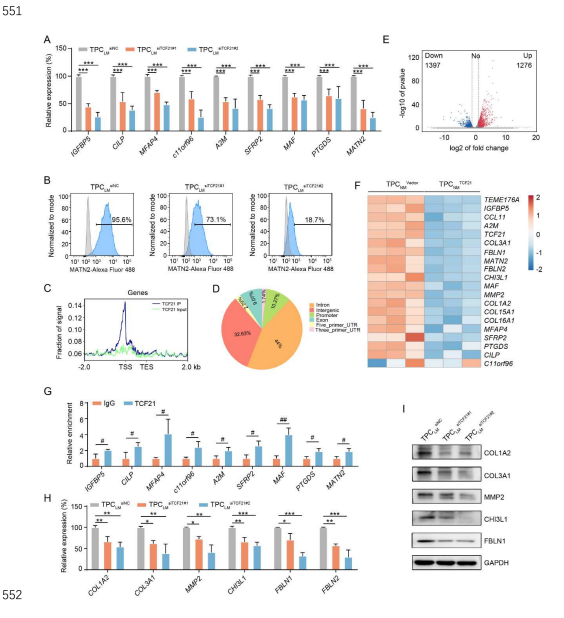
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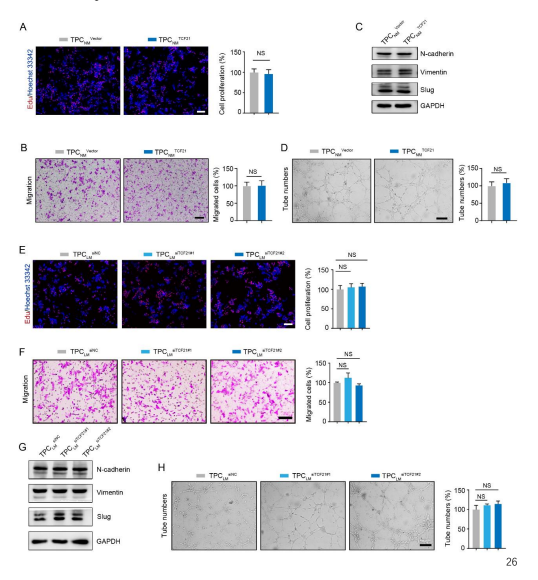
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Supplemental Figure 6. TCF21 stimulates the generation of matrix-pericytes and induces ECM remodeling. (A) RT-qPCR analysis of matrix-pericyte-specific genes in TPC_{LM} transfected with siRNA targeting TCF21 (TPC_{LM}^{siTCF21}) or negative control (TPC_{LM}^{siNC}) (n = 3). (B) FCM analysis of MATN2 expression in TCF21-knockdown TPCs (n = 3). (C) ChIP-seq summary plot of TCF21 enrichment across the indicated genomic distance in TCF21-overexpressing TPCs (n = 3). (D) The distribution of TCF21 peaks on gene elements (n = 3). (E) Volcano Plot of TCF21 regulated genes (n

= 3); Red dots represent the up-regulated genes and blue dots represent the down-regulated genes. **(F)** Heat maps of the differentially expressed genes between TPC_{NM}^{Vector} and TPC_{NM}^{TCF21} (n = 3). **(G)** ChIP-qPCR analysis of TCF21 binding at the promoter of indicated genes in TPC_{NM}^{TCF21} (n = 3). **(H)** RT-qPCR analysis of differentially expressed genes in TPC_{LM}^{siNC} and TPC_{LM}^{siTCF21}. **(I)** Western blotting analysis of COL1A2, COL3A1, MMP2, CHI3L1, and FBLN1 in TPC_{LM}^{siNC} and TPC_{LM}^{siTCF21} (n = 3). Data are presented as mean \pm SEM, *P < 0.05, *P < 0.01, *P < 0.001 by one-way ANOVA followed by Tukey's post hoc test, *P < 0.05, *P < 0.01 by two-tailed unpaired t-test.



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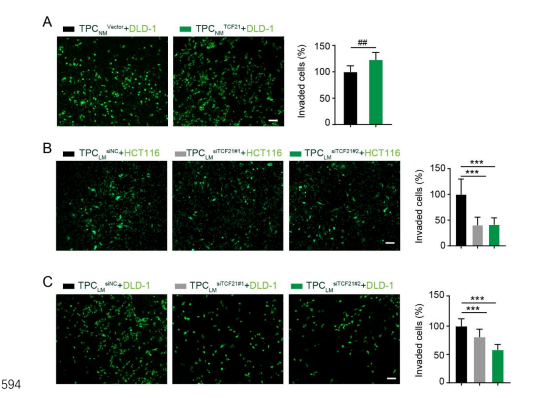
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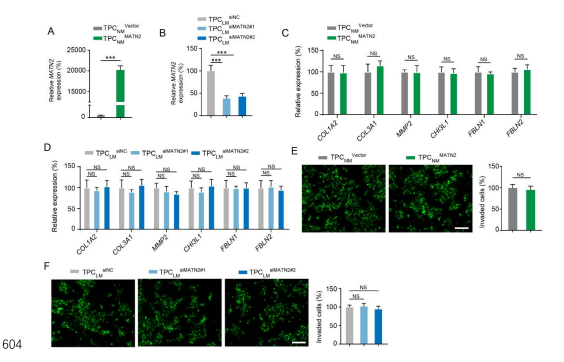
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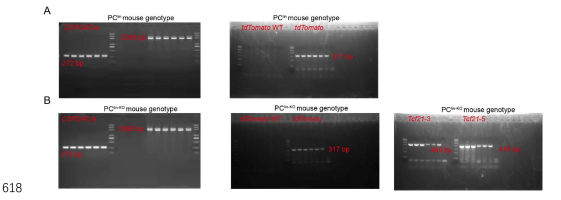
Supplemental Figure 7. TCF21 in TPCs has negligible effects on cell migration and angiogenesis. (A) EdU assay for the proliferation of HCT116 cells primed with conditioned medium from TPC_{NM}^{Vector} and TPC_{NM}^{TCF21} for 48 h (n = 3). Scale bar, 100 μm. (B) Transwell assay for cell migration of HCT116 cells. HCT116 cells were seeded on the upper chamber of the transwell and the bottom compartment was filled with conditioned medium of TPC_{NM}^{Vector} and TPC_{NM}^{TCF21}. After 48 h, the migrated cells were imaged and counted (n = 3). Scale bar, 100 μ m. (C) Western blotting analysis of EMT markers in HCT116 cells primed with conditioned medium of TPC_{NM}^{Vector} and TPC_{NM}^{TCF21} (n = 3). **(D)** Representative images and quantification of tube numbers formed by HMEC-1 cells. HMEC-1 cells suspended with ECM were seeded on the Matrigel coated 96-well plated. After 2 h, ECM were replaced with the conditioned medium of TPC_{NM}^{Vector} and TPC_{NM}^{TCF21}, and the number of formed tubes was calculated 2 hours later (n = 3). Scale bar, 100 μ m. (E) EdU assay for the proliferation of HCT116 cells primed with conditioned medium of TPC_{LM}siNC and TPC_{LM}siTCF21 (n = 3). Scale bar, 100 µm. (F) Representative images and quantification of migrated HCT116 cells. HCT116 cells were seeded on the upper chamber of the transwell and the bottom chamber was filled with the conditioned medium of TPC_{LM}siNC and $TPC_{LM}^{siTCF21}$. After 48 h, the migrated cells were imaged and counted (n = 3). Scale bar, 100 μm. (G) Western blotting analysis of EMT markers in HCT116 cells primed with conditioned medium of TPC_{LM}^{siNC} and $TPC_{LM}^{siTCF21}$ (n = 3). (H) Tube formation assay for HMEC-1 cells treated with conditioned medium of TPC_{LM}siNC and TPC_{LM}siTCF21 as indicated in (D) (n = 3). Scale bar, 100 μ m. Data are presented as mean \pm SEM, NS, not significant. Two-tailed unpaired t-test (A, B, D), one-way ANOVA followed by Tukey's post hoc test (E, F, H).



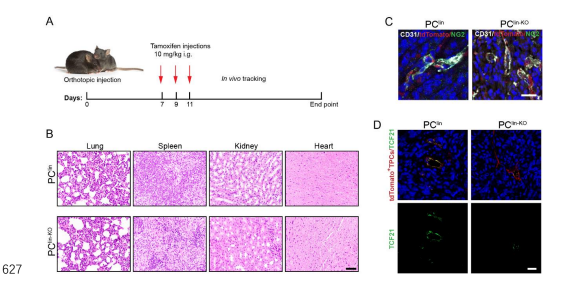
Supplemental Figure 8. TCF21 in TPCs promotes invasion of CRC cells. (A) Representative images and quantification of invaded DLD-1 cells (green). DLD-1 cells mixed with TPC_{NM}^{Vector} or TPC_{NM}^{TCF21} were seeded into the Matrigel-coated transwell. The invaded DLD-1 cells were photographed and counted after 48 h (n = 3). Scale bar, 100 μ m. (B, C) Transwell assay for invasion of HCT116 cells and DLD-1 cells. HCT116 cells (B) or DLD-1 cells (C) were pre-mixed with TPC_{LM}^{SiNC} or TPC_{LM}^{SiTCF21} and subjected to invasion assay as indicated in (A) (n = 3). Scale bar, 100 μ m. Data are presented as mean \pm SEM. ***P < 0.01 by two-tailed unpaired t-test, ***P < 0.001 oneway ANOVA followed by Tukey's post hoc test



Supplemental Figure 9. MATN2 has negligible effects on ECM remodeling and CRC metastasis. (A, B) RT-qPCR analysis of MATN2 in MATN2-overexpressing (A) or -knockdown (B) TPCs (n = 3). (C) RT-qPCR analysis of the ECM-related genes in TPC_{NM}^{Vector} and TPC_{NM}^{MATN2} (n = 3). (D) RT-qPCR analysis of the indicated genes in TPC_{LM}^{siNC} and TPC_{LM}^{siMATN2} (n = 3). (E, F) Representative images and quantification of the invaded DLD-1 cells (green). DLD-1 cells pre-mixed with MATN2-overexpressing (E) or -knockdown (F) TPCs were seeded into the Matrigel-coated transwell. The invaded DLD-1 cells were photographed and counted after 48 h (n = 3). Scale bar, 100 μ m. Data are presented as mean \pm SEM, NS, not significant. ***P<0.001. Two-tailed unpaired *t*-test (A, C, E), one-way ANOVA followed by Tukey's post hoc test (B, D, F).

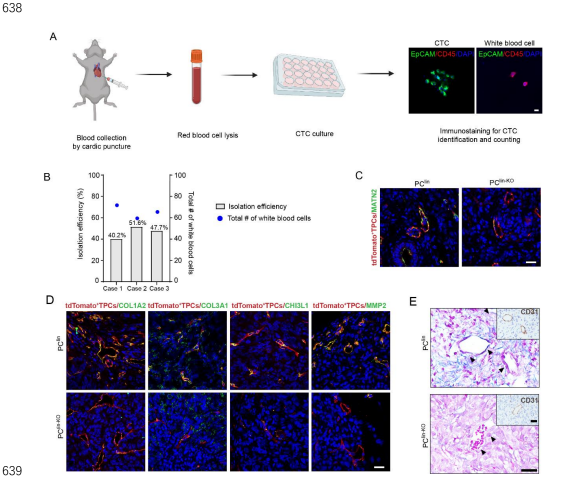


Supplemental Figure 10. Mouse genotyping. (A) PCR analysis of the genotype of PC^{lin} mouse (n = 6). **(B)** PCR analysis of the genotype of PC^{lin-KO} mice (n = 6). All mice were analyzed by PCR genotyping. PCR analysis of Cspg4Cre showed two bands identifying homozygous Cspg4Cre at 2386 bp (knock in) and 272 bp (wild type); PCR analysis of tdTomato showed homozygous tdTomato with a band at 317 bp (knock in) and no signal at 479 bp (wild type); PCR analysis of Tcf21 indicated homozygous Tcf21 flox/flox at 418 bp and 440bp.



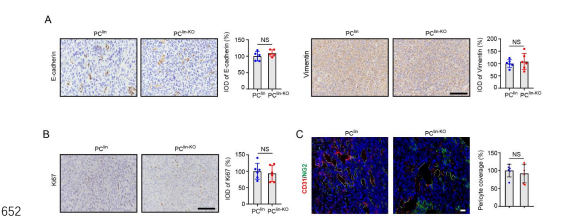
Supplemental Figure 11. Characterization of PClin and PClin-KO mice. (A) Schematic diagram describing the experimental design of *in vivo* experiments. PClin mice and PClin-KO mice were orthotopically injected with MC38-luc-LM3 cells. After 7

days, all mice were treated with tamoxifen (10 mg/kg) through intragastric administration every other day for three times. *In vivo* tracking was performed to detect tumor liver metastasis. **(B)** Representative images of H&E staining of lung, spleen, kidney, heart, and liver derived from PC^{lin} and PC^{lin-KO} mice (n = 6). Scale bar, 50 µm. **(C)** Immunofluorescence analysis of TPCs (tdTomato) in tumor sections by NG2 (green) staining (n = 6). Scale bar, 20 µm. **(D)** Immunofluorescence analysis of TCF21 (green) in TPCs (tdTomato) from PC^{lin} mice and PC^{lin-KO} mice (n = 6). Scale bar, 20 µm.



ECM remodeling and CRCLM. (A) Schematic diagram of the isolation and identification of CTCs. Scale bar, 20 μm. **(B)** Isolation efficiency of CTCs in the blood

spiked with MC38-luc-LM3 cells (n = 3). **(C)** Immunofluorescence staining for MATN2 (green) in TPCs (tdTomato) in primary tumor sections from MC38 allografts (n = 6). Scale bar, 20 μ m. **(D)** Immunofluorescence staining for COL3A1, MMP2, COL1A1 and CHI3L1 (green) in TPCs (tdTomato) from primary tumor sections of MC38 allografts (n = 6). Scale bar, 20 μ m. **(E)** Masson staining for perivascular collagen in primary tumor sections from MC38 allografts. Tumor vessels were labeled with CD31 (n = 6). Black arrows indicate the perivascular collagen fibers. Scale bar, 50 μ m.



Supplemental Figure 13. Pericyte-specific knockout of *Tcf21* has negligible effects on EMT and proliferation of CRC cells. (A) Immunohistochemical staining and quantification of E-cadherin and Vimentin in orthotopic MC38 tumor tissues (n = 6). Scale bar, 50 μ m. (B) Representative images and quantification of Ki67 staining in orthotopic MC38 tumor sections (n = 6). Scale bar, 50 μ m. (C) Immunofluorescence staining and quantification of pericyte coverage as indicated by CD31 (red) and NG2 (green) in tumor sections (n = 6). Scale bar, 20 μ m. Data are presented as mean \pm SEM. NS, not significant. NS by two-tailed unpaired *t*-test.

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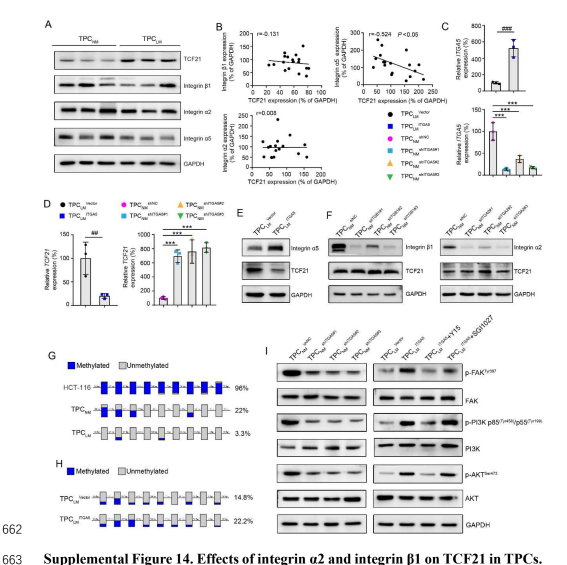
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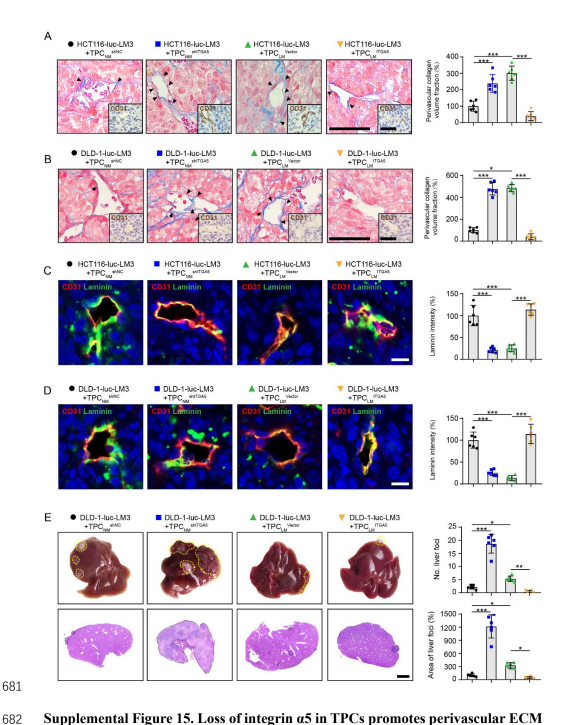
Supplemental Figure 14. Effects of integrin α2 and integrin β1 on TCF21 in TPCs.

(A) Western blotting analysis of TCF21, integrin β 1, integrin α 2, and integrin α 5 in TPC_{NM} and TPC_{LM} (n = 3). (B) Pearson's correlation analysis of the expression of integrin β 1, integrin α 2 and integrin α 5 with TCF21 in TPCs (n = 3). (C) RT-qPCR analysis of the levels of ITGA5 in the integrin α5-knockdown or -overexpressing TPCs (n = 3). (D) RT-qPCR analysis of the levels of TCF21 in integrin $\alpha 5$ -knockdown or overexpressing TPCs (n = 3). (E) Western blotting analysis of integrin $\alpha 5$ and TCF21 in integrin α 5-overexpressing TPC_{LM} (n = 3). (F) Western blotting analysis of integrin $\alpha 2$, integrin $\beta 1$ and TCF21 in integrin $\alpha 2$ or integrin $\beta 1$ -knockdown TPC_{NM} (n = 3). (G)

672 Bisulfite DNA sequencing analysis of TCF21 promoter region in TPCs or HCT116 cells. 673 Blue and gray circles represent methylated and unmethylated CpGs, respectively. The percentage of total methylated CpGs is given on right of each dataset (n = 3). (H) 674 675 Bisulfite DNA sequencing analysis of TCF21 promoter region in integrin α5-676 overexpressing TPC_{LM} (n = 3). (I) Western blotting analysis of FAK/PI3K/AKT signaling axis in integrin α5-knockdown or -overexpressing TPCs with or without FAK 677 inhibitor (Y15) or DNMT1 inhibitor (SGI1027) treatment (n = 3). Data are presented 678 as mean \pm SEM, ##P < 0.01, ###P < 0.001 by two-tailed unpaired t-test, ***P < 0.001 by 679 680 one-way ANOVA followed by Tukey's post hoc test.

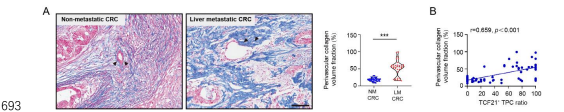
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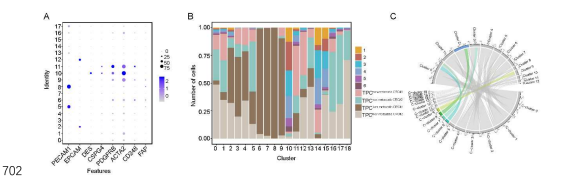


Supplemental Figure 15. Loss of integrin α 5 in TPCs promotes perivascular ECM remodeling and CRCLM. (A, B) Representative images of Masson and CD31 staining in primary tumor sections (n = 6). Black arrows indicate the perivascular collagen fibers. Scale bar, 50 μ m. The quantification of perivascular collagen volume fraction was shown in the right. (C, D) Immunofluorescence staining and quantification of laminin

(green) around the CD31⁺ tumor vessels (red) in HCT116-luc-LM3 xenografts (**C**) and DLD-1-luc-LM3 xenografts (**D**) (n = 6). Scale bar, 20 μ m. (**E**) Representative images and H&E staining of liver metastatic foci (n = 6). Yellow and black dotted lines indicate the metastatic loci. Scale bar, 2 mm. Data are presented as mean \pm SEM. *P < 0.05, **P < 0.01, ***P < 0.001 by one-way ANOVA followed by Tukey's post hoc test.



Supplemental Figure 16. TCF21 in TPCs is associated with perivascular ECM deposition. (A) Masson staining and quantification of perivascular collagen in tumors derived from CRC patients with non-metastasis or liver metastasis (n = 75). Scale bar, 50 μ m. ***P < 0.001 by two-tailed Mann-Whitney test. (B) Pearson's correlation analysis of perivascular collagen volume fraction and TCF21⁺ TPC ratio (n = 75). NM CRC, non-metastatic colorectal cancer, LM CRC, liver metastatic colorectal cancer.



Supplemental Figure 17. Comparison of the scRNA-seq data derived from TPCs and the previous published data. (A) Dot plots for gene expressions in pericytes

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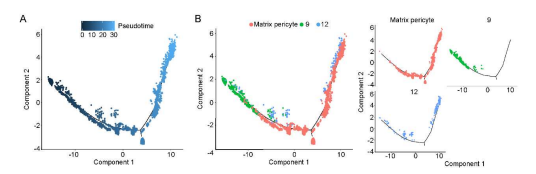
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acquired from previous studies. Raw data of pericyte scRNA-seq were collected from six samples derived from previous studies (GEO accession GSM3140596, GSM3140595, GSM3140594, and GSM3140593) and (GEO accession GSM4159165 and GSM4159164). Total cells derived from previous studies were classified into 18 clusters named L-Cluster 0 to L-Cluster 17. Among them, L-Cluster 3, L-Cluster 9, L-Cluster 10, and L-Cluster 11 were subjected for further analysis as these four populations were positive for DES, CSPG4, PDGFRB, ACTA2, CD248 and FAP (pericyte markers), but negative for PECAM1 (endothelial cell marker) and EPCAM (epithelial cell marker). (B) Analysis of the data of the four populations (L-Cluster 3, L-Cluster 9, L-Cluster 10, and L-Cluster 11) derived from six samples (Sample 1-6) in (A) and our scRNA-seq data (Cluster 0-12). The combined pericytes were categorized into 19 subpopulations, termed C-Cluster 0 to C-Cluster 18. Among them, C-cluster 0-5, C-cluster 10-18 were presented both in our data and the extended data (Sample1-6); however, C-cluster 6-9 were specifically revealed in our data, indicating that the existing pericyte clusters originated from the previous research^{5, 6} were included in our scRNA-seq data and we discovered four new subsets. (C) Comparative analysis of pericyte cluster derived from (B) (C-cluster 0-18) with our data (Cluster 0-12). Among them, C-Cluster 8 was included in Cluster 2 (matrix pericytes).



Supplemental Figure 18. Pseudo-time trajectory for dynamic changes in matrix-

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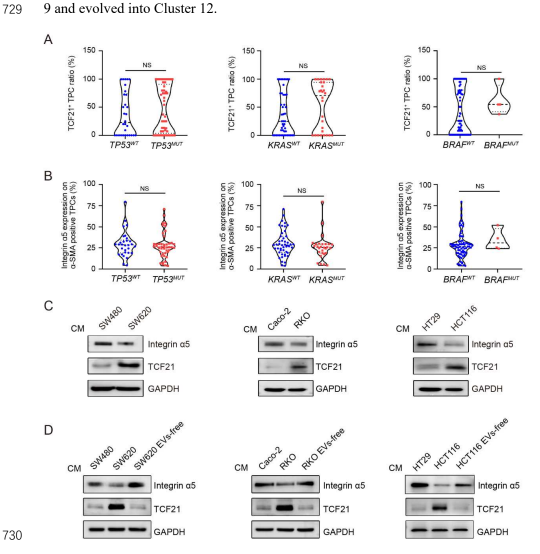
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pericytes. (A) Trajectory analysis plot for matrix-pericytes. Cells are ordered in pseudo-time colored in a gradient from dark blue to light blue (B) The trajectory of the differentiation state of matrix-pericytes. Matrix-pericytes were originated from Cluster 9 and evolved into Cluster 12.



Supplemental Figure 19. Effects of metastatic CRC cells on the expressions of integrin $\alpha 5$ and TCF21 in TPCs. (A) Quantification of TCF21⁺ TPC ratio in CRC patients with or without *TP53*, *BRAF* and *KRAS* mutation (n = 75). (B) Quantification of integrin $\alpha 5$ expression in TPCs derived from CRC patients with or without *TP53*, *BRAF* and *KRAS* mutation (n = 75). (C) Western blotting analysis of integrin $\alpha 5$ and

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TCF21 in TPCs primed with CM from weakly-metastatic (SW480, Caco-2, HT29) and highly-metastatic (SW620, HCT116, RKO) CRC cells (n = 3). (**D**) Western blotting analysis of integrin α 5 and TCF21 in TPCs primed with or without the EVs-free CM of highly-metastatic CRC cells (n = 3). The EVs-free CM of highly-metastatic CRC cells were generated by centrifugation at 1×10^5 g to remove the EVs. EVs, extracellular vesicles; CM, conditioned medium. Each sample on the violin plots represents individual patient data. NS, not significant. NS by two-tailed Mann-Whitney test. *WT*, *wildtype*; *MUT*, *mutant*.

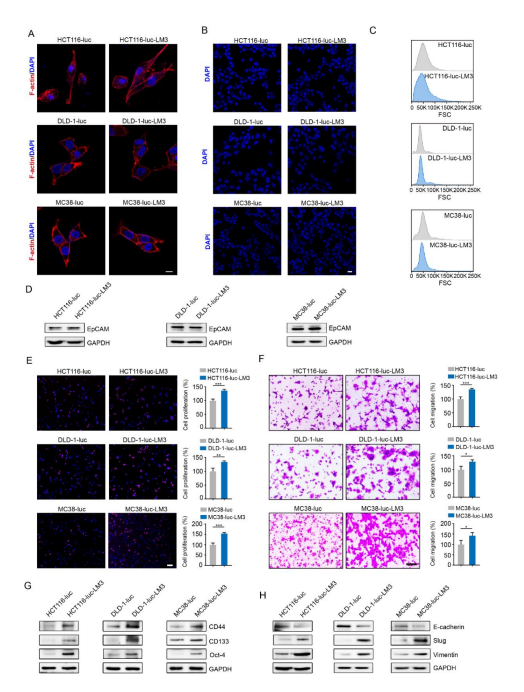
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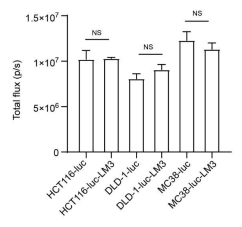
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Supplemental Figure 20. Comparison of parental cells with LM3 cells. (A) Immunofluorescence analysis of cellular morphology (n = 3). Phalloidin-rhodamine was used to identify F-actin. Scale bar, 10 μ m. (B) Immunofluorescence analysis of nucleus size (n = 3). Cell nucleus was measured after DAPI staining of fixed cells. Scale bar, 20 μ m. (C) FCM analysis of the cell size of parental cells and LM3 cells by Forward

scatter (n = 3). **(D)** Western blotting analysis of EpCAM in parental cells and LM3 cells (n = 3). **(E)** EdU assay for the proliferation of parental cells and LM3 cells (n = 3). Scale bar, 100 μ m. **(F)** Transwell assay for the migration of parental cells and LM3 cells (n = 3). Scale bar, 100 μ m. **(G)** Western blotting analysis of CD44, CD133 and Oct-4 in parental cells and LM3 cells (n = 3). **(H)** Western blotting analysis of E-cadherin, slug and Vimentin in parental cells and LM3 cells (n = 3). Data are presented as mean \pm SEM. *P < 0.05, **P < 0.01, ***P < 0.001 by two-tailed unpaired t-test.



Supplemental Figure 21. Comparison of luciferase activities between parental cells and LM3 cells. Bioluminescence detection of parental cells and LM3 cells. Data are presented as mean \pm SEM. NS, not significant. NS by two-tailed unpaired t-test.

769 **Supplemental Table 1.** Clinical characteristics of CRC specimens.

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Character	Overall population
Gender, number of patients (%)	
Female	33 (44.0)
Male	42 (56.0)
Age, median (range)	69 (34 – 91)
Site of primary tumor, number of patients (%)	
Right hemicolon	17 (22.7)
Left hemicolon	58 (77.3)
Histological grade, number of patients (%)	
High/Moderate	47 (62.7)
Low	28 (37.3)
Size (cm), median (range)	4.3 (2 – 8.6)
TNM stage, number of patients (%)	
I-II	41 (54.7)
IV	34 (45.3)
Liver metastasis (%)	
No	41 (54.7)
Yes	34 (45.3)
TP53 mutation (%)	
No	30 (40)
Yes	45 (60)
KRAS mutation (%)	
No	47 (62.7)
Yes	28 (37.3)
BRAF mutation (%)	
No	71 (94.7)
Yes	4 (5.3)

Supplemental Table 2. Correlation analysis between the MATN2⁺ TPC ratio and the clinicopathologic data.

Character	MATN2+	TPC	ratio	MATN2+	TPC	ratio	P value
Character	(≤30%		(>30%)				P value
Gender							
Female	22 (66.7%)		11 ((33.3%)		0.826
Male	29 (69.0%)		13 ((31.0%)		0.820
Age							
< 60	15 (75.0%)		5 (15.0%)		0.422
≥60	36 (65.5%)		19 ((34.5%)		0.433
Location							
Right hemicolon	13 (76.5%)		4 (2	23.5%)		0.395
Left hemicolon	38 (65.5%)	20 (34.5%)				
Differentiatio							
n							
High/Moderate	33 (70.2%)		14 ((29.8%)		0.595
Low	18 (64.3%)		10 ((35.7%)		0.575
Size							
< 5 cm	35 (79.5%)		9 (2	20.5%)		0.011
≥5 cm	16 (51.6%)		15 (48.4%)			0.011
TNM stage							
I-II	40 (97.6%)		1 ((2.4%)		< 0.001
IV	11 (32.4%)		23 ((67.6%)		< 0.001
Liver							
metastasis							
No	40 (97.6%)		1 ((2.4%)		< 0.001
Yes	11 (32.4%)		23 ((67.6%)		· 0.001
TP53 mutation							
No	23 (76.7%)		7 (2	23.3%)		0.189

Yes	28 (62.2%)	17 (37.8%)	
KRAS			
mutation			
No	35 (74.5%)	12 (25.5%)	0.120
Yes	16 (57.1%)	12(42.9%)	0.120
BRAF			
mutation			
No	49 (69%)	22 (31%)	0.000
Yes	2 (50%)	2 (50%)	0.808

Supplemental Table 3. Correlation analysis between the TCF21⁺ TPC ratio and the clinicopathologic data.

Character	TCF21 ⁺	TPC	ratio	TCF21 ⁺	TPC	ratio	P
Character	(:	≤44%)		((>44%)		value
Gender							
Female	16	(48.5%)		17	(51.5%)		0.340
Male	25	(59.5%)		17 (40.5%)			0.340
Age							
< 60	11	(55.0%)		9 ((45.0%)		0.972
≥60	30	(54.5%)		25 (45.5%)		0.972	
Location							
Right hemicolon	6	(35.3%)		11	(64.7%)		0.068
Left hemicolon	35	(60.3%)		23 (39.7%)		0.008	
Differentiation							
High/Moderate	29	(61.7%)		18	(38.3%)		0.113
Low	12	(42.9%)		16	(57.1%)		0.113
Size							
< 5 cm	27	(61.4%)		17	(38.6%)		0.165
≥5 cm	14	(45.2%)		17 (54.8%)			0.105
TNM stage							

I-II	40 (97.6%)	1 (2.4%)	< 0.001
IV	1 (2.9%)	33 (97.1%)	₹ 0.001
Liver			
metastasis			
No	40 (97.6%)	1 (2.4%)	< 0.001
Yes	1 (2.9%)	33 (97.1%)	< 0.001
TP53 mutation			
No	18 (60.0%)	12 (40.0%)	0.440
Yes	23 (51.1%)	22 (48.9%)	0.449
KRAS mutation			
No	29 (61.7%)	18 (38.3%)	0.113
Yes	12 (42.9%)	16 (57.1%)	0.115
BRAF mutation			
No	40 (56.3%)	31 (43.7%)	0.479
Yes	1 (25.0%)	3 (75.0%)	0.478

Supplemental Table 4. Multivariable logistic regression for clinical and demographic factors between CRC patients with or without liver metastasis.

	β	S.E.	Wald	P	OR	95% CI
TCF21 ⁺ TPC ratio (%)	7.112	1.435	24.558	<0.001	1226.464	73.636-20427.781
TP53 mutation	0.074	1.536	0.002	0.962	1.077	0.053-21.851
KRAS mutation	0.448	1.561	0.082	0.774	1.565	0.073-33.387
BRAF mutation	0.875	3.392	0.067	0.796	2.399	0.003-1850.29

Abbreviations: S.E., standard error; OR, odds ratio; CI, confidence interval.

Supplemental Table 5. Clinical characteristics of non-metastatic CRC specimens

Characteristics	Case 1	Case 2	Case 3	Case 4	Case 5	Case 6
Gender/Age (yr)	Male (58)	Male (78)	Male (59)	Female (53)	Male (73)	Female (75)
Date of diagnosis	20190429	20191031	20200331	20210323	20210402	20210420
Tumor type	Colorectal	Colorectal	Colon	Colon	Colon	Colon
rumor type	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma	Adenocarcinoma
Location	Rectum	Sigmoid colon	Sigmoid colon	Sigmoid colon	Sigmoid colon	transverse colon
Tumor size	5 cm	4 cm				
(maximum diameter)	3 CIII	4 CIII	2 cm	4 cm	5 cm	5 cm
Differentiation	Moderate	Moderate	Moderate	Moderate	Moderate	Moderate
TNM stage	T2N0M0	T4aN0M0	T3N1bM0	T4aN1bM0	T3N0M0	T3N1aM0
Clinical stage	I	IIB	IIIB	IIIB	IIA	IIIB
Clinical metastasis	No metastasis	No metastasis	No metastasis	No metastasis	No metastasis	No metastasis
Treatment status	Chemotherapy,	No treatment				

radiotherapy	before surgery				

Supplemental Table 6. Clinical characteristics of liver-metastatic CRC specimens

Female (59) 20200320 Colorectal
Colorectal
Adenocarcinoma
Descending
colon and ileum
7 cm
/ CIII
Low
_

TNM stage	T3N2M1a	T4bN1M1a	T3N2bM1a	T3N2M1a	T4bN1M1a	T4bN1aM1a
Clinical stage	IVa	IVa	IVa	IVa	IVa	IVa
Clinical metastasis	Liver metastasis					
Treatment status	Chemotherapy,	No treatment				
	radiotherapy	before surgery				

Supplemental Table 7. Primers for mouse genotyping

Mice	Primer name	Primer Sequence	Product size	Gene type
1	Rosa26-tF1	CCCAAAGTCGCTCTGAGTTGTTA	W4-4701	
1	Rosa26-tR1	TCGGGTGAGCATGTCTTTAATCT	Wt=479bp	T002249 <i>Rosa26-CAG-</i>
2	tdTomato-tF1	CGGCATGGACGAGCTGTACAAG	VI_2171	LSL-Cas9- tdTomato
2	WPRE-tR2	TCAGCAAACACAGTGCACACCAC	KI=317bp	
3	JS04431-Tcf21-5wt-tF1	GATCCTTCAAATGACTCCAGGCC	WT: 314bp Fl: 418bp	T013083 <i>Tcf21</i> -flox

	JS04431-Tcf21-5wt-tR1	GTTTGCTAACTTGCTGCCACACAC		
4	XM003792- Cspg4-TF1	AAATCTAAGCGCGGGTCTGGC	WT:0bp	
4	XM003792- Cspg4-TR1	TGCGAACCTCATCACTCGTTGC	KI:272bp	T006187 Cspg4-
5	XM003792- Cspg4-TF2	AAATCTAAGCGCGGGTCTGGC	WT:352bp	CreERT2
3	XM003792- Cspg4-TR2	GGACCATGAGTGCAGTCCCCATA	(KI=2386bp)	

Supplemental Table 8. STR profiles of TPCs.

Marker	Allele 1	Allele 2	Allele 3	Allele 4
D19S433	13	14		
D5S818	11	13		
D21S11	29	31.2		
D18S51	13	14		
D6S1043	12	13		
AMEL	X	Y		
D3S1358	16	17		
D13S317	10			
D7S820	8	12		
D16S539	9	11		
CSF1PO	12	14		
Penta D	9	12		
D2S441	10	11		
vWA	14	15		
D8S1179	10	14		
TPOX	8	10		
Penta E	11	12		
TH01	6	8		
D12S391	19	23		
D2S1338	22	23		
FGA	20	23		

Conclusion of cell identification: The results of STR typing showed that there were no multiple alleles at each locus. No cross contamination of human cells was found in the cells.

Supplemental Table 9. Clinical characteristics of the resected CRC patients

Characteristics	Case 1	Case 2	Case 3	Case 4
Gender/Age (yr)	Male (58)	Male (78)	Male (51)	Male (78)
Date of diagnosis	20190429	20191031	20190516	20191010
Tumor type	Colorectal Adenocarcinoma	Colorectal Adenocarcinoma	Colorectal Adenocarcinoma	Colorectal Adenocarcinoma
Location	Rectum	Sigmoid colon	Descending colon and ileum	Sigmoid colon
Tumor size (maximum diameter)	5 cm	4 cm	4 cm	6 cm
Differentiation	Moderate	Moderate	Moderate	Low
TNM stage	T2N0M0	T4aN0M0	T4bN1M1a	T3N2bM1a
Clinical stage	I	IIB	IVa	IVa
Clinical metastasis	No metastasis	No metastasis	Liver metastasis	Liver metastasis
Treatment status	No treatment before surgery			

Supplemental Table 10. STR profiles of HCT116-luc cells and HCT116-luc-LM3 cells.

Locus	HCT116-luc		HCT116-luc-LM3
Amelogenin	X	Y	X Y
D5S818	10	11	10 11
D13S317	10	12	10 12
D7S820	11	12	11 12
D16S539	11		11
vWA	17		17
TH01	8	9	8 9
TPOX	8		8
CSF1PO	7	10	7 10

The number of matched peaks

15

Percent match between the query and the database profile:

100%

Conclusion of cell identification: ①The results of STR typing of the cell DNA of this strain show that there is no multi-allelic phenomenon at each locus. No human cell cross-contamination is found in the cells. ②The submitted HCT116-luc-LM3 cells are 100% match for HCT116-luc cells.

Supplemental Table 11. STR profiles of DLD-1-luc cells and DLD-1-luc-LM3 cells.

Locus		DLD-1-luc	DLD-1-luc-LM3	
Amelogenin	X	Y	X Y	
D5S818	13		13	
D13S317	8	11	8 11	
D7S820	10	12	10 12	

D16S539	12	13	12 13	
vWA	18	19	18 19	
TH01	7	9.3	7 9.3	
TPOX	8	11	8 11	
CSF1PO	12		12	

The number of matched peaks

16

Percent match between DLD-1-luc and DLD-1-luc-LM3:

100%

Conclusion of cell identification: ①The results of STR typing of the cell DNA of this strain show that there is no multi-allelic phenomenon at each locus. No human cell cross-contamination is found in the cells. ②The submitted DLD1-luc-LM3 cells are 100% match for DLD-1-luc cells.

Supplemental Table 12. STR profiles of MC38-luc cells and MC38-luc-LM3 cells.

Locus		MC38-luc		MC38-luc-LM3
18-3	16		16	
4-2	19.3	20.3	19.3	20.3
6-7	14	15	14	15
19-2	13		13	
1-2	19		19	
7-1	26.2		26.2	
8-1	16		16	
1-1	16		16	
3-2	13	14	13	14
2-1	16		16	
15-3	22.3		22.3	
6-4	18		18	
13-1	17.1		17.1	
11-2	16		16	
17-2	15		15	
12-1	17		17	

5-5	17	17	
X-1	27	27	
The number of matched peaks			21
Percent match between the query and the database profile			100%

Conclusion of cell identification: ①The results of STR typing of the cell DNA of this strain show that there is no multi-allelic phenomenon at each locus. No human cell cross-contamination is found in the cells. ②The cells of MC38-luc and MC-38-luc-LM3 are mouse cells from a single source, and there is no human-derived cell contamination. ③The submitted MC38-luc-LM3 cells are 100% match for MC38-luc cells.

Supplemental Table 13. Primers for qPCR.

Genes	Forward (5'-3')	Reverse (5'-3')
ACTB	GTTGCTATCCAGGCTGTGCTATCC	GGTGGCAGTGATGGCATGGAC
MFAP4	TGAAGGCACAAGGAGTTCTCT	GGGTAGATGAGGTACACGCC
MMP2	CGACCACAGCCAACTACGATGATG	GTGCCAAGGTCAATGTCAGGAGAG
FBLNI	TGCTCCATCAACGAGACCTG	AGCACTCCCGATTCTCATGG
COL1A2	CCGTGGCAGTGATGGAAGTGTG	CCTTGTTACCGCTCTCTCCTTTGG
CHI3L1	GGCTTCTTCTGAGACTGGTGTTGG	CGCTTTCCTGGTCGTCGTATCC
COL3A1	TGTACCAGCCAGACCAGGAAGAC	TGTACCAGCCAGACCAGGAAGAC
FBLN2	GACCGAGGACAGTGAGGAGGAAG	CAGGCAGTGATGTGGACAGGATG
IGFBP5	GTACCTGCCCAATTGTGACC	AAGTCCCCGTCAACGTACTC
MATN2	AGAGGTGTGTGGCTGTGGACTAC	GAGCACTGGCAGACGAAGGAATC
ITGA5	GTCGGGGGCTTCAACTTAGAC	ACAGAGGTAGACAGCACCAC
CILP	CTTTGAGAACCTCCGGGCAT	TCGATCCCCCTCAATCTGGT

C11orf96	TCCAGTTACCAGGCGGTGAT	TGCGTCTTGAAGCGAGACTG
A2M	GAGGCAGAAGGACAATGGCT	ATAGGCGGAGAGGGTCACTT
SFRP2	GCCCGACTTCTCCTACAAGC	CTCCTTCATGGTCTCGTGGC
MAF	CGTCCTCTCCCGAGTTTTTCA	GGCTTCCAAAATGTGGCGTA
PTGDS	CCATGTGCAAGTCTGTGGTG	CATGGTTCGGGTCTCACACT
TCF21	TCCTGGCTAACGACAAATACGA	TTTCCCGGCCACCATAAAGG

Supplemental Table 14. Sequences of siRNAs.

SiRNA	Sense (5'-3')	Antisense (5'-3')
siTCF21-1	GGAUUCGAACAAGGAAUUUTT	AAAUUCCUUGUUCGAAUCCTT
siTCF21-2	GCUAACGACAAAUACGAGATT	UCUCGUAUUUGUCGUUAGCTT
siITGB1-1	GAACAGAUCUGAUGAAUGATT	UCAUUCAUCAGAUCUGUUCTT
siITGB1-2	GUGGUUUCGAUGCCAUCAUTT	AUGAUGGCAUCGAAACCACTT

siITGB1-3	GAUCAUUGAUGCAUACAAUTT	AUUGUAUGCAUCAAUGAUCTT
siITGA2-1	CCCGAGCACAUCAUUUAUATT	UAUAAAUGAUGUGCUCGGGTT
siITGA2-2	GCUGGUGACAUCAGUUGUATT	UACAACUGAUGUCACCAGCTT
siITGA2-3	GUGGUUGUGUGAUGAAUTT	AUUCAUCACACACAACCACTT
siMATN2-1	GCAUCCUAAUCUUUGCCAUTT	AUGGCAAAGAUUAGGAUGCTT
siMATN2-2	GCAGUUUGUCACUGGAAUUTT	AAUUCCAGUGACAAACUGCTT

Supplemental table 15. Antibodies for Western blotting

Antibody	RRID	Company
TCF21	AB_10601215	SIGMA
COL1A2	AB_10679394	Abcam
N-cadherin	AB_1310479	Abcam
Vimentin	AB_10562134	Abcam
Slug	AB_777968	Abcam
FBLN1	AB_2553938	Invitrogen
Integrin α5	AB_2233962	Cell Signaling Technology
GAPDH	AB_10622025	Cell Signaling Technology
p-FAK ^{Tyr397}	AB_10891442	Cell Signaling Technology
FAK	AB_2799801	Cell Signaling Technology
$p\text{-PI3K }p85^{(Tyr458)}\!/p55^{(Tyr199)}$	AB_2895293	Cell Signaling Technology
PI3K	AB_2165248	Cell Signaling Technology
p-AKT ^{Ser473}	AB_2315049	Cell Signaling Technology
AKT	AB_2225340	Cell Signaling Technology
EpCAM	Cat. GB12274	Servicebio
E-cadherin	AB_2728770	Cell Signaling Technology
CHI3L1	Cat. AF8379	Beyotime
COL3A1	Cat. AF6531	Beyotime
Integrin alpha 2	Cat. AF2332	Beyotime
Integrin β1/CD29	Cat. AF0207	Beyotime
MMP2	Cat. GB11130	Servicebio
CD44	AB_2750879	Cell Signaling Technology
CD133	AB_2721172	Cell Signaling Technology

Oct-4	AB_823583	Cell Signaling Technology	
HRP conjugated Goat Anti-	Cat. GB23303	Servicebio	
Rabbit IgG (H+L)	Cat. GB23303	Servicedio	
HRP conjugated Goat Anti-	Cat. GB23301	Servicebio	
Mouse IgG (H+L)	Cut. GB25501	Scrviccolo	
HRP conjugated Donkey Anti-	Cat. GB23404	Servicebio	
Goat IgG (H+L)	San SB25 10 1		

Supplemental table 16. Antibody list for immunofluorescence.

Antibody	RRID	Company
MATN2	AB_2811126	Abcam
VWF	AB_298501	Abcam
COL1A2	AB_10679394	Abcam
Laminin	AB_298179	Abcam
α-SMA	AB_2799045	Cell Signaling Technology
Integrin α5	AB_2233962	Cell Signaling Technology
TCF21	AB_10601215	SIGMA
CD31	AB_2161028	R&D Systems
CD45	AB_306361	Abcam
NG2	AB_11213678	MERK
MMP2	Cat. GB11130	Servicebio
CHI3L1	Cat. AF8379	Beyotime
COL3A1	Cat. AF6531	Beyotime
EpCAM	Cat. GB12274	Servicebio

Alexa Fluor 647-Donkey anti-Sheep IgG (H+L)	AB_2535865	Invitrogen
Alexa Fluor 546-Donkey	AB 2534016	Invitrogen
anti-Rabbit IgG (H+L)	118_200 1010	m m ogen
Alexa Fluor 488-Donkey	AB_2535792	Invitrogen
anti-Rabbit IgG (H+L) Alexa Fluor 488-Donkey		
anti-Goat IgG (H+L)	AB_2534102	Invitrogen

Supplemental table 17. Antibody list for immunohistochemistry.

Antibody	RRID	Company
CD31	AB_2161028	R&D Systems
E-Cadherin	AB_2728770	CST
Vimentin	AB_10562134	Abcam
Ki67	Cat. GB11030	Servicebio
HRP- Goat Anti-Rabbit IgG (H+L)	Cat. GB23303	Servicebio
HRP -Goat Anti-Mouse IgG (H+L)	Cat. GB23301	Servicebio
HRP-Donkey Anti-Goat IgG (H+L)	Cat. GB23404	Servicebio