Plasticity of Lgr5-Negative Cancer Cells Drives Metastasis in Colorectal Cancer

Graphical Abstract

Highlights

- The majority of disseminating cells of colorectal cancer are Lgr5^-
- Lgr5^- cancer cells are the main seeds of colorectal cancer metastatic lesions
- Long-term metastatic growth from Lgr5^- cells requires appearance of Lgr5^+ cells
- Lgr5^- metastases have the intrinsic capacity to re-establish the cellular hierarchy

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In Brief

Van Rheenen and colleagues study Lgr5^+ cancer stem cells during colorectal cancer metastasis. They demonstrate that the majority of metastases are seeded by Lgr5^- cells, which upon arrival seed metastases in which Lgr5^+ cells appear. This plasticity can occur independently of stemness-inducing factors and is indispensable for long-term metastatic growth.
Plasticity of Lgr5-Negative Cancer Cells Drives Metastasis in Colorectal Cancer

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SUMMARY

Colorectal cancer stem cells (CSCs) express Lgr5 and display extensive stem cell-like multipotency and self-renewal and are thought to seed metastatic disease. Here, we used a mouse model of colorectal cancer (CRC) and human tumor xenografts to investigate the cell of origin of metastases. We found that most disseminated CRC cells in circulation were Lgr5- and formed distant metastases in which Lgr5+ CSCs appeared. This plasticity occurred independently of stemness-inducing microenvironmental factors and was indispensable for outgrowth, but not establishment, of metastases. Together, these findings show that most colorectal cancer metastases are seeded by Lgr5- cells, which display intrinsic capacity to become CSCs in a niche-independent manner and can restore epithelial hierarchies in metastatic tumors.

INTRODUCTION

Epithelial tissues are hierarchically organized: a small pool of replicative immortal and self-renewing stem cells (SCs) give rise to a large population of more specialized cells that are replicative mortal (Beck and Blanpain, 2013). During development, SCs drive long-term growth, while during adult life they sustain tissues by driving turnover of short-lived specialized cells (Donati and Watt, 2015; Soteriou and Fuchs, 2018). This hierarchical organization is proposed to be maintained in tumors (Batlle and Clevers, 2017; Beck and Blanpain, 2013; Nassar and Blanpain, 2016). Indeed, cell-sorting and transplantation experiments showed that only a small pool of cancer cells, often referred to as cancer SCs (CSCs), have the ability to grow tumors in mice (Al-Hajj et al., 2003; Bonnet and Dick, 1997; Ricci-Vitiani et al., 2007). More recently, lineage-tracing experiments provided proof of the existence of CSCs in unperturbed tumors, including glioblastoma (Chen et al., 2012), squamous skin tumors (Oriessens et al., 2012), colorectal cancer (CRO) (Cortina et al., 2017; Schepers et al., 2012; Shimokawa et al., 2017) and breast cancer (Zomer et al., 2013). In addition to the role in tumor growth and homeostasis, CSCs are thought to be important for metastasis. The essential role of CSCs in the metastatic process is reinforced by studies showing that metastasizing cells acquire SC properties, for example, through epithelial-to-mesenchymal transition (Guo et al., 2012; Mani et al., 2008; Ye et al., 2015). Indeed, selective depletion of CSCs in primary tumors protects from the appearance of distant metastases (de Sousa e Melo et al., 2017). The role of CSCs in metastasis would be particularly crucial if the tumor cell hierarchy is a rigid one-way route, from SCs to more specialized cells (Meacham and Morrison, 2013). However, epithelial hierarchical organization can be dynamic (Medema, 2013). For example, in healthy tissues, upon damage or selective ablation of the SC population, more specialized cells can acquire SC traits (Ritsma et al., 2014; Tetteh et al., 2016; Tian et al., 2011; van Es et al., 2012). This process of cellular plasticity ensures repopulation of impaired SC niches and reestablishment of tissue homeostasis. Whether cellular plasticity also plays an important role in metastasis is unknown. In this study, we dissect the different steps of the metastatic cascade (cell migration, intravasation, metastatic seeding, and outgrowth) and uncover the roles of CSCs, non-CSCs, and cellular plasticity in metastasis.
RESULTS

Lgr5\textsuperscript{eGFP} Labels Functional Stem Cells in Colorectal Tumors

CRC models provide a unique opportunity to clarify the role of CSCs and non-CSCs in the metastatic process, because hierarchical organization is maintained during disease progression and functional CSCs are marked by Lgr5 expression (Cortina et al., 2017; de Sousa e Melo et al., 2017; Schepers et al., 2012; Shimokawa et al., 2017). To study the role of CSCs in metastasis, we combined the Villin\textsuperscript{Cre-ER\textsuperscript{T2}}; APC\textsuperscript{fl/fl}; KRAS\textsuperscript{G12D}; TP53\textsuperscript{ko/ko} colorectal tumor mouse model with Lgr5\textsuperscript{DTR/eGFP} (de Sousa e Melo et al., 2017) and Confetti (Snippert et al., 2010) fluorescent mouse models (Figure 1A). Upon injection of tamoxifen, the entire intestinal tract of these mice becomes tumorigenic, and these animals reach the humane endpoint before metastases are formed. Just before the humane endpoint, we isolated the tumorigenic colons of these mice and established primary CRC organoids that consist of cancer cells expressing RFP-Confetti and of CSCs that additionally express Lgr5-driven eGFP and diphtheria toxin receptor (DTR) (referred as CRC Lgr5\textsuperscript{eGFP} organoids; Figures S1A and S1B). As previously described, orthotopic transplantation of these murine CRC organoids led to the formation of a single colorectal primary tumor that spontaneously metastasizes to liver and lungs (Figures S1C–S1E) (Fumagalli et al., 2017, 2018; Roper et al., 2017; Tauriello et al., 2018). In line with previous findings (Cortina et al., 2017; de Sousa e Melo et al., 2017; Shimokawa et al., 2017), we found that Lgr5 is a good marker of the functional SCs both in organoids and tumors, because selective ablation of the Lgr5\textsuperscript{+} CSCs by diphtheria toxin (DT) (Figures S1F–S1I) caused organoids to collapse and prevented tumor growth and metastasis (Figures 1B–1D). Next, we developed a flow cytometry strategy to isolate Lgr5\textsuperscript{+} CSCs and Lgr5\textsuperscript{−}/C0 cancer cells from both organoids and primary tumors (Figures S1J–S1M). Analysis of DT-treated CRC Lgr5\textsuperscript{eGFP} organoids, in which Lgr5\textsuperscript{+} CSCs are selectively depleted, and post-sort confocal microscopy on untreated CRC Lgr5\textsuperscript{eGFP} organoids confirmed that our gating strategy allowed to reliably isolate Lgr5\textsuperscript{+} and Lgr5\textsuperscript{−}/C0 cancer cells (Figures S1N and S1O). Clonogenic assays confirmed isolation of functional Lgr5\textsuperscript{+} CSCs, as these cells were twice as clonogenic as Lgr5\textsuperscript{−}/C0 cancer cells (Figure 1E). Moreover, isolated Lgr5\textsuperscript{+} CSCs showed higher expression of known intestinal SC markers and lower expression of differentiated markers than the Lgr5\textsuperscript{−} cancer cells (Figure 1E). Of note, we found a very significant overlap in significantly differentially expressed genes between Lgr5\textsuperscript{+} and Lgr5\textsuperscript{−} cells from organoids and orthotopic...
Figure 2. The Majority of Cells Escaping the Primary Colorectal Tumors Are Lgr5⁺

(A) Representative time-lapse intravital images of a non-migratory field within a CRC primary tumor. Note that the resolution of in vivo images is lower than of ex vivo images because of great depth of imaging in living animals. Scale bar, 50 µm.

(B) Time-lapse intravital images of a migratory field showing primary tumor cell migration of Lgr5⁻ and cancer cells (ROI, region of interest 1) and Lgr5⁺ CSCs (ROI2). Dashed lines highlight the migratory cells, continuous lines mark the migratory tracks overtime. Scale bars, 50 µm. See also Videos S1 and S2.

(C) Representative time-lapse intravital images of Lgr5⁺ CSCs (upper panel) and Lgr5⁻ cancer cells (lower panel) moving as cell clusters (see also Videos S3 and S4). Scale bars, 50 µm.

(D) Display of the migratory tracks of Lgr5⁺ CSCs and Lgr5⁻ cancer cells observed in (A) (left) and in (B) (right).

(E) Distribution of total displacement of Lgr5⁺ (green) and Lgr5⁻ (red) migratory cells over a period of 4 h.

(legend continued on next page)
primary tumors (hypergeometric test p value < 10^{-100}), indicating that Lgr5+ and Lgr5− cancer cells maintain their specific cellular identity when analyzed either in vitro or in vivo.

The Majority of Migratory Tumor Cells Are Lgr5 Negative

Using the developed orthotopic transplantation mouse model, we addressed whether Lgr5+ CSCs and Lgr5− cancer cells have differential metastatic behavior. In order to metastasize, cancer cells first need to acquire traits that enable them to leave the primary tumor (Hanahan and Weinberg, 2011). To test which cancer cells escape from the primary site, we filmed the behavior of Lgr5+ CSCs and Lgr5− cancer cells in vivo using multiphoton microscopy (Figure 2; Videos S1, S2, S3, and S4). Consistent with other tumor models (Beerling et al., 2016; Hirata et al., 2015; Patsialou et al., 2013), in some tumor lobes no migration was observed, whereas in other lobes individual and clusters of Lgr5+ CSCs and Lgr5− cancer cells invaded into the non-labeled stroma (n = 9 mice, 73 movies, 1,064 migratory events; Figures 2A–2C; Figure S2A; Videos S1, S2, S3, and S4). Analysis of the migratory cells (i.e., cells that displaced more than half a cell diameter in 4 h; Figures 2D and 2E) revealed that Lgr5+ CSCs and Lgr5− cancer cells mainly escaped as single cells and less frequently as cell clusters (Figure 2F). Moreover, Lgr5− cancer cells might have a slightly higher displacement and velocity than Lgr5+ CSCs (Figures 2G and 2H; Figures S2B and S2C). Surprisingly, of the 1,064 migratory cells that escaped from the primary tumors, the majority were Lgr5− (Figures 2E and 2F; 90.3% ± 2.8%, p < 0.0001), suggesting that most of the escaping cells are in a more differentiated state.

The Vast Majority of Disseminating Cells in the Circulation Are Lgr5 Negative

To determine whether both Lgr5+ CSCs and Lgr5− cancer cells were able to enter the blood circulation, we drew blood from the portal vein of mice bearing metastatic CRC to analyze CTCs directly draining from the primary tumor using flow cytometry (Figure 3A; Figure S3A). In all animals (n = 4) we detected only Lgr5− circulating cancer cells (Figure 3B; Figure S3B). We extended the analysis by collecting systemic oxygen-poor blood from the right ventricle of the heart of a second batch of metastatic CRC mice to analyze CTCs coming from metastases and observed that also in this case the vast majority of circulating cancer cells were Lgr5− (98.4% ± 0.6%, n = 5; Figures S3C–S3E). Importantly, single-cell mRNA sequencing validated that the CTCs displayed an expression profile closely related to Lgr5− cancer cells and did not show enriched expression of other CSC marker genes, such as Prom1 (i.e., CD133), CD44, Aldh1a1, and Bmi1 (Figure 3C; Figures S3F and S3G). Combined these data suggest that the disseminating population of cancer cells is dominated by Lgr5− cancer cells. Indeed, when we analyzed the livers of these mice, all single-cell metastases were Lgr5− (n = 6 mice; Figures 3D and 3E; Figure S3H).

Stem Cell Plasticity Is Required for Metastatic Seeding in the Liver

To test whether escaping Lgr5− cancer cells could initiate metastatic lesions, we analyzed larger metastases in the livers. Although all single-cell and the majority of smaller metastases were devoid of Lgr5+ CSCs, all larger metastases (lesions larger than 80 µm in diameter) contained a population of Lgr5+ CSCs (Figures 3D and 3E; Figure S3H). To test whether Lgr5− cancer cells could give rise to full-blown metastatic lesions, we monitored the real-time outgrowth of Lgr5− cancer cells seeding in the liver, by tracking the same metastatic lesions with repeated multi-day intravitral imaging through an abdominal imaging window (Figure 3F). In all small lesions initially containing only Lgr5− cancer cells, we observed that some of the Lgr5− tumor cells underwent plasticity and acquired Lgr5+GFP expression over time (Figure 3F). Next, we neutralized this plasticity by ablating newly formed Lgr5+ CSCs in early metastatic lesions and observed by intravitral imaging that metastasis composed by only Lgr5− cancer cells stopped growing over time and ultimately regressed (Figures 3G and 3H; Figure S3I). Indeed, after 4 weeks of DT treatment, no metastases could be observed (Figures 3I and 3J). Combined this shows that the appearance of Lgr5+ CSCs is indispensable for the outgrowth of metastases founded by Lgr5− cancer cells.

To compare the efficiency of Lgr5− cancer cells and Lgr5+ CSCs to initiate metastases, we injected equal amounts of sorted Lgr5− CSCs and Lgr5− cancer cells into the mesenteric vein, which are transported through the portal vein to the liver. Four weeks later we found that, in addition to Lgr5+ CSCs, Lgr5− cancer cells were able to form metastases that displayed similar morphology, though with slightly lower efficiency (Figures 3K–3O; Figures S3J and S3K; n = 4 mice). Given that most disseminating cells were Lgr5− and had the ability to initiate metastatic growth, our data reveal that the majority of metastases are seeded by Lgr5− cancer cells.

Lgr5+ Clones Have the Intrinsic Capacity to Restore the Cellular Hierarchy

Because growth of metastases seeded by Lgr5− cancer cells requires cells to become Lgr5+, we next addressed what is driving this plasticity. It has been suggested that cancer cells can acquire SC properties when they are surrounded by a SC niche, such as fibroblasts and immune cells, that provides stemness-inducing signals (Sato and Clevers, 2013). To test whether SC-inducing microenvironmental factors are deterministic for cancer cell plasticity, we set up an organoid-forming assay. In these pure epithelial culture systems, the three-dimensional (3D) microenvironment was mimicked by...
Basement Membrane Extract (BME) and medium providing all the essential SC-inducing factors present in the intestinal SC niche, such as Wnt, R-spondin, and EGF (Sato and Clevers, 2013). By simply removing one or more of these niche factors, it is possible to test whether intestinal epithelial cells and their dynamics are dependent on essential microenvironmental signals (Drost et al., 2015; Fuji et al., 2016). We isolated single Lgr5+ circulating cancer cells from the blood and tested whether they could be cultured in medium without any SC-inducing factors. Interestingly, in this minimal medium we were able to grow organoids from Lgr5+ circulating cancer cells, and also here we observed the appearance of Lgr5+ CSCs over time (Figures 4A and 4B). Moreover, these Lgr5+ cancer cells were functional SCs, as selective ablation of these cells by acute or chronic DT treatment in both organoid-derived and primary tumor-derived cells abolished organoid outgrowth (Figures 4C; Figures S4A–S4F). To exclude that residual growth factors in the BME matrix could trigger cancer cell plasticity in our culture system, we confirmed these data in an engineered 3D matrix consisting of growth factor-free synthetic hydrogel (Figures 4D and 4E). Finally, to test whether plasticity can be enhanced by microenvironmental factors that are released by cancer associated fibroblasts, we cultured Lgr5+ cells in minimal medium supplemented with HGF, FGF, IL4, and IL13 (Chen et al., 2018; Knuchel et al., 2015; Lenos et al., 2018; Todaro et al., 2007, 2014). Indeed, we observed enhanced plasticity (i.e., appearance of Lgr5+ cells) in medium supplemented with HGF and FGF compared with a basal level of plasticity in minimal CRC medium (Figure 4F). These data suggest that clones seeded by Lgr5+ cancer cells have the intrinsic capacity to re-establish the cellular hierarchy, even in conditions devoid of microenvironmental SC-inducing signals, and that this plasticity can be further enhanced by microenvironmental factors such as HGF and FGF.

**DISCUSSION**

Although epithelial hierarchy is often seen as a static, one-directional route from SCs to differentiated cells, our study draws a more dynamic picture of CRC, particularly of the metastatic setting. We showed that, in addition to CSCs, non-CSCs are also capable of giving rise to colorectal liver metastases. Although CSCs are present in the primary tumor and in the migratory population (Figure 2), we did not detect them in circulation. As already shown for healthy intestine (Sato et al., 2011; Serra et al., 2019), CSCs may lose Lgr5 expression and stemness once they are not supported by the surrounding SC niche and therefore may be forced into a Lgr5- state. Regardless of the origin of the disseminating non-CSCs, non-CSCs are in fact the major seeding cells, while conversion to CSCs at the metastatic site is required for efficient metastatic outgrowth. Therefore, cancer cell plasticity, rather than seeding of CSCs, is a key step in the formation of CRC metastases. To confirm the occurrence of plasticity in human tissues, we orthotopically transplanted human tumor organoids characterized by SC Ascl2 reporter that labels Lgr5+ CSCs (i.e., STAR probe; see Oost et al., 2018) (Figure S4G). Similar to their murine counterpart, these experiments showed that the major disseminating population (CTCs) was Lgr5- (Figure S4H). Moreover, human Lgr5+ cells efficiently seed metastases that contained Lgr5+ CSCs (Figure S4I). Plasticity in human cancer cells can happen independently of SC-inducing factors, as Lgr5+ CSCs appeared in organoids that were seeded by Lgr5+ cells and cultured in minimal CRC medium (Figure S4K). These experiments showed that differentiated cancer cells are able to form metastases, are plastic, and can re-establish the cellular hierarchy also in the human setting (Figures S4H–S4K). Cellular plasticity of epithelial cells has already been reported in healthy intestine (Ritsma et al., 2014; Tetteh et al., 2016; 574 Cell Stem Cell 26, 569–578, April 2, 2020.
Tian et al., 2011; van Es et al., 2012) and as recently reviewed by de Sousa e Melo and de Sauvage (2019). Upon targeted depletion of the healthy Lgr5+ SC population, more specialized Lgr5− cells enter the SC niche and are instructed to revert to Lgr5+ SCs, thereby restoring tissue homeostasis (Ritsma et al., 2014; Tetteh et al., 2016; Tian et al., 2011; van Es et al., 2012). The intestinal SC niche consists of supporting cells producing stemness-inducing factors, such as Wnt, R-spondin, and EGF. Similar plasticity processes have been observed in primary CRC tumors, and various microenvironmental signals have been highlighted as responsible for the switch of non-CSCs to CSCs (Lenos et al., 2018; Schwitalla et al., 2013; Vermeulen et al., 2010). However, our data also suggest that cancer cell plasticity can be triggered independently of stemness-inducing factors provided by SC niches. Nevertheless, we also show that factors released by fibroblasts, such as FGF and HGF, can enhance plasticity. Regardless of this enhancement, our data suggest that specifically targeting CSCs and/or the SC-inducing niche, as previously proposed (de Sousa e Melo and de Sauvage, 2019), may not be enough to completely prevent metastatic disease, which is the leading cause of cancer-related death. Instead, endogenous cellular plasticity should be co-targeted to inactivate any potential seeds of metastasis, and future studies should aim to uncover microenvironment-independent mechanisms leading to cell plasticity.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

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- **EXPERIMENTAL MODEL AND SUBJECT DETAILS**
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  - Human organoids

Figure 4. Disseminating Lgr5− Cancer Cells Undergo Niche-Independent Cellular Plasticity

(A) Representative example of organoid formation assay of sorted Lgr5− circulating cancer cells. Dashed lines highlight organoids edges. Arrowhead indicates the appearance of a Lgr5+ CSC. Scale bar, 20 μm.

(B) Quantification of cancer cell plasticity (i.e., emergence of Lgr5+ CSCs) in the organoid-forming assay of sorted Lgr5− circulating cancer cells grown in minimal CRC medium. Data are presented as mean ± SEM (n = 3 independent experiments).

(C) Clonogenicity assay in minimal CRC medium of sorted Lgr5+ CSCs and Lgr5− cancer cells derived from CRC organoid cultures. Single cells were subjected to either vehicle or diphtheria toxin (DT) treatment (n = 3 independent experiments). Data were collected 6 days after plating. Values are presented as mean ± SEM; ***p < 0.0001, calculated using the unpaired t test with Welch’s correction.

(D) Representative example of organoid formation assay of sorted Lgr5− cancer cells cultured in synthetic hydrogel. Dashed lines highlight organoids edges. Scale bar, 50 μm.

(E) Representative images of the effect of vehicle or diphtheria toxin (DT) on sorted Lgr5− CSCs and Lgr5− cancer cells cultured in synthetic hydrogel 6 days after plating (n = 3 independent experiments). Scale bar, 200 μm.

(F) Quantification of cancer cell plasticity (i.e., emergence of Lgr5+ CSCs) of sorted Lgr5− cancer cells grown in either minimal CRC medium (control) or medium containing FGF-2 (10 ng/mL), HGF (75 ng/mL), IL4 (20 ng/mL), IL13 (100 ng/mL), and IL4 and IL13. n = 3 independent experiments. *p < 0.05 and **p < 0.001, calculated using the unpaired t test. Data are represented as mean ± SEM.
AUTHOR CONTRIBUTIONS

A.F. performed the experiments. K.C.O. performed part of the revision experiments with the help of M.A., E.B., and S.J.E.S. L.S. helped with the in vivo experiments. L. Bruijs, L. Bornes, and D.S. helped with the sample preparation of the fluorescence-activated cell sorting (FACS) experiments. L.K. analyzed the single-cell sequencing data. J.M. performed the hydrogel-based assay. D.P. performed the in vitro DT cleaved caspase-3 assay. T.S., S.v.d.E., M.v.B., and D.B. operated the FACS. A.M.S. and J.W.M.M. contributed knowledge and reagents. K.C.O. and M.C.H. provided the organoid line. Experiments were designed with the help of N.V. and H.J.S. J.v.R. together with A.F. designed the experiments and supervised the study. A.F., S.I.J.E., and J.v.R. wrote the manuscript, and the manuscript was approved by all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.
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# STAR★METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Mouse monoclonal anti-ß-catenin | BD Biosciences | Cat# 610154; RRID: AB_397555 |
| Goat polyclonal anti-GFP | Abcam | Cat# ab6673; RRID: AB_305643 |
| Rabbit polyclonal anti-RFP | Rockland | Cat# 600-401-379; RRID: AB_2209751 |
| Mouse monoclonal CD41a, Biotin | Thermo Fisher Scientific | Cat# 13-0411-82; RRID: AB_763484 |
| Mouse monoclonal CD45, Biotin | Thermo Fisher Scientific | Cat# 13-0451-85; RRID: AB_466447 |
| Alexa fluor 488 donkey anti-goat | Abcam | Cat# ab150129; RRID: AB_2687506 |
| Alexa fluor 568 donkey anti-rabbit | Abcam | Cat# ab175470; RRID: AB_2783823 |
| Alexa fluor 647 streptavidin | Thermo Fisher Scientific | Cat# S-21374; RRID: AB_2336066 |
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| Tamoxifen | Merk | Cat# T5648 |
| Advanced DMEM F/12 | Thermo Fisher Scientific | Cat# 12634-010 |
| B27 | Thermo Fisher Scientific | Cat# 17504-044 |
| BME | R&D Systems | Cat# 3533-005-02 |
| Buprenorphine | Multidosis-Astfarma | N/A |
| Collagenase II | Thermo Fisher Scientific | Cat# 17101015 |
| Diphereria Toxin | Merk | Cat# D0564 |
| DNase I | Merk | Cat# 4716728001 |
| Fetal Bovine Serum | Merk | Cat# F7524 |
| FGF | Thermo Fisher Scientific | Cat# PHG0021 |
| HGF | R&D Systems | Cat# 294-HG |
| Hyaluronidase | Merk | Cat# HX0514-1 |
| IL13 | Peprotech | Cat# 214-13 |
| IL4 | Peprotech | Cat# 214-14 |
| Matrigel | Corning | Cat# 356231 |
| N-acetylcysteine | Merk | Cat# A9165 |
| Normal Goat Serum | Monosan | Cat# monx10961 |
| Puramatrix Hydrogel | Corning | Cat# 354250 |
| Purified fibronectin | Merk | Cat# FC010 |
| Purified laminin-111 | Merk | Cat# L2020 |
| Rat Tail High Concentrated Type I Collagen | Corning | Cat# 354249 |
| Recombinant human Noggin | PeproTech | Cat# 120-10C |
| Tissue-Tek OCT | Sakura | Cat# 4583 |
| TrypLE | Thermo Fisher Scientific | Cat# 12605-010 |
| Y-27632 | Abmole | Cat# M1817 |
| Dapi | Thermo Fisher Scientific | Cat# D1306 |

**Deposited Data**

| SINGLE CELL mRNA sequencing | This study | GSE143988 |
|**Experimental Models: Cell Lines** | | |
| Murine_VillinCre-ERT2; APC<sup>Cre<sup>,</sup></sup>, KRAS<sup>LSL-G12D</sup>;<sup></sup>, P53<sup>KO,KO</sup>, R26R-Confetti; Lgr5<sup>OTR-eGFP</sup> | This study | N/A |
| Human CRC | van de Wetering et al., 2015 | p19bT |

**Experimental Models: Organisms/Strains**

| Murine_VillinCre-ERT2; APC<sup>Cre<sup>,</sup></sup>, KRAS<sup>LSL-G12D</sup>;<sup></sup>, P53<sup>KO,KO</sup>, R26R-Confetti; Lgr5<sup>OTR-eGFP</sup> | This study | N/A |
| Human CRC | van de Wetering et al., 2015 | p19bT |

(Continued on next page)
LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Jacco van Rheenen (j.v.rheenen@nki.nl).

All unique/stable reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Generation of a colorectal cancer genetic mouse model

The VillinCre-ER<sup>T2</sup> mouse model (The Jackson Laboratory, cat. no. 20282) was combined with the APC<sup>fl/fl</sup>, KRAS<sup>LSL-G12D/+</sup>, TP53<sup>KO/KO</sup> mouse model in order to generate a VillinCre-ER<sup>T2</sup>; APC<sup>fl/fl</sup>; KRAS<sup>LSL-G12D/+</sup>; TP53<sup>KO/KO</sup>, a Tamoxifen-inducible triple mutant genetic mouse model. The color randomizer R26R-Confetti (gifted by the Clevers laboratory, Hubrecht Institute, NL (Snippert et al., 2010) allele and stem cell fluorescent reporter Lgr5 DTR-eGFP (gifted by De Sauvage laboratory, Genentech, USA (Tian et al., 2011)) were bred in to finally generate the VillinCre-ER<sup>T2</sup>; APC<sup>fl/fl</sup>; KRAS<sup>LSL-G12D/+</sup>; TP53<sup>KO/KO</sup>; R26R-Confetti; Lgr5<sup>DTR-eGFP</sup> genetic mouse model in a mixed background. In these mice Lgr5<sup>+</sup> cells (e.g., the stem cells in the intestine) are labeled with membrane-bound eGFP. Activation of the inducible Cre upon administration of Tamoxifen initiates formation of tumors which are restricted to the intestine and promotes the expression of one of the four Confetti colors, thereby specifically color-coding the tumor cells. Since tumors develop throughout the whole intestinal tract, these mice reach the humane endpoint 5 to 6 days after high dose (200mg/kg) Tamoxifen (Merk cat. no. T5648) injection.

Isolation and culturing of mouse organoids

8 weeks old VillinCre-ER<sup>T2</sup>; APC<sup>fl/fl</sup>; KRAS<sup>LSL-G12D/+</sup>; TP53<sup>KO/KO</sup>; R26R-Confetti; Lgr5<sup>DTR-eGFP</sup> transgenic mice were injected with 200mg/kg Tamoxifen and sacrificed 4 days after induction. Colorectal cancer organoids were isolated from two independent VillinCre-ER<sup>T2</sup>; APC<sup>fl/fl</sup>; KRAS<sup>LSL-G12D/+</sup>; TP53<sup>KO/KO</sup>; R26R-Confetti; Lgr5<sup>DTR-eGFP</sup> genetic mouse model in a mixed background. In these mice Lgr5<sup>+</sup> cells (e.g., the stem cells in the intestine) are labeled with membrane-bound eGFP. Activation of the inducible Cre upon administration of Tamoxifen initiates formation of tumors which are restricted to the intestine and promotes the expression of one of the four Confetti colors, thereby specifically color-coding the tumor cells. Since tumors develop throughout the whole intestinal tract, these mice reach the humane endpoint 5 to 6 days after high dose (200mg/kg) Tamoxifen (Merk cat. no. T5648) injection.

Acceptor mice for transplantation experiments

8 – 14 weeks old male and/or females NOD.Cg-Prkdc<sup>scid</sup>Il2rg<sup>tm1Wjl</sup>/SzJ (NSG, The Jackson Laboratory cat. no. 005557) mice were used as acceptors for subcutaneous injection, orthotopic transplantation and mesenteric vein injection. All experiments were performed in accordance with the Animal Welfare Committees of the Royal Netherlands Academy of Arts and Sciences and the Netherlands Cancer Institute, the Netherlands. Animals were kept at the Hubrecht animal facility in Utrecht or at the Netherlands Cancer Institute facility in Amsterdam, the Netherlands. For every experimental condition, at least 4 mice were used. No animals were excluded from analyses. When experiments consisted of multiple conditions, either males or females of the same litter were randomly assigned to each group.
**Human organoids**

The experiments were performed with patient-derived P19bT human organoids (van de Wetering et al., 2015), mutated in TP53, PIK3CA, BRAF, ERBB3, RNF43) constitutively labeled with H2B-mNeonGreen while cancer stem cells specifically express singleTomato-NLS driven by a ASCL2-responsive minigene (STAR, (Oost et al., 2018)).

**METHOD DETAILS**

**Orthotopic transplantation of colorectal cancer organoids**

Orthotopic transplantation of colorectal cancer organoids was performed as previously described (Fumagalli et al., 2017, 2018). In brief, the day before transplantation organoids were collected and dissociated into small cell clumps. About 250,000 cells were plated in 15 μl drops neutralized Rat Tail High Concentrated Type I Collagen (Corning, cat. no. 354249) and let to recover overnight at 37°C, 5% CO₂ in CRC medium containing 100 μM Y-27632 (Abmo, cat. no. M1817). At the day of transplantation, acceptor mice were sedated using isoflurane inhalation anesthesia (~2% isoflurane/O₂ mixture). Before surgery, the mice were treated subcutaneously with a single dose of buprenorphine (Buprecare, Multidosis-Astfarma, 3μg per mouse). The cecum was exposed through a midline abdominal incision and a collagen drop containing tumor cells was surgically transplanted in the caecal submucosa. The tumor growth was monitored weekly by abdominal palpation.

**Orthotopic transplantation of murine organoids**

Transplantations were performed as described above using 1.5 × 10⁵ human organoid-derived cells embedded in Rat Tail High Concentrated Type I Collagen (Corning, cat. no. 354249).

**Flow cytometry on organoids**

Colorectal cancer organoids were collected and mechanically dissociated with a fire-pointed glass pipet. Subsequently cells were incubated in TrypLE (Thermo Fisher Scientific, cat. no. 12605-010) mix containing 100 μM Y-27632 (Abmo, cat. no. M1817), 4μg/ml DNase I (Sigma-Aldrich, cat. no. 10104159001) for 10 minutes at 37°C and spun down. Cells were resuspended in FACS buffer containing 2% B27 (Thermo Fisher Scientific, cat. no. 17504-044), 1.25 mM N-acetylcysteine (Sigma-Aldrich, cat. no. A9165), 1% Noggin (PeproTech, cat. no. 120-10C), 100 μM Y-27632 (Abmo, cat. no. M1817), 4μg/ml DNase I (Merk, cat. no.4716728001), 0.01% FCS (Merk, cat. no. F7524) in PBS and filtered through a 100 μm strainer (BD Falcon). DAPI 1ng/ml (Thermo Fisher Scientific, cat. no. D1306) was added right before sorting. Cells were sorted on a FACS AriaII Special Ordered Research Product or a FACS Aria Fusion (BD Biosciences). The sort strategy is illustrated in Figures S1J–S1N. A broad FSC/SSC gate was followed by gates excluding doublets and selection of DAPI—living cells. RFP-Confetti + tumor cells were subdivided in (Lgr5)eGFP⁺ and (Lgr5)eGFP⁻ using stringent gating. For each experiment, the quality of the sorting was assessed by examination of the sorted cells at a confocal microscope (as in Figure S1O).

**Flow cytometry on primary mouse material**

Orthotopic colorectal cancer tumors were collected and minced on ice using sterile scalpels. The tumor mass was digested at 37°C in TrypLE (Thermo Fisher Scientific, cat. no. 12605-010) containing 10 μM Y-27632 (Abmo, cat. no. M1817), 4μg/ml DNase I (Merk, cat. no. 4716728001), 20 μg/ml Hyaluronidase (Merk, cat. no. HX0514-1). Digested cell clumps were incubated in TrypLE (Thermo Fisher Scientific, cat. no. 12605-010) mix containing 100 μM Y-27632 (Abmo, cat. no. M1817), 4μg/ml DNase I (Merk, cat. no. 4716728001) for 15 minutes at 37°C and spun down. The cell pellet was resuspended in FACS buffer (composition described above) and filtered through a 35 μm strainer (BD Falcon). DAPI 1ng/ml (Thermo Fisher Scientific, cat. no. D1306) was added right before sorting.

Blood was collected from mice bearing metastatic CRC either via portal vein puncture or right-ventricle cardiac puncture, 7-10 weeks after orthotopic colorectal cancer organoids transplantation. The red blood cells were depleted by NH₄Cl treatment. The remaining circulating tumor cells and immune cells were spun down (4 minutes 500 RCF at RT). Tumor cells and blood cells were blocked in 80% FACS buffer / normal goat serum (Monosan, monx10961) for 10 minutes on ice before labeling with the following antibodies: biotin-conjugated anti-mouse CD41 clone eBioMWReg30 (Thermo Fisher Scientific, cat. no. 13-0411-82) and anti-mouse CD45 clone 30-F11 (Thermo Fisher Scientific, cat. no. 13-0451-85). Secondary labeling was performed using streptavidin-conjugated AF647 (Thermo Fisher Scientific. no. S-32357). Cells were filtered through a 35 μm strainer (BD Falcon). DAPI 1ng/ml (Thermo Fisher Scientific, cat. no. D1306) was added right before sorting.

Cells were sorted on a FACS AriaII Special Ordered Research Product or a FACS Aria Fusion (BD Biosciences). The sort strategy is illustrated in Figures S1J and S1L and Figures S3A and S3D. A broad FSC/SSC gate was followed by gates excluding doublets and selection of DAPI—living cells. Afterward, immune cells and megakaryocytes were excluded in a dump channel. RFP-Confetti⁺ tumor cells were subdivided in Lgr5-eGFP⁺ and Lgr5-eGFP⁻ using stringent gating. The quality of the sorting was assessed afterward by examination of the cells at a confocal microscope.
Flow cytometry on human xenografts

Blood was collected from mice bearing human metastatic CRC either via right-ventricle cardiac puncture, 7-10 weeks after orthotopic colorectal cancer organoids transplantation. Samples were prepared and analyzed at the FACS as described above for experiments with the murine CRC model. H2B-mNeonGreen+ tumor cells were subdivided in STAR-NLS-singleTomato+ and STAR-NLS-singleTomato- using stringent gating. The quality of the sorting was assessed by post-sort purity check and examination of the cells at a confocal microscope.

Immunohistochemistry and fluorescent imaging

Tissues were fixed periodate-lysine-4% paraformaldehyde (PLP)(McLean and Nakane, 1974) buffer overnight at 4°C, incubated in 30% sucrose overnight at 4°C and embedded in Tissue-Tek (Sakura, cat. no. 4583). Organs were cryosectioned and staining were performed on 20-30 μm sections. The stainings were performed with the following primary antibodies: anti-ß-catenin clone 14 (BD Bioscience, cat. no. 610154), anti-GFP (Abcam, cat. no. ab6673), anti-RFP (Rockland, cat. no. 600-401-379). Stained sections were imaged with inverted Leica TCS SP5 and TCS SP8 confocal microscopes (Mannheim, Germany). All images were collected in 12 bit with 25X water immersion (HC FLUOTAR L.N.A. 0.95 W VISIR 0.17 FWD 2.4 mm) or 20X dry (HCX IRAPO N.A. 0.70 WD 0.5 mm) objectives.

In vivo time-lapse imaging of primary tumors

About 250,000 VillinCre-ERT2; APCCre; KRASLSL-G12D/+; TP53KO/KO; RFP-Confetti; Lgr5DTR-eGFP organoid-derived cells were mixed in 100 µl Matrigel (Corning, cat. no. 356231) and injected subcutaneously into the flank of recipient NSG mice. Experiments were performed on 9 NSG mice with organoid lines isolated from two independent VillinCre-ERT2; APCCre; KRASLSL-G12D/+; TP53KO/KO; R26R-Confetti; Lgr5DTR-eGFP transgenic mice. Specifically, 4 NSG were injected with the organoid line derived from the first GEMM donor, 5 NSG with organoid derived from the second GEMM donor. Time-laps intravital imaging was performed on tumors of about 125 mm3. During the entire procedure mice were sedated using isoflurane inhalation anesthesia (1.5% isoflurane/ O2 mixture). The tumor was surgically exposed, and the mouse was placed in a custom designed imaging box with its head in a facemask constantly delivering anesthesia. The imaging box and the microscope were adjusted at 36.5°C using a climate chamber. Intravital images were acquired with an inverted Leica TCS SP5 or TCS SP8 AOBS two-photon microscope (Mannheim, Germany) with a chameleon Ti:Sapphire pumped Optical Parametric Oscillator (Coherent Inc. Santa Clare, CA, USA). The microscopes are equipped with 2 non-descanned and 2 hybrid detectors: GFP and RFP were simultaneously excited at 940 nm and detected with hybrid detectors. Second harmonic generation (Collagen I, stroma). All images were collected in 12 bit and acquired with a 25X water immersion (HC FLUOTAR L N.A. 0.95 W VISIR 0.17 FWD 2.4 mm) objective.

Multi-day imaging of liver metastasis

Lgr5+ cancer cells were seeded into the liver parenchyma of acceptor mice either by mesenteric vein injection (of about 400,000 Lgr5+ sorted cells) or direct liver parenchyma injection (of about 60,000 Lgr5+ sorted cells). For the diphertheria toxin experiment mice were treated every other day with 50 μg/kg Diphertheria Toxin (Merk, cat. no. D0564) administered via intraportal injection every other day. 6-8 days after cells seeding, an abdominal imaging window was applied on the liver of the acceptor mice (Ritsma et al., 2013). Before surgery the mice were treated with a sub-cutaneous injection of buprenorphine (Buprecare, Multidosis-Astfarma, 3mg per mouse). The surgery was performed under aseptic conditions. Animals were sedated with ~2% isoflurane/compressed air mixture. After surgery, the mice were kept at 37°C until fully recovered. For every imaging session, mice were sedated using isoflurane inhalation anesthesia (~1.0% isofluorane/ compressed air mixture). The mice were placed in a custom designed imaging box while kept under constant anesthesia. The imaging box and the microscope were adjusted at 36.5°C using a climate chamber. Between the imaging sessions, mice were let recover in their cage. Intravital images were acquired with an inverted Leica SP8 Dive system (Mannheim, Germany) with a MaiTai eHP DeepSee laser (Spectra-Physics). The imaging areas were retraced in subsequent imaging sessions by storing the stage coordinates, and by visual landmarks such as blood vessels as described in43. The microscope is equipped with a 4Tune non-descanned detector configured with 4 hybrid detectors: Lgr5-DTR-eGFP, Confetti-RFP were simultaneously excited at 940 nm and detected with non-descanned hybrid detectors, together with second harmonic generation (Collagen I, stroma). All images were collected in 12 bit and acquired with a 25X water immersion objective (HC FLUOTAR L 25X/0.95 W VISIR 0.17)

Mesenteric vein injection

FACS-sorted cells were resuspended in 100 µl of sterile PBS and injected in the mesenteric vein of acceptor mice as previously described (van der Bij et al., 2010). Mice were sedated isoflurane inhalation anesthesia (~2% isoflurane/O2 mixture). Before surgery, the mice were treated with a sub-cutaneous dose of buprenorphine (Buprecare, Multidosis-Astfarma, 3mg per mouse). After surgery, the mice were kept at 37°C until fully recovered.
**Intrahepatic injection**

FACS-sorted cells were resuspended in 100 μl of sterile Matrigel (Corning, cat no. 356231) and PBS mix (2.5:1) and injected into the liver parenchyma. The surgical procedure was performed under isoflurane inhalation anesthesia (~2% isoflurane/O₂ mixture). Before surgery, the mice were treated with a sub-cutaneous dose of buprenorphine (Buprecare, Multidosis-Astfarma, 3mg per mouse). After surgery, the mice were kept at 37°C until fully recovered.

**Organoid formation assays**

Lgr5⁺ cancer stem cells, Lgr5⁻ cancer cells from either organoids or primary tumors and Lgr5⁻ circulating tumor cells were isolated by FACS according to their eGFP expression level as described before. Lgr5⁺ cancer stem cells, Lgr5⁻ cancer cells from either organoids or primary tumors were seeded in 20 μl BME (R&D systems, cat. no. 3533-005-02) drops containing 500 cells/drop while Lgr5⁻ circulating tumor cells were seeded in 20 μl BME drops containing 100 cells/drop. Cells were cultured in CRC medium (see above) containing 100 μM Y-27632 (Abmole Bioscience, cat. no. M1817). Cells were imaged daily with an inverted Leica TCS SP5 confocal microscopes (Mannheim, Germany). All images were collected in 12bit with a 25X water immersion (HC FLUOTAR L N.A. 0.95 W VISIR 0.17 FWD 2.4 mm) or with a 20X dry (HCX IRAPO N.A. 0.70 WD 0.5 mm) objective. Enhancement of plasticity by microenvironmental factors was tested by collecting 10000 viable Lgr5⁻ single cells via FACS and plating them in BME (R&D systems, cat. no. 3533-005-02) over three wells with a density of 50-300 cells per well of a glass-bottom 384-well plate (Corning, cat. no. 4581). Organoids were supplemented with either minimal culture medium (control), or medium including 75 ng/ml HGF (R&D Systems, cat. No. 294-HG), 10ng/ml FGF-2 (Thermo Fisher Scientific, cat. no. PHG0021), 20ng/ml IL4 (Peprotech, cat. no. 214-14), 100 ng/mL IL13 (Peprotech, cat. no. 214-13). Organoids were imaged live using a Leica-based spinning disk confocal microscope equipped with an Andor Dragonfly system, using a water 25x objective (HC-Fluotor, N.A. 0.95) with the Argon-laser of 488nm and the Diode-laser of 561nm using a 40μm pinhole and Andor scMOS Zyla 4 2p camera. While kept at 37°C and and 5% CO2 overflow. Organoids were scored at day 0 and after 5 (Figure 4F) or 10 days (Figure S4K) of culture for expression of Lgr5⁺eGFP.

**In vitro Diphertheria Toxin treatment**

Lgr5⁺ cancer stem cells, Lgr5⁻ cancer cells from either organoids or primary tumors were seeded in BME as described above. Diphertheria Toxin (100ng/ml, Merk, cat. no. D0564) or vehicle (demi-water) were added to the CRC culture medium (see above) containing 100 μM Y-27632 (Abmole Bioscience, cat. no. M1817). The medium was refreshed every other day.

**In vivo Diphertheria Toxin treatment**

50 μg/kg Diphertheria Toxin (Merk, cat. no. D0564) or vehicle (demi-water) were administered via intraperitoneal injection every other day.

**Hydrogel assay**

Organoid-derived Lgr5⁺ and Lgr5⁻ were sorted as described above, resuspended in 20% sucrose and mixed with Puramatrix Hydrogel (3mg/ml, Corning, cat. no. 354250), purified fibronectin (0.25mg/ml, Merck, cat. no. FC010), purified laminin-111 (50 μg/ml, Merck, cat. No. 354250), and PBS mix (2.5:1). The mixture was plated as a drop and gelation of the hydrogel was initiated by carefully adding advanced DMEM/F12 medium (adDMEM/F12; Thermo Fisher Scientific, cat. no. 12634-010). Hydrogels were allowed to form for 30 min at 37°C after which the medium was changed to full organoid growth medium with 100 μM Y-27632 (Abmole Bioscience, cat. no. M1817). Organoid medium was refreshed every two-three days.

**Comparison of Lgr5⁺ and Lgr5⁻ metastatic efficiency**

Lgr5⁺ and Lgr5⁻ cells were isolated from 4 different murine primary tumors with stringent gating (sorted strategy described above, Figure S1J and Figure S3A) and post-sort purity was assessed by resorting a fraction of every sorted sample. Only samples above 98% purity used for further in vivo experiments. Cells were inoculated via mesenteric vein injection as described above. Every in vivo biological replicate included: 1. a recipient mouse injected with 50,000 Lgr5⁺ cells; 2. a recipient mouse injected with 50,000 Lgr5⁻ cells; 3. when the purity control is between 98%–100%, a recipient mouse injected with a purity control (Lgr5⁻ cells), established via the percentage of cells falling out from the stringent Lgr5⁻ gate during post-sort purity analysis (i.e., cells that might have mistakenly been sorted as Lgr5⁻). Four weeks after injection the mice were sacrificed and the livers were inspected for presence of metastases under a fluorescence-stereo microscope (Leica). Importantly, none of the mice injected with purity controls showed formation of metastases. Tissues were fixed periodate-lysine-4% paraformaldehyde (PLP) buffer (McLean and Nakane, 1974) overnight at 4°C, incubated in 30% sucrose overnight at 4°C and embedded in Tissue-Tek (Sakura, cat. no. 4583).

**Comparison of STAR⁺ and STAR⁻ metastatic efficiency**

Experiments were performed as described above for murine cancer cells. Human STAR⁺ and STAR⁻ cells were isolated by FACS and post-sort purity was assessed by resorting a fraction of every sorted sample. 50,000 STAR⁺ or STAR⁻ were injected in the mesenteric vein of acceptor mice and a mouse injected with a purity control (STAR⁺ cells), established via the percentage of cells falling out from
the stringent STAR gate during post-sort purity analysis was also included every time that the experiment was performed. Four weeks after injection the mice were sacrificed and the livers were inspected for presence of metastases under a fluorescence-stereo microscope (Leica). Importantly, none of the mice injected with purity controls showed formation of metastases. Tissues were fixed periodate-lysine-4% paraformaldehyde (PLP) buffer (McLean and Nakane, 1974) overnight at 4°C, incubated in 30% sucrose overnight at 4°C and embedded in Tissue-Tek (Sakura, cat. no. 4583).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistics was performed using GraphPad Prism. Paired or unpaired t test (with Welsh correction) was used when data showed normal distribution (verified with normality tests, provided by GraphPad Prism), whereas Mann-Whitney U test was used for data that did not display normal distribution. Adoption of one statistical test or the other is indicated for each experiment in the figure legend.

**Flow cytometry**

Data were manually analyzed using FlowJo 10.6.1 (https://www.flowjo.com/).

**Fluorescent images (fixed samples)**

Images were converted to RGB using Fiji (https://imagej.nih.gov/ij/) and only if necessary, corrected for bleed through, smoothened, cropped, rotated and contrasted linearly. Images were quantified using Fiji.

**Intravital imaging of primary tumors**

Videos were corrected for XYZ drift using a custom-made software (written in Visual Studio 2010 in the .NET framework for Visual Basics. Codes are available on request from J.v.R.) and if needed additionally with match motion compensation software program to correct for rigid and elastic tissue deformation as extensively described by Noordmans and colleagues (Verdaasdonk, 2006). The Z correction of each time point was done manually and the XY correction was done automatically as follows: all images over time were corrected to the image of the first time point. The correction is based on finding the highest spatial correlation between the images. The spatial correlation for pixels in which the gray value in both images are > 0 were analyzed with the Pearson’s correlation coefficient. Then, the image that needs to be corrected was moved one pixel to the left, and the correlation was recalculated. If this led to a higher correlation, all pixels in the image were moved one pixel to the left. The correlation was calculated for a move of the corrected image to the left, right, up or down. This procedure was repeated iteratively until the coefficient no long increased. After the full correction, the correctness of the XY correction was visually inspected and manually adjusted if required. To experimentally test the success of the XYZ correction, the position of randomly picked cells were tracked, and if the movement is less than half a cell diameter in 4 hours, we called the correction successful (see examples in Figure 2 D).

Images were further processed using Fiji. Frames were converted to RGB and only if necessary, corrected for bleed through, smoothened, cropped, rotated and contrasted linearly. Migration was quantified manually using Fiji. Statistical analysis was performed in GraphPad Prism using non-parametric Mann-Whitney U test or unpaired t test with Welch’s correction.

**Intravital imaging of liver metastasis**

Images were processed and analyzed using Fiji. Frames were converted to RGB and only if necessary, corrected for bleed through with AND/OR function, smoothened, cropped, rotated and contrasted linearly.

**Organoid formation assays**

Organoids were counted manually by scoring them using a stereomicroscope. Fluorescent images were processed using Fiji. Frames were converted to RGB and only if necessary, corrected for bleed through with AND/OR function, smoothened, rotated and contrasted linearly. Lgr5-eGFP expression was scored manually.

**Metastasis scoring**

Whole livers were sectioned in 50 μm thick slices and metastasis imaged with an inverted Leica TCS SP5 or a Leica TCS SP8 confocal microscope (Mannheim, Germany). All images were collected in 12 bit with 25X water immersion (HC FLUOTAR L N.A. 0.95 W VISIR 0.17 FWD 2.4 mm). Images were converted to RGB using Fiji and only if necessary, corrected for bleed through, smoothened, cropped, rotated and contrasted linearly. Images were quantified using Fiji. Quantification was blinded concerning to the data group allocation. Statistical analysis was performed with GraphPad Prism.

**Single-cell mRNA sequencing**

Single-cell mRNA sequencing was performed using Sort-seq as described in Muraro et al. (Muraro et al., 2016). Single-cell libraries were sequenced on Illumina NextSeq500 with 75bp paired end reads. Read1 contains the cell barcode and Unique Molecule Identifier, read2 was mapped to the mm10 RefSeq transcriptome using Burrows-Wheeler Aligner with standard parameters. Seurat
(R package Seurat (Stuart et al., 2019)) was used for analyzing the single cell data. Single cell transcriptomes were filtered for cells that had at least 1,000 unique transcripts and subsequently log-normalized to 10,000 transcripts per cell. Clustering was performed on the first 13 principal components and clusters were identified with a resolution of 1. Differential gene expression analysis was performed with the roc test. Cell type identification was performed based on the differentially expressed genes between the clusters.

DATA AND CODE AVAILABILITY

The accession number for the single cell transcriptomic data reported in this paper is GEO: GSE143988.