



Spatiotemporal regulation of clonogenicity in colorectal cancer xenografts

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Cancer evolution is predominantly studied by focusing on differences in the genetic characteristics of malignant cells within tumors. However, the spatiotemporal dynamics of clonal outgrowth that underlie evolutionary trajectories remain largely unresolved. Here, we sought to unravel the clonal dynamics of colorectal cancer (CRC) expansion in space and time by using a color-based clonal tracing method. This method involves lentiviral red-green-blue (RGB) marking of cell populations, which enabled us to track individual cells and their clonal outgrowth during tumor initiation and growth in a xenograft model. We found that clonal expansion largely depends on the location of a clone, as small clones reside in the center and large clones mostly drive tumor growth at the border. These dynamics are recapitulated in a computational model, which confirms that the clone position within a tumor rather than cell-intrinsic features, is crucial for clonal outgrowth. We also found that no significant clonal loss occurs during tumor growth and clonal dispersal is limited in most models. Our results imply that, in addition to molecular features of clones such as (epi-)genetic differences between cells, clone location and the geometry of tumor growth are crucial for clonal expansion. Our findings suggest that either microenvironmental signals on the tumor border or differences in physical properties within the tumor, are major contributors to explain heterogeneous clonal expansion. Thus, this study provides further insights into the dynamics of solid tumor growth and progression, as well as the origins of tumor cell heterogeneity in a relevant model system.

colorectal cancer | tumor growth | intratumor heterogeneity | cancer evolution | cancer stem cells

Solid malignancies result from the accumulation of genetic aberrations that provide cells with a clonogenic advantage over their environment; for example, by promoting proliferation or reducing cell death (1–3). However, our incomplete knowledge of the quantitative effects of these oncogenic events and the fundamental dynamics of tumor expansion have so far precluded a thorough understanding of the dynamics of tumor evolution. For example, it remains unresolved what the effective population size is that drives long-term tumor expansion and progression (4, 5). Do rare cancer stem cells exist, or are all cells capable of driving tumor growth? In addition, the impact of the geometry of tumor expansion on clonogenic outgrowth is a topic of great relevance (6). In contrast to hematological malignancies, cells in solid cancers directly compete for space and nutrients. Furthermore, the dynamics of tissue turnover and the geometry of competing clones are predicted to directly impact on evolutionary trajectories (7, 8). Intratumor heterogeneity, which contributes to resistance to therapies and poor outcome, is a direct consequence of the concepts introduced above and a better understanding of these is essential to improve patient outcomes

(9, 10). Recently, it was suggested in the *big-bang* model of colorectal cancer (CRC) evolution that spatial separation of competing clones results in a largely neutral competition, and that the variation in clone sizes within cancers reflects the age of the clone rather than the relative clonogenic advantage of the unique molecular properties of that lineage (11). However, this model did not consider the possible heterogeneity in clone sizes that could result from a heterogeneous clonogenicity instilled by the specific geometry of the tumor tissue and its microenvironment.

Here we set out to investigate the impact of the environment on clone size variation in primary xenograft models of human CRC. We employed the lentiviral gene ontology (LeGO) method to spatially trace clone lineages within tumors by their unique red-green-blue (RGB) color-coding (12). This improves on previous barcoding studies from which spatial information is absent (13, 14). We found that injection of homogenous populations of cancer cells results in extensive heterogeneity in

Significance

Colorectal cancer (CRC) is a heterogeneous disease, with significant variation in genotype and phenotype within each individual tumor. This intratumor heterogeneity emerges during tumor development due to clonal evolution and in part can explain therapy resistance in CRC. However, a detailed understanding of the spatiotemporal development of tumors underlying cancer evolution and intratumor heterogeneity remains absent. Here, we use lineage-tracing experiments of human CRC cells transplanted into immunocompromised mice, in combination with computational modeling, to study the growth mode of CRC. We found that the clonal position is crucial for clonal outgrowth. This demonstrates that, in addition to the genetic composition, the environment and the geometry of tumor growth play a significant role in shaping tumor evolution.

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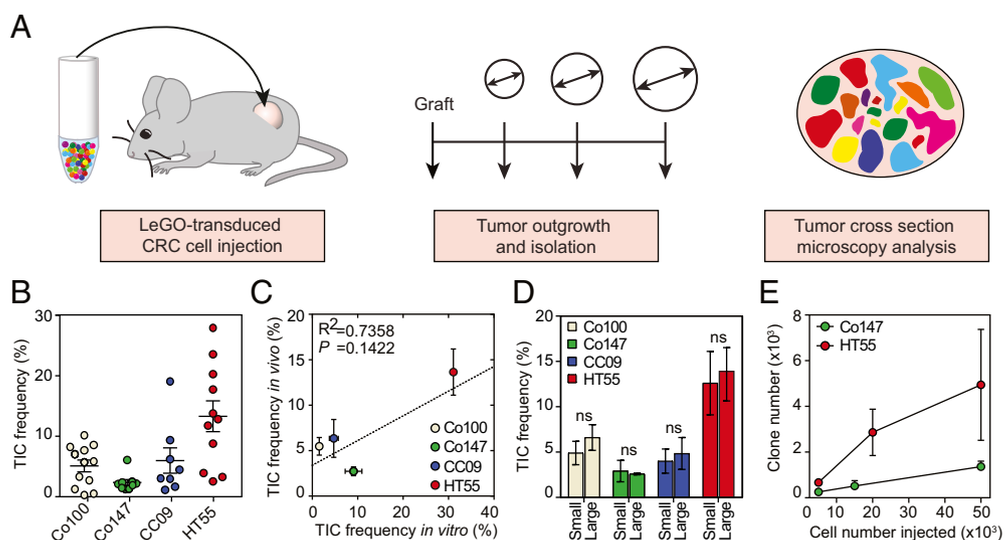
of cancer cells within xenografts is rather limited. These findings are in line with the results of multiregion sequencing analysis that indicate that private mutations are often detected in separate tumor regions and show that the LeGO xenografts are appropriate model systems to study colon cancer growth and progression (11, 15, 16).

Effective Population Size of Colorectal Cancers. Previous work has revealed that not all CRC cells have an equal ability to initiate tumor growth in immunocompromised mice. It has been established that cells that express markers of immature cell types such as *AC133*, or that present with high Wnt pathway activity, have a superior capacity to induce colon cancer xenografts (17–19). These studies have mostly been performed using limiting dilution assays, in which a decreasing number of cells is injected, which then allows for the calculation of the proportion of cells within a population capable of initiating xenograft growth. However, it remains unclear how the reduced cell numbers impact on the ability to initiate tumors; for example, due to the lack of para- and juxtacrine signaling input. Using our model system, we now have the ability to directly determine the proportion of injected cells that contributes to tumor initiation using an equal number of injected cells (Fig. 3A). By high-resolution analysis of xenograft tissue, and quantification of the number of clones, we could estimate the number of injected cells that actively grew out. We found that *in vivo* dispersal of cells is limited (Fig. 2), and this allowed us to identify each clone that resulted from the expansion of an injected cell as a connected region of cells with the same color. We found that the percentage of injected cells contributing to tumor initiation ranged between ~2–20%. The highest proportion of tumor-initiating cells (TICs) was detected in the serum cultured cell line (HT55), and the primary cell cultures displayed more limited clonogenic potential during the initiation phase (Fig. 3B). Additionally, comparison of the TIC frequency in the LeGO model versus the *in vitro* limiting dilution assay revealed only a weak correlation between both methods for determining the clonogenicity of tumor cells (Fig. 3C). We found that the limiting dilution assay could both under- and overestimate the clonogenic cell frequency. This indicates that in some models, injection of a larger cell number suppresses outgrowth of cells, as for example in Co147, while in another model the coinjected cancer cells promote outgrowth of cells (e.g., Co100). Hence, we suggest that the interpretation of data derived from an *in vitro* assay about the clonogenic

capacity of tumor cells should be done with caution. Importantly, the estimated proportion of cells initiating tumor growth was independent of the tumor volume analyzed (Fig. 3D) and actual number of cells injected (Fig. 3E), indicating that clones that contribute to tumor initiation permanently reside in the tumor tissue, and are not lost due to competition for example, making this assay robust to analyze different time points or tumor volumes.

Growth Dynamics of Colorectal Cancer Tissue. To elucidate the underlying dynamics of colon cancer tissue expansion, we mixed LeGO cultures with nontransduced cultures. This had the benefit that LeGO clones were better separated and allowed us to use a semiautomated image analysis pipeline to quantify the clone sizes within the whole xenograft tissues (Fig. 4A and *Material and Methods*). Analysis of hundreds of clones within tumors of different sizes revealed that on average the median clone size increased as expected in an expanding tissue (Fig. 4B). More interestingly, we detected that the heterogeneity in clone sizes was very large, and many clones remained small and did not seem to significantly contribute to tumor expansion (Fig. 4B and *SI Appendix, Fig. S3*). When plotting the relation between the proportion of clones that contribute to which fraction of tumor volume, we indeed detected that a small number of clones is responsible for the majority of the tumor growth (Fig. 4C). Furthermore, in larger tumors, the trend toward a relatively small number of clones driving tumor expansion is increased (Fig. 4C). It has been observed previously, by using genetic clonal tracing strategies in solid tumors, that not all cells contribute equally to cancer growth. In those studies, this heterogeneity was attributed to the intrinsic differences in clonogenic potential of cells, resulting from different cell states, i.e., stem cells vs. differentiated cells (13, 14). We now have the ability to evaluate this by studying the configuration of clones within the tissue. As is immediately apparent from the images from whole LeGO xenograft sections, there is a clear relationship between the position of the clone and its size in all cancer models studied (Fig. 4A and D and *SI Appendix, Fig. S2 A–D*). Larger clones are predominantly located at the xenograft edges, implying that competition for an optimal location instead of the intrinsic properties of clones defines which clones drive expansion in this model. This implies that before clones get into direct competition, i.e., before an established tumor has formed from the

Fig. 3. Clone numbers are stable during tumor growth. (A) Schematic model of the experimental setup for clonal outgrowth quantifications per LeGO-transduced colon cancer cell line. A fixed cell number was injected s.c. for each cell line and tumors were isolated at different tumor volumes. (B) Graph showing the TIC frequency of Co100-, Co147-, CC09-, and HT55-derived tumors. Each dot represents one tumor (a minimum of $n = 8$ per cell line), error bars represent SEM. (C) Graph showing the correlation between the average TIC frequency in an *in vitro* limiting dilution assay versus *in vivo* LeGO experiments for the indicated cell lines. Pearson correlation. Error bars represent SEM. (D) Bar graph showing the TIC frequency of Co100-, Co147-, CC09-, and HT55-derived tumors for two subgroups; tumors indicated as small having an average volume of 170 mm³ and tumors indicated as large having an average volume of 525 mm³. Error bars represent SEM, ns, not significant, Student's *t* test. (E) Graph showing the number of clones as determined by image analysis for Co147 and HT55 xenografts (minimum of $n = 3$ tumors per injected cell number for each cell line) that were derived from injections with different cell numbers, error bar represents SEM.



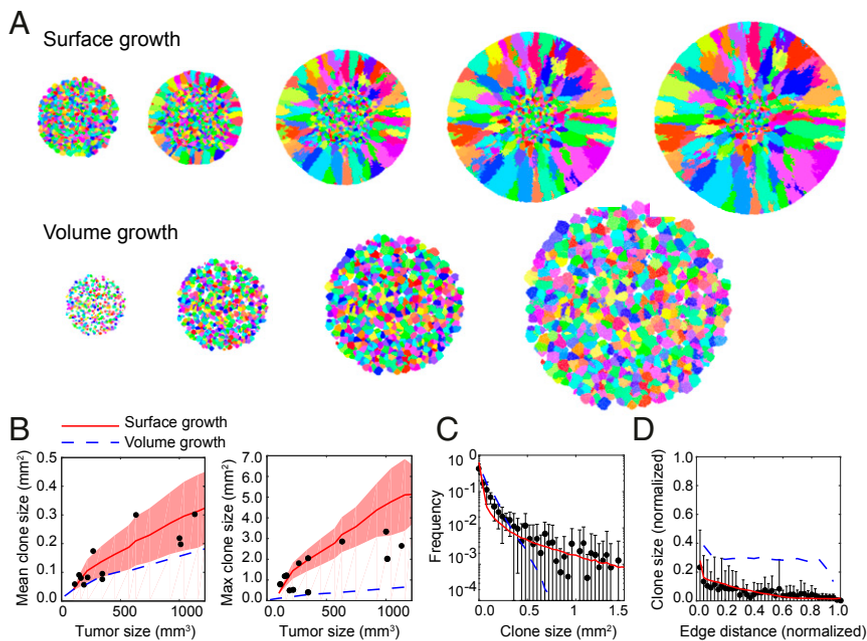


Fig. 5. Clonal expansion is highly dependent on clone location. (A) Cross-sections of tumors of increasing size, simulated with stochastic growth confined to the surface (Top) or throughout the entire tumor (volume growth, Bottom). Tumor sizes (in million cells) from left to right for, respectively, the Top and Bottom: 50, 100, 300, 600, 900, and 25, 150, 600, and 1400. (B–D) Comparison of experimental data (black dots) with model predictions of surface (red lines) and volume growth (dashed blue lines). (B) Mean (Left) and maximum (Right) clone size per xenograft. The red shade displays the SD of numerical mean and maximum clone sizes obtained from cross-sections taken at different positions of simulated tumors growing on the surface. (C) Distribution of sectional clone sizes. Error bars represent SD between xenografts. (D) Average clone size versus distance to tumor edge. Clone sizes and distances are normalized to the maximum of each cross-section. Error bars represent SD of the experimental data.

limited in colon cancer xenografts. This observation has important implications, as it strengthens the notion that clones in colon cancer tissue expand in parallel when residing in an identical environment. Also, this provides geometrical support to the idea that a large proportion of CRCs display predominantly neutral evolution and that no large selective sweeps follow emergence of novel, more aggressive clones in established cancers.

Previous short-term lineage tracing experiments by us and others indicated that clonal expansion is spatially regulated in established CRC xenograft tissue (7, 8). In these studies, clones expressing one or a few different colors were induced in established tumors. The small number of colors in these experimental systems prevented long-term tracing of individual clones because growth and merging of neighboring clones are hard to distinguish with a limited resolution. In the current study, we have overcome this limitation, which has enabled us to study long-term clonal dynamics in xenografts of established CRCs. The combination of long-term lineage tracing and computational modeling revealed that clone size heterogeneity can be fully explained by externally driven growth at the leading edge of the tumor. In contrast, previous barcoding studies in human xenografts indicated that distinct types of stem-like cells in colon cancer xenografts exhibit distinct repopulating features caused by intrinsic functional differences in the self-renewal and tumor-forming capacity of tumor cells (13, 14). Our data provide a radically different explanation for these observations by showing that these different contributions to tumor expansion following xenograft propagation are spatially orchestrated rather than intrinsically defined. Of note, studying clonal dynamics of early phases of premalignant expansion and conversion toward CRC falls beyond the scope of this study, but in these early stages the clonal dynamics are possibly more defined by genomic differences between clones. In addition, our model system lacks a functional immune system and immune effects are not captured in our study. Notwithstanding, we conclude that within the time frame and spatial scale of our experimental setup, the tumor environment is a dominant factor in shaping CRC growth and progression, as expansion mainly occurs at the tumor edge.

Several factors can explain the observed growth at the tumor edge. For example, the enrichment for stroma and secretion of stromal factors can drive clonogenic expansion at the leading edge.

The increased interstitial pressure within the xenograft centers is also likely to contribute. Importantly, manipulation of these factors could yield new therapeutic avenues to improve treatment; for example, Osteopontin would be an interesting target to further investigate (8). Furthermore, targeting the intercellular machinery associated with clonogenic potential, as a cell state enabled by the environment, is another strategy that could be developed to improve the prognosis of patients with solid cancers.

Materials and Methods

Cell Culture. Human primary colon cancer cultures were established as described previously (18). Cultures were isolated from patients with colorectal cancer with approval of the medical ethical committee of the Academic Medical Center and University of Palermo. Primary cell lines are cultured in polyHEMA [Poly(2-hydroxyethyl methacrylate; Sigma) coated flasks (Corning) to allow spheroid growth. Advanced DMEM/F12 (Life Technologies) culture medium is used, which is supplemented with N-2 (Life Technologies), L-glutamine, glucose, HEPES, heparin, insulin, epidermal growth factor (EGF), and basic fibroblast growth factor (bFGF) as described previously (18). The primary human PDAC culture 067 was established as described previously (22) and cultured in Iscove's modified Dulbecco's medium (Life Technologies) supplemented with 8% FBS and L-glutamine. DLD1 (ATCC) and HT55 (Sanger Institute) cells were cultured in DMEM/F12 (Life Technologies) supplemented with 8% FCS (Life Technologies). Capan-2 (ATCC) was cultured in DMEM (Life Technologies) supplemented with 8% FCS. Cell lines were authenticated by short tandem repeat profiling in combination with mutation analysis and have been regularly tested for mycoplasma infection.

Multicolor Marking. Cell lines were simultaneously transduced with three different constructs according to a previously published protocol (12). The following lentiviral gene ontology (LeGO) vectors were used: LeGO-C2 (27339), LeGO-V2 (27340), and LeGO-Cer2 (27338) (Addgene). In short, 50,000 single cells were seeded in a 24-well plate in 500 μ L culture medium in the presence of 8 μ g mL⁻¹ polybrene (Sigma). Lentivirus containing the three vectors was added in a volume that ensured ~60% transduction rate of each vector. Plates were centrifuged for 1 h at 24 °C and incubated overnight at 37 °C. Transduction rates were analyzed by flow cytometry after 3 d. After transduction, cell lines were passaged in a low dilution and for a maximum of five passages before in vivo use.

Flow Cytometry. Flow cytometry was performed on a Fluorescence Activated Cell Sorting (FACS) Aria SORP (BD Biosciences) machine with 405-, 488-, and 561-nm lasers. Data were analyzed with the FlowJo (FlowJo LLC) software.

Vector Expression. Vector integration stability was analyzed by FACS. DLD1 cells were transduced with the LeGO system and then single-cell sorted. Single-cell clone cultures were expanded and passaged twice a week. Upon passaging the expression of Cerulean, Venus and mCherry were analyzed by FACS. Every cell line was analyzed at least twice in a 12-wk follow-up period.

In Vivo Experiments. The Animal Experimentation Committee at the Academic Medical Center in Amsterdam has approved all in vivo experiments (DEC103181 and DEC102348) and all animal experiments were performed according to the national guidelines. Female nude (Hsd:ATHymic Nude-Foxn1^{nu}) mice (6–12 wk old) were obtained from Envigo. Animals were randomly assigned to experimental groups; no blinding was applied for these experiments. Animal exclusion was performed when no tumor growth appeared.

Xenograft Studies. Xenograft tumors were generated by injecting 10,000 (CC09) or 50,000 (Co100, Co147, and HT55) human colon cancer cells in a mixture of medium and Matrigel (Corning) in a 1:1 ratio with a cell density of around 400–1,000 cell/ μ L. Cells were injected s.c. into the flanks of nude mice. Tumor growth was measured manually twice a week using a caliper. Mice were killed based on tumor size at various time points to isolate tumors. After isolation tumors were fixed using 4%-paraformaldehyde in PBS solution overnight at 4 °C followed by preservation in a 20% sucrose solution for 12 h at 4 °C. Tumors were split into two equally sized parts and 10 μ m-thick frozen tissue sections were collected from the tumor center.

In Vivo Transplantation Assay. Center and edge (<0.5 mm from tumor border) located cells were isolated from freshly collected xenografts by using razor blades. Immediately after tissue collection, cells were dissociated by using medium containing collagenase (Roche) and hyaluronidase (Sigma) at 37 °C for 1 h. Before injection, cells were filtered using a 70 μ m cell strainer, and dead cells were removed by 7-AAD staining (BD Biosciences) by using FACS. For each group, 1,000 cells were injected s.c. into the flanks of nude mice ($n = 3$), and tumor growth was measured twice a week.

Copy Number Analysis. DNA was extracted using the NucleoSpin Tissue kit (Bioké) following the manufacturer's procedure. To extract DNA from the inside and outside of tumors, we first mechanically separated the two regions. Shallow sequencing and data analysis were performed as previously described (23).

Limiting Dilution Assay. Cells were dissociated and plated in 96-well plates (Corning) using SH800 Cell Sorter (Sony) in a limiting dilution manner at 1, 2, 4, 8, 16, 24, 32, 64, 128, 256 cells per well. Clonal frequency and significance were determined using the Extreme Limiting Dilution Analysis (ELDA) "Limdil" function (24).

Imaging. Frozen tissue sections were imaged by an EVOS FL Cell Imaging System (Thermo Fisher Scientific). Sections were covered with ProLong Gold Antifade Mountant (Thermo Fisher Scientific) to ensure fluorescent signal preservation. Whole tumor sections were scanned for mCherry, Venus, and Cerulean by using the following LED light cubes; TexasRed (excitation 445/45 and emission 510/42 nm), YFP (excitation 500/24 and emission 524/27 nm), and CFP (excitation 585/29 and emission 624/40 nm). For high-resolution imaging, a SP8-X confocal microscope (Leica) with the Leica Application Suite-Advanced Fluorescence software was used.

Image Analysis. Automated clone size quantification and localization was performed on whole tumor cross-sectional slides imaged by fluorescence microscopy and converted to .tiff file format with a custom written MATLAB program. Boundaries of connected regions with the same color and cross-sections were manually highlighted for accurate tracking of clone position and size. Connected regions with the same color, but separated by >10 cell diameters were considered as separate clones. The number of mixed clones was identified manually.

Spatial Model for Tumor Growth. We adapted the 3D spatial model we recently introduced for tumor evolution, for direct comparison with the xenograft data (6). In short, in this model tumor cells occupy sites of a regular 3D lattice. To simulate growth, iteratively a random cell which has at least one of the neighboring sites (Von Neumann neighborhood) vacant, replicates to a randomly chosen vacant neighbor site. A detailed description of the computational modeling, a description of the different model versions, and how numerical data are compared with the experimental data can be found in *SI Appendix, Computer Models*.

Statistical Analysis. Sample sizes, statistical tests, and definitions of error bars are indicated in the figure legends and calculated using GraphPad Prism 7 or MATLAB. All statistical tests were two-sided. The between-group variances were similar and the data were normally distributed. *P* values of <0.05 were considered significant.

Data Availability. Source data for Figs. 4 and 5 and *SI Appendix, Fig. S3* is provided in *Dataset S1*.

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