An immunocompetent rectal cancer model to study radiation therapy

Graphical abstract

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In brief
Kim et al. demonstrate an immunocompetent rectal cancer implantation model that avoids the use of dextran sulfate sodium. Using this approach, orthotopically transplanted tumors can be used for rapid preclinical screening of radiosensitizers, immunomodulators, and other therapies emerging in cancer treatment and research.

Highlights
- An immunocompetent endoluminal rectal cancer murine model
- Engraftment is based on mechanical disruption
- Tumors recapitulate anatomical origin, histology, and response to radiation
- Ease of use and physiological relevance can improve preclinical therapy modeling
An immunocompetent rectal cancer model to study radiation therapy

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SUMMARY

We describe a mouse model of rectal cancer (RC) involving rapid tumor organoid engraftment via orthotopic transplantation in an immunocompetent setting. This approach uses simple mechanical disruption to allow engraftment, avoiding the use of dextran sulfate sodium. The resulting RC tumors invaded from the mucosal surface and metastasized to distant organs. Histologically, the tumors closely resemble human RC and mirror remodeling of the tumor microenvironment in response to radiation. This murine RC model thus recapitulates key aspects of human RC pathogenesis and presents an accessible approach for more physiologically accurate, preclinical efficacy studies.

MOTIVATION

Rectal cancer (RC) incidence is increasing worldwide, especially among young adults 50 years of age and younger.40–46 The complexity of treatment highlights the need for the development of rapid models that can be used to better understand the biology of RC and to conduct robust preclinical studies. Current approaches utilize induction by dextran sulfate sodium (DSS), which alters the immune physiology of the microenvironment, or models in which tumor origin occurs from the submucosa, which is not the typical site of disease origin. These limitations are particularly important as recent reports suggest that the underlying biology and genetics of inflammatory-associated colorectal tumors are distinct from non-inflammatory tumors.47 How or to what extent DSS treatment influences this is unknown, but the systemic side effects are a concern, and development of a simpler and faster, DSS-free model would be helpful to disentangle these factors. We therefore sought to develop a more anatomically and physiologically relevant RC murine model that is straightforward and rapid. The model described here relies on simple mechanical disruption of the target tissue, rather than DSS treatment, and localizes engraftment to the mucosal tissue, which better reflects the anatomical origin of this tumor type.
INTRODUCTION

Rectal cancer (RC) constitutes approximately one-third of colorectal cancer diagnoses and increasingly affects younger individuals (<50 years old). Rectal cancer is typically treated with radiotherapy (RT) and, if necessary, chemotherapy before radical surgery. Historically, RT sensitivity has largely been considered a cell-intrinsic property resulting from indirect and direct DNA damage, yet tumor genetics alone do not fully explain RT sensitivity. There is growing evidence that immune effects significantly contribute to RT sensitivity. The immune tumor microenvironment (TME) plays a crucial role in tumorigenesis and response to treatment. In turn, RT, by inducing DNA damage and cell death, may induce TME remodeling and modulate immunological responses. Investigating these ideas will require immunocompetent murine models that accurately mimic human RC and its response to RT. These models should also provide an opportunity to develop novel effective therapy regimens involving immunomodulators for the treatment of RC, which has been reported relatively less sensitive to the current immunotherapy compared with right-sided colorectal cancer.

Numerous murine models of colon cancer have been established, but these do not offer an ideal setting in which to examine the TME for RC as the pelvic anatomy and blood supply are unique. A limited number of murine models have been developed in the proper rectal anatomic context, but these involve either chemical induction—a method that Yui et al. utilized successfully to implant colonic organoids orthotopically—which can cause chronic colitis and alter the TME, or intrarectal injection of tumor cells, which results in submucosal-initiated tumors, unlike human tumors that initiate in the mucosa. The advantages and limitations of existing models are summarized in Table 1.

There is an additional need for optimized RT delivery methods within murine RC models. Many studies have used whole-body radiation, which is a common treatment for patients, but this is not representative of localized pelvic RT and adds undesirable systemic effects. Recent studies such as that performed by Nicolas et al. have shown that fractionated radiotherapy in combination with immunotherapy/IL-1α can be used in orthotopic immunocompetent RC models to administer RT treatment with clinical precision, but this is limited by complexity of which medical devices and technology are required. As such, we sought to develop an anatomically and physiologically accurate RC model, along with a widely adaptable pelvic irradiation method that better meets the needs presented above.

RESULTS AND DISCUSSION

We isolated and cultured organoids from genetically engineered (Apc<sup>floX/floX</sup>, LSL-Kras<sup>G12D</sup>, Tp53<sup>flox/floX</sup>) mice (C57Bl/6J) rectal tissue. Organoids were transduced <i>ex vivo</i> with adeno viral vector expressing Cre-recombinase to activate Kras and inactivate Apc and p53 (Apc<sup>-/-</sup>, Kras<sup>G12D</sup>, Tp53<sup>+/−</sup>); hereafter referred to as AKP). This represents a common genetic configuration in human RC. Organoids of the desired genotype were selected by removing WNT ligands and epidermal growth factor from the media, followed by Nutlin-3 selection. The resulting AKP rectal tumor organoids (tumoroids) were prepared as an enema. Mice (C57Bl/6J) were anesthetized, and a small-caliber brush was inserted through the anus to mechanically disrupt the rectal mucosa (Figures 1A and 1B). Thereafter, tumoroids were transanally instilled into the rectum, and the anus was glued shut for 6–8 h to prevent expulsion and facilitate tumoroid engraftment. Step-by-step instructions are provided in Video S1. Of note, the technical aspects of our tumor engraftment procedure are simpler compared with surgical or endoscopic implantation protocols and thus easily adoptable by other investigators. Compared with DSS models that can result in multifocal or segmental lesions and potentially implant tumors anywhere along the colonic tract, our method has less of a systemic inflammatory response and reliably engrafts tumors in the rectum. It is important to note that mechanical disruption reduces anesthesia time compared with methods involving surgical procedures. Animal stress is also reduced compared with DSS methods as the animal does not exhibit a systemic inflammatory state, recovers quickly, and does not require significant delays prior to potential experimental alterations such as treatment with radiation once the tumors engraft. This method allows animals to be engrafted in a manner that is fast, efficient, and well tolerated, enabling larger scale experiments to be performed. Tumoroids engraft at the site of mucosal disruption (Figure 1C) and preserve WNT and KRAS pathways alterations as demonstrated by elevated β-catenin and phospho-ERK (Figures S1A and S1B).

Of 118 mice treated, 20 (17%) died within 2 days after transplantation due to intestinal perforation (confirmed at autopsy). Among the 98 surviving mice, this procedure yielded a 58% engraftment rate (Figure 1A, right panel). We have compared the engraftment rate of different RC models carefully in Table 1. The engraftment rate of this model is close to the immunocompetent model discussed (62%). Higher engraftment rates are seen across immunocompromised models. Tumor engraftment (seen by either a luminal mass or increasing size of a luminal lesion) was apparent on colonoscopy by 2–4 weeks after transplantation (Figure 1D). The median distance of the tumor from the anal verge was 3.5 mm (range, 0–10 mm) (Figure 1E). Engrafted tumors grew and obstructed the rectal lumen within 1–3 months of transplantation and were macroscopically similar to human RC under endoscopy (Figure 1F). In rare cases histopathologic review showed that the engrafted tumors resemble the development and progression from pre-invasive to invasive cancer and to metastatic disease (Figures 1H and 1I) as observed in human RC (Figure 1G). These data establish the feasibility of this RC model and its clinically relevant patterns of tumor progression and metastasis.

To test the application of RT to our model, we administered localized pelvic RT (Figures 2A and 2B) to tumor-bearing mice using a customized Cerrobend holding apparatus (Figures S2A–S2D). Mice were randomized at a median of 5 weeks after transplantation with no difference in average tumor size between cohorts (Figures S1C and S1D). Mice in the RT cohort underwent localized pelvic radiation to 15 Gy in a single fraction. All mice were monitored carefully for health status and subjected to weekly surveillance endoscopy. Euthanasia occurred when the circumferential tumor involvement exceeded 50% of the rectal lumen.
Table 1. Colon and rectal cancer mouse models by anatomic relevance

<table>
<thead>
<tr>
<th>Anatomic location</th>
<th>Tumorigenesis method</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Engraftment rate</th>
<th>Reference</th>
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<td>Heterotopic methods</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Abdominal organ (Renal capsule, peritoneum)</td>
<td>tumor injection</td>
<td>● high engraftment</td>
<td>● ectopic micro-environment</td>
<td>Tanaka et al. [14]; Schoffelen et al. [15]</td>
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<td>Orthotopic methods — colon</td>
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<tr>
<td>Cecum</td>
<td>surgical submucosal implantation of tumor</td>
<td>● high engraftment</td>
<td>● requires surgical technical expertise</td>
<td>100% [16]</td>
<td>Fumagalli et al. [16]</td>
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<td>Predominantly colon</td>
<td>colitis-induced</td>
<td>● inexpensive</td>
<td>● mimics colitis-associated colorectal cancer, not sporadic colorectal cancer</td>
<td>38%–100% [17]</td>
<td>Tanaka et al. [17]; Neufert et al. [18]</td>
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<tr>
<td>Predominantly colon</td>
<td>genetically modified murine model</td>
<td>● mimics sporadic cancer</td>
<td>● tumor location is not specific to rectum</td>
<td>20%–50% [19]</td>
<td>Dow et al. [18]; Xue et al. [19,20]</td>
</tr>
<tr>
<td>Distal colon</td>
<td>genetically modified murine model with focal activation of mutations</td>
<td>● mimics sporadic cancer</td>
<td>● slow tumor growth</td>
<td>71%–96% [21]</td>
<td>Roper et al. [21]; Hung et al. [22]</td>
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(Continued on next page)
Prolapse, a common negative outcome for other similar models, was not observed in any of our implanted mice (n = 118). Mice tolerated RT well without significant weight loss or illness. We also compared the survival rate between pelvic-irradiated mice and whole-body irradiated mice to ensure our localized pelvic RT system provided effective bone marrow shielding and mitigated lethality from whole-body irradiation (Figure S2E). A 10%–40% treatment effect was seen after RT on histological examination and tumor regression grading. This treatment effect is similar to what we see in RC patients. A significant growth delay was noted, endoscopically, after RT, but most mice eventually experienced tumor growth (Figures 2C and 2D). Mice treated with localized pelvic RT had significantly improved survival (Figure 2E).

We sought to evaluate if our model recapitulates changes in the TME in response to RT as it does in human patients. Prior research has shown a significant increase in M2 macrophage polarization in human rectal tumors postRT. Similarly, when examining the TME of mice with AKP endoluminal tumors, we observed a significant increase in macrophage infiltration (detected by macrophage surface marker F4/80) and M2 polarization (CD206+ F4/80+), but not M1 (CD11c+ F4/80+) (Figures 2F–2H). These data suggest that our model reflects some key changes in the TME and its response to RT.

In summary, this model has a wide range of potential preclinical applications, such as the evaluation of radiosensitizers, combinations of RT with immunomodulators, and novel therapies. Furthermore, this model can be used to explore how RT resistance is affected by various oncogenic mutations. For any of these applications, our method provides rapid generation of localized, mucosa-initiated rectal tumors in a model that is reliable, reproducible, and translatable to human RC with limited technical demand.

Limitations of study
Injury to the rectal mucosa without internal visual guidance may result in occasional unintended perforation of the bowel wall. Perforation may occur from a myriad of different factors, but from a technical standpoint, it can be commonly attributed to the following: over-brushing the epithelium, applying excess pressure when causing abrasion, inserting the brush too deep or too quickly into the rectum, insufficient

<table>
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<th>Anatomic location</th>
<th>Tumor model</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Engraftment rate</th>
<th>Reference</th>
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<tr>
<td>Distal colon</td>
<td>colonoscopy-based mural injection of tumor</td>
<td>● high engraftment</td>
<td>● technically challenging</td>
<td>83%&lt;sup&gt;23&lt;/sup&gt; 90%–92%&lt;sup&gt;24&lt;/sup&gt;</td>
<td>Bettenworth et al.&lt;sup&gt;23&lt;/sup&gt;, Roper et al.&lt;sup&gt;24&lt;/sup&gt;</td>
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<td>Orthotopic methods—rectum</td>
<td>Distal colon and rectum</td>
<td>(colitis induction + tumor enema)</td>
<td>● tumors grow from the mucosal lumen</td>
<td>100%&lt;sup&gt;29&lt;/sup&gt; 20%–100%&lt;sup&gt;96&lt;/sup&gt; 70%&lt;sup&gt;25&lt;/sup&gt; 94%&lt;sup&gt;27&lt;/sup&gt; 62%&lt;sup&gt;10&lt;/sup&gt;</td>
<td>O’Rourke et al.&lt;sup&gt;10&lt;/sup&gt;, Kishimoto et al.&lt;sup&gt;25&lt;/sup&gt;, Takahashi et al.&lt;sup&gt;26&lt;/sup&gt;, Ganesh et al.&lt;sup&gt;27&lt;/sup&gt;, Chassaing et al.&lt;sup&gt;28&lt;/sup&gt;</td>
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<tr>
<td>Rectum</td>
<td>intrarectal tumor injection</td>
<td>● tumors form in distal rectum (1–2 mm above anus)</td>
<td>● chemically induced colitis alters the tumor microenvironment</td>
<td>100%&lt;sup&gt;29&lt;/sup&gt;</td>
<td>Kasashima et al.&lt;sup&gt;11&lt;/sup&gt;, Hite et al.&lt;sup&gt;29&lt;/sup&gt;</td>
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Table 1. Continued
Figure 1. An orthotopic rectal cancer model in immunocompetent mice

(A) Diagram of rectal tumoroid preparation for orthotopic transplantation. AKP rectal organoids were cultured in Matrigel, harvested, and engrafted as an enema to C57Bl/6J mice after mechanical disruption of the rectal mucosa.

(B) Critical steps in endoluminal engraftment.
1. Irrigate rectum with PBS.
2. Insert a trimmed P200 pipette tip into the anus as a guide.
3. Insert and irritate mucosa with a small-caliber brush.
4. Confirm a blood smear on the brush, which indicates adequate endoluminal disruption.
5. Pipette tumoroid mixture into the rectum.
6. Seal the anus with Vetbond tissue adhesive.
7. Pinch the anus closed.
8. Allow Vetbond to dry and confirm the anus is sealed. See also Video S1.

(legend continued on next page)
lubrication, improper removal of the brush from the rectum, over-application of tissue adhesive, and/or failure to completely remove tissue adhesive. The procedural mortality rate in our study was 17%. These percentages are reasonable estimates for inexperienced users and should be considered when proposing animal numbers for use and justification. It is important to account for procedural mortality when planning experiments as it impacts mice available for engraftment. Through the course of ongoing experiments beyond the scope of the current paper, we have found that training and experience substantially decreases procedural mortality. As with any new technique, we encourage discussion with the animal use and welfare committee. It is critical to have an approved protocol prior to application of this technique.

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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  - Imaging
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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.crmeth.2022.100353.

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AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

C.L.S. serves on the Board of Directors of Novartis, is a co-founder of ORIC Pharmaceuticals and co-inventor of enzalutamide and apalutamide. He is a science advisor to Agios, Arsenal, Beigene, Blueprint, Column Group, Foghorn, Housey Pharma, Nextech, KSQ, and PMV. S.W.L. receives research funding from Calico and is on the scientific advisory boards and holds equity in Blueprint Medicines, Mirimus Inc., ORIC Pharmaceuticals, Geras Bio, Faeth Therapeutics, Senescea Therapeutics, and PMV Pharmaceuticals. He also recently served as a consultant for Boehringer Ingelheim. J.G.A. reports stock ownership in Intuitive Surgical and has served in a consulting or advisory role for Medtronic, Intuitive Surgical, and Johnson & Johnson. P.B.R. reports prior research funding and is a consultant for EMD Serono, receives research funding from XRAD therapeutics, is a consultant for Faeth Therapeutics, is a consultant for Natera, is on the Medical Advisory Board of the HPV Alliance, and is on the Scientific Advisory Board of the Anal Cancer Foundation. J.J.S. received travel support from Intuitive Surgical Inc. (2015) and served as a clinical advisor for Guardant Health Inc. (2019) and Foundation Medicine Inc. (2022). He also served as a consultant and speaker for Johnson & Johnson Inc. (2022).
Figure 2. Application and evaluation of localized radiation therapy in the orthotopic rectal cancer mouse model

(A) Experimental schema. See also Figure S2.

(B) Radiation dosimetry of the predicted radiation field to the distal pelvis.

(C) Endoluminal tumor growth. Mean circumferential tumor size of the control and RT group mice at the time of treatment and at weeks 1–4 from treatment are plotted. Error bars denote the standard error of mean. ***p < 0.0001 at individual time points by unpaired two-tailed t test.

(D) Representative serial endoscopic pictures of a control mouse and irradiated mouse.

(E) Overall survival of control and RT cohorts. Mice that were sacrificed for having tumor involvement >50% of the lumen were deemed as events. Kaplan-Meier estimates with number at risk depicted; p value for log rank comparison.

(F) Top: representative hematoxylin & eosin (H&E) stained images of rectal tumors from control mouse and irradiated mice. Middle and bottom panels: representative images of immunofluorescent staining for tumor-associated macrophages. F4/80 (green) is a macrophage marker, CD11c (orange) is a marker for M1 polarized macrophages, and CD206 (red) is for M2 polarized; counterstain with Hoechst (blue).

(G) Quantification of tumor-associated macrophages and the M1/M2 subtypes. Each point represents an individual mouse sample. Error bars denote the standard error of mean. p values provided by unpaired two-tailed t test. NS denotes a result that is not statistically significant.

(H) Quantification of CD4+ helper T cells, CD4+FOXP3+ Treg cells, and CD8+ cytotoxic T cells. Data points, error bars, and p values are as represented as in (G).


# STAR METHODS

## KEY RESOURCES TABLE

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, J. Joshua Smith MD PhD (smithj5@mskcc.org).

Materials availability
- This study did not generate new, unique reagents.
- There are restrictions to the availability of ketamine/xylazine due to federal, state, and institutional restrictions as this is a Schedule II substance under the Controlled Substances Act.
- Matrigel basement membrane is a limited resource due to high demand and backorder capacity. Protein concentration may vary slightly between batches. This can be normalized and controlled for using the manufacturer’s batch-specific production information.

Data and code availability
- Endoscopic visualization of colorectal tumors and survival data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
EXPERIMENTAL MODEL AND SUBJECT DETAILS

**Mice**
We used 6- to 8-week-old female C57Bl/6J mice (Jackson Laboratory, stock no. 000664). All animal experiments were conducted under protocols 11-06-012 and 06-07-012 approved by the Memorial Sloan Kettering Cancer Center's Institutional Animal Care and Use Committee in conjunction with the Research Animal Resource Center and American Association (RARC) for Laboratory Animal Science (IACUC).

**METHOD DETAILS**
Step-by-step protocols and details on reagent preparation are provided in Supplemental methods S1. Reagents were used freshly, and nothing more than 6 months old or expired was used.

**Culturing and preparation of tumoroids for endoluminal transplantation**
AKP tumoroids were derived, cultured, and prepared for endoluminal transplantation as described.\(^{10,15}\) Cells were released from Matrigel and resuspended in ice-cold PBS with 5% Matrigel to a concentration of \(2 \times 10^5\) cells/100 \(\mu\)L per mouse.

**Mechanical disruption of the mouse rectal mucosa and tumoroid transplantation**
Critical steps are shown in Video S1. Anesthetized mice (2% isoflurane and oxygen) underwent rectal flushing to expel stool using PBS. A smooth-trimmed P200 pipette tip (lubricated with Vaseline) was inserted into the anus as guide, then a small-caliber brush was inserted through the pipette tip and moved gently in to and out of the rectum 3–5 times to gently disrupt the rectal mucosa. The tumoroid suspension was then slowly injected transanally using a P200 pipette. The anus was sealed using 3 \(\mu\)L of Vetbond Tissue Adhesive to prevent luminal contents from spilling. This bond was removed between 6 and 8 h later.

**Tumor surveillance and measuring tumor growth**
Tumor engraftment was monitored by 1.9-mm rigid 30°C14 small animal endoscope weekly. A video of each endoscopy was taken, and a static picture of the tumor was analyzed to calculate endoluminal involvement. Tumor growth was quantified by the percent of the field of view occupied by the tumor area as described.\(^{26}\)

**Localized pelvic radiation**
Mice were anesthetized by intraperitoneal injection of Ketamine/Xylazine (100 mg/mL; 10 \(\mu\)L/g body weight). Localized pelvic irradiation was delivered using an X-Rad 320 machine (Precision X-ray, Madison, CT) (250kVp/12mA) with customized Cerrobend blocks. The fabrication technique for this apparatus can be shared upon request.

**Histopathology**
Dissected tumor samples were fixed with 4% paraformaldehyde, embedded in paraffin, and sectioned according to standard protocols. For histopathologic evaluation, 5-\(\mu\)m sections were stained with H&E.

**Immunofluorescence**
Tissue sections were deparaffinized, then boiled in pH 6.1 citrate buffer for 20 min for antigen retrieval. Sections were blocked in 10% normal donkey serum and 1% BSA at room temperature for 1 h, immunostained overnight at 4°C with primary antibodies, then for 2 h at room temperature with fluorophore-conjugated secondary antibodies. Cell nuclei were labeled with DAPI or Hoechst, as noted.

**Imaging**
Slides were scanned by a Panoramic Flash slide scanner using a 20 \(\times\)0.8 NA objective. Images were examined and representative areas exported using CaseViewer 2.2. No gamma changes were made to any immunofluorescence images. All brightfield images are unaltered. Immune cells in total tumor regions were quantified using custom macros written in ImageJ (NIH, Bethesda, MD, USA). The number of immune cells in 1mm\(^{-1}\) of tumor regions were calculated from each sample and averaged per group, and a Student’s t test was performed for statistical analysis.

**Monitoring mouse health**
Upon arrival, mice are weighed and observed to confirm robust health prior to implantation. After implantation, mice are observed for 5 min to ensure return to normal levels of body condition, activity, and alertness. Six to 8 h after implantation any remaining Vetbond is
removed. Once a day for three days post-procedure, mice are observed again. Past this period, mice are monitored once a week through general observation and tumors surveilled by endoscopic imaging.

**Adherence to established endpoints**

Mice exhibiting excessive weight loss (>20% decrease from baseline), decreased activity, worsening body condition, reduced alertness, and/or blunted response to stimulation and handling were eligible for euthanasia. Mortality from perforation due to excessive epithelial disruption is most frequently observed <24h post-implantation. For mice developing rectal tumors, if circumferential lumen involvement exceeded 50% or if tumor growth interfered with ambulation, eating, drinking, defecation, or urination, mice were euthanized. While we did not observe rectal prolapse in our mice, likely due to close endoscopic monitoring, it is known that rectal prolapse can occur with large obstructing tumors and mice should be monitored for this closely. Standard, pre-approved institutional methods of CO₂ overdose and asphyxiation were used to euthanize mice as outlined by the *RARC’s Recommended Methods of Euthanasia for Laboratory Animals.*

**QUANTIFICATION AND STATISTICAL ANALYSIS**

The number of biological replicates per experiment and the number of experiments performed for each dataset and the statistical analysis performed are outlined in the corresponding figure legends. Results are depicted as mean ± standard deviation (SD) unless otherwise stated using Microsoft Excel for Mac (v16.56) or GraphPad Prism (v.9.3.1). P-values were calculated using GraphPad Prism (v.9.3.1) and Microsoft Excel for Mac (v16.55). No statistical method was used to predetermine sample size. Sample sizes were estimated according to transplantation success and previous pilot experiments to estimate variability.