Mimicking Metastases Including Tumor Stroma: A New Technique to Generate a Three-Dimensional Colorectal Cancer Model Based on a Biological Decellularized Intestinal Scaffold

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Tumor models based on cancer cell lines cultured two-dimensionally (2D) on plastic lack histological complexity and functionality compared to the native microenvironment. Xenogenic mouse tumor models display higher complexity but often do not predict human drug responses accurately due to species-specific differences. We present here a three-dimensional (3D) in vitro colon cancer model based on a biological scaffold derived from decellularized porcine jejunum (small intestine submucosa+mucosa, SISmuc). Two different cell lines were used in monoculture or in coculture with primary fibroblasts. After 14 days of culture, we demonstrated a close contact of human Caco2 colon cancer cells with the preserved basement membrane on an ultrastructural level as well as morphological characteristics of a well-differentiated epithelium. To generate a tissue-engineered tumor model, we chose human SW480 colon cancer cells, a reportedly malignant cell line. Malignant characteristics were confirmed in 2D cell culture: SW480 cells showed higher vimentin and lower E-cadherin expression than Caco2 cells. In contrast to Caco2, SW480 cells displayed cancerous characteristics such as delocalized E-cadherin and nuclear location of β-catenin in a subset of cells. One central drawback of 2D cultures—especially in consideration of drug testing—is their artificially high proliferation. In our 3D tissue-engineered tumor model, both cell lines showed decreased numbers of proliferating cells, thus correlating more precisely with observations of primary colon cancer in all stages (UICC I-IV). Moreover, vimentin decreased in SW480 colon cancer cells, indicating a mesenchymal to epithelial transition process, attributed to metastasis formation. Only SW480 cells cocultured with fibroblasts induced the formation of tumor-like aggregates surrounded by fibroblasts, whereas in Caco2 cocultures, a separate Caco2 cell layer was formed separated from the fibroblast compartment beneath. To foster tissue generation, a bioreactor was constructed for dynamic culture approaches. This induced a close tissue-like association of cultured tumor cells with fibroblasts reflecting tumor biopsies. Therapy with 5-fluorouracil (5-FU) was effective only in 3D coculture. In conclusion, our 3D tumor model reflects human tissue-related tumor characteristics, including lower tumor cell proliferation. It is now available for drug testing in metastatic context—especially for substances targeting tumor-stroma interactions.
Introduction

Cancer is a leading cause of death worldwide, especially in developed countries, and will gain importance with the progressing age structure of future society. Success in the treatment of this disease depends on the development of appropriate tumor models to improve the translation of novel treatment strategies from research to the patients’ situation. Especially in the field of oncology, attrition rates attain up to 95%. Reasons are that tumorigenesis is not only regulated intrinsically by genetic changes in the tumor cell itself but also by microenvironmental cues and stromal cell-to-cell interactions that instruct tumor development. These conditions are poorly reproduced by conventional two-dimensional (2D) cell culture models currently used in drug development.

Three-dimensional (3D) cell culture approaches such as Matrigel coating and spheroid cultures have gained importance and substantially improved in vitro tumor models. Tissue engineering offers attractive options for innovative 3D scaffolds that correspond more closely to native tumors than standard 2D cell culture systems or animal models. However, a crucial step in cancer progression neglected in most models is the invasion across the basement membrane, a process that starts metastasis formation and mostly excludes patients from complete remission under therapy.

Epithelial cancers—known as carcinomas—can become invasive by mainly two different mechanisms: (1) single tumor cell invasion, mostly preceded by an epithelial-to-mesenchymal transition (EMT), through which cells lose epithelial and gain mesenchymal markers and (2) collective tumor cell invasion, probably driven by cancer-associated fibroblasts and independent of EMT. In this study, fibroblasts associate closely with tumor cells, serve as guidance, digest the matrix, and ultimately deposit extracellular matrix (ECM) components to form a complex, tumor–stroma. Metastasis formation shows a reverse program of EMT called mesenchymal-to-epithelial transition (MET).

To generate tissue-engineered tumors, including tumor–stroma, in a biological tissue architecture, we use a decellularized part of the porcine jejunum derived from the patented Biological Vascularized Scaffold (DE:302014007893; BioVaSc®) structure. We define our scaffold as “small intestinal submucosa with preserved mucosa” (SISmuc). It differs from the commonly used SIS scaffolds by the preservation of the mucosal tissue layer, including crypt, villi, and the basement membrane structures. This allows us to investigate the process of invasion of tumor cells across the basement membrane into deeper layers of the tissue matrix, a procedure that has been successfully used before in a lung cancer model.

In summary, our aim is to generate an artificial colorectal cancer tissue model. This should be achieved (1) by the integration of a biological tissue scaffold, including basement membrane structures, (2) by the interaction of tumor cells with the tumor stroma, and (3) by applying advanced biomechanical technologies. Our new cancer model paves the way for in vitro analyses of drug efficacy that reflect characteristics of human tumor tissue at different stages of malignancy.

Materials and Methods

Cell lines and 2D cell culture

SW480 and Caco2 cells were bought at DSMZ. SW480 cells were cultured in RPMI +10% fetal calf serum (FCS) and Caco2 cells in MEM+20% FCS+1% NEAA+1% Pyruvate. Cells were monitored for pathogen infections at regular intervals. For immunohistochemical 2D stainings, cells were seeded on glass slides until they had reached a confluency of about 70%.

Human fibroblasts

Fibroblasts were isolated from skin biopsies derived from the University Hospital Wuerzburg (ethics proposal: 182/10).

Skin biopsies were rinsed with phosphate-buffered saline (PBS) with calcium and magnesium (0.5 g MgCl₂·6H₂O + 0.5 g CaCl₂/5 L PBS) before all fat and tissue remnants were removed. After a second rinsing with PBS+calcium and magnesium, the skin was cut into strips of 2–3 mm width. Strips were rinsed with PBS and covered with 2 U/mL dispase solution (Invitrogen) over night at 4°C. Then, the dispase solution was removed and 10 mL PBS with calcium and magnesium (0.5 mg MgCl₂·6H₂O + 0.5 g CaCl₂/5 L PBS) was added. Epidermis and dermis were separated mechanically. Dermis strips were transferred to a new Petri dish, chopped into small pieces, and incubated in 5 mL of collagenase solution (500 U/mL) in a 50-mL tube. The Petri dish was re-rinsed with 5 mL of collagenase solution, which was then put into the 50-mL tube containing the chopped pieces of dermis. They were first incubated for 45 min at 37°C and then centrifuged for 5 min at 1200 rpm. Supernatant was removed and the remaining pellet was resuspended in 2 mL of culture medium and transferred to a T75 cell culture flask. The next day, 2 mL of culture medium was added.

SISmuc preparation

SISmuc was prepared from porcine jejunal segments and consists of small intestine submucosa (SIS) and mucosa (muc). All explantations were in compliance with the German Animal Protection Laws (§4 Abs.3) and the institute’s animal protection officer regularly informed the responsible authorities. The animals received proper attention and humane care in compliance with the Guide for Care and Use of Laboratory Animals published by the National Institute of Health (NIH publication no. 85–23, revised 1996) and as approved of by the institutional board of animal protection. After jejunal explantation, the scaffold was prepared as previously described. The SISmuc was generated by removing the mesentery with the vascular tree from the completely decellularized explant.

3D cell culture of in vitro models

For static 3D cell culture, cells were seeded onto the SISmuc (see porcine material), fixed between two tailored metal rings—the cell crowns (Fig. 1). All 3D cell cultures were performed under standard conditions (37°C, 5% CO₂) as described before. For monocultures, 100,000 cells in 500 μL medium were seeded onto the former luminal side of the SISmuc scaffold. For cocultures, 100,000 tumor cells and 100,000 fibroblasts were seeded together in 300 μL medium onto the SISmuc scaffold. After an adhesion time of 2 h, cell crowns with reseeded scaffold were completely filled with medium and cultured for 14 days. Then, the tissue was washed with PBS and fixed in 4% paraformaldehyde (PFA) for 2 h followed by paraffin embedding in an
automated system (STP 120; Thermo Fisher Scientific GmbH) for immunohistochemical analysis. Other samples were frozen without any fixation to isolate mRNA from the cells for quantitative real time-PCR (qRT-PCR) analysis.

For dynamic 3D cell culture, SW480 cells were seeded in monoculture and in coculture with dermal fibroblasts onto SISmuc scaffolds and placed in cell crowns 3 days before starting the bioreactor culture (Fig. 2), thus allowing the cells to attach to the scaffold. Afterward, the scaffolds were transferred to the bioreactor and were cultured under dynamic conditions maintaining a constant medium flow for another 10 days. The speed of the blood flow was adjusted to a speed of $3-4 \times 10^{-3}$ mL/min exerting mechanical shear stress on the constructs (Fig. 2D). For the dynamic cell culture, 1% of Pen/Strep was added to the cell-specific medium.

Bioreactor

The bioreactor (Fig. 2A) was designed to allow the culture of tissues at the interface between two separated fluid circulation systems. For the design of tailored parts, SolidWorks 2010×64 Edition (Dassault Systemes Deutschland GmbH) was used. Construction drawings were sent to GT Labortechnik and parts were manufactured from polysulfone (PSU). To identify an optimal bioreactor setup, different designs were investigated by computational modeling (Fig. 2B, C). For this, COMSOL Multiphysics (Comsol Multiphysics GmbH) was used. Bioreactor geometries were transferred to the simulation software, and the fluid domains were parameterized as water at 37°C. In addition, inlet regions with a specific flow velocity and outlet regions exhibiting ambient pressure conditions were defined. By varying the inlet velocity and by calculating the shear stress distribution, the characterization of a bioreactor design and the identification of the bioreactor operating range were made possible (Fig. 2D).

Stimulation with 5-fluorouracil

Static models were treated with 25 μM (3.25 μg/mL) 5-fluorouracil (5-FU) (Selleckchem) on day 7, 10, and 13.
Dynamic cultured models were treated on day 7 and 11 of dynamic cell culture with 25 μM 5-FU. On day 14 of static or dynamic cell culture, tissue was fixed in PFA and embedded in paraffin for immunohistochemistry. Cell Titer Glo Cell Viability Assay (Promega) was performed with 2D cultured cells following the manufacturer’s protocol.

Fluorescence immunohistochemistry

Glass slides were fixed in 4% PFA for 10 min and SISmuc samples for 2 h. After fixation, glass slides were either directly stained or stored in PBS buffer at 4°C for up to 1 week before staining. SISmuc samples were stained as paraffinized sections (3–5 μm) and primary antibodies (E-cadherin No. 610181, BD Transduction Laboratories; β-Catenin No. ab32572, AbCam; Vimentin No. ab92547, AbCam; PCK No. C 2562-2ML, Sigma-Aldrich Chemie GmbH; Ki67 No. ab16667, AbCam; Collagen IV No. ab6586, AbCam) were diluted 1:100 and incubated over night at 4°C. Secondary antibodies were diluted 1:400 and incubated for 1 h at room temperature. Double stainings were performed by using primary antibodies of two different species and secondary antibodies marked with the fluorescent dyes, Alexa-647 or Alexa-555 (Life Technologies GmbH). Nuclei were counterstained by DAPI solved in Mowiol embedding solution. Pictures were taken by using either a confocal laser scanning microscope (SP-8; Leica Microsystems GmbH) or a digital microscope (BZ-9000; Keyence Deutschland GmbH).

Diaminobenzidine immunohistochemistry

SISmuc samples were stained as paraffinized sections (3–5 μm) with Villin (#V1616C01; DCS) according to the instructions of the EnVision+System-HRP (Diaminobenzidine [DAB]) (Dako Deutschland GmbH).

FIG. 2. Compartmented bioreactor for the culture of tissues at the interface between two fluid circulation systems. (A) The bioreactor comprises a tissue chamber, a peristaltic pump, and a reservoir container for the cell culture medium. From the reservoir container, the peristaltic pump delivers cell culture medium to the tissue chamber harboring up to three tissue constructs in separated compartments. (B) A sectional view of such a compartment depicts the path of the flow allowing defined shear stress conditions. On both sides of the tissue, independent fluid circulations ensure apical- and basolateral-specific culture conditions. Stainless steel electrodes support the electrical characterization of the cultured tissue. (C) The streamline and the resulting shear stress on the tissue surface at an inlet volume flow of 3.8 mL/min were predicted using computational modeling. Except from a ring of ~50 μm at the tissue–bioreactor interface, shear stress values between 3.0 and 5.0 × 10⁻³ N/m² were calculated. (D) By varying the inlet volume flow, the operating range of the bioreactor was characterized to allow the prediction of the average shear stress at a specific velocity of the inlet volume flow. According to the operating range, average shear stress at a volume flow of 3.8 mL was calculated to 3.34 × 10⁻³ N/m². Color images available online at www.liebertpub.com/tec
Quantitative real time-PCR

We isolated total mRNA from cells using the RNeasy Micro Kit (QIAGEN). The synthesis of cDNA was done by using the iScript Kit (Bio-Rad Laboratories GmbH). We used the following primers in a 2-step PCR with the Sso-Fast™ EvaGreen™ Supermix (BioRad):

- **Vimentin**: CTGGATTTCCTTCGTGGA and CGAAA CACCCCTGCAATCTT,
- **E-cadherin**: GACCGGTGCAATCTTCAAAA and CAGGT CTCCCTTGGCTCTG,
- **GAPDH**: TGACGCTGGGGCTGGCATTG and GCTCTT GCTGGGGCTGGTG.

![Conventional 2D cell culture](image)

**FIG. 3.** In conventional 2D cell culture, SW480 cells display a less differentiated and more malignant phenotype than Caco2 cells. In 2D cell culture, phase-contrast pictures show that SW480 cells round up and lose contact with the bottom of the culture flask (A1), whereas Caco2 cells display a well-differentiated monolayer (A2). Immunofluorescence double stainings of vimentin (red right top in B, E) and PCK (green right top in B, E) indicate a higher expression of the mesenchymal marker vimentin in SW480 (B) than in Caco2 cells (E), which was confirmed and quantified by qRT-PCR (D). Immunofluorescence double stainings of the adherence junction protein E-cadherin (green right bottom in C, F) and β-catenin (red right top in C, F) demonstrate a lower expression of E-cadherin in SW480 (C) than in Caco2 cells (F). β-Catenin localizes in the nucleus of several SW480 cells (C, arrows). E-cadherin expression could be also confirmed and quantified by qRT-PCR (D). Scale bars in (B–F): 25 μm. 2D, two-dimensional; PCK, Pan-cytokeratin; qRT-PCR, quantitative real time-PCR. Color images available online at www.liebertpub.com/tec
HPRT: TGACCTTGATTTTTTGCAACC and CGAGCAAGACGTTCAGTCCT. Genes were normalized to HPRT and GAPDH and the ΔΔCT method was used for evaluation of expression ratios. Primer efficiencies were 91–95%.

Results

Caco2 and SW480 colorectal cancer cell lines allow mimicking of different tumor stages

To establish an in vitro tissue-like 3D tumor model that can simulate tumors in different malignant stages, two different cell lines with reportedly more differentiated (Caco2) or undifferentiated (SW480) growth behavior were tested. Phase-contrast pictures in conventional cell culture showed a stronger adherence to the standard culture flasks in Caco2 cells than in SW480 cells. SW480 cells tended to round up and to separate from the surface, whereas Caco2 cells built a tight monolayer (Fig. 3A1, A2). Compared to Caco2 cells, the more malignant SW480 cells showed a higher expression of vimentin in immunofluorescence staining, thus indicating an initiated EMT process going along with a lower expression of the adherence junction marker E-cadherin. This was confirmed quantitatively by qRT-PCR (Fig. 3D). We noted that in some of the SW480 cells, but not in any Caco2 cells, β-catenin was localized within the nucleus (Fig. 3C, arrows), indicating a more malignant state of SW480 cells.16

Generation of 3D tumor models

For 3D in vitro colon cancer tumor model development, we applied the SISmuc as a scaffold with intestinal microarchitecture as used before for artificial tissue and lung tumor models.12,14 First, we determined a cell number of 1 × 10^5 Caco2 cells and a culture period of 14 days on our SISmuc scaffold as being optimal to generate artificial tissue with epithelial characteristics. We could demonstrate differentiation progress by immunohistochemical staining of villin. This protein involved in absorption was basolaterally located after 7 days of culture (Fig. 4A) and moved to the apical surface of the epithelial monolayer after 14 days of culture (Fig. 4B). Furthermore, transmission electron microscopy (TEM) analysis revealed the formation of regular microvilli, close cell-to-cell contacts with apical tight junctions (arrow in Fig. 4d), the presence of storage vacuoles,17 and a close attachment to the preserved basement membrane of the decellularized intestinal scaffold (Fig. 4C–E). The same culture conditions that allowed well-differentiated growth of Caco2 cells were applied to SW480 cells in the following experiments as well as to the coculture approaches with primary fibroblasts.

FIG. 4. On the 3D SISmuc scaffold, differentiation of Caco2 cells is achieved after 14 days of culture. Immunohistochemical DAB-staining of villin on paraffin sections shows a basolateral/cytoplasmic location after 7 days of cell growth (A) and its change to the apical side after 14 days of growth (B, arrows). Ultrastructural analysis of Caco2 cultured on SISmuc for 14 days displays a closed monolayer on top of the SISmuc. The cells show a well-differentiated morphology of microvilli (C, D: mv), the formation of close apical tight junctions (arrow in d) as well as mucin-vacuoles (v in D). The nucleus is marked with n. Moreover, cells attach to the preserved basement membrane (* = lamina lucida, ‡ = lamina densa) of the SISmuc (e). Arrow in (E): collagen fibers. DAB, diaminobenzidine. Color images available online at www.liebertpub.com/tec
The percentage of proliferating cells in 3D cell culture resembles the in vivo situation

In our 3D in vitro tumor model for colorectal carcinoma, we were able to show a clearly decreased proliferation by immunohistochemical Ki67 staining. Compared to almost 100% Ki67-positive cells in conventional 2D cell cultures of Caco2 and SW480 cells (Fig. 5A, B), lower cell proliferation appeared in 3D cultures of both tumor cell lines (Fig. 5C, D). The percentage of proliferating cells in patients’ biopsies from different tumor stages in the tumor center and border ranged from 6% in UICC I, 16% in UICC II, 32% in UICC III, to 35% in UICC IV. To find out if markers from 2D stainings were maintained in 3D and could be aligned to morphological growth patterns indicating invasion or metastasis formation, we investigated tissue morphology by hematoxylin and eosin (HE) staining and stained for molecular EMT markers such as Pan-cytokeratin (PCK) and vimentin, epithelial differentiation (E-cadherin), and nuclear β-catenin localization.

Tissue microenvironment and fibroblast coculture enhance differences of tumor characteristics in Caco2 and SW480 cells

As demonstrated by HE stainings, Caco2 cells formed a monolayer on top of the scaffold (Fig. 6A). Primary fibroblasts exhibited a flat cell morphology and grew inside the scaffold. The former villi topography of the SISmuc was remodeled to a plain surface (Fig. 6B). In contrast to the Caco2 cells, more malignant SW480 cells grew chaotically without the formation of a closed monolayer and migrated more deeply into the former crypt and vessel structures of the gut. Moreover, single SW480 cells completely lost contact with the scaffold (Fig. 6C). The immunofluorescence analysis of vimentin and PCK showed a high expression of PCK in Caco2 cells and vice versa, a high expression of vimentin in fibroblasts (Fig. 6D, E). We noted that in contrast to 2D conditions, vimentin could no longer be detected in SW480 cells cultured on SISmuc (Fig. 6F, compare to inset). In our coculture model, we seeded the same number of tumor cells and fibroblasts at the same time onto the SISmuc scaffold. Interestingly, fibroblasts from the same batch affected Caco2 and SW480 cell growth and morphology differently. After 2 weeks in coculture with fibroblasts, Caco2 cells formed multilayers in single areas. However, HE staining revealed no morphological changes compared to monocultures (Fig. 6G). On the contrary, fibroblast coculture showed a strong morphological impact on SW480 cells and resulted in the formation of tumor-like nodes (Fig. 6H). Cocultures with both cell lines showed less protruding villi structures similar to fibroblast monocultures.

FIG. 5. The amount of proliferating cells in the 3D in vitro tumor model decreases compared to conventional 2D cell culture and correlates better to low-grade (UICC I) and high-grade (UICC IV) tumors. Ki67 immunofluorescence staining of Caco2 cells (A) and SW480 cells (B) in 2D cell culture demonstrates a high proliferation that decreases distinctly in 3D cultures of Caco2 (C) and SW480 cells (D). The number of proliferating cells cultured under 3D culture conditions correlates more precisely with those found in tumors with a low (E, G) and high grade (F, H) of malignancy. Pictures were taken from the tumor border (E, F) as well as from the center of the tumor (G, H). Scale bar in (D) represents 75 µm for (A–D). Scale bar in (F) represents 50 µm for (E–H). Color images available online at www.liebertpub.com/tec
Evidently, neither the seeding order nor a higher ratio of fibroblasts impaired the overall coculture morphologies (data not shown). Depending on the tumor cell type, PCK/ vimentin fluorescence double staining demonstrated that fibroblasts and tumor cells were located in different regions of the SISmuc in 3D cell culture (Fig. 6I, J). Within both cell lines, former crypt (c) and villi (v) structures remain clearly visible. In contrast to this, fibroblasts remodel the matrix to a flat layer without villi structures (B). PCK/vimentin double immunofluorescence staining demonstrates a strong PCK staining of Caco2 and SW480 cells (D, F), and a high expression of vimentin in fibroblasts (E). Vimentin expression is lost in SW480 in 3D culture conditions compared to 2D conditions (F, inset: 2D staining: PCK/vimentin). HE staining of cocultures indicates a preserved monolayer formation of Caco2 cells even though in some parts, multilayers could also be observed (G). In SW480 cocultures, the morphology of tumor cells changed to tumor tissue-like aggregates inside the matrix (H, arrows). Immunofluorescence double staining of PCK/vimentin shows that in Caco2 cocultures, the fibroblasts stay in the bottom compartment with the tumor cells on top (I), whereas in SW480 cocultures, the tumor cells are enclosed by fibroblasts that grow on top, beneath, and between the tumor nodes (J). HE, hematoxylin and eosin. Color images available online at www.liebertpub.com/tec

**FIG. 6.** Fibroblasts induce tumor cell aggregation of SW480 on the SISmuc scaffold. HE staining of paraffin sections of the 3D *in vitro* tumor model shows that Caco2 cells grow as a monolayer (A), while SW480 cells exhibit a less differentiated phenotype with loose cell-to-cell contacts (C). Within both cell lines, former crypt (c) and villi (v) structures remain clearly visible. In contrast to this, fibroblasts remodel the matrix to a flat layer without villi structures (B). PCK/vimentin double immunofluorescence staining demonstrates a strong PCK staining of Caco2 and SW480 cells (D, F), and a high expression of vimentin in fibroblasts (E). Vimentin expression is lost in SW480 in 3D culture conditions compared to 2D conditions (F, inset: 2D staining: PCK/vimentin). HE staining of cocultures indicates a preserved monolayer formation of Caco2 cells even though in some parts, multilayers could also be observed (G). In SW480 cocultures, the morphology of tumor cells changed to tumor tissue-like aggregates inside the matrix (H, arrows). Immunofluorescence double staining of PCK/vimentin shows that in Caco2 cocultures, the fibroblasts stay in the bottom compartment with the tumor cells on top (I), whereas in SW480 cocultures, the tumor cells are enclosed by fibroblasts that grow on top, beneath, and between the tumor nodes (J). HE, hematoxylin and eosin. Color images available online at www.liebertpub.com/tec

Epithelial characteristics of SW480 cells are more prominent in the 3D microenvironment

Next, we correlated our morphological observations to the expression of E-cadherin as a MET-marker and to the expression of β-catenin. As expected, we could show by confocal analysis of E-cadherin/β-catenin immunofluorescence double staining that both colocalize in 2D Caco2 monocultures at the cell junctions. This was not changed by the coculture with fibroblasts in 3D (Supplementary Fig. S1; Supplementary Data
In contrast to that, a weaker expression of E-cadherin and concentration in areas of cell aggregates were found in SW480 monocultures (Fig. 7A, stars). In coculture, E-cadherin accumulated in tumor-like nodes at the cell borders (Fig. 7B).

_b-Catenin expression in mono- and cocultures was more pronounced in SW480 cells than in Caco2 cells (compare Supplementary Fig. S1). Moreover, in SW480 monocultures, _b-catenin was observed in some single cells inside the nucleus (Fig. 7A, arrows), which was also visible in 3D coculture samples (Fig. 7B). In addition, in several other areas of the artificial tumor tissue generated by SW480 cells and fibroblasts, a yellow staining indicative of colocalization appeared in the merged staining of E-cadherin and _b-catenin (Fig. 7B).

Under dynamic culture conditions in the flow bioreactor, this effect is enhanced in the coculture (D) compared to monoculture (C). While in monoculture, SW480 tumor cells fill the former crypts of the SISmuc scaffold and preserve the regular structure of the gut, the former crypt structure is remodeled by the cells in coculture with fibroblasts (D). This effect is more visible in HE staining (E, F) as well as in a double staining of PCK (green in G, H, tumor cells) and collagen-IV (red in G, H, basement membranes). Scale bars in (A–C) represent 75 μm. Scale bars in (E–H) represent 500 μm. Color images available online at www.liebertpub.com/tec

Dynamic cell culture conditions support tumor tissue generation and association with the tumor–stroma

To ensure an optimal _in vivo_ nutrient supply and waste removal, we applied dynamic culture conditions to SW480 mono- and cocultures using a newly constructed flow bioreactor system (Fig. 2). These conditions enhanced SW480 tumor cell growth during a culture period of 14 days. As a result, SW480 filled all the former crypts and built multilayers on the former villi (Fig. 7C, E, G). In contrast to that, SW480 cocultured with fibroblasts remodeled the SISmuc and changed the crypt structure as shown by E-cadherin/ _b-catenin double staining (Fig. 7D) as well as HE staining (Fig. 7F) and PCK/collagen-IV double staining (Fig. 7H). E-cadherin staining displayed quite a homogenous distribution within the areas of cells tightly packed together in monoculture (Fig. 7C) as well as in coculture (Fig. 7D). In contrast to the dynamic SW480 monoculture, where single cells express _b-catenin in the cytoplasm, the majority of cells in the dynamic coculture setting expressed _b-catenin only at the cell borders (Fig. 7D). The formation of a tumor-like tissue with a surrounding tumor stroma was demonstrated by PCK/vimentin double staining to differentiate between tumor tissue and surrounding fibroblasts (Fig. 8). Tumor cell aggregates were
enclosed by fibroblasts and tumor cells completely overgrew the SISmuc in several parts of the scaffold (Fig. 8A, B). The morphology of the generated tissue closely resembled that of a colon adenocarcinoma (Fig. 8C).

**Fibroblasts remodel basement membrane structures under dynamic cell culture conditions**

To analyze basement membrane structures, we stained dynamically cultured SW480 mono- and cocultures for collagen-IV. The tumor cells alone (marked by PCK, green in Fig. 9) did not change the collagen structure of the SISmuc (Fig. 9A–C). Around former crypts, basement membranes are precisely visualized by anticollagen-IV staining and mostly well preserved (Fig. 9B, arrows). In coculture with fibroblasts (Fig. 9D–F), these distinct borders are interrupted (Fig. 9E, arrow) and the red collagen-IV staining shows a thickened structure of the SISmuc (Fig. 9E, F). Fibroblasts are able to remodel the SISmuc alone (Fig. 9G, H) as well as in coculture with SW480 (Fig. 9D–F). In coculture, fibroblasts (stained by vimentin, red in Fig. 9J, K) intermingle with the tumor cells (stained by PCK, green in Fig. 9J, K) and, in some areas, seem to lead them deeply into the collagen structure of the SISmuc (Fig. 9K).

**5-FU treatment reduces tumor cells under static and dynamic conditions in coculture settings with primary fibroblasts**

SW480 cells were nearly insensitive to 5-FU under 2D culture conditions (Supplementary Fig. S2A) and static 3D monocultures (Supplementary Fig. S2C). In contrast to this, the treatment of cocultures with 25 μM 5-FU resulted in decreased tumor cell numbers as clearly shown by a PCK/vimentin staining under 3D static culture conditions (green in Fig. 10A, C), as well as under 3D dynamic culture conditions (Fig. 10C, D). In fibroblasts, we observed no effect of this treatment in monocultures (red in Supplementary Fig. S2C) and in cocultures (red in Fig. 10B, D). In coculture of SW480 with fibroblasts, the dense clusters of tumor cells surrounded by fibroblasts shrank and became less compact after a treatment period of 1 week. In the untreated controls, many living cells were flushed into the medium reservoir, but we found only dead tumor cells in the medium reservoir of bioreactors.
treated with 5-FU (data not shown). In 2D, the coculturing of cells resulted in a fibroblast monolayer with single tumor cells placed on top and no tumor cell aggregates could be observed (Supplementary Fig. S2B, C).

Discussion

We developed a histologically complex 3D intestinal colon cancer model, including a tumor–stroma, on the basis of the biological collagen scaffold SISmuc, which has already been used for a lung tumor test system and which derives from the BioVaSc.

Our tissue-engineered 3D model reflects different aspects of tumors on morphological and molecular levels: (1) the formation of a differentiated monolayer of human Caco2 colon cancer cells in contrast to a more scattered growth pattern of SW480 cells accompanied by a loss of vimentin when cultured in 3D, (2) a reduction of the proliferation correlative to clinical samples, (3) the induction of tumor-like structures of malignantly growing SW480 tumor cells by coculture with fibroblasts destroying basement membrane integrity that was not observed in Caco2 cocultures, and (4) the generation of a tight tissue-like association of tumor cells with each other and with the fibroblasts under dynamic culture conditions in a bioreactor. The resulting artificial tumor tissue with stroma resembles native adenocarcinomas and a clinically relevant treatment with 5-FU could be tested as a proof of principle.

By culturing tumor cells in 3D conditions on our matrix, their proliferation strongly decreases. Although an increase
in the percentage of proliferating cells can be seen in some metastases where the Ki67 index has been identified as a prognostic marker, the number of proliferating cells is usually lower in most tumors. Concerning cytostatic drugs, our model should thereby reduce false-positive results in preclinical testing that are thought to be derived at least, in part, from artificially high proliferation rates up to 90 or even 100% in 2D culture.

In 1998, Kammerer et al. tested a similar decellularized human colonic matrix for the development of tumor models. They stressed the potential of decellularized tissue scaffolds as they included parts of the decellularized human colon as basis for tumor and stroma generation. However, they did not succeed with further developments. In the present study, we used a jejunal segment of porcine origin, which has several advantages. We developed a standardized tumor model based on the SISmuc, which is a non-vascularized part of the BioVaSc. This model can be vascularized in future settings by reseeding human endothelial cells into the preserved vessel structures as done before. The jejunal compartment is superior to colon tissue as it exhibits a higher vascularization and contains significantly less bacteria. This facilitates the decellularization process in bioreactors and the creation of a sterile scaffold, a fact of major importance as the use of sodium deoxycholate for decellularization hinders the addition of antibiotics. Moreover, the jejunum is much longer than the colon and thus fewer animals have to be sacrificed.

Since evolution has produced a high conservation of ECM proteins from porcine to human beings, it is even possible to implant such constructs into patients after recellularization of our scaffold with human cells. This indicates that our porcine matrix is suitable for human tumor model generation. Furthermore, porcine gut is easily obtainable from commercial pig breeders, which facilitates the production of pharmaceutical test systems.

Caco2 cells are well established for applications in 3D cell culture models that simulate the intestinal barrier and are used for drug uptake studies. As shown before, they exhibit a differentiated phenotype in a suitable microenvironment. To investigate to which extent our decellularized tissue scaffold reflects tumor characteristics—especially concerning invasion and metastasis—we focused on the more malignant SW480 cell line. In fact, the 3D microenvironment showed a stronger impact on SW480 cells than on Caco2 cells as marked by a downregulation of vimentin indicating MET induction—a hallmark for metastasis generation.

Remarkably, coculture with fibroblasts induced only SW480 cells to form tumor-like cell clusters with fibroblasts within the matrix, whereas Caco2 cells grew on top of the matrix separated from the fibroblasts beneath. In some areas, we observed the generation of multilayers as found in hyperplastic epithelial regions: one of the first steps in tumorigenesis. This indicates different cell-specific interactions of both tumor cell types with the same fibroblast population and stresses the impact of signaling circuits between the tumor–stroma and the tumor cells on tumor formation. It is widely recognized that the tumor–stroma plays a key role in tumor growth and invasion by the secretion of degrading enzymes and by storing cytokines. This changes signaling and thereby drug response in different invasive states. In fact, colon cancer tissue usually displays a very large proportion of fibrotic tissue. These issues further emphasize the importance to include the stroma component in preclinical tumor models.

To test the robustness of the system, different seeding conditions were compared such as different ratios of tumor cells to fibroblasts, different orders of seeding, first the fibroblasts and then the tumor cells or vice versa, as well as different donors of primary fibroblasts. Similar morphological results were observed in all the different experimental settings (data not shown). For standardization, we seeded both kinds of cells at the same time to generate the test model.

On a molecular level, we investigated the differentiation status and the degree of EMT by E-cadherin/β-catenin and PCK/vimentin fluorescence double staining, respectively. In healthy epithelia, β-catenin is linked to E-cadherin at the cell border connecting it to the actin cytoskeleton. In primary tumors, E-cadherin expression is reduced and translocates to the cytoplasm at the invasion front. β-Catenin dissociates from E-cadherin and also translocates to the cytoplasm as well as into the nucleus. In this study, it
induces the expression of other genes important for EMT and inhibitors of apoptosis, as reviewed concisely by Lee et al.30 Nuclear localization of β-catenin correlates with a poor prognosis in the clinic.31 In colorectal carcinogenesis, many pro-oncogenic factors encourage β-catenin translocation from the adherens junction to the nucleus, and anti-oncogenic factors inhibit nuclear β-catenin signaling. This stresses its vital part in tumor development.52 When metastases extravasate into distant healthy tissues to form a macroscopic secondary tumor, cancer cells have to cross the basement membrane and revert to an epithelial state in a process that is called MET and a strong E-cadherin expression is found in these areas.33

Our 3D SW480 colon cancer model showed strong E-cadherin expression particularly in areas of dense tumor cell aggregate formation. This effect was more pronounced in a flow bioreactor under dynamic conditions enhancing close cell associations. Furthermore, we observed a strong downregulation of vimentin expression in 3D compared to 2D culture—under static as well as dynamic conditions. Since SW480 cancer cells are known to develop more epithelial characteristics at a higher confluency,34 this effect could be related to a higher confluency in the 3D cell culture. Thus, the induction of tighter cell to cell contacts in 3D cell culture conditions could trigger MET induction, as it takes place in tissues into which metastases originating from circulating single tumor cells grow. The MET process characterized by E-cadherin re-expression is thought to be a critical step during metastasis formation, as reviewed by Wells et al.35

As mentioned before, at least two different mechanisms exist for tumor cell invasion where fibroblasts play a central part. First, fibroblasts are important for single-cell invasion because this needs the EMT of tumor cells induced by myofibroblasts of the adjacent tumor–stroma.35,36 Second, they are also essential for collective invasion processes37 as they are able to remodel the scaffold and to guide tumor cells through this matrix. Accordingly, in deep layers of the collagen scaffold, we could observe patterns of fibroblasts in front of bulky tumor cell formations. In contrast to single-cell invasion, collective invasion is considered to be independent of EMT in tumor cells and to be associated with fibroblasts. The mesenchymal marker vimentin is not reactivated in SW480 cells in the coculture setting, indicating no EMT associated with fibroblasts. The mesenchymal marker vimentin is not reactivated in SW480 cells in the coculture setting, indicating no EMT associated with fibroblasts. Double staining of Ki67 and the epithelial marker PCK enables the assessment of tumor cell proliferation in a coculture setting. This model can become more sophisticated by using the standardized biological scaffold BioVaSc-TERM,14 which allows vascularization of tumor models. In serial bioreactors, this would also enable the investigation of the metastatic spread of single tumor cells into the artificial “blood” circulation following the process of MET, which is a critical step in metastasis.

SW480 colon cancer cells harbor an activating KRAS mutation that occurs in 30–40% of colorectal cancer,47 making them insensitive to epidermal growth factor receptor (EGFR) inhibitors.48 In the clinic, patients with these mutations are excluded from targeted therapies available against the EGFR. Therefore, novel therapeutic strategies are urgently needed for efficient intervention and could be gained by analysis of signaling changes on a targeted treatment in certain mutational backgrounds. These data could be integrated in bioinformatics network analyses as done before in the combined in vitro/in silico lung tumor model.12

Conclusion

In this study, we established a modular tissue-engineered 3D colon cancer model, including the adjacent tumor environment on a decellularized matrix. We were able to foster cell growth and metastasis-like tissue formation by coculturing fibroblasts and tumor cells in a newly developed bioreactor. This test system enables analyses of drug responses in advanced tumor stages, particularly of substances that target tumor–stroma interactions.

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Disclosure Statement

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