Establishment and characterization of cell lines from chromosomal instable colorectal cancer

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Abstract

AIM: To generate novel tumor models for preclinical validation of biomarkers that allow drug response prediction and individual therapeutic decisions.

METHODS: Cell line establishment was conducted by both direct in vitro culturing and in vivo xenografting followed by in vitro culturing procedure. A comprehensive characterization was subsequently performed. This included quality control, consisting of the confirmation of human and colorectal cancer (CRC) origin by DNA fingerprint and epithelial cell adhesion molecule (EpCAM) staining, as well as mycoplasma and human virus testing. Phenotypic analysis was done by light microscopy and multicolor flow cytometry. Histopathological examination ($\beta$-catenin and cytokeratin staining) was conducted in direct comparison to parental tumor tissues. Extensive molecular-pathological profiling included mutation analysis for CRC-associated driver mutations, assessment of chromosomal and microsatellite instability, and the grade of CpG island methylation. Additionally, an array-based comparative genomic hybridization analysis was performed. Drug responsiveness was assessed for a panel of classical and novel substances in clinical use for the treatment of solid cancers. Finally, tumorigenicity of the cell lines was tested by xenografting into immunocompromised nude mice.

RESULTS: Herein we describe the establishment of three ultra-low passage cell lines from two individual patients suffering from sporadic CRC. One cell line was derived directly from an early stage case (HROC18), whereas two cell lines could be established both direct from patient material and after xenografting from a late stage tumor (HROC32). All cell lines were free of contaminating mycoplasma and viruses. Molecular-pathological analysis allowed all cell lines to be classified as chromosomal instable (CIN$^+$). They were aneuploid, with CpG island promoter methylation and microsatellite instability being absent. The following mutational profile was observed both in the cell lines and the parental tumor tissue: HROC18: APC$\text{mut}$, p53$\text{mut}$, K-ras$\text{wt}$; HROC32:
INTRODUCTION

The landmark paper by Vogelstein et al[1] ushered in a new era of understanding genetic alterations during colorectal tumor development and led to a new model of colorectal tumorigenesis. Since then, great leaps have been made in categorizing the main molecular classes of colorectal carcinoma (CRC) by pathogenetic mechanisms, namely those having chromosomal and microsatellite instability, as well as by CpG island methylation phenotype (CIMP)[2]. These mechanisms allow the definition of five molecular subtypes of CRC[3]. Initially, Fearon and Vogelstein proposed that inactivation of the adenomatous polyposis coli (APC) tumor suppressor gene occurs as a first step with the activation of KRAS mutations[3]. The subsequent malignant transformation is driven by additional mutations in the transforming growth factor PIK3CA and TP53 pathways[4,5]. As a consequence of these fundamental findings, further mutational analyses have shown that, in an individual cancer, only a limited number of pathways are dysregulated by “driver” mutations from a circumscribed number of about 80 candidate cancer genes (i.e., cancer genome landscapes)[6]. As tumors are highly heterogeneous and normal mutation rates are insufficient to account for the required number of mutations, Loeb et al[7] proposed a mutator phenotype of cancer cells that allows them to increase their rate of new mutations.

This chromosomal instability phenotype (CIN+) is observed in 65%-70% of sporadic CRC and indicates an accelerated rate of gains or losses of whole or large portions of chromosomes, resulting in a karyotypic variability[8]. The consequence of CIN+ is an imbalance in the number of chromosomes (aneuploidy), subchromosomal amplifications, and a high frequency of heterozygosity loss[9].

Transfer of these insights into clinical practice led to the development of therapies targeting epidermal growth factor (EGF)-receptor (EGFR) blockade (with the pre-requisite of KRAS mutational analysis)[10,11], and it is highly probable that further “individual” molecular testing of patients’ tumor tissue will become standard to guide and improve anticancer therapy regimens in the near future.

In this study, we describe the establishment and functional characterization of novel CIN+ cell lines with their corresponding xenograft model. A broad analysis of tumor biology, genetic alterations, and assessment of chemosensitivity towards an extended range of chemotherapeutic drugs is shown. Considering these aspects, the described cell lines provide an excellent tool for further development of individual therapy regimens.

MATERIALS AND METHODS

Tumor preparation and cell line establishment

Primary CRC resection specimens of HROC18 and HROC32 were received fresh from surgery, with informed written patient consent. Both patients underwent surgery upon initial diagnosis; hence, neither of them received any kind of prior therapy (i.e., surgery, chemotherapy, or radiotherapy). All procedures were approved by the Ethics Committee of the University of Rostock (reference number HV 43/2004) in accordance with generally accepted guidelines for the use of human material. Tumor samples were cut into small pieces. Parts of the tumor were immediately frozen in freezing medium [fetal calf serum (FCS) containing 10% DMSO; Sigma Aldrich, Munich, Germany] at -80°C for subsequent xenografting. Other pieces were frozen in liquid nitrogen for molecular analysis. The cell line establishment protocol was adapted from that of Maletzki et al[12]. Tumor cells were released from surrounding tissue by scraping and passage through a nylon mesh (100 μm). Single cell suspensions were seeded on collagen-coated 6-well plates in Quantum tumor medium [supplemented with 10%
**In vivo tumorigenicity**

In vivo tumorigenicity of HROC18 and HROC32 cells was tested following subcutaneous (sc) injection of $5 \times 10^6$ cells (with or without Matrigel®, BD Biosciences, Heidelberg, Germany) into the right and left hind flank of six-week-old female NMRI Foxn1® mice. Mice were bred in the University of Rostock’s animal facilities and maintained in specified pathogen-free conditions. Their care and housing were in accordance with guidelines as put forth by the German Ethical Committee. Tumor growth was monitored for at least 10 wk. Outgrowing tumors were removed for histological examination.

In a parallel experiment, in vivo xenografting of a HROC32 patients’ tumor was performed. Subcutaneous tumor implantation was performed as described. Established xenografts ($\geq 1500$ mm$^3$) were removed and subjected to in vitro culture protocols as described above. In the case of HROC18, the quantity of tumor material was insufficient for in vitro xenografting; hence, cell line establishment was only done on patient tumor material.

**Histology and immunohistochemistry of original tumors and xenografts**

Histopathological examination of HE-stained primary tumors and the corresponding xenografts was performed according to standard protocols for clinicopathological CRC staging\(^{[22]}\), with additional staging information being compiled from patients’ clinical charts. Supplementary immunostainings from paraffin-embedded primary tumors were done for β-catenin (Cell Signaling Technology, Boston, United States) and MNF116 (cytokeratin; Dako, Hamburg, Germany).

**Mutational and methylation profile of tumor-associated target genes and determining the level of chromosomal instability**

Molecular classification was done according to Ostwald et al\(^{[3]}\) on an ABI 3500 genetic analyzer (Applied Biosystems, Darmstadt, Germany). Data analysis was carried out by taking advantage of SeqScape® Software v2.7 (Applied Biosystems). These data, as well as staging information compiled from the clinical charts, are summarized in Table 1. Mutations in tumor-associated APC, p53, KRAS, and BRAF$^{V_{600E}}$ genes were analyzed as described. DNA-methylation in CIMP-sensitive promoters was traced by MethyLight with a modified marker panel originally published by Ogino et al\(^{[21]}\). The degree of chromosomal instability was assessed using SNP Array 6.0 from Affymetrix (Cleveland, OH) according to the manufacturer’s instructions.

**DNA typing**

Genomic DNA was isolated from cell lines at different passages, matched tumor and normal tissue, and corresponding B cells using the Wizard® Genomic DNA Purification Kit provided by Promega (Mannheim, Germany). Highly polymorphic short tandem repeat (STR) DNA marker (CSF1PO, TPOX, TH01, vWA, D16S539, D13S317, and D7S820) and the gender marker amelogenin were amplified in standard polymerase chain reaction (PCR) reactions and analyzed on an ABI Prism 3100 system. PCR primers were based on the original publication\(^{[20]}\).

**Generation of peripheral B cultures from patients**

Peripheral blood lymphocytes were purified by Ficoll density-gradient centrifugation. B-lymphoid cell lines (B-LCLs) were generated by Epstein-Barr virus-transformation as described previously\(^{[23]}\). Outgrowing B-LCL cultures were harvested, expanded, characterized, and frozen.

**In vitro growth kinetics, ploidy, and cell cycle analysis**

Doubling time of HROC18 and HROC32 cells was determined from serial passages. Therefore, $5 \times 10^6$ viable cells were seeded into 25 cm$^2$ flasks and the number of viable cells (defined by trypan blue exclusion) determined daily for seven days. Cultures were fed every 3 or 4 d. Ploidy and cell cycle analysis was performed

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**Table 1 Clinical and pathological characteristics of patients as well as cell line establishment protocol**

| Tumor-ID | Age/ gender | Tumor location | TNM-stage | UICC stage | Tumor type | Molecular type | β-catenin translocation | Corresponding Xenograft | Direct cell line establishment | Cell line from xenograft | Paired B-LCL |
|----------|-------------|----------------|------------|------------|------------|----------------|------------------------|------------------------|---------------------------|------------------------|-------------|
| HROC18   | 65/F        | Caecum         | G2T2N0     | I          | Primary adenocarcinoma | spStd | +       | -                     | +                      | -                        | Yes         |
| HROC32   | 83/F        | Ascending colon| G2T4N2     | N          | Primary adenocarcinoma | spStd | +       | +                     | +                      | +                        | Yes         |

F: Female; spStd: Sporadic standard; B-LCL: B lymphoid cell line; +: Positive; –: Negative.
Mycoplasma contamination was tested by the 16S-rRNA-Mycoplasma and human viral infection (University of Rostock Medical Center). Institute of carbohydrate antigen 19-9 was kindly accomplished by the Measurement of carcinoembryonic antigen (CEA) and plate reader (Tecan Infinite immediately measured at a wavelength of 450 nm on a stopped (100 incubation in the dark until color change. Reactions were (3,5,3',5'-tetramethylbenzidine) solution (100 min). Reaction visualization was done by adding TMB and subjected to streptavidin-HRP conjugates (1:500; 30 again, incubated with biotinylated antibody (1:200, 1 h), incubated (1 h, room temperature). Plates were washed (1 × PBS) and plates following five washing steps (1 × PBS) and blocking (2% BSA, 1 h, room temperature), standards and samples were transferred in duplicate wells and incubated (1 h, room temperature). Plates were washed again, incubated with biotinylated antibody (1:200, 1 h), and subjected to streptavidin-HRP conjugates (1:500; 30 min). Reaction visualization was done by adding TMB (3,5,3',5'-tetramethylbenzidine) solution (100 μL/well) and incubation in the dark until color change. Reactions were stopped (100 μL of 2 mol/L H2SO4 solution) and plates immediately measured at a wavelength of 450 nm on a plate reader (Tecan Infinite® M200, Crailsheim, Germany). Measurement of carcinoembryonic antigen (CEA) and carbohydrate antigen 19-9 was kindly accomplished by the Institute of Clinical Chemistry and Laboratory Medicine (University of Rostock Medical Center).

Flow cytometric phenotyping of primary cell lines
Cell surface marker expression on CRC lines was traced by flow cytometry with and without interferon-gamma (IFN-γ) pre-treatment using a panel of FITC-, PE, or APC-conjugated Abs: CD11b, CD15, CD20, CD24, CD25, CD28, CD34, CD43, CD44, CD45, CD45ra, CD45rb, CD50, CD54, CD55, CD56, CD58, CD62L, CD66acde, CD69, CD71, CD80, CD86, CD102, CD154, CD178, CD326, MHC I (Immunotools, Friesoythe, Germany), MHC II (Miltenyi Biotec, Bergisch-Gladbach, Germany), and HLA-A2 (cell culture supernatant clone BB7.2). For HLA-A2, a secondary FITC-conjugated anti-mouse Ab was applied (Dako). Samples were analyzed using CellQuest software (BD Biosciences).

HLA typing
A two-digits encompassing resolution typing on patient-derived cell lines of HLA loci (HLA-A, -B and, -C, and HLA-DR and -DQ) was performed by Transfusion Medicine at the University of Rostock.

Cytokine secretion pattern
Spontaneous cytokine release by tumor cells was analyzed from duplicates (3 × 10⁶ cells/well) in 24-well plates. Cell-free supernatants were harvested at different time-points and cytokine production (IL8, and IL10, Immunotools) free supernatants were harvested at different time-points from duplicates (3 × 10⁴ cells/well) in 24-well plates. Cell-cycle analysis was carried out by applying Modfit software (Verity Software House, Topsham, ME, United States). Matched B-LCLs were used as diploid controls.

Wound healing assay
Tumor cells were seeded in duplicates (6 × 10⁶ cells/well, 6-well plates) and grown to 100% confluency. Monolayers were scratched with a 10 μL pipette tip to induce a wound. Wounded edges were imaged using a Zeiss inverted microscope. Images were taken daily for a period of 9 d under a × 10 objective lens. Migration property was calculated by measuring scratch diameter under a microscope within each well.

In vitro chemosensitivity analysis
Cells were seeded into 96-well microtiter plates at 5 × 10³ or 1 × 10⁴ cells/well. When cells reached 30%-40% confluency, cultures were exposed to increasing drug concentrations of 5-fluorouracil (5-FU), cisplatin, irinotecan, gemcitabine, taxol, trastuzumab, rapamycin, erlotinib (courtesy of the University of Rostock's hospital pharmacy), or nilotinib (kindly provided by Novartis AG). Substances were selected based on the following criteria: (1) clinical approval for first-, second-, or third-line therapy of CRC patients (5-FU, irinotecan)[24]; (2) experimental and/or (pre-) clinical evidence of efficacy against CRC tumors (gemcitabine, taxol, cisplatin, and trastuzumab)[25,26]; and (3) promising novel targeted drugs, especially in case of metastatic disease (rapamycin, erlotinib, and nilotinib)[27].

After three days of exposure, media were removed and replaced by fresh medium supplemented with therapeutics. Following another three days, medium was removed, plates carefully washed, and they were then stained with crystal violet (0.2%, 10 min). Finally, drug effects from triplicate wells were determined at the level of 50% inhibition (IC50) in comparison to control, measured at 570 nm (reference wavelength: 620 nm).

Statistical analysis
Statistical analysis was done upon treatment with chemotherapeutics and targeted inhibitors. IC50 values were calculated using Sigma Plot 12.5 software (Systat Software Inc., San Jose, CA) applying the four parameter logistic function standard curve analysis for dose response. Values are given as absolute numbers.

RESULTS
Clinicopathologic characteristics of the primaries and cell line establishment
Parental tumors for HROC18 and HROC32 were primary colorectal carcinomas of the right colon. Histologically, both tumors were moderately well-differentiated tubular adenocarcinomas; one presenting at an early stage with infiltration into the muscular bowel wall, but without gene-based PCR amplification method from whole cell lysates. For determining potential polyomavirus infection (JC/BK and SV40), gDNA was isolated using the Wizard genomic DNA purification kit (Promega). All procedures were performed as described by Maletzki et al[26].
regional or distant metastasis (parental to HROC18), and the other at late stage with penetration of the peritoneum, regional lymph node metastases, and synchronous distant metastases to the liver and brain. Molecularly, both tumors were classified as sporadic standard (spStd) type and with nuclear beta-catenin translocation (Figure 1 and Table 1).

For cell line establishment, in vitro and in vivo approaches were combined as previously described[18]. With this method, two individual patient-derived cell lines could be established (HROC18 and HROC32). Furthermore, a xenopatient-derived cell line from HROC32 was also obtained. In the case of HROC18, the obtained tumor material was insufficient for xenografting in addition to primary cell line establishment. All three lines exhibited stable outgrowth as defined by passaging > 50 times. Hence, three cell lines obtained from two individual tumor patients were available for subsequent analysis. For clarity, patient-derived cell lines were marked with the suffix P and the xenograft-derived cell line with suffix X.

Morphology and viral contamination
As determined by phase contrast microscopy, cells adhered tightly to the cell culture flask. Morphologically, all cell lines showed an epithelial-like phenotype without contaminating fibroblasts. HROC18 cells proliferated as polygonal cell clusters in shape with more regular dimensions (Figure 2, upper panel). Patient-derived HROC32 and their matched xenopatient-derived cells proliferated as tightly-packed multi-cellular islands (Figure 2, middle and low panels) that did not grow to complete confluence. Of note, no apparent morphological differences were observed between these two lines, indicative for the same dominant clone growing out of the initial tumor culture. In all cases, morphology did not change during long-term cell culture (up to 60 passages). As determined by semi-quantitative PCR, cell lines were found to be free of mycoplasma (Figure 3A) or other potential contaminants like human polyoma viruses (SV40 and JC/BK, data not shown).

Growth kinetics, ploidy analysis, and phenotype
Next, doubling times were determined over a culture period of seven days. Experiments revealed no significant differences between individual cell lines. Doubling times were 32 ± 10.6 h for HROC32P, 33.8 ± 5.4 h for HROC32X, and 35 ± 7.2 h for HROC18 cells. Cells did not grow faster after serial passages (data not shown).

Their epithelial origin was confirmed by high epithelial cell adhesion molecule expression (EpCAM; each > 98%; Table 2). More detailed phenotypic characterization revealed comparable expression levels for the transferrin receptor CD71, but heterogeneous expression of adhesion and co-stimulation markers (CD54, CD58, and CD102). All cell lines expressed MHC class I and were additionally found to be HLA-A2 positive (Table 2). MHC class II expression was absent on all lines, but could be induced by IFN-γ pre-treatment (up to 80%; Table 3).

Secretion profile
Secretion of the CRC tumor markers CA19-9 and CEA, as well as chemotactic and pro-angiogenic factor IL8, was determined from supernatants collected at different times of culture (Figure 3B). HROC18 and HROC32 secreted detectable levels of both tumor markers, with HROC18 showing high CA19-9, but low CEA secretion. On the other hand, the secretory picture of HROC32 was reversed, with high CEA and quite low CA19-9.
levels. IL8 was predominantly detected in HROC18-derived supernatants. IL10 was not detectable at all.

Molecular signature, DNA fingerprint, and identification of novel tumor suppressor genes

First, DNA fingerprint was performed to verify genetic identity with the patients. A panel of highly polymorphic STR DNA markers covering four validated STR loci on different chromosomal locations was included into this analysis. Using this approach, identity with the patient was confirmed for all three cell lines even at late passage, hence cross-contamination with other (tumor) cells could be ruled out. Additionally, mutational profiling was done for a series of frequently affected tumor suppressor/proto-oncogenes (i.e., TP53, APC, KRAS, and BRAF). Examinations were done on all three cell lines and compared to the parental tumor. Notably, no differences were found in any of the matched samples (Table 4). HROC18 and HROC32 showed characteristic molecular features associated with the spStd type. HROC18 displayed TP53 and APC gene mutations. HROC32 cells presented with wild type APC, but mutations in the TP53 and KR-AS genes. In line with the characteristic molecular signature, no methylation in CIMP-sensitive promoters, as traced by MethyLight, was found. Similarly, no BRAF or PTEN mutations were detected.

More detailed analysis of potentially novel tumor suppressor genes identified individual differences between both tumor models. In detail, expression patterns of MBNL1, IIMPL2L, FHIT, WWOX, and MACROD2 were...
examined by endpoint PCR. Alterations were detected as heterozygous point mutations, as demonstrated for FHIT (HROC18) and WWOX. IMMP2L and MACROD2 were homozygously deleted in both tumor models. MBNL1 showed no alteration with normal expression values.

Finally, we performed a very high resolution genomic analysis employing SNP Array 6.0. In this analysis, instability on the chromosomal level was confirmed. Both

Table 3  Human leukocyte antigen: Typing of colorectal carcinoma cell lines

| DNA     | HLA class I | HLA class II | HLA class III |
|---------|-------------|--------------|---------------|
|         | A<sup>+</sup> | B<sup>+</sup> | Cw<sup>+</sup> | DRB1<sup>+</sup> | DQB1<sup>+</sup> |
|         | 1 allele | 2 allele | 1 allele | 2 allele | 1 allele | 2 allele | 1 allele | 2 allele | 1 allele | 2 allele |
| HROC18  | 01 | 02 | 08 | 39 | 07 | - | 03 | 13 | 02 | 03 |
| HROC32P | 01 | 02 | 08 | 51 | 07 | 14 | 03 | 11 | 02 | 03 |

<sup>*</sup>The number of the alleles. Information on both alleles is provided; in case of homozygosity the “second” allele is marked by “-”. HLA: Human leukocyte antigen.

Table 4  Comparative molecular characterization of patients’ tumor and corresponding colorectal carcinoma cell lines

| Tumor-ID | TPS3 | APC | KRAS |
|----------|------|-----|------|
|          | Exon 5 | Exon 6 | Exon 7 | Exon 8 | Cd1 | Cd2 | Cd12 | Cd13 | B-raf<sup>V600E</sup> |
| HROC18   | Patient tumor | wt | wt | wt | mut | mut | wt | wt | wt | wt |
| HROC32   | Cell line | wt | wt | wt | mut | mut | wt | mut | wt | wt |
|          | P-Cell line | wt | wt | wt | mut | wt | mut | wt | wt | wt |
|          | X-Cell line | wt | wt | wt | mut | wt | mut | wt | wt | wt |

wt: Wild type; mut: Mutated; Cd: Codon.

Figure 3  Mycoplasma polymerase chain reaction, ploidy status, and secretion profile of tumor cells. A: Displayed are the results for 16S-rRNA-gene-based polymerase chain reaction for mycoplasma. All three lines were found free of contaminating mycoplasma. Samples amplified in the absence of tumor DNA served as a negative control, whereas a highly contaminated cell line was applied as a positive control, run in duplicates; B: Cytokine secretion pattern as determined by enzyme-linked immunosorbent assays. Cytokine concentrations were determined by comparison with a standard curve generated from serial dilutions of individual standards. Quantitative analysis of carcinoembryonic antigen (CEA), carbohydrate antigen 19-9 (CA19-9), and interleukin (IL) 8 secretion after three and five days of culture, respectively. Results show the mean ± SD of three independent experiments.
Figure 4 Single nucleotide polymorphism array 6.0 for assessment of chromosomal instability in cell lines. Analysis was performed according to manufacturer’s instructions. A: HROC18 cells; B: HROC32P cells.
Migration of tumor lines

A scratch wound healing assay was performed, with cell migration upon wounding being assessed over a period of 9 d. As can be taken from Figure 5, HROC18 cells migrated within five days throughout the initial wound (migration speed: 34.8 μm/d). By contrast, HROC32P cells exhibited lower levels of migration (Figure 5), finally taking 9 d to close the wound (migration speed: 24.8 μm/d).

In vitro drug response

Drug response testing on individual cell lines has become increasingly important due to the large heterogeneity of individual tumors in terms of therapy response. Hence, a series of clinically approved and potentially novel substances was included into in vitro analysis (Table 5). Responsiveness was examined following two 72-hour cycles, closely resembling the clinical regimen.

Generally, HROC18 and HROC32 (P + X) cells responded to clinically approved CRC drugs (i.e., 5-FU, cisplatin, and irinotecan). HROC18 cells showed the highest sensitivity towards 5-FU (IC50 value: 0.48 ± 0.28 μmol/L). HROC32X tended to be more resistant against these drugs than its parental line (HROC32P), especially following irinotecan treatment (IC50 values: 3.83 ± 1.40 μmol/L vs 0.15 ± 0.07 μmol/L). Of note, all lines were susceptible towards targeted therapies. Among these therapeutics, the mammalian target of rapamycin (mTOR) inhibitor rapamycin (IC50 value of each cell line: < 0.20 μmol/L) was most potent. Nilotinib, an inhibitor of the DDR, KIT, and PDGFR receptor kinases, showed IC50 values between 5.4 and 7.5 μmol/L, and erlotinib,
a selective EGFR inhibitor, inhibited proliferation with IC₅₀ values in the range of 0.87-12.2 μmol/L. By contrast, taxol did not show significant growth inhibition at all, which is in line with the recent literature in CIN tumors[29]. Likewise, trastuzumab had no impact on cellular growth.

**In vivo tumorigenicity**

To evaluate in vivo growth behavior, mice were subcutaneously implanted with 5 × 10⁶ patient-derived HROC18 and HROC32 tumor cells, respectively. Growth kinetics were different between these two cells, with HROC32P growing more rapidly than HROC18 (Figure 6). Thirty-five days after injection, HROC32 tumors became apparent and continued to grow until the experimental endpoint (day 77, tumor diameter: 10.2 ± 1.9 mm). By contrast, HROC18 tumors presented with low tumorigenic potential and tumor sizes suspended at 6.2 ± 0.3 mm in diameter (day 77). The addition of Matrigel to the tumor cell suspension did not improve tumor growth (data not shown).

Next, experiments were performed in NOD-SCID mice. Due to a lack of NK cells, these mice grafted human cells better than Foxn1nu mice. Once again, low or absent tumor growth was observed for HROC18 cells, confirming their low tumorigenic potential in vivo (data not shown).

**DISCUSSION**

Identifying subsets of patients who will benefit from a particular (targeted) therapy requires improved integration of pre-clinical models and cutting-edge clinical examination. In this case, individual tumor models have become increasingly important research tools[29-32]. Such models helped to identify the large morphologic and molecular heterogeneity that exists among individual tumor cases; yet these findings have contributed to a better understanding of CRC biology and guided the way for individualized pre-clinical drug testing. Hence, this study was conducted to establish novel CRC cell lines obtained from patients with primary spStd type tumors. Tumors were resected at early and late stages, thereby representing two different models that expand our collection on individual tumor models. This collection includes primary material (frozen and paraffin-embedded tumors, normal tissue, and lymphocytes) together with xenografts, patient-derived and xenopatient-derived cell lines, as well as matched B-LCLs[18,33,34].

Cell lines generated in this study were directly obtained from patient tumor material, thereby perfectly reflecting the original molecular signature of the tumor. For cultivating primary tumor cells, a simple, yet rapid, method described by Maletzki et al[38] was applied. This method was found to be reliable and highly reproducible, yielding an approximately 10% success rate of primary tumor cell line establishment, which is comparable to that reported by others[35,36]. In order to improve cell line establishment and obtain more individual tumor models for functional comparative studies, tumor fragments were additionally engrafted in immunodeficient mice without prior in vitro exposure[35,37]. With this approach, up to 60%-80% success can be achieved (own unpublished data). Cell lines established following xenografting often show even better in vitro growth behavior than their parental counterpart cell line. They even adequately recapitulate molecular hallmarks of parental tumors and exhibit good in vivo tumorigenic potential, which is crucial for therapeutic interventions. However, there are also some limitations that must be kept in mind: (1) xenopatient-derived cell lines undergo a kind of in vivo preselection and may thus not represent the full tumor heterogeneity; (2) due to selective pressure, growth behavior, and consequently drug response, may differ; and (3) in vivo xenografting is very time-consuming, lasting up to 6 mo with follow-up cell line establishment potentially adding another year.

In this study, HROC32 cells gave rise to in vitro growth after xenografting, and consequently three cell lines were available from two individual tumor cases. The genetic identity with the patient was verified for these lines by means of STR fingerprinting. All three lines showed characteristic molecular features associated with the spStd type (i.e., complex chromosomal aberrations, TP53 gene mutations, and absence of GpG island promoter methylation). In line with our previous study on microsatellite unstable tumor cell lines, molecular profiles matched between cell lines (both X and P) and the parental counterpart[39]. KRAS, a well-established predictive biomarker for EGFR-targeted therapies[38], was mutated only in the late stage CRC case (HROC32). Moreover, this tumor harbored a mutation in codon 12 that is associated with worse outcome and early relapse after chemotherapy compared to codon 13 or other KRAS gene mutations[39].

Confirming their origin, cell lines were deliberated as epithelial tumor cells (> 80% EpCAM+) with heterogeneous expression of adhesion molecules (CD15, CD54, CD58, and CD102) and tumor antigens (CD66acde). Transferrin receptor CD71 was highly expressed on tumor cells, indicative for a high proliferative index. Due to their natural immunosuppressive behavior, co-stimulatory molecules (CD80 and CD86) were not detectable at all, while MHC class I-expression was preserved. HROC18 and HROC32 cells were additionally found to be HLA-A2 positive. This provides a ready basis for subsequent immunological studies aiming at the identification of immunogenic epitopes from novel candidate antigens. Given the fact that about half of Caucasians and Asians are HLA-A2 positive, this finding is of particular relevance.

Common characteristics for adaptation to cell culture conditions include changes in morphology towards a rather undifferentiated phenotype and an accelerated rate of growth[18,40]. However, both parameters remained unchanged in HROC18 and HROC32 (X + P) lines, at least over 60 passages. Directional cell migration, as determined by a classical wound healing assay[41], was
quite individual, with HROC18 cells migrating faster than HROC32 cells. Since these findings may give a hint towards in vivo growth behavior and has potential consequences for the efficacy of antineoplastic drugs in vivo, the tumorigenic potential was studied. However, tumorigenicity was always found to be low. Together, these findings are indicative for outgrowth of one dominant cell clone already in the initial in vivo and in vitro culture.

In many CRC patients, therapy resistance and subsequent relapse occurs early after initial objective response. In terms of targeted therapies, it was found that single cells resistant to any targeted agent are already present at the start of therapy. In most cases, these cells clonally expand once the therapy is applied, causing tumor recurrence. Recently, a mathematical model describing the evolutionary dynamics of lesions in response to treatment was developed that attempts to predict the efficacy of drugs and certain combinations. Hence, a large individual variability is actually found even in molecular closely-matched tumor samples. Therefore, maintenance of the original tumors’ molecular signature is in demand for determining (initial) drug sensitivity experimentally and for predicting response. Since this condition is completely fulfilled in our freshly established cancer cell lines, a large panel of chemotherapeutics and targeted drugs was tested. In principle, cell lines were susceptible towards commonly used chemotherapeutics (i.e., 5-FU, cisplatin, and irinotecan), while the targeted therapies evaluated did not show significant growth inhibition, they were indicative for intrinsic resistance (of the in vitro outgrown cell clone). Of note, HROC18 cells were generally more resistant than HROC32 cells (both X and P), underscoring the necessity of personalized therapy. A rather unexpected, yet very interesting, finding was the responsiveness towards rapamycin. Rapamycin, produced by 

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torosus 

lyngbyensis, 

was the first mTOR inhibitor discovered that has been defined as a “longevity enhancer and cancer preventative agent” in TP53 gene deficiency. Though evidence from the literature indicates the resistance of CRCs to the antitumorigenic effects of rapamycin, we still observed significant growth inhibition in our cell lines. The underlying resistance mechanism is supposed to be mediated by feedback activation of PI3K/Akt and Ras-MAPK signaling and, accordingly, cell survival. Recently, the addition of sorafenib has been found to abrogate rapamycin resistance in cells carrying oncogenic KRAS and PIK3CA. Hence, this combination may deliver another option for selected patients and underscores the role of mTOR inhibitors as combinatorial agents in cancer therapy despite their so far limited clinical success.

In principle, an ideal preclinical tumor model is of low passage, shows close similarity to its parental tumor (≥ 95%), harbors specific genomic alterations of interest (for targeted approaches), and fulfills more practical issues like good in vitro and in vivo growth characteristics. Our collection of individual tumor models meets these criteria; hence these models provide a virtually unlimited source of tumor material for predicting treatment and response. This will help to improve the ability for personalizing tumor therapy.

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COMMENTS

Background

The chromosomal instability (CIN) subtype occurs in 65%-70% of all sporadic colorectal cancer (CRC). Preclinical validation of novel biomarkers that allow drug response testing and prediction is ideally being performed in patient-derived, individual tumor models. However, ultra-low passage and well-characterized CRC models are still rare.

Research frontiers

Individual tumor models provide excellent tools for both basic and translational research. Even more importantly, they are a prime choice for the development of personalized tumor therapies.

Innovations and breakthroughs

In this study, the authors describe the successful establishment of two different tumor models showing characteristic features associated with the CIN phenotype. Due to their individual character, morphology, phenotype, and drug response were quite heterogeneous. This substantially extends the collection of individual tumor models, which include all known CRC subtypes. By establishing and comprehensively characterizing obtained tumor models, useful instruments for subsequent development of individualized tumor therapies are supplied.

Applications

Patient-derived tumor models constitute a virtually unlimited source of tumor material, both for further analyzing their biological characteristics and for predicting drug response. This will help to improve the ability for personalizing tumor therapy in the near future.

Terminology

Chromosomal instability: A typical feature of most solid cancers involving structural and numerical alterations of the normal chromosome numbers. Ultra-low passage tumor cell lines: Culture of primary (patient-derived) tumor cells which grow continuously, but with only a low number of cell divisions outside the body. This guarantees very low or absent molecular changes in comparison to the parental tumor.

Peer review

This descriptive study represents the methodology and characterization of CRC cell lines both from patient material and xenografts of original patients’ tumors. By applying a large number of methods for analyzing tumor biology, genetic alterations, and chemosensitivity, the authors present an interesting way for generating individual tumor models.

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