Label-Free Imaging Analysis of Patient-Derived Cholangiocarcinoma Organoids after Sorafenib Treatment

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Abstract: Monitoring tumor growth dynamics is crucial for understanding cancer. To establish an in vitro method for the continuous assessment of patient-specific tumor growth, tumor organoids were generated from patients with intrahepatic CCA (iCCA). Organoid growth was monitored for 48 h by label-free live brightfield imaging. Growth kinetics were calculated and validated by MTS assay as well as immunohistochemistry of Ki67 to determine proliferation rates. We exposed iCCA organoids (iCCAOs) and non-tumor intrahepatic cholangiocyte organoids (ICOs) to sub-therapeutic concentrations of sorafenib. Monitoring the expansion rate of iCCAOs and ICOs revealed that iCCAO growth was inhibited by sorafenib in a time- and dose-dependent fashion, while ICOs were unaffected. Quantification of the proliferation marker Ki67 confirmed inhibition of iCCAO growth by roughly 50% after 48 h of treatment with 4 \( \mu \)M sorafenib. We established a robust analysis pipeline combining brightfield microscopy and a straightforward image processing approach for the label-free growth monitoring of patient-derived iCCAOs. Combined with bioanalytical validation, this approach is suitable for a fast and efficient high-throughput drug screening in tumor organoids to develop patient-specific systemic treatment options.

Keywords: label-free live imaging; brightfield microscopy; tumor organoids; primary liver cancer; intrahepatic cholangiocarcinoma; sorafenib

1. Introduction

Bile duct cancers, or cholangiocarcinomas (CCAs), are rare but aggressive tumors arising from the biliary tree. CCAs have a very poor prognosis [1] and are categorized by anatomical location as intrahepatic (iCCA), perihilar (pCCA), or distal (dCCA). Liver resection or transplantation are the only curative treatment options for CCA [2]. However, the five-year survival is less than 10% [3] due to the high recurrence rate of 85% within the first three years after resection [4].

Today, non-resectable CCAs are treated with a combination of gemcitabine and cisplatin, extending median survival significantly by about 3.6 months [5]. Sorafenib, a multi-kinase inhibitor, has shown to be effective in the treatment of solid cancers such as renal cell carcinoma and hepatocellular carcinoma [6]. Although several studies showed a potential therapeutic beneficial effect, the therapeutic benefit in CCAs is still largely unknown [7–9].
Importantly, the interpatient heterogeneity of CCAs requires individual, patient-tailored systemic therapy to optimize the therapeutic outcome while simultaneously reducing adverse effects. However, appropriate procedures involving the patient-derived tumor tissue to screen for the most effective drug treatment are rarely available, if at all.

In addition to intrahepatic cholangiocyte organoids (ICOs) derived from the biliary compartment of the liver, an innovative CCA organoid model (CCAO) was recently established from resected tumor specimens and core tumor needle biopsies [10–12]. They were shown to preserve the majority of the mutational landscape and represent the epithelial cancer cell compartment of the original tumor to a large extent, leading to similar histopathology upon xenografting [10,11,13–15]. Moreover, they are suitable models for high-throughput drug screening approaches using luminescence or spectrophotometric endpoint readouts for cell viability [10,13,16].

Understanding biological systems such as (tumor) organoids and their dynamic behavior requires adequate techniques to investigate cellular dynamics under close-to-physiological conditions with minimal extrinsic perturbation [17,18]. To this aim, live imaging by advanced light microscopy together with subsequent advanced image analysis offer suitable approaches [19]. Different advanced light microscopy techniques are currently available that each hold unique advantages but also limitations with regards to in-depth resolution and invasiveness into the observed system [20].

To study organoid dynamics live at a global (entire culture) scale, brightfield microscopy presents a suitable solution for observing global changes in organoid size and morphology in multiple organoids simultaneously [21]. Brightfield microscopy offers a label-free, minimally invasive approach that can be easily adapted for high-throughput imaging. The reduced level of detail in the acquired image data allows for comparatively easy data handling and subsequent image analysis [22,23].

More detailed in-depth observations on the single-cell level can be achieved by fluorescence microscopy techniques such as confocal laser scanning microscopy (CLSM) [24] or light sheet-based fluorescence microscopy (LSFM) [25,26]. The vast amounts of data produced with these techniques, however, require considerable resources for post-processing and data evaluation [27,28].

In this project, we combined brightfield microscopy and an organoid-specific brightfield image analysis pipeline [21] to monitor the growth of patient-derived iCCAO treated with sorafenib, a drug currently not used for therapeutic intervention in cholangiocarcinoma.

2. Materials and Methods

2.1. Patient-Derived Liver Tissue and Organoid Initiation

Organoids were generated from one iCCA tissue sample collected after resection of the tumor for curative treatment (iCCAOs), and one healthy donor liver tissue sample (ICO) collected at the Erasmus MC—University Medical Center Rotterdam. The use of these tissue samples was approved by the Medical Ethics Committee of the Erasmus MC—University Medical Center Rotterdam (MEC-2013-143 and MEC-2014-060). All patients consented to the use of materials for research purposes. Organoids from iCCA biopsies and from healthy liver tissue were cultured as described previously by Broutier et al. [10] and Huch et al. [29]. The iCCAOs used in this study were characterized by Broutier et al. (originally labelled CC-1), and tumorigenicity was confirmed by xenografting and whole-exome sequencing [10].

2.2. Cell Proliferation Assay

The relative proliferation of iCCAOs was determined by CellTiter 96© Aqueous One Solution Cell Proliferation Assay (MTS) according to the supplier’s instructions (CellTiter 96© AQueous One Solution Cell Proliferation Assay, Technical Bulletin, version: revised 12/12, TB254, Promega, Walldorf, Germany). Organoids were dissociated into single cells by incubation in trypsin-EDTA (0.25%, Thermo Fisher Scientific, Dreieich, Germany) at 37 °C for five to ten min and vigorous pipetting. In total, 5 × 10^3 organoid cells per well
were seeded into 96-well plates (5 µL Matrigel™, Corning, Wiesbaden, Germany + 100 µL human liver expansion medium [30]). Cells were grown for seven days to form organoids before treatment with sorafenib (Santa Cruz Biotechnology, Heidelberg, Germany) for 72 h. Before the addition of 20 µL of CellTiter 96® AQueous One Solution Reagent, the expansion/treatment medium was exchanged with 100 µL of basal medium [30]. Samples were incubated for one hour at standard culture conditions (37 °C, 5% CO₂, humidified). The assay medium was transferred into a white/clear bottom 96-well plate and absorbance was measured on the microplate reader Tecan® Infinite® M200 with the corresponding software i-control™ (Tecan, Crailsheim, Germany).

2.3. Organoid Growth Analysis Pipeline—Organoid Size Measurements on Time-Resolved 2D Brightfield Image Data

The organoid growth analysis pipeline described here widely corresponds to the brightfield pipeline published in Hof et al. 2021 [21] with minor changes in sample preparation, image acquisition, and data analysis. In brief, organoids (iCCAOs and ICOs) were seeded into 96-well plates (5 µL Matrigel™ + 100 µL human liver expansion medium [30]), and grown for 72 h prior to image acquisition. The medium was exchanged and supplemented with the resp. compounds, where indicated, before starting the image acquisition. Organoids were imaged for 48 h with a recording interval of 0.5 h (=97 time points, counting 0 h as time point 1) under controlled standard culture conditions (37 °C, 5% CO₂, humidified).

Brightfield time-lapse recordings of organoids were acquired on the microscope AxioObserver.Z1 equipped with objective lens Plan-Apochromat 5×/0.16, camera AxioCam MR R3, and software ZEN blue (version 3.2, Carl Zeiss, Jena, Germany). Organoid cultures were imaged with 10 z-planes (z-spacing: 65 µm), 2 × 2 tiles (overlap: 15%, scan mode: meander), a pixel size of 1.29 × 1.29 µm², an image size of 52,379 × 23,432 pixels, a bit depth of 12, an exposure time of 15 milliseconds (light source intensity: 3.22 V), and a binning mode of 1 × 1.

The recorded time-lapse image stacks were exported, converted from .czi to generic .tif format with software ZEN blue, and pre-processed with Fiji (ImageJ version 1.51n https://imagej.nih.gov/ij/download.html (accessed on 28 September 2022); Java version 1.8.0_6, 64-bit [31]). The dimensionality of the raw data sets was reduced from 4 (2 × 2) tiles with 10 z-planes each to one stitched image with one z-plane per time frame using the plugin Grid/Collection stitching [32]. The resulting image stacks were segmented based on projected luminal areas of the organoids by using the Fiji plugin Morphological Segmentation (MorphoLibJ [33]). Segmented luminal areas were measured with the Fiji plugin Region Morphometry (MorphoLibJ [33]). The luminal areas of individual organoids were normalized to the average luminal area over the first 5 time points. Organoids that were not detected in all 97 time frames were manually excluded from the analysis.

2.4. Immunohistochemistry

Organoids were fixed in paraformaldehyde and embedded in histogel (Thermo Fisher Scientific, Dreieich, Germany) followed by embedding in paraffin. Sections of 1 µm thickness were deparaffinized in the descending alcohol series. Epitopes were demasked for 30 min in citrate buffer, pH 6.0, using a pressure cooker. Following cooling for 30 min in ice water, endogenous peroxidases were blocked for 30 min with 3% H₂O₂ at 4 °C in the refrigerator. Sections were washed in PBS for 5 min, immediately followed by a 90 min BSA blocking step (5% BSA + 0.5% Tween20) in a humid chamber at RT. After applying the Vector-Kit SP-2001 according to the manufacturer’s manual (Vector Laboratories, Newark, CA, USA), slices were incubated overnight in a humid chamber at 4 °C with the mouse anti-human-Ki67 primary antibody (1:100, Dako M7240, Agilent Technologies Germany GmbH & Co., KG, Waldbronn, Germany). Following 3 washes with PBS for 5 min each, slices were incubated with the biotinylated goat anti-mouse secondary antibody (1:200, Dianova, Hamburg, Germany) for 1 h at RT in a humid chamber. After three additional
washes with PBS, incubation was continued for 30 min with the ABC reagent (Vector Kit
PK-6100; Vector Laboratories, Newark, CA, USA), and slices were counterstained with
nuclear fast red for 3 min, followed by washing with PBS and subsequent incubation
with Histogreen (Linaris GmbH, Dossenheim, Germany) for 2 min in a humid chamber.
After the ascending alcohol series, the sections were embedded in Entellan (Merck GmbH,
Darmstadt, Germany). For image acquisition, the microscope AxioImager.Z1 (Carl Zeiss,
Jena, Germany) was applied.

For the detection of SOX9, the protocol as described was essentially the same, except
that the primary rabbit anti-SOX9 antibody (1:200, ab185966, Abcam, Cambridge, UK) was
used in combination with the biotinylated donkey anti-rabbit antibody (1:200, Dianova,
Hamburg, Germany) as the secondary antibody.

Antigen demasking for immunofluorescence was achieved by 30 min incubation in
Tris-EDTA buffer (pH 9.0). Subsequently, unspecific binding was blocked in 5% goat
serum/PBS (ccpro, Oberdorla, Germany) for 20 min followed by 60 min in 5% BSA/0.5%
Tween20. For the co-detection of E-cadherin and ZO-1, the mouse anti-E-cadherin (1:200,
610182, BD Biosciences, Heidelberg, Germany) and the rabbit anti-ZO-1 (1:100, 44-2200,
InVitrogen, Carlsbad, CA, USA) antibodies were offered overnight in 1% BSA in PBS at 4°C
in a humid chamber. Following stepwise incubation with the goat anti-mouse Cy3 antibody
(1:200, 115-165-003, Dianova, Hamburg, Germany) for 1 h, two washings for 10 min each
in PBS, incubation with the goat anti-rabbit AlexaFluor 488 antibody (1:200, A11008, Life
Technologies, Ober-Olm, Germany) for 1 h at RT, three washings for 5 min each in PBS,
and nuclear counterstaining with DAPI (1:1000, Carl Roth GmbH + Co., KG, Karlsruhe,
Germany) for 5 min, slices were finally embedded in 50% glycerol solution and lacquer.
Images were taken using the microscope AxioObserver.Z1 (Oberkochen, Germany).

EpCAM and MDR1 were detected using the primary rabbit anti-EpCAM (1:200,
ab213500, Abcam, Cambridge, England) and the rabbit anti-MDR1 (1:250, ab170904, Ab-
cam, Cambridge, England) antibodies, resp., overnight at 4°C in combination with the
secondary goat anti-rabbit AlexaFluor 488 antibody (1:200, A11008, Life Technologies,
Ober-Olm, Germany).

2.5. RNA Sequencing Analysis

Total RNA was isolated using TRIzol™ Reagent according to the supplier’s instruc-
tions (TRIzol™ Reagent User Guide, version: 28 January 2020, Thermo Fisher Scientific,
Dreieich, Germany). In brief, organoids (iCCAOs and ICOs) were grown in 24-well plates
(8 × 5 µL Matrigel™ droplets + 500 µL human liver expansion medium (PMID: 27560176))
for at least 7 days. Organoid-containing wells were rinsed with warm PBS (37°C). Organoid
material from two 24-wells was resuspended and combined in 1.5 mL of TRIzol™ Reagent.
Volumes of reagents in the following steps were adjusted accordingly (×1.5). Lysates
were vortexed vigorously for 30 sec and passed through a 20-G needle ten times before
proceeding with the TRIzol™ protocol. RNA pellets were air-dried for at least 20 min and
resuspended in 20 µL of UltraPure™ DNase/RNase-Free Distilled Water (Thermo Fisher
Scientific, Dreieich, Germany). Samples were incubated at 55°C for 15 min.

cDNA was synthesized from 1 µg of total RNA isolate using the Maxima First Strand
cDNA Synthesis Kit for RT-qPCR according to the supplier’s instructions (Maxima First
Strand cDNA Synthesis Kit for RT-qPCR Product Information, version: 2012, Thermo Fisher
Scientific, Dreieich, Germany).

Transcripts of interest were amplified using KRAS-specific primers (Hs KRAS isoform
A/B, forward primer: 5′-GAG AGA GGC CTG CTG AAA ATG-3′, reverse primer: 5′-CCC
CGG CTC TCG GTT ATA AG-3′, biomers.net, Ulm, Germany) and Phusion™ High-Fidelity
DNA Polymerase according to the supplier’s instructions (annealing temp.: 56.0°C, 3-step
protocol, Phusion™ High-Fidelity DNA Polymerase, Product Information Sheet, version:
28 January 2020, Thermo Fisher Scientific, Dreieich, Germany). Reaction volumes were
scaled up to 100 µL containing 1 µL of undiluted cDNA template. To verify amplicon
sizes, 20 µL of the PCR product were subjected to standard agarose-gel electrophoresis
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(2% agarose/TAE, 100 V, 30 min, TAE buffer: 40 mM Tris-HCl (pH 7.6), 20 mM acetic acid, 1 mM EDTA in ultrapure H₂O).

In all, 80 µL of PCR product were cleaned-up with NucleoSpin® Gel and PCR Clean-up according to the supplier’s instructions (5.1 PCR clean-up, NucleoSpin® Gel and PCR clean-up, user manual, version: February 2017/Rev. 4, Machery-Nagel, Dueren, Germany). The PCR product was eluted in 20 µL of Buffer NE. A-tailing of cleaned-up blunt-end PCR product was achieved by incubation with Taq DNA Polymerase (Thermo Fisher Scientific, Dreieich, Germany) at 72 °C for 20 min (A-tailing reaction mix/10 µL: 1 µL of 10× PCR Buffer/-Mg, 0.25 µL of 50 mM MgCl₂, 0.25 µL of 10 mM dATP, 8.25 µL of template cDNA, 0.25 µL of Taq DNA Polymerase).

Next, 3 µL of the A-tailed PCR product was ligated into the pGEM®-T Easy Vector System I according to the supplier’s instructions for 10 µL reactions (pGEM®-T and pGEM®-T Easy Vector Systems, Technical Manual, version: revised 12/18, TM042, Promega, Walldorf, Germany) at 16 °C overnight.

The ligation product was transformed into Subcloning Efficiency™ DH5α™ Competent Cells according to the supplier’s instructions with slight modifications using the entire volume (10 µL) of ligation product (Subcloning Efficiency™ DH5α™ Competent Cells, Product Information Sheet, version: 17 January 2006, Thermo Fisher Scientific, Dreieich, Germany). In brief, cells were heat-shocked for 30 sec at 42 °C. Transformed cells were incubated in 950 µL of SOC medium (New England BioLabs, Frankfurt am Main, Germany) at 37 °C and 225 rpm for one h.

Transformed cells were LB/ampicillin/IPTG/X-Gal plates (100 µg/mL ampicillin in LB agar, coated with 100 µL of 100 mM IPTG, and 20 µL of 50 mg/mL X-Gal, Carl Roth, Karlsruhe, Germany; according to pGEM®-T and pGEM®-T Easy Vector Systems, Technical Manual, version: revised 12/18, TM042, Promega, Walldorf, Germany). Plates were incubated for 16-24 h. White colonies were selected and incubated in 5 mL of LB medium + 100 µg/mL ampicillin (Carl Roth, Karlsruhe, Germany) at 37 °C and 225 rpm overnight. Amplified plasmid DNA was extracted with NucleoSpin® Plasmid according to the supplier’s instructions (5.1 Isolation of high-copy plasmid DNA from E. coli, NucleoSpin® Plasmid, User manual, version: March 2019/Rev. 11, Machery-Nagel, Dueren, Germany). Plasmid DNA was eluted in 30 µL of Buffer AE.

Insert sizes in amplified plasmid DNA were determined by restriction digest at 37 °C for one h (Restriction digest mix: 12.5 µL of nuclease-free H₂O, 2 µL of rCutSmart™ Buffer, 5 µL of plasmid DNA, 0.5 µL of NotI-HF®, New England BioLabs, Frankfurt am Main, Germany). The digestion mix was subjected to standard agarose-gel electrophoresis to verify samples (2% agarose/TAE, 100 V, 30 min, TAE buffer: 40 mM Tris-HCl (pH 7.6), 20 mM acetic acid, 1 mM EDTA in ultrapure H₂O).

Verified samples were diluted to 100 ng/µL in 20 µL final sample volume (Buffer AE, Machery-Nagel, Dueren, Germany) and sent for sequencing by Eurofins Genomics (SupremeRun Barcodes, GATC Services, Ebersberg, Germany). Sequencing data were analyzed with Geneious Prime® (version 2021.1.1, Biomatters, Auckland, New Zealand).

2.6. Data Display and Statistical Analysis

Image data were processed with Fiji (ImageJ version 1.51n; Java version 1.8.0_6, 64-bit [31]) and displayed with PowerPoint® (Microsoft; München, Germany).

Numerical data were cured with Excel® (Microsoft; München, Germany), visualized with OriginPro® 2020 (64-bit, SR1 9.7.0.188, Academic, OriginLab Corp., Northampton, MA, USA), and displayed with PowerPoint®. Normality or skewness in data distribution (descriptive statistics) was determined by Shapiro–Wilks testing in OriginPro® 2020. Statistical analyses (hypothesis testing) were performed in OriginPro® 2020. Data sets with small sample size and normal distribution were subjected to two-sample t-testing. Data sets with small sample size and skewed distribution were subjected to Mann–Whitney testing. Further details are given in the respective results sections.
Sequencing data were analyzed and visualized with Geneious Prime® (version 2021.1.1, Biomatters, Auckland, New Zealand).

Statistics for the quantitative analysis of proliferation measured by Ki67 immunohistochemistry were performed using SPSS. For the non-normally distributed data set (Shapiro–Wilk), a Kruskal–Wallis test and post-hoc Bonferroni analysis of significance were performed. Values were considered as significantly different at the $P$-level as indicated.

3. Results

3.1. Cell Proliferation Assay Shows Dose-Dependent Susceptibility of iCCAOs to Multi-Kinase Inhibitor Sorafenib

A first screen with 133 compounds from an FDA-approved screening library showed intermediate to strong inhibitory effects of the multi-kinase inhibitor sorafenib on the established iCCAO line (results not shown). Additionally, previous studies also proposed a potentially beneficial effect of sorafenib in CCAs [6–9]. We therefore performed an exemplary MTS proliferation assay on the iCCAO line. We aimed to determine two concentrations of sorafenib for studying treatment effects at the morphological scale. This required efficacy without inducing high levels of apoptosis, which would impede long-term observations of morphological alterations under treatment conditions.

We determined 4 $\mu$M sorafenib to be the lowest effective concentration which significantly inhibits organoid growth over a treatment period of 72 h. We saw a drastic change in the morphological appearance of organoids (Figure 1A) and reduced mean proliferation by about 30% (Figure 1B). Accordingly, we decided to use 4 $\mu$M for brightfield microscopy experiments to further monitor and investigate changes in organoid growth and morphological behavior under treatment conditions. Although 2 $\mu$M sorafenib did not significantly decrease the MTS proliferation assay read-out, we still observed a reduction in mean proliferation of about 20% (Figure 1B). Therefore, we decided to use this concentration to investigate whether minor changes in morphological or growth behavior were elicited under prolonged treatment at half the lowest significantly effective concentration of 4 $\mu$M.

3.2. Organoid Growth Analysis Demonstrates Reduced Growth of iCCAOs upon Sorafenib Treatment

To study the effects of sorafenib treatment on organoid growth over time at a global morphological level by label-free, minimally invasive, and high-throughput means, we applied our recently published brightfield microscopy and corresponding image analysis pipeline [21]: here referred to as the organoid growth analysis (pipeline). We quantified organoid sizes in time-resolved brightfield data (48 h, recording interval: 0.5 h) of entire organoid cultures (on average 41–139 organoids per 96-well/5 $\mu$L Matrigel™) treated with 2 $\mu$M and 4 $\mu$M sorafenib.

By monitoring and quantifying the expansion of iCCAOs and ICOs under sorafenib treatment, we saw that growth in tumor organoids was generally more affected by sorafenib than growth in healthy organoids (Figure 2A,B). Compared to untreated organoids, relative sizes in tumor organoids after 48 h of treatment were significantly reduced to 71% at 2 $\mu$M sorafenib and 49% at 4 $\mu$M sorafenib. The vehicle control (DMSO 0.02%) showed a minor effect on tumor organoid growth with a size difference of 7%. When directly comparing the tumor organoid sizes at 2 $\mu$M and 4 $\mu$M sorafenib, we saw a dose-dependent size reduction of 31% (Figure 2B, Table 1).
Figure 1. iCCAOs show dose-dependent decreases in cell proliferation under sorafenib treatment. Organoids were grown from single cells for seven days before treatment with sorafenib for 72 h. The vehicle control (DMSO 0.02%) corresponds to the vehicle concentration in 4 µM sorafenib. Vehicle controls for other conditions were excluded from display. (A) Phase-contrast images of organoids after treatment. Microscope: AxioVert 40 CFL; objective lens: EC-Achro-Plan 10×/0.5; camera: AxioCam ICm 1; pixel size: 1.29 × 1.29 µm²; scale bar: 1000 µm. (B) Cell proliferation was quantified by MTS assay [proliferation \( \approx \) absorbance]. Proliferation values were normalized to the mean viability value of the untreated control. Statistics were performed on mean proliferation values \( n = 9, \) normal distribution, two-sample \( t \)-test.

ICOs showed a minor size reduction to 98% at 2 µM sorafenib but were highly affected by 4 µM sorafenib with a size reduction to 74% compared to untreated organoids. Surprisingly, the vehicle control (DMSO 0.02%) for ICOs showed a size increase of 15%. A direct comparison of organoid sizes at 2 µM and 4 µM sorafenib in ICOs further revealed a dose-dependent size reduction of 24% (Figure 2B).

Taken together, our results indicate that sorafenib inhibits growth more extensively in tumor organoids than in ICOs at low concentrations.
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**Figure 2.** Sorafenib treatment limits iCCAO growth more extensively than ICO growth. (A) Overview of pre-processed (z-projected, stitched) brightfield images for subsequent segmentation and post-processing in the organoid growth analysis pipeline. Organoids were grown for 72 h prior to treatment. Post-treatment, organoid growth was monitored over 48 h with a recording interval of 0.5 h (≈ 97 time points, counting 0 h as time point 1). To capture the vast majority of organoids within one well (one Matrigel™ droplet), 2 × 2 tiles and 10 z-planes (z-spacing: 65 µm) were recorded with a magnification of 5×. Figure 2 shows representative stitched average intensity z-projections of n = 4 recordings for each organoid line and each condition. Microscope: AxioObserver.Z1; objective lens: Plan-Apochromat 5×/0.16; camera: AxioCam MR R3; pixel size: 1.29 × 1.29 µm²; scale bars: 1000 µm. (B) Quantification of organoid growth under sorafenib treatment. Data were generated with the organoid growth analysis pipeline. Graphs show median lines of normalized projected area (NPA) values for 97 consecutively measured time points (equivalent to recorded/analyzed time points: 48 h, 0.5 h recording interval ≈ 97 time points, counting 0 h as time point 1). Projected areas (PAs) for each organoid were normalized to their individual mean PA value over the first five time points. An overview and a comparison of relative final organoid size in NPA and percentage values are given in Table 1. For details on data distribution refer to Figures S1 and S2. Statistics were performed on median NPA values [n = 97, skewed distribution, Mann–Whitney test, *** p < 0.001].
3.3. Determination of Proliferation and Apoptosis in iCCAOs by Immunohistochemistry

The determination of organoid growth inhibition by the organoid growth analysis pipeline as shown in Figure 2 was validated on the cellular level by immunohistochemical staining of the proliferation marker Ki67. As compared to untreated controls, the vehicle DMSO did not impact the number of Ki67-positive cells in the organoids, which was in the range of 80–90%. Treatment with 4 µM sorafenib for 48 h inhibited cell proliferation significantly by 50% (Figure 3A,B). This corroborates data determined by the organoid growth analysis pipeline, and moreover demonstrates that the growth of organoids is due to cell proliferation and not, e.g., to physical influences artificially “blowing up” the organoid size.

Table 1. Relative organoid sizes after 48 h of sorafenib treatment compared to untreated control. Percentages (right columns) were calculated from normalized projected area (NPA) values (left columns) obtained by organoid growth analysis (data shown in Figure 2). NPA values were rounded to two decimal places. Percentages were rounded to zero decimal places.

| Condition          | iCCAOs | ICOs |
|--------------------|--------|------|
|                    | Determined NPA Value | NPA Relative to Control in % | Determined NPA Value | NPA Relative to Control in % |
| Untreated          | 2.27   | 100  | 2.57 | 100 |
| DMSO 0.02%         | 2.10   | 93   | 2.95 | 115 |
| SOR 2 µM           | 1.60   | 71   | 2.53 | 98  |
| SOR 4 µM           | 1.11   | 49   | 1.91 | 74  |
| SOR 4 µM/SOR 2 µM  | —      | 69   | —    | 76  |

Figure 3. Sorafenib inhibits cell proliferation in iCCAOs in a concentration-dependent manner. (A) iCCAOs were treated for 48 h as indicated with the vehicle control (DMSO 0.02%), or with 2 and 4 µM sorafenib. Proliferating cells were identified by immunohistochemical detection of the nuclear antigen Ki67 (arrows). (B) For quantitative evaluation, positive cells in each 10 visual fields were detected using ImageJ and expressed as percentage number of total cells. Data were not normally distributed as verified by the Shapiro–Wilk test. The Kruskal–Wallis test and post-hoc Bonferroni analysis of significance were performed. Statistical analysis was accomplished using SPSS. Horizontal lines indicate the P-level of significance.

To show whether sorafenib inhibition of proliferation was associated with an increase in cell death, we stained iCCAOs for Caspase 3 activation. Yet, stained cells were found in the organoid tissue only occasionally. Stained cells were often visible in the organoid lumen, indicating extrusion of dying cells. Since these formed irregular conglomerates, quantification was not possible (Figure S3). Thus, whether cell death was due to normal turnover or due to sorafenib treatment cannot be decided unequivocally.
3.4. Expression of Functional Markers in iCCAOs after Sorafenib Treatment

The impact of sorafenib on functional features of iCCAOs was assessed by immunohistochemical detection of structural and functional markers after treatment with 4 µM sorafenib. The epithelial organization of the organoids was retained after treatment as indicated by the apical expression of the tight junction protein ZO-1 and the basolateral expression of the adherens junction protein E-cadherin (Figure 4A). This was supported by the expression of EpCAM, which was exclusively expressed in epithelial cells and played a role in tumorigenesis and metastasis [34], in both the control and sorafenib-treated organoids (Figure 4B). MDR1 is a bile export protein expressed in the liver at the apical membrane of hepatocytes and cholangiocytes [35]. This expression pattern was observed in the organoids after sorafenib treatment as demonstrated by the expression of MDR1 on the luminal side both in the controls and treated organoids (Figure 4C). The expression of SOX9, a transcription factor involved in proliferation, self-renewal, and tumorigenicity of cancer stem cells [36], was slightly, albeit not significantly, reduced by the treatment with sorafenib (Figures 4D and S4). Taken together, the results suggest that protein expression patterns were largely similar in organoids with and without treatment with sorafenib.

Figure 4. Sorafenib treatment does seemingly not affect protein expression of epithelial markers and proliferation marker SOX9 in iCCAOs. (A) iCCAOs were treated for 48 h as indicated either with the vehicle control (DMSO 0.02%), or with 4 µM sorafenib, resp. Fluorescent immunohistochemistry revealed apical and basolateral expression of ZO-1 (green) and E-cadherin (red), resp. Nuclei are labeled with DAPI (blue). Insets show digital magnifications (orig. magnification: 20×) of selected areas of the pictures behind. Arrows indicate luminal expression of ZO-1. (B) iCCAOs were treated for 48 h as indicated. Fluorescent immunohistochemistry showed membranous expression of EpCAM (green). Nuclei are labeled with DAPI (blue). (C) iCCAOs were treated for 48 h as indicated. Fluorescent immunohistochemistry revealed apical expression of MDR1 (green, white arrows). (D) iCCAOs were treated for 96 h as indicated. SOX9 expression was localized in the nuclei (insets, black arrows in digital magnifications (orig. magnification: 10×) of selected areas of the pictures behind) as shown by non-fluorescent immunohistochemistry.

3.5. RNA Sequencing Analysis Reveals Common KRAS Mutation in iCCAOs

To investigate whether the different sensitivities of ICOs and iCCAOs to sorafenib resulted from underlying mutational differences, we performed an RNA sequencing analysis of KRAS. This gene is commonly mutated in iCCA [37] and is involved in MAPK/ERK signaling, which is inhibited by sorafenib at the level of BRAF [38].

iCCAOs expressed both isoforms of KRAS (Iso 4A, Iso 4B) described in the literature [39] (Figure 5A,B). Both isoforms exhibited the commonly occurring G12D mutation (GGT > GAT), in which glycine was replaced by aspartic acid constituting the activation
of KRAS. This was also observed in the whole-exome sequencing of this organoid line performed earlier [10]. Additionally, we sequenced one transcript of Iso 4B (Iso 4B.I), which did not exhibit the G12D mutation but a rarely described silent D173D (GAT > GAC) mutation. These results indicate that both mutations are monoallelic.

ICOs also contained both isoforms of KRAS and expressed the wild-type form of Iso 4A (Figure 5). Only in one transcript of Iso 4B (Iso 4B.I), did we observe the aforementioned silent D173D mutation, indicating that this mutation is also monoallelic here.

Our results indicate that the mutational status of KRAS could partially explain why tumor organoids are more sensitive to sorafenib treatment than healthy organoids.

4. Discussion
4.1. Quantifying Organoid Growth in Patient-Derived Tumor Organoids by Label-Free Time-Lapse Brightfield Microscopy

We evaluated organoid growth under low-dose sorafenib treatment using our recently published brightfield microscopy-based image segmentation pipeline [21] (here referred to as: the organoid growth analysis pipeline). Our results show that this pipeline can be easily adapted to iCCAOs. This label-free imaging approach offers great potential for high-throughput screening of patient-derived organoid cultures [22,23,40,41]. However,
detailed data from other disciplines such as proteomics [42] and transcriptomics [43] are needed to ascribe the morphologic changes observed over time to specific underlying cellular processes. For example, we need to know whether shrinking of the organoid is a mere mechanical phenomenon caused by deflation of the luminal volume, or if cell death is the driving underlying mechanism. Yet, the increase in organoid size can at least in part be attributed to proliferation as shown by the proliferation marker Ki67 (cf. Figure 3).

Regarding the technical aspects of the pipeline, the pre- and post-processing operations of the data still require a lot of manual work and need to be further optimized [21]. This will help to reduce the manual workload, and further push the pipeline towards a fully automated high-throughput image analysis approach. One tangible idea to improve the pre-processing steps is to adapt image acquisition, background subtraction, filtering, and binarizing to produce a better signal-to-noise ratio for subsequent image segmentation. In this project, we segmented and analyzed coherent, sequential time-resolved image stacks of organoid cultures observed for up to 48 h (recording interval: 0.5 h). However, we noticed that single time points are segmented more accurately than coherent time series (the smaller the series, the better the segmentation). Hence, another strategy to improve the segmentation process is to segment each time point separately and fuse the segmented time points afterwards for a subsequent analysis.

It is debatable whether the level of detail on organoid size change produced by the organoid growth analysis pipeline is required for evaluating treatment effects in patient-derived organoid cultures. To increase throughput and decrease the acquired amount of data, the temporal resolution can be decreased. However, we argue that even minor fluctuations in organoid size or morphology, which can be monitored especially under low-dose treatment conditions, enable detailed conclusions on individual treatment efficiency.

4.2. Understanding the Observed Morphological Treatment Effects in Patient-Derived Tumor Organoids

By quantifying organoid sizes in the time-resolved data of entire organoid cultures under low-dose sorafenib treatment, we observed that sorafenib had a higher impact on tumor organoid growth (for this particular patient) compared to growth in healthy organoids. To investigate whether these findings might result from underlying mutations in the MAPK pathway, which is targeted by sorafenib, we sequenced the KRAS gene [44]. KRAS is frequently mutated in iCCA [45]. While ICOs expressed the wild-type form of KRAS, we identified a common G12D mutation associated with constitutive activation of KRAS [46] and poor prognosis [37] in iCCAOs. Our results suggest that iCCA patients with this specific mutational background in KRAS might benefit from systemic low-dose sorafenib treatment as previously reported [47,48].

However, here, we only looked in detail at one specific component (KRAS) in a very complex system. Broader analyses by multi-omics approaches (e.g., next generation sequencing, mass spectrometry) [49,50] and integrating clinical patient characteristics are required to draw further conclusions on predictive parameters and paint a bigger picture on how the mutational background relates to morphological appearance/behavior and susceptibility to treatment [51,52].

5. Conclusions

The organoid growth analysis pipeline is a promising technique to measure drug response over time and could become a high-throughput predictive tool upon further validation both in vitro for more organoid lines and drugs, and by comparison to patient responses in vivo. The integration of results from multi-omics studies [51,52] could lead to more insight into the underlying substantiating network of predictive parameters to maximize interpretation of the acquired imaging data. Once this network of predictive parameters is established, high-throughput imaging approaches such as our label-free brightfield imaging pipeline can be used to obtain vast information on treatment effects without the need for further cross-validation by other disciplines.
Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/cells11223613/s1, Figure S1: Data distribution of iCCAO growth upon sorafenib treatment; Figure S2: Data distribution of ICs growth upon sorafenib treatment; Figure S3: Immunohistochemical detection of cleaved Caspase 3 in iCCAOs with and without sorafenib treatment; Figure S4: Image analysis of the immunohistochemical detection of SOX9 in iCCAOs with and without sorafenib treatment; Table S1: KRAS sequencing analysis nucleotide alignment.

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