Abstract. Three-dimensional (3D) cultures are indispensable for capturing tumor heterogeneity in colorectal cancer (CRC) in vitro. Although 3D cultures (such as sphere-forming assay and organoid culture) can partially preserve the morphological and molecular characteristics of primary CRC, whether these 3D cultures maintain the long-term stemness of cancer stem cells (CSCs) remains largely unknown. In the present study, spheres and organoids were generated side by side using individual primary CRC specimens, then respectively processed as serial passages. The results revealed that during serial passages, the percentage of CSCs (such as cluster of differentiation-133+ and Wnt+ cells) in organoids and the tumor-initiating capacity of organoid-derived cells were constant, while they gradually increased in the sphere-derived cells. Furthermore, during serial passages, resistance to chemotherapeutic agents (including 5-fluorouracil and oxaliplatin) in sphere- and organoid-derived cells was evaluated. The results indicated that the percentage of chemoresistant cells was constant in serial organoid cultures; however, it gradually increased in the serial sphere-forming assays. Taken together, the results of the present study comprehensively demonstrate that, with regard to long-term culture in vitro, organoid culture may be useful in maintaining tumor heterogeneity and the levels of chemoresistant cells, while the sphere formation assay enriches for CSCs and chemoresistant cells.

Introduction

Colorectal cancer (CRC) is the third most common cause of cancer-associated mortality worldwide (1). A number of studies have revealed that CRC is heterogeneous, manifesting with variegated cellular morphologies, histopathology and chemoresistant capacity; these biological characteristics of CRC have contributed to the failure of conventional therapeutics (2-4). In order to improve the prognosis of patients with CRC, preclinical tumor models are essential for simulating primary tumors, exploiting novel therapeutic approaches and predicting drug responses in vivo (5). Even though preclinical models have been widely used, their advantages and limitations remain largely unknown.

It has been suggested that cancer stem cells (CSCs) may be responsible for tumor heterogeneity (6). In CRC, it has been revealed that the tumor bulk exhibits cellular hierarchy with CSCs at the apex, possessing self-renewal abilities, multi-differentiation potential and an inherent chemoresistant capacity (7). In addition, tumor recurrence in patients with CRC is closely associated with the presence of CSCs in tumors (8). These findings indicate that CSCs may be promising therapeutic targets. Therefore, the long-term maintenance of stem cell-like properties in preclinical models is critical for capturing the real primary tumor conditions.

Three dimensional (3D) cultures, such as sphere formation assays and organoid culture, can be used as platforms that support the long-term expansion of primary tumor cells (9). However, whether these 3D models can preserve the original properties of parental tumors remains unclear. Sphere formation assays, for instance, have been reported to expand CSCs during serial passages, and thus they are not a suitable platform for investigating drug activity (10). Organoid culture, on the other hand, has been exploited for predicting drug efficacy in vivo (11-14). However, it is still unclear whether the stem cell-like properties would be maintained long-term in organoid culture.
The present study generated sphere and organoid cultures side by side using individual CRC specimens and demonstrated that: i) The sphere formation assay was enriched for CSCs, while the organoid culture only maintained CSCs; and ii) the frequency of chemoresistant CRC cells in each of the generations during the serial organoid passages were almost same; however, the serial sphere formation assay increased the frequency of chemoresistant cells.

Materials and methods

Collection of CRC specimens and preparation of the single cell suspension. Surgical human colorectal adenocarcinoma samples were obtained with written informed consent and approval from the Institutional Review Board of Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology (Wuhan, China; IRB ID: 20141106); the experiments were conducted according to the principles of the Declaration of Helsinki. In total, 20 tumor specimens from CRC patients were included in the present study, and the patients were assigned case numbers CRC1-20. The patient clinical characteristics are listed in Table SI. The CRC specimens were disassembled into single primary CRC cells as described previously (15). Briefly, fresh specimens were minced into small sections with scissors. The completely minced pieces were then incubated in serum-free Dulbecco’s modified Eagle’s medium (DMEM/F12 (Thermo Fisher Scientific, Inc., Waltham, MA, USA) containing 1.5 mg/ml collagenase IV (Gibco; Thermo Fisher Scientific, Inc.), 20 µg/ml hyaluronidase (Sigma-Aldrich; Merck KGaA, Darmstadt, Germany) and 1% penicillin/streptomycin (Thermo Fisher Scientific, Inc.) at 37°C for 1 to 2 h. To eliminate red blood cells, the cells were incubated in red blood cell lysis buffer (eBioscience; Thermo Fisher Scientific, Inc.) on ice for 10 min, then washed twice with PBS. The single primary CRC cells were then resuspended in PBS for use in subsequent organoid culture, sphere forming assay and animal studies.

Organoid culture. Organoid culture was processed as previously described (15,16), with several modifications. For culture establishment, single primary CRC cells were embedded in Matrigel (growth factor reduced; phenol free; BD Biosciences, Franklin Lakes, NJ, USA) and seeded into 24-well culture plates at the indicated dosage (1,000 cells/30 µl Matrigel/well). Following Matrigel polymerization, the cells were overlaid with human colorectal cancer organoid culture medium and incubated at 37°C in humidified air containing 5% CO₂. The composition of the human colorectal cancer organoid culture medium was as follows: DMEM/F12 (Invitrogen; Thermo Fisher Scientific, Inc.) supplemented with 1X B27 (Invitrogen; Thermo Fisher Scientific, Inc.), 20 ng/ml recombinant human EGF (Sigma-Aldrich; Merck KGaA), 10 nM Gastrin (Sigma-Aldrich; Merck KGaA) and 500 nM A83-01 (Tocris Bioscience, Bristol, UK) (16). The medium was changed every 3 days. To calculate the forming efficiency, following 7 days post-embedding, organoids with diameters >100 µm were scored under an inverted microscope (Leica Microsystems, Wetzlar, Germany). The forming efficiency (%) = scored organoid number / total embedding CRC cells. For serial passages, following 7 days post-embedding, whole organoids were trypsinized using 0.025% trypsin/EDTA at 37°C for 15-20 min. Single organoid-derived cells were then resuspended with PBS and employed for a new round of organoid culture in Matrigel.

Sphere formation assay. The sphere formation assay was conducted as previously described (17,18). Single primary CRC cells were resuspended in standard stem cell medium (SCM), which consisted of the following: DMEM/F12 (Invitrogen; Thermo Fisher Scientific, Inc.) supplemented with 1X B27 (Invitrogen; Thermo Fisher Scientific, Inc.), 20 ng/ml human recombinant epidermal growth factor (Sigma-Aldrich; Merck KGaA) and 20 ng/ml basic fibroblast growth factor (Sigma-Aldrich; Merck KGaA) (17). The spheres were cultured at 37°C in humidified air containing 5% CO₂. To calculate the forming efficiency, following 7 days post-plating, spheres with diameters >50 µm were scored under an inverted microscope (Leica Microsystems). The forming efficiency (%) = scored sphere number / total plating CRC cells. For serial passage experiments, at 7 days post-plating, spheres were trypsinized as aforementioned. Single sphere-derived cells were then replated in SCM for a new round of sphere forming assay.

Animal experiments. Female, 4- to 6-week-old NOD/SCID mice (n=135) were purchased from Beijing HFK Bioscience Co., Ltd. (Beijing, China) and were maintained according to protocols approved by the Institutional Animal Care and Use Committee, Huazhong University of Science and Technology (Hubei, China; IACUC ID: 2014S652). During experimentation, all mice were kept in the animal center of the Tongji Medical college, Huazhong University of Science and Technology under specific pathogen-free (SPF) conditions (License Number: SYXK#2016-0057). For maintenance of mice, a stable breeding environment was provided: a 12-h/12-h light/dark, a temperature of 21°C and a relative humidity 50% and allowed access to water and food ad libitum. For the establishment of patient-derived xenografts (termed PDXs), n=40 mice were used, and the detailed information of PDX generation is presented in Table SI. For transplantation assays (n=5 mice were used), the single cells derived from CRC6 specimens and the corresponding organoids and spheres generating form them were resuspended in a PBS/Matrigel (BD Biosciences) mixture (1:1 volume) (15,17). The cells were then injected into the subcutaneous tissue of mouse flanks using 27-gauge needles at at dose of 125,000 cells/point. For histopathological analyses, the xenografts derived from primary cells (termed PDX, 1 mouse), organoid cells (termed ODX, 2 mice) and sphere cells (termed SDX, 2 mice) were harvested on day 30 post-implantation. The mice were anesthetized and sacrificed according to the AVMA guidelines, and hematoxylin and eosin (H&E) staining was performed as previously described (15). For limited dilution assays (n=90 mice were used), 10,000, 1,000 and 100 single cells derived from organoids at 1st - 3rd generation, or 1,000, 100 and 10 single cells derived from spheres at 1st - 3rd generation were implanted per injection (5 mice for each dose at each group). For the limited dilution assays, the tumor volumes were examined every 3 days, and the mice with excessive weight loss (humane endpoint) post-injection were excluded.
from the study and euthanized (n=7), and these included: 1 mouse injected with organoid cells at 1st generation; 2 mice injected with organoid cells at 2nd generation; 4 mice injected with organoid cells at 3rd generation. Following sacrifice, tumors were removed from the mice and weighed to evaluate tumor development. The frequency of tumor-initiating cells and statistical significance were examined using the Extreme Limiting Dilution Analysis software (bioinf.wehi.edu.au/software/elda/index.html).

**Immunofluorescence.** The immunofluorescence procedures for organoids and spheres were performed as previously described (19). Prior to staining, organoids or spheres were fixed in 4% PFA at 4°C for 20 min. For immunofluorescence staining, the following antibodies were used to detect antigens: Mouse-anti-human cluster of differentiation (CD)-133 (1:100; cat. no. 66666-1-lg; ProteinTech Group, Inc., Chicago, IL, USA) and rabbit-anti-human cytokeratin (CK)-20 (1:100; cat. no. 13063; Cell Signaling Technology, Inc., Danvers, MA, USA). Briefly, the sections were blocked with 5% (w/v) bovine serum albumin (Invitrogen; Thermo Fisher Scientific, Inc.) in PBS, then incubated with the primary antibodies at 4°C overnight, followed by staining with secondary antibodies conjugated to Streptavidin-Cy3 (1:100; cat. no. SA1010; Thermo Fisher Scientific, Inc.) or Alexa Flour 488 (1:100; cat. no. 705-546-147; Jackson ImmunoResearch Laboratories, Inc., West Grove, PA, USA) at room temperature for 2 h. Finally, nuclei were stained with 4,6-diamidino-2-phenylindole (DAPI, Sigma-Aldrich Merck KGaA) at room temperature for 10 min. Images of organoids and spheres were captured under a fluorescence microscope (TRRFM; Olympus Corp., Tokyo, Japan) or a confocal microscope (FV1000; Olympus Corp.).

**Reverse transcription-quantitative polymerase chain reaction (RT-qPCR).** Total RNA was extracted using RNAiso Plus (Takara Bio, Inc., Otsu, Japan), and then cDNA was synthesized using the Mixima First Strand cDNA Synthesis kit (Thermo Fisher Scientific, Inc.). RT-qPCR analysis was conducted using an ABI PRISM 7300 Sequence Detection System instrument with Maxima SYBR-Green/ROX qPCR Master Mix (Thermo Fisher Scientific, Inc.) according to the manufacturer's instructions. The thermocycling conditions for RT-qPCR were: step1: 50°C for 2 min; step 2: 95°C for 2 min; step 3 (x40 cycles): 95°C for 15 sec and 60°C for 1 min. The Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as an internal control, and the gene expression was measured using the 2ΔΔCq method (20). The sequences of the primers were as follows: GAPDH forward, 5'-TCGTGGAAGAGCTCATGACC-3' and reverse, 5'-TCCACCACTGTTGCTGTA-3'; CD44 forward, 5'-AGCAACCAAGAGGCAAGGCAAAGAAA-3' and reverse, 5'-GTTGTTGGTGAATGGTGCTG-3'; CD133 forward, 5'-TCTTTGACCAGACTGAGCCAAAA-3' and reverse, 5'-CTATGTTTCTCAACCGCCTCTT-3'; ABCG2 forward, 5'-TCTATATGTTGGAATGGTGCTG-3' and reverse, 5'-TTTCAGGCGGTGAAACTCTTT-3'.

**Lentiviral reporter assays.** The TCF/LEF reporter, which drives the expression of GFP (TOP-GFP) lentivirus, was purchased from SBO Medical Biotechnology Co. (Shanghai, China) (18). Primary cancer cells were infected as previously described (21). Primary cells were infected with TOP-GFP lentivirus at an MOI of 25 for 72 h. To separate the GFP+/GFP- cells, the top (GFP+) and bottom (GFP-) 5-10% cells were purified out by flow cytometry. To evaluate Wnt activity, the intensity and frequency of GFP+ cells were detected using a fluorescence microscope (TRRFM; Olympus Corp.) or flow cytometry according to the manufacturer's instructions of a FACS Aria II Cell Sorter (BD Biosciences) (18).

**Flow cytometric analysis and purification of CD133+/CD133- cells.** Fluorescence-activated cell sorting (FACS) was performed according to the manufacturer's instructions using a FACS Aria II Cell Sorter (BD Biosciences) (18). To measure CD133 expression in the organoids and spheres, the cells were stained with phycoerythrin (PE)-conjugated mouse anti-human CD133 (1:1; cat. no. 130-090-756; Miltenyi Biotec GmbH, Bergisch Gladbach, Germany), then analyzed using a BD FACS Aria II flow cytometer (18). To separate CD133+/CD133- cells in primary CRC, tumor specimens were processed into single cells as described above (15). The cells were then stained with PE-conjugated mouse anti-human CD133 at 4°C for 15 min. For purification, only the top (CD133+) and bottom (CD133-/-) 10-20% cells were purified out (18).

**Cell death analysis.** The cell death of organoid- and sphere-derived cells was assessed using the Cell Counting kit-8 (CCK-8; Dojindo Molecular Technologies, Inc., Rockville, MD, USA), as well as with 5-fluorouracil (5-Fu) and oxaliplatin (OXA) treatment, which were purchased from Sigma-Aldrich (Merck KGaA). Briefly, the cells were seeded in complete medium at 30,000 cells/well in 96-well plates. Following 12 h, the cells were treated with either 5-Fu (1 µM) or OXA (1 µM). After 72 h, 10 µl CCK-8 solution was added to each well. The plates were then incubated at 37°C for 1 h, and cell viability was determined by scanning with a microplate reader (Thermo Fisher Scientific, Inc.) at 450 nm.

**Statistical analysis.** Statistical significance was calculated with SPSS Statistics 18.0 software (IBM Corp., Armonk, NY, USA). All measurement data are presented as the means ± standard deviation of at least 3 independent experiments. The measurement data were analyzed using the Student's t-test or one-way analysis of variance followed by a Tukey's post hoc test to determine the significant differences of means in two or multiple groups (n>2) comparisons. The enumeration data were analyzed using the Fisher's exact test to determine the significant differences of rates in two categories comparisons. P<0.05 was considered to indicate a statistically significant difference.

**Results**

Organoids effectively simulate the tumor heterogeneity of CRC tumors in vitro. Sphere formation assay (22) and organoid culture (23) are 3D models that support the long-term expansion of primary tumor cells in vitro. In this study, to investigate whether these 3D models accurately simulate the tumor heterogeneity of CRC tumors, we generated spheres and organoids side by side using freshly surgical CRC specimens. H&E staining revealed
that the organoids harbored parental tumor-like budding structures, while the spheres were observed as solid globes (Fig. 1A). Notably, the organoids formed more complex structures in the cells derived from high-grade tumors (such as Dukes’ C and D), whereas the spheres retained similar global shapes (Fig. 1B). Since cells expressing CD133 have been reported to enrich for CSCs in CRCs (24) and cytokeratin 20 (CK20) is a differentiated cell marker (25), the present study conducted further CD133 and CK20 staining for spheres and organoids derived from primary CRCs. The results revealed that the expression of CD133 and CK20 more closely resembled that of primary tumors in the organoids than in the spheres (Fig. 1C). Quantitative analysis demonstrated that the organoids harbored similar percentages of CD133+ cells when compared with the corresponding primary tumors; however, sphere-forming assays significantly increased the percentage of CD133+ cells (Fig. 1D), implying that organoid culture may maintain the cellular heterogeneity of parental CRCs. These results demonstrate that, compared with the sphere-forming assay, organoid culture may more effectively simulated the tumor heterogeneity of primary CRCs in vitro.

Organoid culture more effectively simulates primary colorectal tumors than sphere-forming assays. In general, tumor cells are able to generate xenografts in immuno-compromised mice subcutaneously, and the PDXs can recapitulate original tumor heterogeneity (5). In order to explore the tumor heterogeneity of CRC tumors, it is also crucial to establish successful cell culture in vitro (5,26,27). However, whether cells cultured in 3D models preserve the ability to generate parental tumor-like xenografts (i.e., PDXs) remains unclear. Consistent with the findings of previous studies (5,7,25), the results of the present study demonstrated that primary CRC cells and their corresponding organoids and spheres were all capable of generating tumor xenografts in NOD/SCID mice (Fig. 2A). To determine whether ODXs and SDXs exhibit the same tumor heterogeneity of primary tumors, the present study performed CK20 (25) staining for primary CRC tumors, PDX, ODX and SDX. As shown in Fig. 2B, the present study revealed that the expression pattern of CK20 in ODX more closely resembled primary tumors and the corresponding organoids and spheres than SDX (Fig. 2B), suggesting that organoid culture more accurately reproduced the tumor heterogeneity of primary tumors than the sphere formation assay. In order to examine the efficiency of generating organoids or spheres from primary CRC tumors, the present study performed side-by-side organoid culture and sphere-forming assays for CRC specimens (Table S1).
results revealed that organoids in 15 of the 20 CRC specimens were successfully generated (success rate, 75%), whereas spheres were only generated for 5 of the 16 CRC specimens (success rate 31%; Tables I and S1). Notably, the primary CRC cells formed more organoids than spheres when the same cell dosage was applied (Fig. 2C and D). Taken together, these results demonstrate that organoid culture possesses a higher success rate and better efficiency to simulate primary colorectal tumors than sphere-forming assay.

Organoid culture maintains the number of CSCs during serial passages. The cell surface marker, CD133, is widely used to enrich CSCs in CRCs (16,23). In the present experimental system, using FACS, CRC cells expressing high levels of CD133 (CD133+) and those expressing little or no CD133 (CD133-) were assessed and sorted. CD133+ CRC cells were found to produce more spheres and organoids than CD133- cells, implying that sphere- and organoid-forming cells were

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Tumor specimens were obtained from colorectal cancer patients and processed into single cell suspensions. Cells were then seeded in 3D culture conditions to generate spheres, organoids in vitro and implanted into NOD/SCID mice to generate PDXs in vivo. P-values are listed comparing two categories using Fisher's exact test.
enriched for CSCs (Fig. 3A). Serial sphere formation assays are generally applied to enrich and expand CSCs (27). However, the dynamics of CSCs in serial organoid cultures remain unclear. The results of the present study revealed that the levels of CD133 and CD44, the two cell surface markers widely used to enrich CSCs in CRCs (24,29), remained constant in serial organoid cultures, while they significantly and gradually increased in serial sphere formation assays, indicating that the dynamics of CSCs may differ between organoid culture and sphere-forming assays during serial passages (Fig. 3B). To visualize CSCs and differentiated cells within individual organoids and spheres during serial passages, the present study performed CD133 and CK20 immunofluorescence staining for organoids and spheres (Fig. 3C and D). Quantification analysis revealed that the proportion of CD133+ cells remained relatively constant in organoids during serial passages (Fig. 3E). Flow cytometric analysis of CK20+ cells in organoids and spheres during serial passages (Fig. 3F) showed that the proportion of CK20+ cells remained relatively constant in organoids during serial
passages; however, there were increased levels of CD133$^+$ cells and decreased levels of CK20$^+$ cells in spheres during serial passages (Fig. 3E and F). Taken together, these results demonstrate that organoid culture produces steady ratios of CSC subsets in long-term cultures, whereas sphere-forming assays may enrich CSCs.

Organoid culture retains the frequency of tumor-initiating cells (TICs), while sphere-forming assays enrich TICs during serial transplantations. TICs, also known as CSCs, are a rare population within CRC (30). In this study, to investigate whether the frequency of TICs was altered during serial transplantations, we performed limited dilution assays to evaluate the

Figure 4. Organoid culture retains the frequency of tumor-initiating cells while sphere-forming assays enrich tumor-initiating cells during serial transplantations. (A and B) Limited dilution assays for serial organoid- and sphere-derived cells. The frequency of tumor-initiating cells and statistical significance were examined using Extreme Limiting Dilution Analysis software with Lindill function of the Stamod package. (http://bioinf.wehi.edu.au/software/elda/index.html). (C) The cells were isolated from organoids during serial cultures, and implanted subcutaneously in NOD/SCID mice. Tumor volumes were measured in the mice with 1,000 cell injections starting from 21 days post-implantation. Data are presented as the means ± standard deviation. (D) The cells were isolated from spheres during serial passages, and implanted subcutaneously in NOD/SCID mice. Tumor volumes were measured in the mice with 1,000 cell injections starting from 12 days post-implantation. Data are presented as the mean ± standard deviation. ***P<0.001. The cells used in (A-D) were derived from CRC6. CRC, colorectal cancer; NS, not significant.
tumorigenic capacity of organoid- and sphere-derived cells in each generation (15,30). As shown in Fig. 4A, during serial transplantations (1st to 3rd), organoid-derived cells possessed a constant number of TICs. On the other hand, serial sphere-forming assays had significantly increased numbers of TICs with serial passaging (Fig. 4B). Notably, it was observed that organoid-derived cells initiated tumor generation at almost the same time in each generation, while sphere-derived cells initiated tumor generation earlier than the former generation (Fig. 4A and B). Consistent with this finding, there were no significant differences among the tumor volumes of organoid-derived xenografts in serial transplantations (Fig. 4C), while sphere-derived cells developed increasingly larger tumors in serial transplantations (Fig. 4D). Taken together, these results indicate that organoid culture maintains the frequency of TICs (i.e., CSCs) in long-term culture, while sphere-forming assays enrich TICs.

Serial organoid cultures preserve the proportion of Wnt+ CRC cells. It has been demonstrated that a high Wnt activity functionally designates the colon CSC population (18,31-33). To further evaluate the alternations in Wnt activity during serial passages, in the present study, we infected purified primary CRC cells with lentivirus, in which the TCF/LEF reporter drove the expression of GFP (TOP-GFP) and represented the cellular levels of Wnt activity (18,31,32). The transfected cells were then utilized in organoid culture and sphere-forming assay; when the organoids and spheres were growing, the expression of GFP was evaluated. The results revealed that the intensity of GFP expression was stronger in the spheres than the organoids (Fig. 5A), indicating that spheres were more highly enriched for CSCs than organoids during culture. The present results also demonstrated that during serial passage, the number of cells expressing GFP (i.e., GFP+ or Wnt+ cells) gradually increased in serial sphere-forming assays, but remained relatively constant in serial organoid cultures (Fig. 5B and C). Consistent with these findings, FACS analysis further demonstrated that the intensity of Wnt+ cells remained unaltered in organoids during serial passaging, while they markedly increased in spheres with serial passaging (Fig. 5D). Collectively, these data indicated that organoid cultures preserved relatively constant numbers of Wnt+ cells and the level of Wnt signaling in the cells during long-term culture, whereas the sphere-forming assay enriched Wnt+ cells and increased the intensity of Wnt signaling in the cells.

Organoid culture preserves chemoresistant cells over time. As the presence of CSCs and the activation of Wnt signaling have been demonstrated to be associated with the chemoresistance of tumor cells (31-33), we hypothesized that serial
sphere-forming assays, which increased the proportion of CSCs and improved the levels of Wnt signaling during serial passages, may also increase the frequency of chemoresistant cells; whereas, organoid culture may only maintain the chemoresistant capacity in the cells during serial passages. To examine this hypothesis, the present study performed

Figure 6. Organoid culture preserves chemoresistant cells over time. Representative images of (A) organoids and (B) spheres treated with chemotherapeutic agents (1 µM OXA or 1 µM 5-Fu) during serial passages. Scale bars, 200 µm. (C) Organoid-forming efficiency in the serial organoid cultures and (D) sphere-forming efficiency in the serial sphere formation assays upon the administration of chemotherapeutic agents (1 µM OXA or 1 µM 5-Fu). **P<0.01 and ***P<0.001. Flow cytometric quantification analysis of Wnt cells in the serial (E) organoid cultures and (F) sphere formation assays upon administration of chemotherapeutic agents (1 µM OXA or 1 µM 5-Fu). **P<0.01 and ***P<0.001. Quantification of the IC_{50} of primary cells, organoid and spheres of CRC upon (I) OXA (1 µM) and (J) 5-Fu (1 µM) treatment. The primary CRC cells used in (A-J) were derived from CRC6. (K) The cell viability of CRC1, CRC5, and CRC9 derived organoids or spheres treated with (K) OXA (1 µM) and (L) 5-Fu (1 µM) during serial passages. *P<0.05, **P<0.01 and ***P<0.001. IC_{50}, 50% inhibitory concentration. CRC, colorectal cancer; OXA, oxaliplatin; 5-Fu, 5-fluorouracil; ABCG2, adenosine triphosphate binding cassette subfamily G member 2; NS, not significant.

sphere-forming assays, which increased the proportion of CSCs and improved the levels of Wnt signaling during serial passages, may also increase the frequency of chemoresistant cells; whereas, organoid culture may only maintain the chemoresistant capacity in the cells during serial passages. To examine this hypothesis, the present study performed
serial organoid cultures or sphere-forming assays combined with conventional chemotherapeutic agents (oxaliplatin or 5-fluorouracil), and then detected the efficiency of organoid or sphere formation. As shown in Fig. 6A-D, the results indicated that, upon the administration of chemotherapeutic agents, the efficiency of organoid and sphere formation decreased; however, along with serial passages, this impairment was partially rescued in serial sphere-forming assays, but remained unchanged in organoid culture. Furthermore, in each generation, the present study detected the frequency of Wnt+ cells and the expression levels of adenosine triphosphate binding cassette subfamily G member 2 (ABCG2), one of the resistant markers of CRC (34), in organoids and spheres treated with chemotherapeutic agents. The results verified that, the frequency of Wnt+ cells, as detected by FACs, and the expression level of ABCG2, as determined by RT-qPCR, significantly increased in spheres and organoids following treatments with chemotherapeutic agents; however, during serial passaging, these markers remained unaltered in the organoid culture, while they gradually increased with serial sphere formation (Fig. 6E-H). The half-maximal inhibitory concentration (IC50) analysis upon OXA and 5-Fu treatment suggested that, since the 1st generation, organoid cells exhibited a similar IC50 to that of primary cells, while the sphere forming assay significantly increased the original IC50 (Fig. 6I and J). These data, obtained from CRC6, indicated that organoid culture preserved the characteristics of chemoresistance in primary CRC cells. To verify the universality, a chemoresistance assay in serial passages was respectively performed in CRC1, CRC5 and CRC9. The subsequent CCK8 assay further demonstrated that the frequency of chemoresistant cells gradually increased in serial sphere-forming assays, while it remained relatively constant in serial organoid cultures (Fig. 6K and L). Overall, the results indicated that, when compared with sphere-forming assays, organoid culture can preserve the frequency of chemoresistant tumors cells in the long-term culture, and may serve as a better pre-clinical model for drug screening and discovery.

Discussion

Preclinical models are essential for assessing tumor progression and designing therapeutic approaches (5). For decades, using numerous in vitro and in vivo preclinical models, the identification of oncogenes and tumor suppressor genes, aberrant signaling pathways and interactions in tumor microenvironment, have yielded several novel therapeutic strategies, including monoclonal antibodies, chemotherapy, small molecule inhibitors and immune therapy (5). However, previous studies have indicated that the inability of preclinical models to simulate tumor heterogeneity contributed to the failure of treatments (5,26,27). To avoid senseless trials and achieve better efficacy, the advantages and/or limitations of various preclinical models must be elucidated, and a relatively robust platform for drug testing must be determined.

To date, the widely used preclinical models have comprised of PDX, 2D monolayer cells culture (i.e., adherent cell culture and clonal culture) and 3D culture models (i.e., sphere formation assay and organoid culture) (5). For the PDX model, it has been demonstrated that the histopathology and genome mutational landscapes of patients can be highly preserved in the established individual PDXs (35), and the drug design procedures based on PDXs can partially predict patient response (36). Nonetheless, several limitations of PDX models impair its utilization, including the following: i) Limited success rate of initiation; ii) the lack of tumor-host interactions and immunity in xenograft animals; iii) it is highly labor intensive and time consuming; and iv) there are many ethical issues involving the use of animals (5,37). For 2D monolayer cells culture, there are several practical advantages: i) Immortalized cancer cell lines are easy to propagate, and can easily process transfection and genomic modification; and ii) the investigation concerning oncogenic pathways relies heavily on cell lines based on their stable manipulation (5). However, 2D monolayer cell culture possesses very low successful initiation rates for primary cells (<10% in CRC), and most notably, 2D culture fails to represent the tumor heterogeneity of primary cancers, which finally leads to the loss of the original clinical characteristics of tumor cells during long term passaging (26,27).

To overcome the limitations of PDX and 2D cell culture, in recent years, numerous novel 3D culture models have been applied. Among the 3D culture models, the two most popular ones are, the sphere formation assay and organoid culture (5,26,27). 3D culture models are considered to more accurately simulate the tumor heterogeneity of primary tumors when compared with 2D cell culture, and be less time-consuming and expensive than PDX (5). In general, sphere formation assays can be applied to enrich cancer cells with limited or unlimited self-renewal capacity for primary tumor cells and cell lines (38); on the other hand, organoid culture has been utilized in a range of cancer studies including drug testing, disease modeling, and co-culture models of cancer and stromal cells (11,14,39-41). Although the two 3D culture models can support the continuous expansion of primary tumor cells, whether they maintain the self-renewal capacity of cells in long-term culture, which is critical to simulate the tumor heterogeneity of primary tumors, remains largely uncertain.

It has been revealed that the tumor bulk harbored cellular hierarchy with CSCs at the apex, possessing self-renewal capacity and multi-differentiated potential (42). CSCs were considered to be closely associated with tumor heterogeneity in cancers, particularly in CRCs (43). Based on the expression of cell surface markers (such as CD133 and CD44) and the activity of Wnt signaling in CRC cells, it has been demonstrated that CSCs are enriched in the cells that highly express such markers or possess high activity of Wnt signaling (43). Furthermore, transplantation assays are applied to test the self-renewal capacity of CSCs in vivo (42,44). In the present study, the proportion of CSCs (including CD133+, CD44+ and Wnt+ cells) was enriched in serial sphere formation assays, but maintained constant in serial organoid culturing. Notably, following transplantation assays, the results demonstrated that serial sphere forming assays expanded CSCs (including CSCs and TICs), while serial organoid passage maintained the original levels. In addition, the results further demonstrated that cellular heterogeneity in organoids more closely resembled that of the primary tumor and the corresponding PDX than in spheres.

Sphere formation assay and organoid culture are both candidates of preclinical drug testing models (9). In previous
studies, a sphere-forming assay has been used to evaluate CSCs, the putative inherent chemoresistant cells (38,45). However, several studies have suggested that, during the sphere formation assay, as the CSCs become enriched, the drug activity of cancer cells became unstable (10,28). On the other hand, organoid culture, with high efficiency to establish patient-derived organoids (PDOs), enabled patient-specific drug testing and the development of personalized treatments (11-14). Although PDOs have been applied in drug testing in various types of solid tumors, whether organoid culture maintains the chemoresistant capacity of primary tumor cells in the long-term culture remains largely unknown. The present results demonstrated that the frequency of chemoresistant cells was constant in serial organoid cultures, whereas it gradually increased in the serial sphere-forming assays.

In conclusion, taken together with previous findings, the results of the present study demonstrated that organoid culture is the better preclinical model for reproducing tumor heterogeneity and testing drug responses, especially in long-term cultures.

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Availability of data and materials

All data generated or analyzed during this study are included in the article.

Authors’ contributions

HZ, CY and JQ contributed to the conception and design of the study. HZ performed the majority of the experiments. JQ and HZ wrote the manuscript. JQ revised the manuscript. YH, LM, XL and DT analyzed the experimental data. HZ, KH and QL performed the animal studies. All authors have read and approved the final manuscript.

Ethics approval and consent to participate

Surgical human colorectal adenocarcinoma samples were obtained with written informed consent under guidelines approved by Institutional Review Board of Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology, (Hubei China; IRB ID: 20141106), conducted according to the principles of the Declaration of Helsinki. The mice used in the present study were maintained according to guidelines approved by Institutional Animal Care and Use Committee, Huazhong University of Science and Technology (Hubei China; IACUC ID:20145652), and were anesthetized and sacrificed according to AVMA guidelines.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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