Research paper

Profiling of conditionally reprogrammed cell lines for in vitro chemotherapy response prediction of pancreatic cancer

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ABSTRACT

Background: The establishment of patient-derived models for pancreatic ductal adenocarcinoma (PDAC) using conventional methods has been fraught with low success rate, mainly because of the small number of tumour cells and dense fibrotic stroma. Here, we sought to establish patient-derived model of PDAC and perform genetic analysis with responses to anticancer drug by using the conditionally reprogrammed cell (CRC) methodology.

Methods: We performed in vitro and in vivo tumourigenicity assays and analysed histological characteristics by immunostaining. We investigated genetic profiles including mutation patterns and copy number variations using targeted deep sequencing and copy-number analyses. We assessed the responses of cultured CRCs to the available clinical anticancer drugs based on patient responsiveness.

Findings: We established a total of 28 CRCs from patients. Of the 28 samples, 27 showed KRAS mutations in codon 12/13 or codon 61. We found that somatic mutations were shared in the primary-CRC pairs and shared mutations included key oncogenic mutations such as KRAS (9 pairs), TP53 (8 pairs), and SMAD4 (3 pairs). Overall, CRCs preserved the genetic characteristics of primary tumours with high concordance, with additional confirmation of low-AF NPM1 mutation in CRC (35 shared mutations out of 36 total, concordance rate=97.2%). CRCs of the responder group were more sensitive to anticancer agents than those of the non-responder group ($P < 0.001$).

Interpretation: These results show that a pancreatic cancer cell line model can be efficiently established using the CRC methodology, to better support a personalized therapeutic approach for pancreatic cancer patients.

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1. Introduction

Pancreatic ductal adenocarcinoma (PDAC) has the worst prognosis among all cancers, with an overall 5-year survival of approximately 10%, and it is expected to become the second leading cause of cancer deaths by 2030 [1]. Even in patients who undergo complete resection, long-term survival is poor owing to tumour recurrence [2].

Studies on many types of cancers have shown that next-generation sequencing (NGS)-based genomic analysis sheds light on profiling of genetic aberrations, identification of novel therapeutic targets and clinical practices for personalized treatment that remarkably increase survival [3–6]. Unlike other cancers, however, these applications have been largely limited in PDAC because of the difficulty in obtaining sufficient amount of high quality specimens, owing to the
Research in context

Evidence before this study

Pancreatic ductal adenocarcinoma is one of the most aggressive malignancies. Next-generation sequencing-based genomic analyses and genetic aberration profiling have been employed for many tumour types to identify novel therapeutic targets. However, in most cases, only small amounts of biopsy specimens are available for analysis. To address these problems, various conventional preclinical cancer models are being developed. This strategy has been largely limited in pancreatic ductal adenocarcinoma because of the difficulty in obtaining sufficient amounts of high-quality specimens, owing to the low eligibility of the patients for surgery due to rapid disease progression, and the predominance of stromal cells in the tumour.

Added value of this study

We rapidly established patient-derived pancreatic cancer cell lines from tumour samples and then performed genetic analyses using these cells. We found that somatic mutations were maintained in cell lines and shared mutations included key oncogenic mutations such as KRAS, TP53, and SMAD4. Overall, patient-derived cell lines preserved the genetic characteristics of the primary tumours with high concordance. Cells can be isolated from patients at all stages of pancreatic cancer using the limited tissue available from a fine needle tumor biopsy as the starting tissue. Furthermore, ex-vivo examination of patient-derived cell lines revealed drug responsiveness predictions, which matched the clinical results.

Implications of all the available evidence

Our findings suggest that patient-specific preclinical cancer cell lines can be efficiently established and maintain original tumour characteristics. These matched patient-derived cells might support a personalized therapeutic approach for pancreatic adenocarcinoma.

low eligibility of patients for surgery because of the rapid disease progression, and the predominance of stromal cells in the tumour [7]. Hence, in most cases, only small amounts of biopsy specimens are available for analysis. To address these problems, various preclinical cancer models have been developed including cancer cell line models and patient-derived xenografts [8,9]. However, reduced stability, high cost, and lengthy preparation time hindered their active use in PDAC, emphasizing the immediate requirement of a more efficient model.

Recently, a new technique that establishes patient-derived cell lines without the use of extrinsic genetic immortalization, named “conditional reprogramming” (CR) enabled rapid and stable cell culture [10]. The conditionally reprogrammed cells (CRCs)–patient-derived tumour cells generated by CR—could be grown indefinitely under defined conditions and the karyotype was maintained even after numerous passages [11–15]. Moreover, the entire procedure can be started with a very small amount of primary tissue (~5 mm), such as biopsy specimens. Considering these benefits, we assumed that CR could be used as a beneficial platform for applying genomic analysis and predicting patient response in PDAC.

Here, using the CRC methodology, we aimed to develop PDAC cell lines that phenotypically represent primary human PDAC specimens and to identify patient-derived cell line models that can predict patient therapeutic responses in pancreatic cancer.

2. Methods

2.1. Patients and tissue samples

Patients diagnosed with PDAC were enrolled for establishment of the tumour model and genetic analysis. Tumour specimens (~1 cm) were obtained from the resected tissues of patients who underwent surgery for PDAC. For patients with unresectable PDAC, endoscopic ultrasonic (EUS)-guided biopsy or percutaneous biopsy was performed to obtain the tumour specimens (Fig. 1). Tumour tissues and paired peripheral blood samples were collected simultaneously. All tissues were placed in a medium containing antibiotics. Using forceps and a scalpel, the residual fat tissue was removed. Tumour tissues were minced into 1–2 mm small fragments with sterile scissors. Deselected specimens were placed in the medium. Primary cells were isolated within 1–2 h of tumour resection. If the specimens could not be processed immediately to prepare CRCs, the tumour tissues were frozen in liquid nitrogen for long-term storage.

2.2. Ethics

This study was approved by the Institutional Review Board of Severance Hospital, Seoul, South Korea (No. 4-2015-0297). Written informed consent was obtained from all patients for sample collection and molecular analysis. All experiments were performed in accordance with the relevant guidelines.

2.3. Conditionally reprogrammed cell line culture and cell characterization

The experimental procedure was performed as described previously [10,11,14,16]. Cells were characterized by immunofluorescence and Soft agar colony formation assay, and tumourigenicity was evaluated according to the manufacturer’s protocol.

2.3.1. Conditionally reprogrammed cell lines

The tissue was re-suspended in collagenase (1 mg/mL, Sigma, St. Louis, MO, USA) in culture medium and incubated for 30 min at 37°C with agitation to dissociate the tumour tissue from the collagenous stroma. We added 5x F medium for neutralization, followed by centrifugation at 1500 rpm for 3 min. The supernatant was filtered through a cell strainer (70 μm nylon, Falcon). The filtered tumour cells were seeded on a feeder layer of lethally irradiated (30 Gy) J2 murine fibroblasts in F medium. The F medium consisted of 70% Ham’s F-12 nutrient mix (Hyclone, Logan, UT, USA) and 25% complete Dulbecco’s modified Eagle’s medium, supplemented with 0.4 μg/mL hydrocortisone (Sigma), 5 μg/mL insulin, 8.4 ng/mL cholera toxin (Sigma-Aldrich, St. Louis, MO, USA), 10 ng/mL epidermal growth factor, 5% fetal bovine serum (Hyclone), 24 μg/mL adenosine (Sigma), 10 μg/mL gentamycin (Life Technologies), and 250 ng/mL Amphotericin B (Thermo Fisher Scientific, Waltham, MA, USA). Cells were cultured in the presence of the Rho-associated kinase (ROCK) inhibitor, Y-27632 at a final concentration of 5 μM (Enzo Life Sciences, Farmingdale, NY, USA). Cells were incubated at 37°C in a humidified atmosphere with 5% CO2. Tumour cells on the plates were visibly apparent by morphology relative to stromal cells (e.g., fibroblasts). Contaminating stromal cells were removed by differential trypsinization or selective scraping of the plates as necessary. The cell lines were pretreated with 500 ng/mL mycoplasma removal agent (MP Biomedicals, Santa Ana, CA, USA). The generated cell lines were regularly checked to ensure that they were not infected with mycoplasma.

2.3.2. Immunofluorescence assay

Cells were fixed in 4% paraformaldehyde for 15 min at room temperature (RT) and washed three times with PBS, followed by blocking with 5% normal goat serum for 1 h and incubation with primary
antibodies at room temperature (1 h). The cells were then washed three times with PBS, and incubated with secondary antibody for 30 min. The following primary antibodies were used: α-Amylase (sc-25562, rabbit polyclonal; Santa Cruz Biotechnology, dilution 1:100), cytokeratin 19 (A53-B/A2: sc-6278, mouse monoclonal; Santa Cruz Biotechnology, dilution 1:100), insulin (sc-8033, mouse monoclonal; Santa Cruz Biotechnology, dilution 1:100), and vimentin (D21H3: #5741, mouse monoclonal; Cell signaling technology, dilution 1:100). The cells were stained using Alexa Flour-conjugated secondary antibodies from Invitrogen.

3.3.3. Soft agar colony formation assay
To evaluate tumourigenicity, in vitro tumourigenesis and anchorage-independent growth of CRCs were evaluated by colony formation assay in soft agar. Soft agar colony formation assays were performed using the double-layer soft agar method. In each well of a 6-well plate, 5 × 10⁴ cells were plated on the top agar (0.5% agarose gel) over a base agar (1% agarose gel; Santa Cruz Biotechnology, dilution 1:100), and vimentin (D21H3: #5741, mouse monoclonal; Cell signaling technology, dilution 1:100). The cells were stained using Alexa Flour-conjugated secondary antibodies from Invitrogen.

3.4. Immunofluorescence staining for tissue slide
Immunostaining was performed on paraffin-embedded human pancreatic cancer tissues with antibody to S100A2 (#S6797, sigma-aldrich), GATA6 (#AF1700, R&D systems) and esophagus cancer tissue was using GATA6 negative control. The paraffin-embedded slides were deparaffinized by immersion three times in xylene for 5 min. Rehydrate were performed two times 100% EtOH for 5 min, 95% EtOH for 5 min and 90%~30% EtOH for 1min followed by washes two times with H2O for 5 min. Antigen retrieval was performed 0.01 M citrate buffer (pH 6.0) in boiled water bath for 30 minutes and slides were allowed to step-down cooling in the same buffer for 30 min. Endogenous peroxidase activity was blocked 0.3% H2O2 with MeOH at room temperature for 20 minutes. Slides were then incubated with 10% normal horse serum and 0.2% Triton X-100 for 1 hour to reduce background non-specific staining and permeabilized the tissues, followed by three times washing with PBS buffer for 5 min. Each slides was incubated in first primary antibody GATA6 (10 ug/ml) overnight at 4°C. After washing three times, slides were incubated with secondary antibody Alexa Fluor 594 (1:400, Invitrogen) for 1 hr at room temperature. After washing three times, the subsequent reaction was repeated blocking step. Blocked slides were incubated in second primary antibody S100A2 (100 ug/ml) for 1 hr at room temperature. After washing three times, slides were incubated secondary antibody Alexa Fluor 488 (1:400, Invitrogen) for 1 hr at room temperature. After three times washing, the slides were mounted using anti-fade mounting solution contained with DAPI (H-1200, vector Laboratories Inc. Burlingame, CA 94010). The immunofluorescence images were acquired Olympus BX53 microscope and the signal intensity of each image was analysed using ImageJ analysis software (NIH).
3.5. Targeted deep sequencing and RNA sequencing

We validated the genetic similarity of the established CRCs with the original PDAC tissue by comparing 83 targeted gene sequences to develop representative genomic data of PDAC (Supplementary Table 1). Targeted sequencing was performed using the Cancer-SCAN panel (83-gene panel at ~900 × 1) [17]. Initially, DNA obtained by the microdissection of formalin-fixed paraffin-embedded PDAC tissue and DNA from pancreatic CRCs of the same patient were sequenced. DNA was extracted using the QIAamp DNA Mini Kit and sequenced using an Illumina HiSeq2500 system (Illumina, San Diego, CA, USA). Quality of the DNA was evaluated using a Nanodrop spectrophotometer (Thermo Fisher Scientific). Preprocessing of read sequences was conducted by quality filtering using fastQC. We followed the Genome Analysis Toolkit (GATK) best practices workflow for improving the quality of variant calls. Sequencing reads in the targeted region were aligned and compared to the human reference genome (UCSC hg19) using BWA-MEM version 0.7.10 and Picard tools version 1.119 (http://broadinstitute.github.io/picard/). In the case of CRCs, alignment was performed using the combination of human and mouse reference (UCSC mm10) sequences, to remove false positive variants from mouse contamination through feeder layer. According to the GATK best practice, the location of insertions and deletions was recalibrated based on the dbSNP database, version 150, of known variants. Cross-contamination ratio was estimated using ContEst [18] (Supplementary Fig. 1) to identify cross-contamination between the tumours and CRC samples.

3.5.1. Somatic single-nucleotide variant (SNV) and indel calling

SNVs and indels were identified in normal-tumour pair and normal-CRC pair samples by GATK, version 4.1.2.0 Mutect2 [19]. SNVs in 83 targeted genes were annotated with genetic features using SnipEff version 4.3, to obtain the common somatic mutation list based on the Catalog of Somatic Mutation in Cancer database (COSMIC) for annotated mutations [20,21]. Additionally, we used PROVEN version 1.1.5 and SIFT version 6.2.1 to predict the effects of mutations on biological functions [22,23]. To retain high-confidence somatic variants, we applied the following filtering criteria provided in Mutect2: [1] variants with alternative allele counts less than 5, [2] variants with total allele counts (read depth) less than 20, and [3] variants with allele frequency (AF) less than 0.05. The candidates who fulfilled the above criteria were manually checked using Integrative Genomics Viewer (IGV) version 2.3.81 to evade false-positive variants (Supplementary Fig. 2) [24]. The mutations detected in both, tumour and paired CRCs of the same patient were defined as concordance mutations. The concordance rate was calculated as the formula below for primary-CRC pairs.

\[
\text{concordance rate} = \frac{\text{count of shared mutations}}{\text{count of total mutations}}
\]

While primary-only mutation was detected as a somatic mutation in primary tumours, the mutation failed to pass the filtering criteria in the paired CRCs. A mutation is defined as a CRC-only mutation, when it is detected in CRCs as a high-confidence somatic variant, but is not present in the tumour. In order to confirm that a mutation existed in both tumour and matching CRCs, we checked the presence of alternative allele using IGV in the location where the somatic mutation occurs (Supplementary Fig. 2). We checked the tumour and CRCs-only mutations individually using IGV and included them in the concordant mutation category if a low-frequency allele was present in the counterpart.

3.5.2. Copy number variant (CNV) calling

CNV calling was performed using CNVkit version 0.8.6.9 [25]. The corrected on- and off-target log2 values of bin-level copy ratios with associated weights were concatenated using the fix command. The call command with the center mode option was used to recalculate the copy number of the default center-centering the copy number neutral area slightly above or below the expected log2 value of zero. After these adjustments, we used the threshold method to calculate the absolute integer copy number of each segment.

3.5.3. RNA sequencing and characterization of molecular subtype

RNA sequencing using TruSeq Stranded Total RNA Sample Prep Kit with Ribo-Zero H/M/R was performed to confirm the similarity between the original PDAC tissue and CRCs even at the gene expression level. We trimmed the FASTQ files to obtain clean cropped read with length 100 using Trimmomatic [26]. In the case of CRCs, the FASTQ files used Disambiguate removed mouse contamination through feeder layer [27]. STAR and Cufflinks are mapped the read to transcriptome as a reference and calculated normalized gene-level expression values such as FPKM (Features Per Kilobase Million) [28,29]. The analysis of the molecular subtype of CRCs and original tissue proceeded in the same way as described in the previous paper in 2017 [30,31]. We divided the samples into two groups, known as Moffitt subtypes, based on 50 genes expression. Based on 62 genes expression, we also identified three groups known as Collisson subtypes (classical, quasi-mesenchymal, and exocrine).

3.6. Establishment of patient-derived xenographs (PDX) using CRCs

Exponentially growing PDAC CRCs were tryptosinized, dispersed into single cells, and suspended in 200 μL of Matrigel (BD Biosciences). To evaluate tumourigenicity in vivo, 2 × 106 cells prepared in 0.2 mL of Matrigel were injected subcutaneously into the flank region of 5-week-old male NOD/SCID mice with severe combined immunodeficiency (National Cancer Institute, USA). The mice were sacrificed by CO2 inhalation; tumours were harvested once they reached a volume of approximately ~1500 mm3 (20 mm). PDx tumour tissue was digested for: 1) fixation in 10% buffered formalin for paraffin embedding/histology, 2) cryopreservation, 3) PDx cell line development and passaging, and 4) optimal cutting temperature compound embedding. Inclusion criteria are 5-week-old, male, and NOD/SCID mice) and exclusion criteria is that animal might be dropped from the study and euthanized before the predetermined time point if the size of a subcutaneous tumour exceeds 20mm. The primary outcome measure that was used to determine the sample size was the diameter and volume of tumour mass. In order to evaluate the response to chemotherapeutic agents in xenograft models from the CRCs, we performed the randomization of 12 mice into two groups (treatment group, n=6; control group, n=6). We injected gemcitabine in chemotherapeutic group (Gemcitabine, 50 mg/kg, twice/week). Animals were housed at the Yonsei University animal care facility in accordance with the institutional guidelines. All experiments were performed in accordance with the relevant animal use guidelines.

3.7. Cell viability assay and drug sensitivity

The cell viability assay was performed to determine the drug sensitivity of each cell line. CRCs of patients with different prognoses were selected to test the response to anticancer agents. The responder group was defined as follows: [1] patients who had complete remission (CR)/partial response (PR)/ stable disease for at least 4 months or [2] patients who had CR or PR at best. Cells were seeded in 96-well plates at a density of 5,000 cells per well. To identify compounds that are toxic to CRCs, after attachment, the cells were exposed to gemcitabine, 5-FU, oxaliplatin, irinotecan, nab-paclitaxel, and combinations of these drugs for 72 h. CellTiox Green (Promega, Heidelberg, Germany) was added to the growth medium in accordance with the manufacturer’s instructions, and the cells were followed by live cell imaging using the IncuCyte microscope (Essen Bioscience) to identify toxicity. The data from the images were
quantified using the IncuCyte software (version 2011A rev2) and the data plots of cell toxicity extraction and proliferation were plotted using the GraphPad Prism software. Values are the means of triplicate wells from three independent experiments for each drug concentration.

### 3.8. Key resource table

| Reagent or resource | Source | Identifier |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Rabbit polyclonal anti-a-Amylase | Santa Cruz Biotechnology | Cat# SC-25562, RRID:AB_623872 |
| Mouse monoclonal anti-Cytokeratin19 | Santa Cruz Biotechnology | Cat# SC-6278, RRID:AB_627851 |
| Mouse monoclonal anti-Insulin | Santa Cruz Biotechnology | Cat# SC-8033, RRID:AB_627285 |
| Mouse monoclonal anti-Vimentin | Cell signaling technology | Cat# 5741, RRID:AB_10695459 |
| Mouse monoclonal anti-S100A2 | Sigma-Aldrich | Cat# 56797, RRID:AB_261539 |
| Goat polyclonal anti-GATA6 | R and D Systems | Cat# AF1700, RRID:AB_2108901 |
| Alexa Fluor 384 | Thermo fisher scientific | Cat# A27016, RRID:AB_2536080 |
| Alexa Fluor 488 | Thermo fisher scientific | Cat# A28175, RRID:AB_2536161 |
| DAPI | Vector Laboratories | Cat# H-1200, RRID:AB_2336790 |
| Oligonucleotides |        |            |
| PCR primers for KRAS |        |            |
| Forward primer: AGGCCTGCTGAAAATGACTGAA | Bioneer corp. | N/A |
| Reverse primer: GGTCTCTGCACCAATATGCA | Bioneer corp. | N/A |
| Oligonucleotides |        |            |
| Y-27632 dihydrochloride | Enzo Life Science | Cat# ALX-270-333 |
| Nutrient mixture F-12 HAMS | hyclone | Cat# SH30026.01 |
| DMEM/high glucose | hyclone | Cat# SH30243.01 |
| Hydrocortisone | Sigma | Cat# H-0135 |
| Human Epithelial growth factor | Gibco | Cat# PHG0313 |
| Insulin | Sigma | Cat# I-9278 |
| AmphotericinB | Thermo fisher scientific | Cat# A2942 |
| Gentamycin | Gibco | Cat# 15750-060 |
| Chloramxin | Sigma | Cat# C-8052 |
| Adenine | sigma | Cat# A2786 |
| Agar | Thermo fisher scientific | Cat# BD214220 |
| Collagenase I | Gibco | Cat# 17100-017 |
| Collagenase IV | Gibco | Cat# 17104-019 |
| QIAamp DNA Mini kit | Qagen | Cat# 51304 |
| RNeasy Mini kit | Qagen | Cat# 74104 |
| Alcian blue stain kit | Vector laboratories | Cat# H-3501 |
| Matrigel | BD Biosciences | Cat# 356234 |
| zylene | Duscan | Cat# H29118 |
| Ethanol | Millipore corp. | Cat# 100983 |
| Methanol | Millipore corp. | Cat# 106012 |
| Hydrogen peroxidase | FUJIFILM | Cat# 081-04215 |
| TritonX-100 | Sigma | Cat# 78787 |
| Citrate tribasic dihydrate | Sigma | Cat# 54641 |
| Horse serum | Thermo fisher scientific | Cat# 16050130 |
| Fetal bovine serum | Thermo fisher scientific | Cat# 16000-044 |
| ReliaPrep FFPE gDNA Miniprep system | Promega | Cat# A2352 |
| Thiazolyl Blue Tetrazolium Bromide | Sigma | Cat# M2128 |
| Gemcitabine | Lilly | Cat# 7502 |
| Cell lines |        |            |
| YPAC-2 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-5 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-6 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-7 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-10 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-13 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-15 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-16 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-17 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-20 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-21 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-23 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-25 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-26 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-27 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-28 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-29 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-30 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-31 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-32 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-33 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-34 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-35 | Severance Hospital, Seoul, Korea | N/A |
| YPAC-37 | Severance Hospital, Seoul, Korea | N/A |

(continued)
3.9. Statistical analysis

All statistical analyses were performed using GraphPad Prism 7 software. IC 50 value were statistically analysed using Student’s t test, and the graphs show the mean ± SD. Differences between groups were considered to be significant at a P value of <0.05. We did not include additional statistical tests for data distributions.

3.10. Role of funders

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4. Results

4.1. Establishment of CRCs from clinical specimens

We established a total of 28 CRCs from patients who were pathologically diagnosed with PDAC. The following were the original sources for tissue acquisition: surgical resection (n=12), EUS-guided biopsy (n=15), and percutaneous liver biopsy (n=1) [Supplementary Table 2]. Subsequently, we performed genomic analysis of 9 patient samples, in which the original FFPE tissue contained sufficient amount of DNA (>1 μg) after microdissection. Drug screening was performed in 7 patients who showed a defined chemotherapeutic response.

4.2. Characterization of CRCs

Epithelial colonies were readily observed at 2 days and rapidly proliferated to reach confluence in approximately 7-14 days (Fig. 2a). Immunofluorescence for CRCs was performed using a monoclonal antibody against cytokeratin 19. The cytoplasm of cancer cells was clearly stained with this antibody (Fig. 2b and Supplementary Fig. 3). Soft agar colony formation assay showed in vitro tumourigenesis (Fig. 2d). All colony assays can be divided into three groups by their growth (large size and rapid growing, moderate, small size and slow growing). YPAC-2, 5, 26, 35, and 46 showed large colony formation and rapid growing feature. In contrast, YPAC-10, 15, 16, 23, 25, 29, 39, 43, 44, 46, Swiss 3T3-J2 cell (fibroblast, mouse) from Georgetown University, N/A; NOD/SCID from National Cancer Institute, USA, N/A; BALB/c Nude from Institute of Medical Science, University of Tokyo, N/A.

![Fig. 2.](image-url) Characterization of conditionally reprogrammed cell lines (representative image). (a) Conditionally reprogrammed cell formation was confirmed after 15 days of culture, and within one month. (Scale bar, 200 μm) (b) Characterization of the intensely fluorescent cells in conditionally reprogrammed cells. (Scale bar, 100 μm) (c) KRAS mutations were analysed by polymerase chain reaction (p.Gly12Val (GGT → GTT)). (d) Soft agar colony formation assay showed in vitro tumourigenesis.
30, and 34 showed small colony formation and slow growing feature. Large colony group showed relatively larger tumour size at their diagnosis, many proportion of unresectable cancer stage, metastatic status. CRCs with aggressive cancer characteristics showed large colony formation and rapid growth (Supplementary Fig. 4 and Supplementary Table 6). With respect to the key oncogenic mutations, PCR was used to perform KRAS mutation (Fig. 2c and Supplementary Fig. 5). Of the 28 samples, 27 showed KRAS mutations in codon 12/13 (n=25, KRAS mutation was confirmed by targeted sequencing. Sanger sequencing was not performed for 1 patient sample because of low amount of DNA) or codon 61 (n=1).

4.3. Genetic characterization of patient-derived models

4.3.1. Sample pair matching

To verify the origin of the patient-derived models, all the 9 CRCs and their matched primary tissues used in this study were checked for their germline single-nucleotide polymorphism (SNP) similarity (Fig. 3a). CRCs were matched with their paired blood and original tumour tissue. In addition, the proportion of exonic variations was well retained among the primary pancreatic cancer tissues and the corresponding cancer CRCs (Fig. 3b). Analysis of the distribution of base substitutions in both pancreatic cancer tissues and CRCs revealed over-representation of C>T transition; this is in agreement with the mutational spectrum presented in previous studies [32].

4.3.2. Genetic variants

In the initial call, 36 somatic mutations were detected from each set of the 9 primary tumours and 9 CRCs. We found that 30 of these mutations were shared in the primary-CRC pairs (Fig. 4a, Supplementary Table 3 and 4), leaving two primary-only, and four CRC-only mutations. The shared mutations included key oncogenic mutations such as KRAS (9 pairs), TP53 (8 pairs), and SMAD4 (3 pairs). The overall number of non-synonymous mutations per patient was consistent with that presented in a previous report (median of 2 SNVs, 4.27 per Mb) [33]. The two primary-only mutations were, a synonymous mutation in ATRX (YPAC-2) and an intron variant in NPM1 (YPAC-17), implying a low phenotypic impact. Furthermore, deeper assessments identified that the four CRC-only mutations were not de novo but had pre-existed in alleles from matching primary tumours at low AF, which only became callable with an increase in VAFs (see Methods, Supplementary Fig. 2). While primary tumour specimens normally contain non-tumourous cells (e.g., stromal cells), frequently in high numbers (e.g., 30-80%), increased tumour cell purity in CRCs enables more accurate profiling of somatic mutations [34]. Overall, the 9 CRCs preserved the genetic characteristics of the primary tumours with high concordance, with additional confirmation of low-AF NPM1 mutation in CRC (35 shared mutations out of total 36, concordance rate=97.2%).

The presented AFs of the 36 shared mutations confirmed the clarity of genomic profiles in the CRCs (Fig. 4B). The list of genes with mutations was preserved and the overall VAFs were increased in

Fig. 3. Sample pair matching. (a) The concordance ratio of germline single-nucleotide polymorphisms (SNPs) of objects on the y-axis to the x-axis is defined as the similarity score and the closer the maximum value is to 1, higher is the similarity, represented by blue colors. Nine conditionally reprogrammed cell lines (CRCs) were matched with their paired blood (normal) and original tumour tissue. Furthermore, we confirmed the genetic coincidence and originality between the original tissue and patient-derived model. Colors indicate the similarity of paired samples, with high similarity represented by blue and low similarity represented by yellow. (b) The proportion of base substitutions was maintained in a similar pattern in each of the primary pancreatic cancer tissue and corresponding cancer CRCs, confirming that they were from the same individual. (For interpretation of the references to color in the text, the reader is referred to the web version of this article.)
Fig. 4. Genetic characterization. (a) The two connected circles indicate the somatic mutation detected in both the primary and the CRC, and the single circle implies that the expressed mutation was detected only in either the primary or the CRC. The effects on biological functions due to allele change were distinguished by color, wherein light grey color represents the tolerant effect of the mutation such as synonymous mutation and intron variants. The dark grey color indicates the damaging effect of the mutation such as missense variant and frameshift. The nine CRCs represent the concordance of somatic mutation with the exception of the tolerant mutation of YPAC-2 marked as the dashed line. (b) The x-axis represents the variant allele frequency (VAF) of the mutations in the primary tissue; the y-axis represents the VAF of the mutations in the CRC. In addition, the allele frequency plot shows the gene name and mutation type for each mutation, which are color-coded, and tolerant mutations are marked in grey. (c) The graph indicates comparison between the copy number in the primary tumour (left on the x-axis) and CRCs (right on the x-axis) for 83 targeted genes. The copy number on the y-axis with a value 2.5 or higher is marked as amplification, colored with red and deletion as blue for 1.5 or less of copy number. The unaltered copy numbers in the primary tumour and CRCs are marked as conserved and represent the orange area. Copy number variants (CNVs) are marked with bold dotted lines. (For interpretation of the references to color in the text, the reader is referred to the web version of this article.)
CRCs (average VAF=60.9%) compared to those in primary tumours (24.5%). Specifically, in KRAS mutations (i.e., confident drivers and expected to be clonal in all tumour cells), the VAFs were less than 30% in 5 out of 9 primary tumours (12.3-26.7%), which strongly suggests low tumour purity in the specimen. In the matching CRCs, the VAFs were increased to approximately 50% (Fig. 4B). Similar or a significant increase of VAFs was also observed in other key mutations including TP53 (25.9% to 89.1%), SMAD4 (17.7% to 79.1%), CDKN2A (29.9% to 74.8%), and ARID1A (14.4% to 50%). We noted that the VAFs of some mutations (e.g., TP53) reached approximately 100%, which can be obtained either by homozygous mutation, loss of heterozygosity (e.g., loss of wild-type copy), or multi-copy amplification. While many studies have observed these copy number changes during tumour progression [35], there could also be a possibility for selective advantage during the generation of CRCs, which requires further investigation to separate these effects.

Likewise, all the 27 CNVs (18 deletions and 9 amplifications) initially called in primary tumours were retained in CRCs (Fig. 4C, Supplementary Table 5). Moreover, the estimated copy numbers (CNs) of the 83 targeted genes remained comparable in CRCs (average CN of 2.00 vs. 2.00). Of note, we found that the degree of copy number changes in key oncogenic CNVs including CDKN2A deletion (1.9 vs. 0.6) and AKT2 amplification (1.9 vs. 2.2) was increased in CRCs. The increased somatic mutation VAFs suggest that CNVs can be better clarified by constructing CRCs.

On WES, The mutation burden of 173 TCGA-PAAD samples was distributed from 1 to 115 (average 37.03). Comparing to TCGA mutation burden, we found that the mutation burden in 10 CRC samples distributed from 1 to 115 (average 37.03). Comparing to TCGA mutation type was matched with original tumour (YPAC-2, 5). On average, 50% of the total samples have KRAS mutations and 60% (6/10) of the total samples have TP53 mutations, the key gene of Pancreatic Adenocarcinoma, as shown in the TCGA-PAAD analysis (Supplementary Fig. 8).

### 4.3.3. Molecular subtype

We confirmed the preservation of subtypes between CRCs and their corresponding primary tumours by using four primary-CRCs pairs (YPAC-5, 17, 23, 26) (Fig. 5). All samples showed a concordance between subtype based on moiffit et al. and half of samples maintained their molecular subtype based on collision et al. (30, 31). YPAC-5, 26 were classified as basal type and YPAC-17, 23 were classified as classical type through transcriptome analysis. In transcriptome analysis, key gene expression such as GATA6, HNF4A of classical type was higher in YPAC 17, 23 (Fig. 5a). To validate the critical gene expression, we investigated the immunofluorescence expression of GATA6 and S100A2 between original tissue slide and CRCs (Fig. 5b). S100A2 and GATA6 expression in tumour tissue was consistent with the expression of matched CRCs for YPAC-23 and 26. Classical type YPAC-23 and its original tumour tissue simultaneously showed low signal intensity of S100A2 and high signal intensity of GATA6. Compared to YPAC-23, basal type YPAC-26 and its original tumour tissue simultaneously showed high signal intensity of S100A2 and low signal intensity of GATA6.

### 4.3.4. In vivo tumourigenesis

In vivo, three CRCs (YPAC-2, 5, 21) were implanted in three NOD/SCID mice, respectively (Supplementary Fig. 9). Implanted CRCs showed tumour engraftment and the grafted tumours were palpated in the flank of mice after 2-3 months (Fig. 6a, YPAC-2). YPAC-21 was from a patient with cystic adenocarcinoma tumour tissue. Mouse xenografted tumours were also mucinous (Fig. 6b and Supplementary Fig. 10), similar to the original tumour phenotype. Histological analysis showed the presence of atypical cells. They were distributed in mucin-filled cysts lined by stacked layers of dysplastic epithelial cells with high nucleus-to-cytoplasm ratio. Hematoxylin and eosin staining showed that the implanted tumour tissues shared similar morphological features with the parent tumour tissue (Fig. 6c). After targeted deep sequencing and sanger sequencing, we confirmed the KRAS mutation type was matched with original tumour (YPAC-2, 5) (Supplementary Fig. 11). In order to evaluate the response to chemotherapeutic agents in xenograft models from the CRCs, we implanted...
the pancreatic cancer patient-derived CRCs in the flank of mouse (vehicle, n=6; gemcitabine, n=6). And, we injected normal saline in vehicle group and gemcitabine in chemotherapeutic group. We found the anti-tumour effect of gemcitabine using xenograft model derived from the CRCs (Supplementary Fig. 12).

4.4. Drug sensitivity assay

We compared responses to anticancer agents in CRCs, with the clinical response of pancreatic cancer patients. Therapeutic options for pancreatic cancer patients include combination chemotherapy regimens with gemcitabine/nab-paclitaxel or FOLFIRINOX (5-fluorouracil, leucovorin, irinotecan, and oxaliplatin). Based on responsiveness to chemotherapy, we divided the patients into a responder group (YPAC-25, 26, 30) and non-responder group (YPAC-2, 5, 16, 31). Other patients showed intermediate responsiveness, therefore we did not perform drug sensitivity test. Results of the cell viability assay indicated that the anti-proliferative effect of anticancer agent was significantly higher in YPAC-25, 26, 30 than in YPAC-2, 5, 16, 31. Drug sensitivity of CRCs corresponded with individual patient’s anticancer treatment response. CRCs of the responder group were more sensitive to anticancer agents than those of the non-responder group (Fig. 7a-e). The IC50 value of each drug was statistically lower in the responder group than in the non-responder group (Fig. 7f and g). In a clinical setting, YPAC-2 patient showed increased tumour size in the first chemotherapy response just after 2 months. However, YPAC-25 patient showed good response with decreased tumour size for over 12 months compared to YPAC-2 patient (progression free survival,

Fig. 6. In vivo tumourigenesis. (a) Implanted CRCs (YPAC-2) showed successful tumour engraftment, and a 15 mm sized transplanted tumour on the side of the mouse was palpated after three months. (N=3, Supplementary Fig. 9) (b) CRCs of a patient who was diagnosed with cystic adenocarcinoma showed similar phenotype in mouse CRC implantation. White circle, cystic adenocarcinoma; Black dotted circle, implanted cystic tumour; black dotted square, mucin-filled cysts lined by layers of dysplastic epithelial cells (c) Histological examination showed that CRCs and xenograft were phenotypically similar when compared with primary cancer tissue. CK19, cytokeratin 19. (Scale bar, 200 μm)
360 vs. 65 days) (Supplementary Fig. 13). Ex-vivo examination of patient-derived cell lines also showed similar drug responsiveness, which matched with clinical results.

5. Discussion

We established cell culture models derived from tumour samples of PDAC patients and then performed genetic analyses using these cells. The key oncogenic mutation present in the tumour tissue was reliably identified in the derived cell line. CRCs can be isolated from patients at all stages of pancreatic cancer using the limited tissue available from a fine needle tumour biopsy as the starting tissue.

To date, various cancer models have been developed for cancer research. At first, cancer cell lines also exhibit tumour characteristics but long-term culture is difficult and individual patient-derived matched model is difficult to develop. A few models have been developed using primary PDAC to establish stable cancer cell lines [36-38], and varying success rates (7.4-74.1%) have been reported [39,40]. However, commercialized cancer cell line models have repeatedly shown technical and biological limitations including cellular senescence, reduced stability over time, and insufficient recapitulation of the primary tumour [7,9,41,42]. Moreover, generating patient-derived xenograft models (PDX) commonly used for cancer research requires a cost- and time-intensive endeavor (more than 4-6 months). Though the recently developed 3D cancer model, called organoid, recapitulates the original tumour, it stimulates LGR5 (+) stem-like cells during the formation of initial tumours [41]. Recent studies have reported that for organoid culture—although the culture needs many niche factors—the composition of the surrounding niche factors still changes in pancreatic cancer [43,44]. Above all, niche factors can also affect the result of drug screening through chemical reaction and, considerable financial resources are required for high-throughput drug screening using the organoid culture [8].

In previous studies, the success rate of CRC culture was found to differ according to cancer type (from 24.2 to 50%) [10,45-47]. For pancreatic cancer, Liu et al. developed four cell lines (2 tumour and 2 normal) from 8 pancreatic tissue samples [10]. Natalya et al. reported three cell line models from surgical tissue for pancreatic cancer, but the number of patients was too small to characterize the method [46]. The present study was the largest to develop PDAC CRCs. Further, previous studies have been conducted using surgical tissues that account for only 10% to 20% of PDAC. In this study, we performed experiments using endoscopic biopsies of patients with inoperable PDAC, indicating the novelty of this study. Histology-based morphological analysis revealed similarities between mutations in matched samples, primary tumours, and CRCs, with increased AF observed in CRCs. KRAS, TP53, and SMAD4 are representative somatic mutations of pancreatic cancer [48]. In samples subjected to sequencing and PCR, these typical mutations were detected. Furthermore, the key oncogenic mutations—present at a low AF in the tumour—showed an increased AF in CRCs. This may

![Fig. 7. Conditionally reprogrammed cell lines reveal heterogeneity of chemotherapeutic response. The results of the cell viability assay indicated that the anti-proliferative effect of the anticancer agent was significantly higher in YPAC-25, 26, and 30 than in YPAC-2, 5, 16 and 31. Drug sensitivity of CRCs correlated with individual patient anticancer treatment response. CRCs of responder group were more sensitive to anticancer agents than those of the non-responder group. Viability assay were performed at least two times in triplicate experiments. ***, P < 0.0001. AUC, Area Under The Curve [IC50 value was statistically analysed using Student’s t test, and the graphs show the mean ± standard deviation]
be a useful tool for detecting minor allele deficiency mutations in cancer.

Even if the CRC protocol is obviously less lab-intensive and more affordable, tumour organoid culture is well-established method for in vitro maintenance of patient-derived samples. For side-by-side comparison to the success rate in both methods, previous studies for CRCs showed 50–100% success rate, even if the number of sample was small (3–5 cases) and there was no enough information about the tissue acquisition route [10,46,49]. In case of organoids, the success rate of between 66% and 100% for samples from fine needle biopsy is reported by previous researchers [44,50,51]. But, there have been no large-scale studies of fine needle biopsy in CRCs. Second, there are few studies to investigate molecular feature of CRC methods in PDAC rather than organoid culture, previously. In other cancer type, they showed the maintenance of molecular feature in CRC matched with original tumour and preserved key mutation and tissue heterogeneity [52]. Beglyarova et al. demonstrated the MYC–ERCC3 interaction as a target for PDAC and a new mechanistic approach for the disruption of critical signaling in MYC-dependent cancers using the patient-derived pancreatic cancer CRCs [46]. Relatively, several studies reported the preservation of similar molecular characteristics in organoids compared to matched cancer tissue [44,51]. Tuveson et al. found that pancreatic cancer patient-derived organoids therapeutic profiles paralleled patient outcomes. They proposed that combined molecular and therapeutic profiling of organoids may predict clinical response and enable prospective therapeutic selection [44].

Precision medicine focuses on delivering the most appropriate therapy to a patient through drug screening based on the clinical and molecular features of their disease. In the present study, CRCs showed similar characteristics to those of the original tumour samples. We tested the feasibility of using CRCs derived from metastatic pancreatic cancers as drug screening tools and validated the robustness of our approach. The drug sensitivity test revealed that anti-cancer drug sensitivity was reliably associated with patient prognosis. By evaluating the drug sensitivity of a large panel of clinical agents using CRCs, this platform might identify a targetable new drug [46].

Recent developments in genomic medicine for PDAC treatment have provided various options for personalized treatment [30,31,53–58]. Studies have been reported wherein patients were divided into several groups based on their responsiveness to chemotheraphy and survival [53–56,59–61]. In PDAC research, major limitations in implementing treatment strategies based on mutational status are the availability of tumour samples for detailed sequence analysis. In addition, [1] difficulty in primary cell culture owing to limited life span, [2] difficulty in tumour sequencing owing to the dominance of stromal cells, and [3] absence of an appropriate pre-clinical model preserving the molecular characteristics of tumour cells are also serious limitations. Therefore, generating CRCs for amplifying tumour epithelial cells could serve as an important pre-clinical strategy. Established CRCs of pancreatic cancer predominantly comprising neoplastic cells thereby enable scientists to study low frequency nucleotide variants and copy number alterations that would be difficult to discern in primary tumour tissue with low neo-plastic cellularity. Furthermore, most primary cell cultures are difficult to maintain because they have a limited lifespan due to their gradual decrease in proliferation, eventually leading to senescence. In the present study, we maintained YPAC-2 for over 2 years. Therefore, the CRC methodology has wide application potential and can be adapted for live bio-banking and basic research, as well as for diagnostic and therapeutic purposes.

There are several limitations of CRC methodology compared with PDX or organoids. Compared with PDX, some CRC cells derived from malignant tumours were often non-malignant, without tumour-derived mutations. It is also a kind of issue in organoids with normal epithelial cell contamination [62,63]. Researchers try to reduce of normal like organoid using specialized method such as media selection, picking method according to morphology in organoids [64]. In this study, we used refined method as follows: IF such as pancreatic cancer marker, PCR by most common gene KRAS, tumour formation assay, and targeted sequencing. Another limitation of CRC is that it cannot recapitulate with tumour microenvironment. It is difficult to evaluate the impact of stromal cells on tumour-cell growth and their effect on the tumour-cell response to treatments. Regarding organoids, recent research for co-culture system with cancer associated fibroblast and/or immune cell is promising [65–68]. Recent study reported organoid culture after CRC establishment which can be a solution of above problem in cancer research [52].

Nonetheless, there are several strengths in CRCs methods. At first, rapid expansion is possible for high throughput drug screening because of the characterization of 2D culture. Additionally, these cultures are more suitable for high throughput rather than low-throughput drug screening. Overall it takes 4-6 weeks to provide enough cells for drug screening. CRC can be reproduced for long periods of time without genomic alterations. Conditionally reprogrammed cells retain cell lineage commitment and maintain the heterogeneity of cells present in a biopsy. Compared to organoids, it does not depend on the extracellular matrix matrigel, which can hamper drug penetration and be adverse in drug screens.

In conclusion, using the conditional reprogramming cell culture technique, we established patient-specific preclinical cancer cell lines. Sequencing analysis revealed that the CRCs maintained the mutations present in the original tumour tissue. The use of matched patient-derived cells provided a unique ex vivo model for personalized cancer therapy of PDAC.

6. Author’s Contributions

H.S.L., E.K., S.K., and S.B. designed the study and directed the entire study. E.K. and S.K. performed analyses of sequencing data. H.S.L., J.L., C.H.P., S.J.P., and J.M.H. performed in vitro experiments. H.S.L. and S.J. performed statistical analyses. H.K.H., C.M.K., H.K., J.H.J., I.R.C., M.J.C., J.Y.P., S.W.P., S.Y.S., provided samples and clinical data. H.S.L. and E.K. wrote the manuscript. H.K.H., C.M.K., H.K., J.H.J., I.R.C., M.J.C., J.Y.P., S.W.P., S.Y.S. critically reviewed the manuscript. All authors reviewed and approved the final manuscript.

7. Data sharing

Data presented in this study can be downloaded through the Sequence Read Archive (SRA) under BioProject ID: PRJNA673595. The authors declare that all data supporting the findings of this study are available with its supplementary information files. Extra data is available from the corresponding authors upon request. The data can be available to others in de-identified form after all primary and secondary endpoints have been published and in the presence of a data transfer agreement.

Declaration of Competing Interest

The authors declare no conflict of interest.

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Supplementary materials

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