Abstract. Desmoplastic malignant pleural mesothelioma (DMM) is a rare histological variant of malignant pleural mesothelioma, which is a highly aggressive neoplasm of the mesothelium. DMM is associated with distant metastases and short survival. Effective treatments for DMM are not established and the development of histotype-tailored treatments is difficult due to the rarity of the disease. Although patient-derived cancer models are crucial tools for the development of novel therapeutics, they are difficult to obtain for DMM; no DMM cell lines or xenografts are available from public biobanks and only two cell lines have been reported. Thus, the present study aimed to establish a novel cell line of DMM as a resource for drug screening. A cell line of DMM was established, designated as NCC-DMM1-C1, using surgically resected tumor tissues from a 73-year-old male patient with DMM. Characteristics of NCC-DMM1-C1 cells were examined, such as growth, spheroid formation and invasion capability. Drug targets and anti-cancer drugs with anti-proliferative efficacy were examined using a comprehensive kinase activity assay and drug screening of 213 anti-cancer agents, respectively. NCC-DMM1-C1 exhibited fast growth, spheroid formation and invasion capability, suggesting that the NCC-DMM1-C1 cells retained the aggressive features of DMM. NCC-DMM1-C1 cells and the tumor tissue shared common activity profiles of kinases, which included FES, Wee1, platelet-derived growth factor receptor-β and Src. The drug screening revealed that bortezomib, fostamatinib, gemcitabine, homoharringtonine and vinorelbine had anti-proliferative effects, which have not been previously reported for DMM. It was concluded that NCC-DMM1-C1 cells may be a useful tool for the study of DMM.

Introduction

Desmoplastic malignant pleural mesothelioma (DMM) is a rare histological variant of malignant pleural mesothelioma (MPM), accounting for 5-10% of all cases of MPM. DMM is categorized as a subtype of sarcomatoid tumor (1) and characterized by dense, collagenized tissue (>50%) separated by atypical cells arranged in a storiform or ‘patternless pattern’ in the tumor specimen (2). The pathological diagnosis of DMM is difficult, with fibrous pleuritis and reactive mesothelial hyperplasia as potential differential diagnoses (3). The role of immunohistochemistry is important for the diagnosis of DMM and broad-spectrum staining of cytokeratins is crucial to diagnose DMM correctly. In addition, p16 deletion is useful to distinguish DMM from benign pleuritic (2). The characteristic computed tomography (CT) findings include unilateral pleural effusion, thickening of the mediastinal pleura, circumferential and nodular pleural thickening of >1 cm and interlobar fissure thickening (4-6). MPM is refractory to chemotherapy and radiation therapy, and the combination of pemetrexed and cisplatin has become the standard first-line chemotherapy regimen for MPM based on the results of a randomized phase III trial (7). However, this regimen has only improved median survival from 9.3 months for treatment with cisplatin alone to 12.1 months (7) and it lacks the impact on survival that second-line therapy is expected to have (8). Thus, further investigations for novel therapeutics have long been desired for MPM and DMM (9).

Patient-derived cancer cells that retain the genetic and phenotypic profiles of their original tumor tissue are crucial for the elucidation of molecular mechanisms underlying the malignant features of tumors and for the development of novel therapies (10). In particular, patient-derived cell lines provide a model for screening numerous anti-cancer agents and investigate their modes of action in a high-throughput...
DNA was assessed in...

DNA was extracted from the cells and the tumor tissue using (Promega Corporation), as previously described (17). Genomic repeats (STRs) in 10 loci using the GenePrint 10 System. The cell line was authenticated by examining the short tandem repeats of adequate patient-derived cancer models is one factor hindering the development of novel therapies for DMM and the establishment of patient-derived DMM models is an urgent requirement.

In the present study, a novel cell line, designated NCC-DMM1-C1, was established from tumor tissue of DMM and characterized. Its utility in high-throughput screening of anti-cancer agents for anti-proliferative effects was then evaluated. To the best of our knowledge, the present study is the third report describing the establishment of a patient-derived cell line of DMM.

Materials and methods

Patient history. The patient was a 73-year-old male who visited the National Cancer Center Hospital (Tokyo, Japan) with a major symptom of progressive dyspnea. The patient had been exposed to asbestos from the age of 25 to 60 years. Enhanced CT detected pleural thickening and pleural effusion on the right side (Fig. 1A). Pathological diagnosis using pleural biopsy suggested DMM. Pleurectomy/decortication was performed and the definitive diagnosis from surgically resected tumor tissues was DMM (pT2 pN1 cM0 pStage II). The tumor tissue obtained at the time of surgery was used to establish the cell line. After seven months, multiple liver metastases occurred and the patient developed jaundice and died from pneumonia and respiratory failure due to DMM recurrence.

Cell culture procedure. Cell culture was performed according to a previous study by our group (17). Informed consent was obtained from the patient. In brief, the resected tumor tissue was mechanically dissected with scissors in tissue culture plates (Thermo Fisher Scientific, Inc.). The cells were maintained in Dulbecco's modified Eagle's medium (DMEM)/F12 (Sigma-Aldrich; Merck KGaA) supplemented with 10% heat-inactivated fetal bovine serum (FBS; Gibco; Thermo Fisher Scientific, Inc.), 100 U penicillin G and 100 µg/ml streptomycin (Gibco; Thermo Fisher Scientific, Inc.). After reaching sub-confluence, the cells were treated with 0.25% trypsin-EDTA (Gibco; Thermo Fisher Scientific, Inc.) and transferred to a new tissue culture plate.

Figure 1. Clinical imaging. Computed tomography indicated irregular pleural thickening involving both parietal and visceral pleurae and pleural effusion in the right chest area. The yellow arrow indicates the right-sided desmoplastic malignant pleural mesothelioma.

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Authentication and quality control of the established cell line. The cell line was authenticated by examining the short tandem repeats (STRs) in 10 loci using the GenePrint 10 System (Promega Corporation), as previously described (17). Genomic DNA was extracted from the cells and the tumor tissue using a DNeasy Blood & Tissue kit (Qiagen GmbH) and quantified using a NanoDrop 8000 Spectrophotometer (Thermo Fisher Scientific, Inc.). STRs were amplified and sequenced using a 3500xL Genetic Analyzer (Applied Biosystems; Thermo Fisher Scientific, Inc.). STR profiling was performed using the GeneMapper Software (Applied Biosystems; Thermo Fisher Scientific, Inc.). The similarity of STRs among the reported cell lines was examined using the CLASTR function of Cellosaurus (18). Based on the Tanabe algorithm (19), the STR match ratio between the established cell lines and the corresponding original tumor was calculated. The score met the standard match threshold of 80% (19) based on Tanabe algorithms and non-empty marker mode that computed the score to handle cases in which allele data is missing for the query or the reference. Mycoplasma DNA was assessed in the tissue culture medium using an e-Myc Mycoplasma PCR Detection Kit (Intron Biotechnology).

Single nucleotide polymorphism (SNP) array. Copy number alterations were examined using an Infinium OmniExpressExome-8 v.1.4 BeadChip (Illumina, Inc.) following the manufacturer's protocol. Detailed descriptions of these procedures are provided in previous reports (20). Abnormal copy number regions were detected using the circular binary segmentation algorithm (21,22) using the R package DNAcopy from Bioconductor (23). Amplifications were defined as regions for which the copy number was >1. Deletions were defined as regions of which <1 copy was present in the tumor cells. Among identified genes with copy number alterations, a search for ‘cancer-related genes’ was performed in the Cancer Gene Census in the Catalogue Of Somatic Mutations In Cancer database (GRCh 37 v.91) (24).

Histological evaluation. All specimens for H&E staining were cut into 4-µm slices and placed on the slides. After deparaffinization, the nuclei were immersed in hematoxylin staining solution (Muto Chemical) for 15 min, followed by washing out the solution with tap water. Subsequently, cytoplasm and stromal matrix were stained with eosin (Muto Chemical). The slides
were washed in water and mounted. Immunohistochemical staining was performed on formalin-fixed, paraffin-embedded specimens obtained through surgery and on NCC-DMM1-C1 cells. NCC-DMM1-C1 cells were detached by treatment with trypsin and cell suspensions were solidified using iPGell (Genostaff) according to the manufacturer’s protocol. Cell masses were fixed with 10% formalin and embedded in paraffin. Cell blocks were cut into sections that were processed for immunolabeling and H&E staining. Sections were incubated with antibodies against various proteins. The expression of cytokeratin, D2-40, HEG1 and WT1 was evaluated using the following primary antibodies: Cytokeratin AE1/3 (AE1/AE3; 1:200 dilution; cat. no. sc-81714; Santa Cruz Biotechnology, Inc.), D2-40 (D2-40; 1:200 dilution; cat. no. PDM558; Diagnostic Biosystem), sialylated heart development protein with EGF-like domains 1 (SKM9-2; prediluted; cat. no. 418231; Nichirei Biosciences, Inc.) and WT1 (6F-H2; M3561, 1:50 dilution; Agilent Technologies, Inc.). Antigen detection was performed using a Dako Autostainer and EnVision Detection System (Dako; Agilent Technologies, Inc.) according to the manufacturer’s protocol. The slides were counterstained with hematoxylin.

Cell proliferation assay. The cells were seeded into 24-well culture plates at a density of 2.5x10^4 cells/well as described previously (17). The cell viability was monitored using Cell Counting Kit-8 (CCK-8; Dojindo Molecular Technologies, Inc.). The absorbance of each well at 450 nm was recorded at multiple time-points using a microplate reader (Bio-Rad Laboratories, Inc.). Growth curves were constructed by plotting the absorbance as a function of culture time and these were used to estimate the population doubling time. The cells were maintained in a humidified atmosphere of 5% CO₂ at 37°C.

Spheroid formation assay. Spheroid formation was performed by placing 1x10^4 cells in a 96-well Clear Flat Bottom Ultra-Low Attachment Microplate (Corning, Inc.) in DMEM/F12 medium containing 10% FBS, as described previously (17). After three days of plating, spheroid formation was monitored using a BZ-9000 fluorescence microscope (Keyence). Spheroid formation assays were performed in duplicate.

Transwell cell invasion assay. The invasive capability of NCC-DMM1-C1 cells and MG63 osteosarcoma cells (The Japanese Collection of Research Bioresources) (25) was measured as previously described (17). Cells (1x10^5 and 2x10^5) were seeded in the upper chamber. Following incubation, the cells in the three separate areas were counted under a microscope (Keyence) at x100 magnification.

Tyrosine kinase activity assay. Tyrosine kinase activity was examined for multiple kinases using the PamChip TK peptide microarray system (PamGene International B.V.) as previously described (26). The protein lysate (5 µg) was extracted from the tissue cultured cells and tumor tissue and hybridized to the membrane array, which included 144 peptides. The average signal intensity of the 144 hybridized peptides based on the end levels of the phosphorylation curve was used for the activity analysis. All data analyses were performed using BioNavigator v.6.3.67.0 (PamGene International B.V.).

### Table I. Results of short tandem repeat analysis.

| STR locus (chromosome) | NCC-DMM1-C1 | Normal tissue |
|------------------------|-------------|---------------|
| Amelogenin (X Y)       | X, Y        | X, Y          |
| TH01 (3)               | 9           | 9             |
| D21S11 (21)            | 29, 32.2    | 29, 32.2      |
| D5S818 (5)             | 9, 11       | 9, 11         |
| D13S317 (13)           | 12          | 12            |
| D7S820 (7)             | 8           | 8             |
| D16S539 (16)           | 9, 12       | 9, 12         |
| CSF1PO (5)             | 12          | 12            |
| vWA (12)               | 18, 19      | 18, 19        |
| TPOX (2)               | 8, 11       | 8, 11         |

Active kinases were predicted using PhosphoSitePlus (27), the UniProt database (28) and the Human Protein Reference Database (29).

### Screening for anti-proliferative effects of anti-cancer agents.

The anti-proliferative effects of 213 anti-cancer agents (Table S1) were examined using the CCK-8 assay, as described previously (17). The cells were seeded at 5x10^3 cells/well in DMEM/F12 medium supplemented with 10% FBS using the Bravo Automated Liquid Handling Platform (Agilent Technologies, Inc.). The following day, anti-cancer agent compounds (10 µM; Selleck Chemicals) and 0.1% DMSO were added using the Bravo Automated Liquid Handler. After 72 h, survival rates were assessed using the CCK-8 reagent according to the manufacturer’s protocol. The response rate was calculated relative to that of the DMSO control. Dose-response experiments were performed to validate the candidate anti-cancer agents that were identified during the pilot screening. The compounds were dispensed into 384-well plates with serial dilution at 10 different concentrations, ranging from 0.1 to 100,000 nM, using an Echo 555 Acoustic Liquid Handler (Labcyte Inc.) and IC₅₀ values were determined. CCK-8 absorbance was measured using an Epoch multimode multiplate reader (BioTek Instruments). Absorbance values were plotted as a function of the compound concentrations to obtain IC₅₀ values using GraphPad Prism 8.1.0 software (GraphPad Inc.). This screening was performed in duplicate.

### Results

**Authentication and quality control of the established cell line.**

NCC-DMM1-C1 was maintained for >16 months and passaged >30 times under tissue culture conditions. The characterization of NCC-DMM1-C1 cells was performed by STR profiling at the 20th passage. The same patterns of the peaks in all loci (>30 times under tissue culture conditions. The characterization of NCC-DMM1-C1 cells was performed by STR profiling at the 20th passage. The same patterns of the peaks in all loci (i.e., D5S818, D13S317, D7S820, D16S539, vWA, TH01, TPOX, CSF1PO, and the sex chromosomal marker amelogenin) were observed in both the normal tissues and the NCC-DMM1-C1 cells. The normal tissues were composed of the adjacent tissue to the tumor tissue which did not include the
tumor cells near the chest wall. The authentication indicated that the normal tissues and the NCC-DMM1-C1 cells were derived from the same origin (Table I, Fig. S1). Calculated based on the Tanabe algorithm (19), the STR match ratio between the NCC-DMM1-C1 cells and the corresponding original tumor was 100%. The score met the standard match

Figure 2. Single nucleotide polymorphism genotyping of NCC-DMM1-C1 cells. (A and B) The chromosome numbers are displayed on the x-axis; chromosomes 1 to 22 are presented from left to right using green and black colors, and chromosomal segments are displayed in red dots. The copy number values are indicated on the y-axis. Multiple deletions were identified in NCC-DMM1-C1 cells. (B) Cytogenetic band of chromosome 9 and the copy number variation data of chromosome 9. The genomic positions are indicated on the x-axis and the copy number values are indicated on the y-axis. The red dots indicate chromosomal segments. The red bar indicates the corresponding site of cyclin-dependent kinase inhibitor 2A.
According to Cellosaurus, the profile did not match that of cell lines deposited in public cell banks. It was therefore indicated that NCC-DMM1-C1 is a novel cell line. The DNA sequence of *Mycoplasma* was not detected in the tissue culture medium (data not shown) and it was concluded that NCC-DMM1-C1 was not contaminated with *Mycoplasma*.

**Cell line characteristics.** SNP array experiments revealed the presence of copy number variants mostly involving partial deletions of chromosomal arms (9p) (Fig. 2A) and cyclin-dependent kinase inhibitor 2A was located in the loci of deletions (Fig. 2B). NCC-DMM1-C1 cells exhibited a polygonal or spindle-shaped cytoplasm, arranged in a partially cohesive sheet-like structure, with enlarged and variably sized nuclei (Fig. 3A). The NCC-DMM1-C1 cells formed spheroids in low-attachment substrates (Fig. 3B) and exhibited constant growth (Fig. 3C). The population doubling time based on the growth curve was 25.9 h (Fig. 3C). The invasion capability of NCC-DMM1-C1 cells was higher than that of MG63 osteosarcoma cells (Figs. 3D and S2).

**Histological evaluation.** H&E staining of the sectioned tissue indicated that the spindle-shaped cells had a storiform pattern and proliferated densely (Fig. 4A). Immunohistochemistry revealed diffusely positive immunoreactivity for cytokeratin AE1/AE3 (Fig. 4B) and D2-40 (Fig. 4C) and negative immunoreactivity for WT1 (Fig. 4D). H&E staining of the NCC-DMM1-C1 cells indicated the same pattern as that of the tissue section with a storiform pattern and proliferated density and the composition of one type of cells, i.e. tumor cells with no other cells (Fig. 4E). Immunohistochemistry revealed diffusely
positive staining for cytokeratin AE1/AE3 (Fig. 4F), negativity for D2-40 (Fig. 4G), weakly but diffusely positive staining for HEG1 (Fig. 4H) and mostly positive staining for WT1 (Fig. 4I).

Kinase activity assay. Pearson's correlation scatter plot revealed high similarity of kinase activity between NCC-DMM1-C1 cells and tumor tissue ($r^2=0.86$; Fig. 4, Table SII). The 25 most highly phosphorylated peptides were identified in both NCC-DMM1-C1 cells and tumor tissue (Table SIII). These peptides included FES, Wee1, platelet-derived growth factor receptor (PDGFR)-β and Src (Table SIII).

Sensitivity to anti-cancer agents. The IC$_{50}$ values of 13 of the anti-cancer agents applied were calculated, including four anti-cancer agents used for DMM and nine drugs that had relatively high inhibitory effects on cancer cell proliferation at 10 µM (Fig. S3, Table SIV). The calculated IC$_{50}$ values are summarized in Table II and the representative growth curves of five anti-cancer agents with IC$_{50}$ values <100 nM (bortezomib, gemcitabine, homoharringtonine, fostamatinib and vinorelbine) are displayed in Fig. 5.

Discussion

DMM is an aggressive neoplasm with poor prognosis. The effects of surgery, chemotherapy and radiotherapy have remained to be fully defined. Thus, it is necessary to improve the current knowledge regarding the molecular mechanisms underlying disease progression of DMM and develop assay systems to evaluate novel anti-cancer agents. However, due to the rarity of the disease, only a small number of DMM cell lines have been reported and cell lines and xenografts for DMM are not readily available from public cell banks. The present study reported on the establishment a novel DMM cell line, designated as NCC-DMM1-C1 and determined its characteristics. Furthermore, anti-cancer drugs were screened using NCC-DMM1-C1 cells to evaluate whether the cell line may be used to screen drug candidates or not.

NCC-DMM1-C1 cells exhibited a spindle cell morphology and demonstrated constant growth and aggressive invasion, which reflect the characteristics of the original tumor. The spheroid formation capability contributes to the understanding of the behavior of DDM in a 3D environment. While NCC-DMM1-C1 cells had these features reflecting the original tumor, the immunohistochemical staining for D2-40 and WT1 was different between the tumor tissue and NCC-DMM1-C1 cells. NCC-DMM1-C1 cells stained negative for D2-40 expression and the tumor tissue demonstrated positive staining for D2-40 expression. Furthermore, NCC-DMM1-C1 cells demonstrated mostly positive WT1 expression and the tumor tissue demonstrated negative WT1 expression. Tumor tissue of DMM is composed of collagen tissue occupying >50% of the tumor and tumor cells (2) and NCC-DMM1-C1 was composed of tumor cells only. In addition, the expression of D2-40 and WT1 in NCC-DMM1-C1 cells was different from that of the tumor tissue due to heterogeneity. The cell components and the heterogeneity may have led to the distinct differences in D2-40 and WT1 expression.

The drug screening assay identified bortezomib, gemcitabine, homoharringtonine, fostamatinib and vinorelbine as anti-cancer agents with low IC$_{50}$ values in NCC-DMM1-C1. Gemcitabine is a DNA-damaging agent and has been investigated as a first-line single treatment in chemotherapy-naïve patients or as a second- or third-line combination therapy in patients with MPM (8,30,31). Vinorelbine is a microtubule-damaging agent and its efficacy has been reported in patients with progressive disease after treatment with pemetrexed-platinum chemotherapy (8,31,32). Bortezomib is a proteasome inhibitor and its cytotoxic effects were revealed in a preclinical study using six cell lines of MPM (33). However, two subsequent Phase II studies did not reach the desired endpoint in MPM (34,35). Further clinical trials are necessary to clarify the clinical utility of these three anti-cancer drugs for DMM. Homoharringtonine, a natural plant alkaloid extracted from Cephalotaxus harringtonia used

| CAS no. | Name of drug            | IC$_{50}$ (µM) |
|---------|-------------------------|----------------|
| 15663-27-1 | Cisplatin              | 5.38           |
| 95058-81-4  | Gemcitabine            | 0.053          |
| 901119-35-5 | Fostamatinib          | 0.097          |
| 443913-73-3 | Vandetanib            | 13.75          |
| 366017-09-6 | Mubritinib            | 0.43           |
| 179324-69-7 | Bortezomib            | 0.055          |
| 26833-87-4  | Homoharringtonine      | 0.072          |
| 65271-80-9  | Mitoxantrone           | 0.26           |
| 267243-28-7 | Canertinib            | 0.76           |
| 357166-30-4 | Pemetrexed            | 0.49           |
| 149647-78-9 | Vorinostat            | 1.26           |
| 71486-22-1  | Vinorelbine            | 0.04           |
| 252916-29-3 | Orantinib             | 30.06          |

Table II. Summary of IC$_{50}$ values.

IC$_{50}$, half-maximal inhibitory concentration.
in Chinese Traditional Medicine (36), has efficient inhibitory activity against myelocytic leukemia (37,38). Fostamatinib is a pro-drug inhibitor of spleen tyrosine kinase, which is a key mediator of Fc and B-cell receptor signaling in inflammatory cells and has efficacy in non-Hodgkin lymphoma and chronic lymphocytic leukemia (39). The treatment utility of homoharringtonine and fostamatinib in MPM has not been previously reported and it is worth elucidating their effects on MPM using NCC-DMM1-C1 cells.

Comprehensive kinase activity assays revealed that kinase activity was similar between NCC-DMM1-C1 cells and the original tumor tissue, suggesting that the NCC-DMM1-C1 cell line may be useful for examining the effects of kinase inhibitors in vitro. It was also determined that the kinases FES, Wee1, PDGFR-β and Src were highly activated in both NCC-DMM1-C1 cells and the tumor tissue. Tsao et al (40) demonstrated that Src was expressed and activated in MPM cell lines and 46 MPM tumor specimens. Additionally, the MPM cell lines were sensitive to an Src inhibitor, as determined using in vitro cytotoxicity assays. Wee1 is a tyrosine kinase that phosphorylates and inactivates cyclin-dependent kinase 1 and is involved in G2 checkpoint signaling (41). Xu et al (41) reported that a kinome-wide CRISPR/Cas9 knockout screen identified Wee1, the loss of function of which sensitizes cells to standard combination cisplatin and pemetrexed chemotherapy in MPM cell lines and PDXs. Several groups have reported that the PDGF/PDGFR pathway is involved in mesothelioma carcinogenesis (42,43). Melaiu et al (44) reported that, in MPM cells, PDGFRB silencing causes a decrease in the proliferation rate and reduces the colony formation capacity, which highlights the utility of PDGFR-β as a drug target.

In the present study, high FES activity was observed, which has not been previously reported in MPM, to the best of our knowledge. Experiments to evaluate kinase activity cannot be perfect and to compensate for the drawbacks of each experiment, it is required to use different types of experiments prior

Figure 6. Growth curves of NCC-DMM1-C1 cells treated with anti-cancer agents. The proliferation of NCC-DMM1-C1 cells treated with anti-cancer agents, which had marked anti-proliferative effects at a fixed concentration, was assessed using growth curves. (A) Bortezomib, (B) fostamatinib, (C) gemcitabine, (D) homoharringtonine and (E) vinorelbine. Data are presented as mean ± standard deviations based on two biological replicates.
to clinical trials. Therefore, kinase activities identified in the present study need to be validated using other methods such as western blot analysis. The present findings confirm that kinase activity may be used as a drug target and biomarker to assess the utility of kinase inhibitors.

In conclusion, in the present study, a novel cell line was established from tumor tissue of DMM, designated as NCC-DMM1-C1, and its characteristics and utility in high-throughput drug screening were demonstrated. However, the results of the present study were obtained from only a single cell line; further cell lines from different patients are required to generate reliable results. In addition, the present results regarding the candidate anti-cancer drugs should be validated in other patient-derived cancer models, including organoids and xenografts, prior to clinical trials. In addition, the candidate anti-cancer drugs were identified using the drug screening based on a single agent. Reflecting the practical clinical situation, drug screening based on several agents of the combination treatment will be performed in the future. Furthermore, evaluation of highly phosphorylated kinases and the relevant molecular pathway should be investigated by other proteomics techniques such as mass spectrometry. The present findings, along with future validations, will provide a deeper understanding of DMM, suggesting the possible utility of NCC-DMM1-C1 cells in drug development.

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Availability of data and materials

The cell line of the current study is available from the corresponding author on reasonable request. The datasets generated during the current study are available from the Gene Expression Omnibus (GEO) repository (45). The present data of the peptide microarray are accessible through GEO (series accession no. GSE185107; https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE185107) and the data of the SNP array are accessible through GEO (series accession no. GSE185549; https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE185549).

Authors’ contributions

RN and YYoshimatsu established the cell line. NM and YYa performed the pathological diagnoses and immunohistochemical staining. SW, Y Yoshida and AS prepared tumor tissues and obtained clinical information of the donor patient. RN, YYoshimatsu and TO performed all experiments, including authentication, drug screening assay and kinase assay. TK was responsible for study conception and design. RN, YYoshimatsu and TK prepared the manuscript, including the figures and tables. RN, Y Yoshimatsu and TK edited the manuscript. All authors read and approved the final manuscript prior to submission. RN and TK checked and confirm the authenticity of all the raw data.

Ethics approval and consent to participate

The use of clinical materials for this study was approved by the ethics committee of the National Cancer Center (approval no. 2015-108) and written informed consent was obtained from the patient. The informed consent was for collection of tissues from the patient for the present study and publication of data obtained using the tissues. All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration.

Patient consent for publication

Although detailed clinical data were not included in the present study, consent for publication was obtained from the patient included in the present study.

Competing interests

The authors declare that they have no competing interests.

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