Plectin promotes tumor formation by B16 mouse melanoma cells via regulation of Rous sarcoma oncogene activity

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Abstract

Background: Melanoma is a malignant tumor characterized by high proliferation and aggressive metastasis. To address the molecular mechanisms of the proto-oncogene, Rous sarcoma oncogene (Src), which is highly activated and promotes cell proliferation, migration, adhesion, and metastasis in melanoma. Plectin, a cytoskeletal protein, has recently been identified as a Src-binding protein that regulates Src activity in osteoclasts. Plectin is a candidate biomarker of certain tumors because of its high expression and the target of anti-tumor reagents such as ruthenium pyridinecarbothioamide. The molecular mechanisms by which plectin affects melanoma is still unclear. In this study, we examined the role of plectin in melanoma tumor formation.

Methods: We used CRISPR/Cas9 gene editing to knock-out plectin in B16 mouse melanoma cells. Protein levels of plectin and Src activity were examined by western blotting analysis. In vivo tumor formation was assessed by subcutaneous injection of B16 cells into nude mice and histological analysis performed after 2 weeks by Hematoxylin-Eosin (H&E) staining. Cell proliferation was evaluated by direct cell count, cell counting kit-8 assays, cyclin D1 mRNA expression and Ki-67 immunostaining. Cell aggregation and adhesion were examined by spheroid formation, dispase-based dissociation assay and cell adhesion assays.

Results: In in vivo tumor formation assays, depletion of plectin resulted in low-density tumors with large intercellular spaces. In vitro experiments revealed that plectin-deficient B16 cells exhibit reduced cell proliferation and reduced cell-to-cell adhesion. Since Src activity is reduced in plectin-deficient melanomas, we examined the relationship between plectin and Src signaling. Src overexpression in plectin knockout B16 cells rescued cell proliferation and improved cell-to-cell adhesion and cell to extracellular matrix adhesion.

Conclusion: These results suggest that plectin plays critical roles in tumor formation by promoting cell proliferation and cell-to-cell adhesion through Src signaling activity in melanoma cells.

Keywords: Melanoma, Src, Plectin, Tumor genesis, Cell adhesion

Background

Melanoma is an aggressive tumor derived from skin melanocytes [1, 2]. Patient prognosis is usually poor because the rapidly proliferating melanoma progresses from radial to vertical growth and frequently metastasizes [1, 2]. Recently developed therapeutic interventions target melanoma metastasis by blocking immune checkpoints using...
cytotoxic T-lymphocyte-associated antigen 4 (CTLA-4) or programmed death 1 (PD-1) inhibitors [3]. However, these reagents occasionally cause adverse immune-related events such as large intestinal inflammation, interstitial pneumonia, and type 1 diabetes [4]. Furthermore, some tumors fail to respond to therapy and acquired resistance during immune checkpoint therapy can occur. In addition, melanoma secretes immune suppressing agents such as cortisol and lymphotoxic intermediates [5]. Therefore, novel therapeutic approaches need to be developed.

Rous sarcoma oncogene (Src), a non-receptor tyrosine kinase, plays a physiological role in several cellular processes [6–9] including proliferation, adhesion, migration, and actin filament-organized kinase [7, 8]. Src signal transduction is initiated by the phosphorylation of substrates such as protein tyrosine kinase 2 beta (Pyk2) [8, 10] or actin-binding proteins. Src activation suppresses stress fiber formation, leading to the conversion of spindle-shaped cells into round cells with dot-like actin structures [11]. Src, which was originally identified as the first protooncogene, is involved in the malignancy, growth, and invasion of several tumors, including melanoma [12–17].

Plectin is a large protein consisting of more than 4000 amino acids. Plectin binds to actin, microtubules, and intermediate filaments, and regulates the cytoskeleton, cell shape, and chromosomal structure [18, 19]. Plectin is also involved in the organization of keratin intermediate filament networks within cells. Disruption of Plectin-linked cytoskeletal networks leads to Src and extracellular signal-regulated kinase (Erk) activation [20]. On the other hand, it has also been demonstrated that plectin is essential for Src activation during osteoclastogenesis [21, 22]. In osteoclasts, plectin acts as a scaffold for the recruitment of Src and Pyk2 substrates [22]. Plectin is a candidate biomarker for certain types of tumors because its expression in tumor specimens is higher than that in surrounding normal tissues [23–25]. Recently, a novel anti-tumor agent targeting plectin, ruthenium pyridinecarbothioamide, was shown to suppress tumor invasion of several tumors, including melanoma [12–17].

Western blot analysis
Cells were lysed with a 1% Triton X-containing lysis buffer, as described previously [22]. Lysates were boiled with sodium dodecyl sulfate (SDS) sample buffer for 5 min at 95°C [22]. Samples were separated on 7.5, 10, 12 and 15% SDS-PAGE gels and transferred to polyvinylidene difluoride (PVDF) membranes. After blocking with 5% bovine serum albumin (BSA), the membranes were incubated with primary antibodies. The antibodies used were anti-Plectin-1 (#12254), anti-Src pY416 (#2101), anti-Pyk2 (#3292), anti-Pyk2 pY402 (#3291), anti-p38 MAPK (#8690), anti-phospho-p38 MAPK (Thr180/Tyr182) (#9211), anti-p44/42 MAPK (Erk1/2) Antibody (#9102), anti-phospho-p44/42 MAPK (Erk1/2) (Thr202/Tyr204) (#4370) (Cell Signaling Technology, CST, Danvers, MA), anti-Src (ab-1) (Merck, Darmstadt, Germany), anti-DDDDK-tag (Flag-tag, Fla-1), anti-GAPDH mAb-horseradish peroxidase-conjugated (HRP)-DirectT (Medical & Biological Laboratories, MBL, Tokyo, Japan), and β-actin (A2228, Sigma Aldrich Chemicals, St. Louis, MO). The membranes were then incubated with HRP-conjugated anti-mouse or anti-rabbit secondary antibodies. Finally, the blots were imaged using a LAS4000 (Fujifilm Wako) with Immobilon ECL Ultra Western HRP Substrate (Merck).
Immunocytochemical analysis
A total of $1 \times 10^5$ cells were plated on a cover glass and cultured for 1 d. Cells were fixed with 3.7% formaldehyde and washed with phosphate buffered saline (PBS). Cells were permeabilized with PBS containing 0.2% Triton X100 and blocked with 5% BSA for 2h or 4h. Cells were then incubated with anti-vimentin (E-5, sc-373,717, Santa Cruz Biotechnology, TX), anti-Ki-67 rabbit monoclonal antibody (ab92742, Abcam Cambridge, UK), anti-Flag-tag in 5% BSA at 4°C overnight. Target proteins were visualized using Alexa 488-conjugated or Alexa 555-conjugated secondary antibodies (Thermo Fisher Scientific). The actin cytoskeleton was visualized using rhodamine phalloidin (Thermo Fisher Scientific). Nuclei were visualized using 4',6-diamidino-2-phenylindole (DAPI). Slides were imaged using a BZ-X810 microscope (Keyence, Osaka, Japan) and analyzed using ImageJ software (NIH). Cells with a maximum diameter more than 5 times greater than their minor diameter were defined as “elongated cells” [29].

Animals and tumor injection
Ten-week-old male BALB/cAJcl-nu/nu mice were purchased from CLEA Japan, Inc. (Tokyo, Japan). All mice were used in accordance with the guidelines of the Animal Care and Use Committee of Kyushu Dental University, based on the Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines. This study was approved by the Animal Care and Use Committee of Kyushu Dental University (Approval #19–020). The mice were housed under specific pathogen-free conditions. Anesthesia was performed by intraperitoneal injection of an anesthetic agent cocktail (0.3 mg/kg body weight (b.w.) medetomidine, 4.0 mg/kg b.w. midazolam, and 5.0 mg/kg b.w. butorphanol) [30]. Anesthetized mice were injected subcutaneously with $1 \times 10^5$ cells in a 100μL volume [28]. After 2 weeks, mice were anesthetized with an anesthetic agent cocktail and perfused with 4% paraformaldehyde in PBS [31]. Tumors were then dissected, measured with calipers, and weighed using an electronic balance scale. Volumes were calculated using the formula $\text{volume} = \frac{\text{width} \times \text{length} \times 0.52}{2}$, as previously described [32, 33]. Sections of 10μm were prepared using a cryostat. The sections were stained with hematoxylin and eosin (H&E) and imaged using a BZ-X810 microscope. Cell numbers and tumor areas were quantified with ImageJ software. Cell density was calculated as the number of the cells divided by tumor area. The area between cells was determined by subtracting total cell area from total tumor area on the histological section. Sections were also stained with 0.1% azocarmine G solution, aniline blue and orange G solution (Azan staining) and imaged using a BZ-X810 microscope.

Cell proliferation and viability assays
Cell proliferation was assessed with the Cell Counting Kit-8 (CCK-8, Dojindo, Kumamoto, Japan), according to the manufacturer’s protocol [28]. Cells were plated at a density of 5000 cells/well in 96-well plates. For direct viable cell counts by the trypan blue pigment exclusion method, cells were plated at a density of 2.5 $\times 10^4$ cells/well in 24-well plates [34]. Three days later, cells were detached with trypsin (Fujifilm Wako) mixed with Trypan Blue counted with a TC20 Automated Cell Counter (Bio-Rad).

Reverse transcription and quantitative PCR (qPCR)
Total RNA was isolated using the FastGene™ RNA Basic Kit (Nippon Genetics, Tokyo, Japan) and reverse-transcribed into cDNA using the High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific) [35]. Real time quantitative (q)PCR was performed using a Quantstudio 3 system (Thermo Fisher Scientific) with specific primers for murine Cyclin D1 ($\text{Ccnd1}$); (forward, 5′- gtggtacgaccagaggcatac - 3′; 5′- taacctcagaagggcttcaa - 3′), human Cyclin D1 ($\text{CCND1}$); (forward, 5′- tacaccccgacattccgt - 3′; 5′- tcggattttgggaaggaattg - 3′), murine β-actin ($\text{Actb}$); (forward, 5′- aagggccaaccgtgaaaa - 3′; reverse, 5′- gtggtacgaccagaggcatac - 3′), and human β-actin ($\text{ACTB}$); (forward, 5′- caccattggaatcagcgcgt - 3′; reverse, 5′- agtcttttggtgtgggtctacct - 3′) [36, 37]. Gene expression levels were calculated using the ΔΔCT method and normalized to that of β-actin.

Spheroid formation
Cells were seeded at a density of 5000 cells in 15μL of media on the inner side of a dish lid. The lid was then placed upside down on a PBS-filled plate. Cells were

[See figure on next page.]

**Fig. 1** Src signaling is impaired in plectin deficient cells. A. PLECTIN levels in control B16 and 2 clones of plectin deficient B16 cells (PKO1 and PKO2) were determined by western blotting analysis. B. SRC was immunoprecipitated with an anti-Src antibody (IP:SRC) and interaction with PLECTIN was detected by western blotting analysis. C. Expression and phosphorylation of PYK2, or GAPDH was determined by western blotting assay 24 h after plating. SRC was immunoprecipitated with an anti-Src antibody (IP:SRC) and interaction with PYK2 was detected by western blotting analysis. D. SRC was immunoprecipitated with an anti-Src antibody (IP:SRC) and interaction with PYK2 was detected by western blotting analysis. E. Cells were stained with Rhodamine-phalloidin and vimentin/anti-mouse Alexa fluor 488. Long actin fibers are indicated by arrows. Cell shapes are outlined with a dotted line in the bright-field image. Scale bar = 50 μm. F. Quantification of the number of elongated cells in samples shown in (E) (n = 5). *; p < 0.05 vs control.
Fig. 1 (See legend on previous page.)
cultured for 7 d at 37 °C, with 5% CO₂. Images of spheroids were taken with a Leica EZ4 HD microscope (Leica, Germany) and analyzed using ImageJ software. The volumes were calculated using the formula width² × length × 0.52, similar to that used for the tumor.

**Dispase-based dissociation assay**

Cells were seeded at a density of 1.0 × 10⁶ cells/well in 6-well plates. Twenty-four hours after confluency, the cells were incubated with 1440 PU/ml dispase II (Fujifilm Wako) solution for 40 min at 37 °C to detach the monolayer from the well bottom. The dispase solution was then carefully transferred into a new 6-well plate using a 1 ml pipette. Next, the cell monolayer was mechanically stressed by pipetting up and down 7 times [38, 39].

**Cell adhesion assay**

96-well plates were coated by overnight incubation with 10 μg/mL fibronectin at 4 °C. Wells were then blocked with 2% BSA for 2 h at room temperature (RT). Cells were plated at a density of 3 × 10⁴ cells per well in serum-free medium and allowed to attach for 2 h at 37 °C with 5% CO₂. Non-adherent cells were washed away with serum-free medium. The remaining adherent cells were fixed using 1% glutaraldehyde/PBS for 10 min and stained with crystal violet in 2% methanol/PBS for 20 min at RT with shaking. Crystal violet was solubilized with 1% SDS and quantified by absorbance measurement at 595 nm using a microplate reader (Bio-Rad, Hercules, CA). Adherent cells were counted using ImageJ software.

**Src overexpression**

Src mutating tyrosine 527 to phenylalanine (constitutively activated Src) with a Flag-tag was previously described [40, 41]. Constitutively activated Src was transfected B16 cells with Lipofectamine 2000 (Thermo Fisher Scientific) and cultured for 1 d. Then, cells were seeded and performed each experiment.

**Data analysis and statistics**

Statistical significance of differences between groups was analyzed using one-way ANOVA followed by a Tukey-Kramer post-hoc test. Statistical significance was set at p < 0.05. Data are expressed as mean ± standard deviation of the mean (n = number of samples). All experiments were independently performed at least twice and similar results were obtained.

**Microarray data mining**

Dataset GDS1375 from the NCBI Gene Expression Omnibus (GEO) was analyzed for plectin and Src gene expression in melanoma patient samples, as previously described [28, 42]. Means were compared using Kruskal-Wallis followed by Dunn’s post hoc test.

**Results**

**Src signaling is impaired in plectin-deficient melanoma cells**

To examine the role of plectin in melanoma, we used CRISPR/Cas9 gene editing to delete plectin in murine B16 melanoma cells (PKO cells). Western blotting confirmed
successful deletion of plectin (Fig. 1A). We have previously shown that in osteoclast cells plectin interacts with Src and facilitates Src activation [22]. We next examined whether plectin also interacts with src in melanoma cells. By co-immunoprecipitation, we confirmed that plectin and Src are bound in control wildtype B16 cells but not in plectin knockout cells (Fig. 1B). Furthermore, Src activation, determined by the level of phospho-Pyk2 (Tyr402), was decreased in plectin-deficient cells (Fig. 1C). Moreover, binding of Src to Pyk2 was also disrupted in PKO cells (Fig. 1D). Spindle-shaped cells with thick, long actin fibers and abnormal cell spreading are characteristic of disturbed Src signaling [29]. Consistent with abnormal Src activity in PKO cells, we observed that PKO cells were elongated and less round than control cells (Fig. 1E, F). Vimentin, an intermediate fiber regulated by plectin, forms networks around the nuclei in control B16 cells. In contrast, vimentin localization was diffused in plectin deficient cells (Fig. 1E and F).

To determine whether these observations were cell-type specific, we also generated plectin deficient human HMV-II human melanoma cells by shRNA mediated knockdown (Fig. S1A). Similar to our observations in B16 cells, plectin knockdown reduced Src signaling (Fig. S1B) and produced cells with abnormal elongated morphology (Figs. S1C and S1D). Thus, the depletion of plectin impairs Src signaling in melanoma cells.

**Plectin regulates cell proliferation and adhesion**

To determine the function of plectin in tumor formation in vivo, control or PKO cells were subcutaneously injected into the backs of nude mice (Fig. 2A). There were no significant differences in tumor weight or volume between control and PKO cells (Figs. S2A and S2B). However, the density of PKO tumors was lower than that of the control tumors (Fig. 2B). Histological analysis showed that the area between cells and subsequently
the cell density within the PKO tumor was greater than in control tumors (Fig. 2C and D). Azan staining, which usually stains collagen and extracellular matrix blue, showed that the intercellular spaces did not contain any collagenous extracellular matrix (Fig. S2C). No other extracellular elements other than some red blood cells were detected in the tumors. Thus, intercellular spaces may be filled with interstitial fluid similar to edemas or the spontaneous skin blistering that occurs in patients with plectin mutations who develop epidermolysis bullosa simplex [43–45]. Taken together, these in vivo data suggest that plectin may affect cell number and intercellular connections.

Next, the effects of plectin on cell proliferation was examined in vitro. Direct cell counting as well as a CCK-8 assay showed that the number of viable cells were significantly reduced after plectin depletion in B16 cells (Fig. 3A and B) or in HMV-II cells (Figs. S3A and S3B). mRNA levels of Cyclin D1, a regulator of cell cycle progression, were also decreased in plectin-deficient cells (Figs. 3C and S3C). Immunofluorescence analysis showed that the number of Ki-67 positive cells, a marker of cell proliferation, was also decreased in plectin-deficient cells (Figs. 3D and S3D).

To determine the role of plectin in the formation of intercellular cell connections we next performed a spheroid formation assay using PKO cells. Similar to our

Fig. 4 Cell adhesion is impaired in plectin knockout cells. A Cells were cultured at a density of $5 \times 10^3$ cells per 15 μl drop on the inner side of a dish lid to form spheroids for 7 d. Images were acquired with a stereoscopic microscope and volumes quantified with ImageJ. Scale bar = 250 μm ($n = 5$). B Confluent cells were detached with dispase and mechanically fragmented by pipetting. The left panel shows an image of the fragmented cell monolayer and the right panel shows quantification of the number of cell fragments. Scale bar = 20 mm ($n = 6$). C Cells were plated on a fibronectin-coated dish and counted the number of adherent cells after 2 h. The adherent cells were stained with crystal violet (left 3 panels). After taken photo, the adherent cells were lysed and measured the absorbance shown at graph (right panel) ($n = 6$). *; $p < 0.05$ vs control.
observation in tumor formation in vivo, plectin-deficient cells formed larger spheroids than the control cells (Figs. 4A and S4A). A dispase-based dissociation assay showed that PKO cell monolayers were more sensitive to mechanical fragmentation indicative of weakened cell-to-cell adhesion (Fig. 4B). In addition, cell attachment to fibronectin was reduced in plectin-deficient cells compared to that in control cells in an adhesion assay (Figs. 4C and S4B). This suggests that plectin plays an important role in both cell proliferation and in the formation of cell-to-cell or cell-to-matrix adhesion of melanoma cells.

Overexpression of Src rescues the plectin knockout phenotype

Given that Src activity is reduced in PKO cells, we next sought to determine whether plectin affects melanoma cell behavior via Src signaling. We therefore examined the effect of overexpressing a constitutively activated Src construct in PKO cells. Src overexpression was able to increase p-Pyk2 levels in wildtype and PKO cells (Fig. 5A). Overexpression of Src in PKO cells reduced the abnormally high percentage of elongated cells occurring in PKO cell cultures (Fig. 5B and C). In addition, Src
overexpression restored the vimentin network in PKO cells to a peri-nuclear localization (Fig. S5). Overexpression of Src also prevented the decreased cell proliferation in PKO cells as determined by the number of viable cells and Ki-67 positive cells (Fig. 6A, B and C). In the spheroid formation assay, Src overexpression in PKO cells led to the formation of small, tightly packed spheroids similar to control cells instead of the large loose phenotype observed in PKO cells without Src overexpression (Fig. 6D). In the dispase-dissociation assay, Src overexpression was able to prevent the fragmentation of PKO cells suggesting that the loss of cell-cell adhesion in PKO cells is Src-dependent (Fig. 6E). Furthermore, cell adhesion to fibronectin was also improved when Src was overexpressed in PKO cells (Fig. 6F).

**Discussion**

In this study, we demonstrated that plectin was essential for Src activation in melanoma cells. This is similar to our previous observations in osteoclasts [22]. In plectin-deficient melanomas, proliferation and adhesion are impaired. Overexpression of constitutively activated Src partially rescued the defective proliferation and adhesions suggesting that plectin regulates the proliferation and adhesion of melanoma cells through Src-dependent and Src-independent mechanisms.

The relationship between plectin and Src activation appears to be complex and cell-type dependent. For example, although plectin negatively regulates Src in keratinocytes [20], plectin plays an essential role in activating Src signaling to organize the cytoskeleton in osteoclasts [22]. In the focal adhesion complex, plectin participates in mechanosensing to activate Src signaling downstream of the vimentin intermediate filament network [46]. Our data confirmed that plectin interacts with Src in a manner that promotes cell adhesion similar to our previous observations in osteoclasts.

In tumor cell proliferation, Src mediates growth factor signaling by activating Ras GTPase and mitogen-activated protein kinases (MAPKs) [8]. Src and cyclin-dependent kinase 1 (Cdk1) have common substrates related to cell proliferation [47, 48]. Plectin is known to be involved in cell proliferation independently of Src. Cdk1 interacts with plectin and regulates the rearrangement of the cytoskeleton during interphase to promote mitosis [19]. Plectin directly binds and organizes actin, microtubules, and intermediate filament networks at each step of the cell cycle [19]. Therefore, cell division is disrupted in the absence of plectin.

Src is a major regulator of both cell adhesion and aggregation. Src signaling promotes focal adhesion formation via Pyk2 and Rho GTPase activation for adhesion to the extracellular matrix [49–51]. Plectin has been shown to directly organize vimentin networks, cell polarization and focal adhesions for cell adhesion in fibroblasts [46, 52]. We observed disturbed vimentin network organization in plectin deficient cells that could be restored by Src activation. Src directly phosphorylates and organizes vimentin around the nuclei [53]. Plectin may thus likewise be critical for vimentin organization through its Src dependent functions. Src-independent regulation of vimentin or actin networks by plectin can however not be excluded. Src and Pyk2 signaling also regulates cell-to-cell contacts by phosphorylating cadherins during desmosome and tight junction formation [54, 55]. Similarly, plectin also directly organizes the keratin network or vimentin network to maintain tight junction and adherence junction [39, 56–58]. Plectin's role in the cytoskeletal integrity and cell-cell contacts is evident in human patients with plectin mutations where disruption of keratin networks within the skin leads to extensive blistering or epidermolysis bullosa simplex [43–45].

P38 MAPK and extracellular signal-regulated kinase (Erk) 1/2 have been shown to be the mediators of plectin function in cytoskeleton organization [20, 46]. In our study we found that phosphorylation of p38 and Erk was enhanced in PKO cells (Fig. S6) similar to the plectin-deficient keratinocytes [20]. In melanoma, Erk and p38 are mutually activated and induce cell migration [59]. Erk activation also causes ECM degradation and decreased Cadherin expression [60]. Elevated Erk activation could contribute to the weakened cell adhesion in plectin deficient B16 cells. In keratinocytes, Src increases Erk phosphorylation, while in melanomas, Erk phosphorylation is decreased [22].
Fig. 6 (See legend on previous page.)
is increased despite low Src activity. Further studies are needed to reveal the extent and mechanisms of MAPK regulation by plectin in the cytoskeleton.

Analysis of a GEO dataset showed that plectin-and Src-related genes are highly expressed in human primary melanoma in patients compared to human melanocytes (Fig. S7). Based on our experimental data, plectin may be required for malignancy and primary tumor formation in melanoma. During metastasis, plectin may play a positive role in the invasion and migration into the vascular system. Src signaling and plectin are essential for organizing actin structures in invadosomes [61, 62]. Plectin may also be necessary for cell anchoring to the target tissue during metastasis or separation of cells from the primary tumor mass. Therefore, the role of plectin in melanoma progression may differ at each phase.

An anti-tumor agent targeting plectin, ruthenium pyridinacarbothioamide, has been recently developed for the treatment of melanoma [26, 27]. Our present study strongly supports a role for plectin in melanoma during early stage events, such as primary tumor formation. However, the effects of plectin inhibition during the late stage of melanoma remain unclear. Furthermore, data from the Human Protein Atlas (https://www.proteinatlas.org/ENSG00000178209-PLEC/pathology/melanoma) and The Cancer Genome Atlas (TCGA) showed that melanoma patients with high plectin expression survived significantly longer than those with low expression levels. This suggests that the suppression of plectin may promote melanoma after the formation of the primary tumor because the primary tumor had already been formed upon diagnosis. Further studies, especially on invasion and metastasis, are required to clarify the role of plectin in melanoma progression.

Conclusions
Plectin plays a key role in robust tumor formation in melanoma by regulating cell proliferation and aggregation through Src signaling.

Supplementary Information
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Authors’ contributions
T. M. and S. K. designed the study. K. M. and T. M. performed the experiments. A.G. and M.N. analyzed tumor tissue. K. M. T. M., S. K. wrote the first draft. W. A. provided feedback. M.K., Y.T-S., T. Y., H.H., I. Y. and S. K. performed literature review. The author(s) read and approved the final manuscript.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request. The dataset in Fig. S5 is available from GSE1375 dataset (GEO accession number 1375) of NCBI Gene Expression Omnibus (GEO) (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE1375).

Declarations
Ethics approval and consent to participate
All mice were used in accordance with the guidelines of the Animal Care and Use Committee of Kyushu Dental University, based on the Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines. This study was approved by the Animal Care and Use Committee of Kyushu Dental University (Approval #19-020).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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