Filamin B Enhances the Invasiveness of Cancer Cells into 3D Collagen Matrices

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ABSTRACT. Numerous types of cancer cells migrate into extracellular tissues. This phenomenon is termed invasion, and is associated with poor prognosis in cancer patients. In this study, we demonstrated that filamin B (FLNb), an actin-binding protein, is highly expressed in cancer cell lines that exhibit high invasiveness, with a spindle morphology, into 3D collagen matrices. In addition, we determined that knockdown of FLNb in invasive cancer cells converts cell morphology from spindle-shaped, which is associated with high invasiveness, to round-shaped with low invasiveness. Furthermore, di-phosphorylation of myosin regulatory light chain (MRLC) and phosphorylation of focal adhesion kinase (FAK) are inhibited in FLNb-knockdown cancer cells. These results suggest that FLNb enhances invasion of cancer cells through phosphorylation of MRLC and FAK. Therefore, FLNb may be a new therapeutic target for invasive cancers.

Key words: cancer cells, filamin B, focal adhesion kinase, invasion, myosin regulatory light chain

Introduction

Some cancer cells are able to migrate into surrounding tissues. This phenomenon is called invasion and is a critical characteristic of cancer cells that are associated with poor prognosis in cancer patients. To migrate into tissues, cancer cells must reorganize actin filaments, a process regulated by actin binding proteins (Yamazaki et al., 2005).

Filamins are a group of actin binding proteins that consist of three isoforms: filamin A (FLNa), filamin B (FLNb), and filamin C (FLNc). They have some similar functions, including regulating the formation of the three-dimensional actin cytoskeleton network (Feng and Walsh, 2004; Stossel et al., 2001; Zhou et al., 2010). However, isoform-specific functions for filamin have been identified. For instance, FLNa plays a key regulatory role in apical extrusion (Kajita et al., 2014); FLNb contributes to angiogenesis by regulating the migration of endothelial cells (del Valle-Perez et al., 2010); FLNc is specifically expressed in cardiomyocytes and skeletal myocytes and promotes structural integrity (Fujita et al., 2012). In addition, FLNa is reported to enhance the invasive ability of human melanoma cells (Cunningham et al., 1992; Zhang et al., 2014). On the other hand, FLNa has been shown to prevent invasion and metastatic potential in human breast cancer cells (Xu et al., 2010). Though the role of FLNa in regulating invasion has been characterized, the function of FLNb in this process is not well understood.

Contractile force is essential for cell migration. Cellular force is generated through actomyosin contractility, which is regulated by di-phosphorylation of myosin regulatory light chain (MRLC) (Mizutani et al., 2006). We have previously reported that the invasive ability of A549 lung cancer cells is regulated by di-phosphorylation of MRLC (Ishihara et al., 2013). In addition, FLNa promotes the phosphorylation of MRLC in human breast epithelial cells (Gehler et al., 2009). However, the role of FLNb in regulating MRLC phosphorylation is not well studied.

Focal adhesions are adhesive structures that connect cells to their underlying substrate and consist of a large variety of proteins. Turnover of focal adhesions is important for cell migration (Bugide et al., 2014; Ezratty et al., 2005). Focal adhesion kinase (FAK) regulates the turnover of focal adhesions and is activated via phosphorylation at Y397, which subsequently promotes focal adhesion disassembly (Hamadi et al., 2005). FAK also contributes to invasion of cancer cells (Sulzmaier et al., 2014). Furthermore, a previous study indicated that FLNa inhibits turnover of focal adhesions (Xu et al., 2010). However, the role of FLNb in regulating focal adhesion turnover is not well understood.
In this study, we demonstrate that FLNb is highly expressed in invasive cancer cells and contributes to their invasive ability. Furthermore, our data indicate that FLNb is critical for di-phosphorylation of MRLC and phosphorylation of FAK in invasive cancer cells.

Materials and Methods

Cell culture

A549 human lung adenocarcinoma cells and HT1080 human fibrosarcoma cells were purchased from American Type Culture Collection (ATCC; Manassas VA) and Riken Cell Bank (Tsukuba, Japan), respectively. Subclonal A549 cells (P-3 cells) and irradiation-tolerant P-3 cells (P-3IR cells) were generated as we previously reported (Ishihara et al., 2010). Cells were cultured using Dulbecco’s Modified Eagle’s Medium (DMEM; Sigma, St. Louis, MO) supplemented with 10% fetal bovine serum (Bist-Tec; Equitech Bio Inc., Kerrville, TX, Biowest; Biowest, Nuaille, France) and 1% antibiotic/antimycotic solution (Invitrogen, Carlsbad, CA). Cells were kept in a humidified incubator at 37°C with 5% CO₂.

Reagents and antibodies

Cellmatrix Type I-P (1.6 mg/mL; Nitta Gelatin Inc., Osaka, Japan) was used for preparing collagen gels. Primary antibodies used for the detection of filamins were anti-FLNa (Millipore, Billerica, MA, MAB1680) and anti-FLNb (Abcam, Cambridge, UK, ab97457) antibodies. Primary antibodies for western blotting included anti-di-phosphorylated myosin regulatory light chain (pp-MRLC, Cell Signaling Technology, Beverly, MA, #3674), anti-total amount of myosin regulatory light chain (total-MRLC, Cell Signaling Technology, #3672), anti-total amount of focal adhesion kinase (total-FAK, BD biosciences, San Jose, CA, 610087), anti-phosphorylated focal adhesion kinase at Y397 residue (FAK (pY397), BD biosciences, 611806), and anti-GAPDH antibody (Ambion, Foster City, CA, AM4300). Secondary antibodies of immunofluorescence staining included Alexa-Fluor 546 goat anti-mouse antibody (Molecular probes, Carlsbad, CA, A-11003) and Alexa-Fluor 546 goat anti-rabbit antibody (Molecular probes, A-11010). Secondary antibodies for western blot analysis included HRP anti-mouse antibody (Bio-Rad, Hercules, CA, 70-6516) and HRP anti-rabbit antibody (Cell Signaling Technology, #7074). Alexa-Fluor 488 phallolidin (Molecular probes, A-12379) was used for staining F-actin.

siRNA transfection

We generated siRNA duplexes using in vitro Transcription T7 kit (Takara Bio Inc., Otsu, Japan). The siRNA against FLNb targeted the sequence 5'-ATTATCTCTGACCTGTAGG-3' (antisense sequence). FLNb siRNA or scrambled siRNA (as a negative control) were transfected into cells using Lipofectamine RNAiMAX Reagent (Invitrogen). Knockdown of FLNb was confirmed three days after transfection using RT-PCR and/or western blotting.

Collagen gel overlay method and time-lapse imaging

The collagen gel solution (150 μL) was added to a hand-made glass dish with a radius of 8 mm, the bottom of which was made of commercial cover glass (Matsunami Glass Ind., Ltd., Osaka, Japan). Either P-3IR or HT1080 cells (3.0×10⁵ cells) were seeded on the gel. Then, 24 h after seeding, 125 μL of collagen solution was poured on top of the cells and incubated for 1 h in a humidified incubator at 37°C with 5% CO₂. The pouring of collagen provided the cells with a 3D culture environment. The dish was filled with growth medium and sealed with silicone grease. Time-lapse microscopy was performed by using a phase-contrast microscope (TE300, Nikon, Tokyo, Japan) with a ×10 objective. During the experiment, the dish was kept at 37°C in an acrylic resin box. Time-lapse images were captured every 5 min using Image-Pro Software (Media Cybernetics Inc., Silver Spring, MD). The manual tracking plugin in Image J software (National Institutes of Health, Bethesda, MD) was used to track cell migration and analyze migration velocity. Student’s t-test was used for statistical analysis.

Immunofluorescence staining

Cells were cultured for 24 h in a 3D collagen gel using the above described collagen gel overlay method. Cells were fixed using 4% formaldehyde in phosphate-buffered saline (PBS), permeabilized with 0.5% Triton X-100 in PBS, and blocked using 0.5% of bovine serum albumin (Sigma) in PBS. The cells were incubated with primary antibodies (FLNa, 1:2000; FLNb, 1:250) in PBS for 1 h at room temperature. After three washes with PBS, the cells were incubated with secondary antibodies (Alexa-Fluor 546 goat anti-mouse antibody, 1:250, for FLNa staining; Alexa-Fluor 546 goat anti-rabbit antibody, 1:250, for FLNb staining) for 1 h at room temperature. After three washes with PBS, anti-bleaching reagent was added to the cells. Fluorescence images were obtained using confocal laser scanning microscopy (Cl confocal system; Nikon) with a ×60 objective.

Western blotting

P-3 or P-3IR cells (1.0×10⁵) were seeded onto plastic dishes (Corning, Tewksbury, MA) with a radius of 17.5 mm that were previously filled with 450 μL of collagen gel. Twenty-four hours after seeding, 350 μL of collagen gel solution was poured onto the top of the cells and incubated in a humidified incubator at 37°C with 5% CO₂. HT1080 cells (1.0×10⁶) were seeded onto a plastic dish with a radius of 17.5 mm previously filled with 500 μL of collagen gel. Twenty-four hours after seeding, 450 μL of collagen gel solution was poured onto the cells and incubated in a humidified incubator at 37°C with 5% CO₂. Then, the cells were cultured for 24 h. Cell extractions were prepared as we previously described (Ishihara et al., 2014). Cells were fixed with 10% trichloroacetic
acid in PBS for 3 min on ice and then treated with 0.1% collagenase-L (Nitta Gelatin) in PBS for 90 min at 37°C with 5% CO₂ to digest the collagen gel. Then, cells were isolated from the collagen solution using centrifugation. Cells were lysed using SDS sample buffer (0.25 M Tris-HCl, 5% of dithiothreitol, 2.3% of sodium dodecyl sulfate, 10% of glycerol, 0.01% of bromophenol blue, pH=6.8). Lysates were sonicated and boiled at 95°C for 5 min. The lysates were separated using SDS-PAGE (20 mA per gel, 70 min). Further, 8.0% polyacrylamide gels were used in the detection of FLNa, FLNb, FAK (pY397), total-FAK, and GAPDH, whereas 12.5% polyacrylamide gels were used in the detection of pp-MRLC, total-MRLC, and GAPDH. After SDS-PAGE, transfer to PDVF membranes (Millipore) was performed (92 mA per gel; 60 min for the detection of pp-MRLC, total-MRLC, and GAPDH; 150 min for the detection of FLNa, FLNb, FAK (pY397), total-FAK, and GAPDH). Next, membranes were blocked using 0.5% skim milk in TBS-Tween (for the detection of FAK (pY397), pp-MRLC, and GAPDH) or 5% skim milk in TBS-Tween (for the detection of FLNa, FLNb, total-FAK, total-MRLC, and GAPDH) for 60 min. The membranes were then incubated with primary antibodies (GAPDH, 1:50000–1:100000; FLNa, 1:500; FLNb, 1:200–1:500; FAK (pY397), 1:100–1:300; total-FAK, 1:100–1:300; pp-MRLC, 1:300; total-MRLC, 1:300) in TBS-Tween at 4°C overnight. After three washes with TBS-Tween, the membranes were incubated with secondary antibodies (HRP anti-mouse antibody, 1:15000–1:100000, for the detection of FLNa, FAK (pY397), total-FAK, and GAPDH; HRP anti-rabbit antibody, 1:10000–1:50000, for the detection of FLNb, pp-MRLC, and total-MRLC) in TBS-Tween for 1 h at room temperature. After three washes with TBS-Tween, signals were detected with Immobilon Western Chemiluminescent HRP substrate (Millipore). The band intensity was quantified using Image J software (National Institutes of Health). The levels of phosphorylated MRLC and FAK were determined using the ratio of the band intensity of pp-MRLC/total-MRLC and FAK (pY397)/total-FAK, respectively. Statistical analysis was performed using a Student’s t-test.

**RT-PCR**

A plastic dish with a radius of 17.5 mm filled with 450 μL of a collagen gel solution was prepared. P-3 or P-3IR cells (1.0×10⁵) were seeded onto the dishes. Twenty-four hours after seeding, 350 μL of collagen gel solution was poured onto the cells and incubated in a humidified incubator at 37°C with 5% CO₂ for 48 h; we were unable to obtain sufficient number of HT1080 cells in the collagen gel overlay for this experiment owing to low growth of the cells. RNA was isolated using TriPure Isolation reagent (Roche, Basel, Switzerland). Reverse transcription reaction was performed using a ReverTra Ace qPCR RT kit (TaKaRa Bio Inc.). Then, polymerase chain reaction was performed using Taq DNA polymerase with ThermoPol buffer (TOYOBO, Osaka, Japan). We used following primers in the reactions: GAPDH, 5’- ACCACAGTCCATGCATCAC-3’ (upper) and 5’-TCCACCACCTGTGCTGA-3’ (lower); FLNb, 5’-GGTGCTCAGGCGAAGCGCA-3’ (upper) and 5’-CAGGGCTCCCCAGGGTCTTG-3’ (lower). The products were detected using electrophoresis with a 2% agarose gel.

**Results**

**FLNb expression is associated with invasiveness in cancer cells**

To investigate the relationship between filamin expression and invasiveness, we examined the protein expression levels of FLNa and FLNb in cancer cell lines with different invasive capacities. For these experiments, we utilized P-3 and P-3IR cells (Ishihara et al., 2010). Using a collagen gel overlay culture-method to analyze invasiveness in vitro, we previously reported that P-3IR cells, which have a spindle-shaped morphology, are highly invasive than P-3 cells (Ishihara et al., 2010). Therefore, we compared the protein expression level of filamins in P-3 cells with P-3IR cells that were grown in collagen gel overlay culture. Using immunofluorescence staining and western blot analysis, we determined that the expression of FLNa was not significantly different between P-3 and P-3IR cells, whereas the expression of FLNb was higher in P-3IR cells than in P-3 cells (Fig. 1A, B and Fig. S1). Therefore, FLNb protein expression level is associated with a high invasive ability in human lung adenocarcinoma cells.

**FLNb is critical for invasiveness in irradiation-tolerant lung cancer cells**

Next, we investigated whether FLNb regulates invasiveness in P-3IR cells. We transfected P-3IR cells with scrambled siRNA, as a negative control, or siRNA targeting FLNb. Knockdown of FLNb was confirmed by RT-PCR and western blot analysis (Fig. 2A, B). We analyzed invasiveness of the cells in vitro by observation of morphology and migration using a collagen gel overlay culture-method. P-3IR cells transfected with scrambled siRNA invaded into the 3D collagen gel by generating protrusions (Fig. 2C). In this condition, cells migration was restricted to the horizontal direction (Movie S1). Therefore, we were able to analyze cell migration velocity using phase-contrast images. In contrast, FLNb knockdown cells exhibited low invasive ability and had a round-shaped morphology (Fig. 2C). Migration velocity in P-3IR cells transfected with scrambled siRNA was significantly greater than cells expressing siRNA targeting FLNb (Fig. 2D). These data indicate that FLNb contributes to the invasiveness of P-3IR cells into 3D collagen matrices.
FLNb plays an important role in the invasiveness of fibrosarcoma cells

We examined the role of FLNb in regulating invasiveness in HT1080 human fibrosarcoma cells. HT1080 cells are known to invade into 3D collagen matrices while adopting spindle-shaped morphologies (Wolf et al., 2003), similarly to P-3IR cells. Knockdown of FLNb in HT1080 cells was confirmed using RT-PCR and western blot analysis (Fig. 3A, B). As seen with P-3IR cells, HT080 cells transfected with scrambled siRNA adopted an invasive phenotype, generating spindle-shaped protrusions into the 3D collagen gel (Fig. 3C). For HT1080 cells, migration was restricted to the horizontal direction for the collagen gel overlay (Movie S2). However, FLNb knockdown cells displayed a rounded-shape morphology and had low invasive activity into the 3D collagen gel (Fig. 3C). The migration velocity of HT1080 cells transfected with scrambled siRNA was sig-
significantly faster than cells expressing siRNA targeting FLNb (Fig. 3D). Thus, FLNb plays an important role for the invasive phenotype of HT1080 cells, in addition to P-3IR cells.

**Discussion**

Our data indicate that FLNb contributes to invasiveness in cancer cells. We previously reported that P-3IR cells displayed an invasive phenotype with a spindle-shaped morphology, while P-3 cells exhibited low invasiveness and a round-shaped morphology in 3D collagen matrices (Ishihara et al., 2010, 2013). In this study, we demonstrated that P-3IR cells have a higher protein expression level of FLNb than P-3 cells. Furthermore, knockdown of FLNb induced a low-invasive phenotype and round morphology in P-3IR cells. Because P-3IR cells are radiation-tolerant cancer cells, we also investigated the contribution of FLNb to the invasiveness of non-irradiated HT1080 cells, which also indicates high invasiveness in a collagen gel. FLNb knockdown in HT1080 and P-3IR cells showed low invasiveness. Taken together, these data suggest that FLNb promotes invasive capability and a spindle-shaped morphology in cancer cells.

We also demonstrated that FLNb is a critical regulator of di-phosphorylation of MRLC and phosphorylation of Y397 in FAK. Di-phosphorylation of MRLC is important for generating contractile actomyosin forces (Mizutani et al., 2006). Furthermore, these contractile forces are crucial for the invasive ability of cancer cells (Friedl and Wolf, 2010). Thus, FLNb may enhance contrac-

**FLNb regulates di-phosphorylation of MRLC and phosphorylation of FAK**

We previously reported that di-phosphorylation of MRLC regulates invasiveness in P-3IR cells (Ishihara et al., 2013). Therefore, we assessed whether FLNb regulates di-phosphorylation of MRLC in invasive cancer cells. Western blot analysis indicated that knockdown of FLNb in HT1080 cells reduced the level of di-phosphorylated MRLC (Fig. 4A, B). Therefore, FLNb regulates MRLC di-phosphorylation in HT1080 cells.

In addition, we examined the role of FLNb in regulating FAK phosphorylation in HT1080 cells, which is known to contribute to invasion (Sulzmaier et al., 2014). We assessed the phosphorylation of residue Y397 in FAK, as this site is crucial for the activation of FAK and is associated with tumor progression (Sulzmaier et al., 2014). The level of phosphorylated FAK was significantly lower in FLNb knockdown HT1080 cells than in control cells (Fig. 4C, D). Therefore, phosphorylation of FAK at Y397 is regulated by FLNb expression in HT1080 cells.
tile forces generated by actomyosin via regulation of di-phosphorylation in MRLC. In addition, phosphorylation of FAK is reported to regulate focal adhesion turnover (Hamadi et al., 2005). Since turnover of focal adhesions is important for cell migration (Bugide et al., 2014), FLNb may promote invasion in cancer cells through FAK-mediated regulation of focal adhesion turnover. Previous work has reported that di-phosphorylation of MRLC and phosphorylation of FAK at Y397 residue ([FAK (pY397)]), total amount of FAK (total-FAK), FLNb, and GAPDH in HT1080 cells as detected by western blot analysis is shown. (D) The relative phosphorylated level of FAK ([FAK (pY397])/total-FAK ratio) was calculated from (C). The mean values are shown with standard deviation (shown as error bars) from 3 independent experiments. *p<0.05.

Fig. 4. Filamin B (FLNb) enhances phosphorylation of myosin regulatory light chain (MRLC) and focal adhesion kinase (FAK) in HT1080 cells. (A) Protein expression of di-phosphorylated MRLC (pp-MRLC), total amount of MRLC (total-MRLC), FLNb, and GAPDH in HT1080 cells as detected by western blot analysis is shown. (B) The relative phosphorylated level of MRLC (pp-MRLC/total-MRLC ratio) that was quantified from (A). The mean values are shown with standard deviation (shown as error bars) from 3 independent experiments. *p<0.05. (C) Protein expression of phosphorylated FAK at Y397 residue ([FAK (pY397)]), total amount of FAK (total-FAK), FLNb, and GAPDH in HT1080 cells as detected by western blot analysis is shown. (D) The relative phosphorylated level of FAK ([FAK (pY397])/total-FAK ratio) was calculated from (C). The mean values are shown with standard deviation (shown as error bars) from 3 independent experiments. *p<0.05.

Our data suggest that FLNb is a potential therapeutic target for invasive cancers, including lung cancer and fibrosarcoma. In this study, we demonstrated that downregulation of FLNb is sufficient to inhibit the invasive ability of lung carcinoma cells and fibrosarcoma cells. Therefore, inhibition of FLNb may improve the prognosis of patients with invasive cancer.
Acknowledgments. The authors are grateful to Kosaku Kato for performing preliminary experiments. This study was supported by JSPS KAKENHI Grant Numbers 26430104, 26106702, 25127701, 24370069 to H.H., 26106704 to T.M., 25287106 to K.K., 26860964 to S.I. This study was also partly supported by Special Expenditures for “Reverse Translational Research from Advanced Medical Technology to Advanced Life Science” to H.H. and S.I. granted by MEXT, Japan.

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(Received for publication, March 10, 2015, accepted, April 22, 2015 and published online, April 29, 2015)