Periostin induces epithelial-to-mesenchymal transition via the integrin α5β1/TWIST-2 axis in cholangiocarcinoma

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Abstract. Periostin (PN) (also known as osteoblast-specific factor OSF-2) is a protein that in humans is encoded by the POSTN gene and has been correlated with a reduced survival of cholangiocarcinoma (CCA) patients, with the well-known effect of inducing epithelial-to-mesenchymal transition (EMT). The present study investigated the effect of PN, through integrin (ITG)α5β1, in EMT-mediated CCA aggressiveness. The alterations in EMT-related gene and protein expression were investigated by real-time PCR, western blot analysis and zymogram. The effects of PN on migration and the level of TWIST-2 were assessed in CCA cells with and without siITGA5 transfection. PN was found to induce CCA cell migration and EMT features, including increments in Twist-related protein 2 (TWIST-2), zinc finger protein SNAI1 (SNAIL-1), α-smooth muscle actin (ASMA), vimentin (VIM) and matrix metalloproteinase 9 (MMP-9), and a reduction in cytokeratin 19 (CK-19) together with cytoplasmic translocation of E-cadherin (CDH-1). Additionally, PN markedly induced MMP-9 activity. TWIST-2 was significantly induced in PN-treated CCA cells; this effect was attenuated in the ITGα5β1-knockdown cells and corresponded to reduced migration of the cancer cells. These results indicated that PN induced CCA migration through ITGα5β1/TWIST-2-mediated EMT. Moreover, clinical samples from CCA patients showed that higher levels of TWIST-2 were significantly correlated with shorter survival time. In conclusion, the ITGα5β1-mediated TWIST-2 signaling pathway regulates PN-induced EMT in CCA progression, and TWIST-2 is a prognostic marker of poor survival in CCA patients.

Introduction

Cholangiocarcinoma (CCA) is a primary liver cancer arising from bile duct epithelial cells and characteristically manifests as an abundant stromal reaction (1). CCA is associated with a poor patient prognosis and a low survival rate due to its high capability to invade and metastasize (2). Increased α-smooth muscle actin (ASMA)-positive fibroblasts have been indidentified in the stroma of CCA tissues and higher levels have been correlated with shorter patient survival times (1). These fibroblasts were found to have upregulated expression of several secreted proteins, and periostin (PN) is a crucial one exclusively overexpressed in stromal fibroblasts and significantly correlated with poor patient prognosis (3). PN (also known as osteoblast-specific factor OSF-2) is a protein that in humans is encoded by the POSTN gene and has been reported to enhance several tumorigenic properties in various types of cancers including ovarian (4), breast (5), colon (6), head and neck (7), and pancreatic cancer (8). Enhancement was shown to occur through interactions with membrane receptor integrin (ITG)αvβ3 or αvβ5 in ovarian cancer (4), ITGα6β4 in pancreatic cancer (9), and ITGα5β1 in CCA (10). Mino et al revealed that the malignant potential of PN is expressed through the induction of epithelial-to-mesenchymal transition (EMT) in CCA via ITGαv (11). Although PN acts through ITGα5β1 in CCA cell invasion (10), understanding of the role of ITGα5β1 in the EMT phenotype of CCA cells is still limited.

EMT has been strongly implicated in several types of cancer in regards to a key impact on cell invasion and metastasis (12,13). Characteristics of cells undergoing EMT include an increase in mesenchymal markers such as vimentin (VIM),
N-cadherin (CDH-2), ASMA, and fibronectin (FN-1) and a reduction in epithelial markers, in particular, E-cadherin (CDH-1) and cytokeratin (CK) (14,15). Three transcription factors, zinc finger protein SNAI1 (SNAI1-1), SLUG (SNAI1-2) and TWIST have been demonstrated to function in the regulation of EMT in cancers (14). Zinc finger protein SNAI1 (SNAI1-1) has been identified as a key molecule in transforming growth factor (TGF)-β1-activated EMT in pancreatic cancer (16,17). TWIST was shown to induce breast cancer cells to undergo EMT (18). However, the role of TWIST-2 in EMT that follows PN activation of ITGα5β1 has not been well elucidated. In the present study, recombinant PN activated EMT which led to CCA migration and TWIST-2 activation, and additionally, the potential clinical use of TWIST-2 as a marker for poor prognosis in human CCA was revealed. These findings also highlight the potential impact of targeting the PN/ITGα5β1/TWIST-2 pathway driving CCA migration to attenuate the progression of disease.

Materials and methods

CCA cell line culture. Human CCA cell lines KKU-100, KKU-139 and KKU-213 were cultured in DMEM medium (Invitrogen; Thermo Fisher Scientific, Inc.) supplemented with 10% fetal bovine serum (FBS), 100 U/ml penicillin, 100 µg/ml streptomycin (Invitrogen; Thermo Fisher Scientific, Inc.), and an anti-fungal agent. Cells were cultured in a 5% CO2 incubator at 37˚C, and passaged with 0.25% trypsin-EDTA. Cells with more than 90% viability were used throughout this study.

Migration assay (wound healing assay). KKU-100, KKU-139, and KKU-213 tumor cell lines (50,000 cells/well) were cultured in 6-well plates until they reached approximately 90% confluence. A reference midline was drawn under the plate. Cells were scraped off along the line using a sterile 200-µl pipette tip and the detached cells were washed away with serum-free medium. The remaining cells were then treated with medium containing either 100 ng/ml recombinant PN or medium without PN. The scraped area indicated by the reference line was recorded at the beginning of treatment and again at 24 h. The efficiency of migration into the scraped area was taken as a measure of wound healing and was calculated by the following formula: % wound healing=[(wound space at 0 h-wound space at 24 h)/wound space at 0 h] x100.

Transwell migration assay. A total of 5x104 KKU-213 cells in DMEM containing 1% FBS with or without 100 ng/ml PN was plated in the upper chamber of a 24-well Corning Transwell plate (Corning #3428 Transwell) and 600 µl of 1% FBS DMEM was added to the lower chamber. After culture in a humidified incubator at 37˚C for 12 h, the upper chamber was fixed in 70% ethanol for 30 min and stained with 0.5% crystal violet for 15 min. After drying, migrating cells were counted under an inverted microscope (original magnification, x400).

Measurement of EMT gene expression in PN-treated CCA cell lines by real-time PCR. Total RNA was extracted using a Perfect Pure RNA Cultured Cell Kit (5 PRIME; Thermo Fisher Scientific, Inc.) according to the manufacturer’s instructions. The cDNA was synthesized from 1 µg of total RNA using SuperScript™ III First-Strand Synthesis System for RT-PCR (M-MLV; Invitrogen; Thermo Fisher Scientific, Inc.) according to the instructions. The sequences of all genes in this study were retrieved from PubMed (www.ncbi.nlm.nih.gov) and the primers were designed using Primer 3 software (Version 0.4.0) (Table I). The PCR cycles consisted of: 95˚C for 10 min, followed by 50 cycles of 95˚C for 15 sec and 60˚C (except 58˚C for TWIST-2) for 15 sec. Crossing point data were automatically obtained by the fit-point of Lightcycler® 480 software version 1.5 (https://lifescience.roche.com/en_th/products/lightcycler14301‑480‑software‑version‑15.html). ACTB (β-actin gene) was used as an internal control. The expression level of each gene in the PN-treated CCA cells was compared to that of the control cells without PN treatment (2^(-ΔΔCp)) as follows (19): 2^-ΔΔCp=(Cpuntreated sample-CpACTB)-(Cptreated sample-CpACTB) (To note Cp is the crossing point as calculated by Lightcycler 480 software). The expression of EMT genes in TGF-β-treated cells was used as the positive control for EMT.

Western blot analysis of EMT markers. Cells with or without PN treatment were collected after refrigerated centrifugation of the cell suspensions at 400 x g for 5 min. The cell pellets were lysed in 1X sample buffer containing 50 mM Tris-HCl pH 6.8, 2% (w/v) SDS, 10% (v/v) glycerol, 5% (v/v) β-mercaptoethanol and 0.05% (w/v) bromophenol blue. Cell lysates were boiled for 1 min and centrifuged at 8,000 x g for 1 min to remove the undissolved proteins and cell debris. Cell extracts were then separated using 10% SDS-PAGE and transferred onto PVDF membranes (Amersham). The membranes were blocked in 5% skim milk containing TBST for 1 h at room temperature (RT). The immunodetection steps for TWIST-2, ASMA, VIM, CK-19, MMP-9, MMP-13, and ITGα5 were performed using mouse anti-TWIST-2 (dilution 1:250; cat. no. ab57997; Abcam), mouse anti-ASMA (dilution 1:400; cat. no. A5228; Sigma-Aldrich; Merck KGaA), mouse anti-VIM (dilution 1:200; cat. no. sc-6260; Santa Cruz Biotechnology, Inc.), mouse anti-CDH-1 (dilution 1:50; cat. no. sc-71008; Santa Cruz Biotechnology), mouse anti-CK-19 (dilution 1:200; cat. no. sc-6278; Santa Cruz Biotechnology), mouse anti-MMP-9 (dilution 1:100; cat. no. sc-217333; Santa Cruz Biotechnology, Inc.), anti-MMP13 (dilution 1:1,000; cat. no. sc-30073; Santa Cruz Biotechnology, Inc.), and mouse anti-ITGα5 (dilution 1:500, sc-376199; Santa Cruz Biotechnology). The incubation periods of primary antibodies for TWIST-2, ASMA, VIM, CK-19, and ITGα5 were overnight at 4˚C, and those of MMP-9 and MMP-13 were 2 and 1 h, respectively, at RT. The secondary antibodies were HRP-conjugated goat anti-mouse IgG antibody (dilution 1:2,000; cat. no. 62-6620; Zymed, Thermo Fisher Scientific, Inc.) for mouse primary antibody, and HRP-conjugated goat anti-rabbit IgG antibody (dilution 1:3,000; Ab6717; Abcam) for rabbit primary antibody. The immunoreactive signals were visualized by ECL (Thermo Fisher Scientific, Inc.) under Gel Document Syngene (Syngene). The β-actin level was used as an internal control with mouse anti-β-actin (dilution 1:10,000; sc-47778; Santa Cruz). The bands were quantified by ImageJ software (Version 1.52a; National Institutes of Health, Bethesda, MD, USA). The obtained staining intensities of the proteins of interest were then normalized against the intensity of the staining of β-actin protein (ACTB).
Zymography. The conditioned-media of CCA cells that had been treated with PN, or that had been left untreated, were centrifuged at 400 x g for 5 min to remove cell debris, and supernatants were then collected. The supernatants were concentrated with Vivaspin 20 (MWCO 3,000) by centrifugation at 3,750 x g at 4˚C for 2 h. The protein concentrations were measured using the Coomassie Plus Assay Kit (Thermo Fisher Scientific, Inc.). The samples were loaded using 12% SDS-PAGE containing 1 mg/ml gelatin (Ajax Finechem) as a substrate. The electrophoresis was performed at 200 V for 5 min in pre-cooled SDS-PAGE running buffer, followed by prolonged electrophoresis in a 4˚C refrigerator for 1 h 20 min. Afterward, the zymogram gels were twice washed for 30 min with 2.5% Triton X-100 to renature the enzymes. Zymographic activities were processed with developing buffer at 37˚C for 18 h. The results were visualized as clear bands after staining with 0.006% Coomassie blue for 2 h at RT. The bands were quantified by ImageJ software. The cut-off at 1.5 for upregulated gene expression was used as it typically causes enough upregulated protein production to affect cell phenotype (3).

Immunocytochemistry for localization of CDH-1. KKU-213 and KKU-139 CCA cells (5,000 cells/well) were grown in 96-well plates to 40% confluence and treated with 100 ng/ml PN for 48 h. Cells were washed with 1X PBS pH 7.4 and fixed with 4% paraformaldehyde at RT for 10 min. For cell permeabilization, 0.5% Triton X-100 in 1X PBS was added and incubated at RT for 5 min. The non-specific binding was blocked with 10% (w/v) FBS in 1X PBS for 1 h 30 min. The samples were loaded using 12% SDS-PAGE containing 1 mg/ml gelatin (Ajax Finechem) as a substrate. The electrophoresis was performed at 200 V for 5 min in pre-cooled SDS-PAGE running buffer, followed by prolonged electrophoresis in a 4˚C refrigerator for 1 h 20 min. Afterward, the zymogram gels were twice washed for 30 min with 2.5% Triton X-100 to renature the enzymes. Zymographic activities were processed with developing buffer at 37˚C for 18 h. The results were visualized as clear bands after staining with 0.006% Coomassie blue for 2 h at RT. The bands were quantified by ImageJ software. The cut-off at 1.5 for upregulated gene expression was used as it typically causes enough upregulated protein production to affect cell phenotype (3).
IgG-Cy3 (dilution 1:2,000; cat. no. 115-166-071; Jackson ImmunoResearch). The nuclei were stained with Hoechst at 1:1,000 (Invitrogen Thermo Fisher Scientific, Inc.). After washing, the cells were fixed with 4% paraformaldehyde at RT for 10 min and mounted in 50% glycerol. The positive staining cells were visualized under a fluorescence inverted microscope (Olympus; x400 original magnification). The patterns and localizations (cytoplasmic or nuclear) were recorded.

**Immunofluorescence staining of ITGα5β1.** Cells were plated onto 12-mm glass coverslips in DMEM containing 10% FBS in a 24-well plate at 2x10^4 cells/well and cultured overnight. Cells were fixed with 4% (w/v) paraformaldehyde for 15 min at RT. The cells were blocked with 1X PBS containing 0.5% BSA for 30 min at RT. The cells were stained with primary antibody, mouse anti-human ITGα5 (dilution 1:100, C-9, Santa Cruz Biotechnology, Inc.), for 2 h at RT in 1X PBS containing 0.5% BSA, and incubated with Cy3-conjugated anti-mouse IgG secondary antibody (dilution 1:2,000; cat. no. 115-166-071; Jackson ImmunoResearch Laboratory Inc.). Based on the fact that the ITGα5 subunit can bind only to the ITGb1 subunit (20), changes in ITGα5 were taken as equivalent to changes in ITGα5β1. The nuclei were stained with Hoechst 33342 (1:1,000; Invitrogen; Thermo Fisher Scientific, Inc.) for 1 h at RT. The stained samples were mounted in 50% glycerol and immunofluorescence images were obtained on an inverted microscope and a confocal microscope (LSM800, Carl Zeiss, Jena, Germany; x63 original magnification).

**Transient knockdown of ITGα5 in CCA cells.** Silencing the expression of ITGα5β1 was performed by transfection with siITGα5 (Santa Cruz Biotechnology, Inc.) according to the manufacturer's instructions. Briefly, 1.5x10^5 cells/well were seeded onto 6-well plates and cultured with complete medium at 37°C in a 5% CO₂ incubator for 24 h. A siITGα5 transfection solution was prepared by separately mixing 100 µl of OptiMEM® I (Invitrogen; Thermo Fisher Scientific, Inc.) with either 10 µl of 10 pmole siITGα5 or 5 µl of Lipofectamine™ 2000 (Invitrogen; Thermo Fisher Scientific, Inc.). The solutions were then mixed and incubated for 20 min at RT. In addition, 200 µl of OptiMEM® I mixed with 5 µl of Lipofectamine™ 2000 was simultaneously used as a mock solution. Cells were washed three times with antibiotic-serum-free medium. The siITGα5 transfection solution (cat. no. STSCS-29372, Santa Cruz Biotechnology, Inc.) was slowly dropped into each well and incubated at 37°C in a 5% CO₂ incubator for 6 h. Then the complete medium was replaced. The efficiency of ITGα5 silencing was confirmed by real-time PCR and immunofluorescence staining.

**Immunohistochemistry of ITGα5β1 and TWIST-2 in CCA tissues.** Paraffin-embedded tissue samples from 50 CCA cases were collected from the Cholangiocarcinoma Research Institute, Faculty of Medicine, Khon Kaen University (Khon Kaen, Thailand) under the approval of the Human Research Ethics Committee, Khon Kaen University (HE490143). The median age of the patients was 58 years with a range of 37-75 years. The percentage of females was 38% (19/50) and males was 62% (31/50). The samples were recruited during 2006-2011. The tissues were baked at 60-65°C for 4 h, deparaffinized, soaked in 10 mM citrate buffer pH 6, and boiled at 95°C for 40 min. Endogenous peroxidase was blocked with 3% H₂O₂ in methanol for 30 min. The tissues were incubated with 2% BSA in 0.05 M Tris-HCl pH 7.6 to block non-specific binding, then mouse anti-human TWIST-2 (dilution 1:200; ab57997; Abcam) was applied and the tissues were incubated at 4°C overnight. The tissues were then incubated with Envision® System horseradish peroxidase (Dako) for 30 min. The signal was developed using 0.05% 3,3'-diaminobenzidine (DAB) and counterstained with hematoxylin. The scoring was assessed using the percentage of positive staining cells (P) and the intensity of the staining signal (I). For P, samples containing 0-25, 26-50, 51-75, and 76-100% positive cells were classified as grades 0, 1, 2, and 3, respectively. For I, unstained, slightly stained, intermediately stained, and strongly stained cells were classified as 0, 1, 2, and 3, respectively. The multiplied score (P x I) as a total score of 0-9 was determined for each stained slide by 2 investigators double-blinded to the clinical data. A total score of more than 4 was classified as high TWIST-2 level whereas scores less than or equal to 4 were in the low-level group. This cut-off value was derived from the median of the scoring of all cases. In this research study, the normal adjacent area which was approximately 5-10 mm distant from the border of the cancerous area in the same slide of the CCA tissue section was used to represent normal bile duct.

For ITGα5β1, the level in CCA tissues was detected using the same protocols as for TWIST-2 except that the antigen retrieval was performed using samples microwave-heated with 10 mM sodium citrate buffer with 0.05% Tween 20, pH 6.0 for 10 min, and the mouse anti-human ITGα5 antibody (dilution 1:100, sc-376199; Santa Cruz Biotechnology, Inc.) was used as the primary antibody.

**Statistical analysis.** All quantified data are expressed as mean ± standard deviation (SD), except for survival time for which median values were used. Statistics for the in vitro studies were compared with controls using the Student's t-test. One-way analysis of variance (ANOVA) with all pairwise multiple comparison by Tukey test was utilized to analyze the significant results of more than 2 groups. The statistical analysis of two variables and the interaction between the groups was performed by two-way ANOVA. The correlation of TWIST-2 level with clinicopathological data was verified by Cox regression multivariate analysis, and the Kaplan-Meier Log Rank test was used for survival analysis. A P-value of <0.05 was considered to be statistically significant.

**Results**

**PN-induced alterations in the EMT phenotype and EMT-related markers in CCA cells.** The results revealed with statistical significance that KKU-213 CCA cells treated with 100 ng/ml PN had a greater ability to migrate than the untreated control cells, which had a migration ability similar to CCA cells treated with 10 ng/ml TGF-β (Fig. S1). Uregulated genes in the PN-treated cells were defined as those with relative gene expression 1.2-fold over those in the untreated cells. Downregulated genes had a relative gene expression equal to or less than 0.8-fold. The results showed
that PN treatment induced the expression of both \textit{SNAIL}-1 and \textit{TWIST}-2, with statistical significance only for \textit{TWIST}-2 (Fig. 1A). Notably, \textit{SNAIL}-1 was activated by both PN and TGF-\(\beta\), whereas \textit{SLUG} was not changed. Upregulation of \textit{ASMA} and \textit{VIM} mesenchymal markers was observed in the PN-treated CCA cells (Fig. 1B). Downregulation of the \textit{CK}-19 epithelial marker was significant, whereas no significant change was observed in the \textit{CDH}-1 mRNA level (Fig. 1C).

The Western blot results confirmed that the PN-induced EMT of CCA cells was accompanied by significant downregulation of \textit{CK}-19, and a significant increase in the production of \textit{TWIST}-2 and \textit{VIM} (Fig. 1D).

Interestingly, although the difference in the expression level of \textit{CDH}-1 mRNA was not significant, the protein level revealed by western blot analysis was significantly reduced in the PN-treated cancer cells (Fig. 1E). This was supported by the finding in TGF-\(\beta\)-treated cells. Moreover, \textit{CDH}-1, expressed predominantly on the cell membranes of both KKKU-213 and KKKU-139 CCA cells, was translocated after PN treatment to the cytoplasm (Fig. 1F). Thus, the membrane localization of \textit{CDH}-1 in the untreated cells was found on approximately 84 and 85\% of the KKKU-213 and KKKU-139 cells (Fig. 1G). In cells treated with PN, the membranous protein staining pattern of \textit{CDH}-1 was reduced to 16 and 26\% of KKKU-213 and KKKU-139 cells respectively, and the percentages of cells displaying a cytoplasmic staining pattern was increased to approximately 84 and 74\%.

\textbf{Effect of PN on the induction of MMPs in CCA cells.} No alterations of \textit{MMP}-1, \textit{MMP}-7, \textit{MMP}-10, or \textit{MMP}-14 were detected in PN-treated CCA cells, whereas 100 ng/ml PN induced levels of \textit{MMP}-9 and \textit{MMP}-13 to 1.99\pm0.21-
SONGBUA et al.: PN INDUCES ITGα5β1/TWIST-2-MEDIATED EMT IN CCA

1.81±0.17-fold that of the untreated control cells (Fig. 2A-F). However, the western blot analysis results showed that neither pro-MMP-9 (92 kDa) nor active MMP-9 (84 kDa) were induced by PN whereas the intermediate form of MMP-13 (Inter-MMP-13) at 56 kDa showed a significant increase (Fig. 2G). The zymography results showed that after exposure to PN, CCA cells secreted MMP-9 into the conditioned-medium with a marked increase in activity compared to the activity observed in the control condition (Fig. 2H). Densitometric analysis showed that MMP-9 activity was significantly increased in the PN-treated CCA cells to 7-fold that of the control cells. No significant change in MMP-13 in the PN-treated CCA cells was observed.

ITGα5β1-mediated CCA cell migration and MMP expression. The siITGα5-transfection significantly downregulated intrinsic ITG5 mRNA expression with an efficiency of approximately 35-50% (Fig. 3A). The efficiency of knockdown was significant in cells up to 96 h post-siRNA treatment (Fig. 3B). The wound-closure assay showed that PN significantly induced CCA cell migration after 24 h of treatment, while in the siITGα5-transfected cells, the migration was slower than that in the mock-transfected cells (Fig. 3C). In cells in which ITGα5β1 was diminished, the induction of expression of both MMP-9 and MMP-13 by PN treatment was less than that in the mock-transfected cells. This difference was significant for MMP-9 (Fig. 3D and E).

ITGα5β1/TWIST2-mediated CCA cell migration. The results showed that siITGA55β1 treatment successfully inhibited the production of membrane ITGα5β1 in CCA cell lines (Fig. 4A). Moreover, the siITGα5-transfected cells had a significantly lower response rate in the PN-induced cell migration assay compared to the scrambled-siRNA-treated cells (Fig. 4B). These results were observed in all three CCA cell lines. TWIST-2 levels in these ITGα5-transient-knockdown CCA cells were not induced by PN and this was in accord with the decreased level of cell migration induced by PN (Fig. 4C).
Expression of ITGα5β1 and TWIST-2 in CCA tissues and the clinicopathological relevance. The demographic data of the enrolled patients are shown in Table II. Most of the cases (60%) were male and more than 90% of the enrolled cases were diagnosed with late-stage disease. Various positive stain-intensity scores of TWIST-2 were found in CCA tissues (Fig. 5A-C) whereas normal bile duct showed no TWIST-2 expression (Fig. 5D). Most of the CCA cases, (33/50, 66%), showed high TWIST-2 levels. Multivariate Cox regression analysis revealed that elevated TWIST-2 level and a categorization as a poorly differentiated cancer type were independent parameters associated with poor prognosis. Other parameters showed no significant correlations. Interestingly, CCA patients with high TWIST-2 had shorter mean survival times (211±14 days) compared to those with low TWIST-2 (637±503 days). The overall survival analysis by Kaplan-Meier test showed, with statistical significance, that patients exhibiting high TWIST-2 levels had shorter survival times than those with low TWIST-2 using both 3-year and 5-year survival as the cut-off values (Fig. 5E). The level of ITGα5β1 in CCA clinical samples was also explored, and the results showed elevated expression in 70% (7/10) of the CCA cases. The levels ranged from low to high expression (Fig. 5F).

Discussion

Epithelial-to-mesenchymal transition (EMT) is a transient phenomenon but is an important step in cancer cell activation by many intracellular and extracellular stimuli, including soluble substances from both cancer cells and cancer stromal fibroblasts in the cholangiocarcinoma (CCA) microenvironment (21-23). The aberration of secretomes in ASMA-positive stromal fibroblasts led to the discovery of peristin (PN) as a major protein having a tumorigenic impact in intrahepatic CCA (3), and a variety of EMT markers have been proposed as indicators for the poor...
SONGBUA et al.: PN INDUCES ITGα5β1/TWIST‑2‑MEDIATED EMT IN CCA

8

patient prognosis (24). The present study confirms the effect of PN on the EMT phenotype in CCA cells and is consistent with recently published research indicating that this protein acts in an autocrine manner (11).

The high level of ITGα5β1 in CCA cell lines and its role as a receptor for PN have been reported in our previous publication (10). In the present study, it was revealed that PN treatment of CCA cell lines induced an EMT pattern of gene expression that included upregulation of ASMA and VIM, an increase in MMP‑9 activity, and downregulation of CK‑19 through the ITGα5β1 and TWIST‑2 signaling pathway. Additionally, the level of TWIST‑2 was correlated with patient survival time, with statistical significance. These findings highlight the potential impact of inhibiting the PN/ITGα5β1/TWIST‑2-pathway that drives CCA migration, thereby attenuating the progression of this disease.

The expression pattern of EMT markers in PN‑induced CCA cell lines revealed alterations and translocation of membranous CDH‑1 to the cytoplasm of cells. This hallmark of EMT gene expression has also been found in TNF‑α and HGF‑α‑treated CCA cells (21,23). Although CDH1 mRNA in peristin‑treated cells was not significantly changed compared to the control untreated cells, the protein level was reduced. The level of protein, the final product, is considered to be more relevant than the mRNA level. Moreover, there is an evident showing that PN regulated EMT and EMT transcriptional factor expression by repressing microRNA‑38 expression in lung cancer (25). In the same way, CDH‑1 was shown in breast cancer to be downregulated by miR‑9 resulting in increased cell motility and invasiveness (26). These findings highlight the potential of PN to regulate gene expression at the post‑transcriptional or translational level which supports the finding of unaltered mRNA but an altered protein level of CDH‑1 in the study herein. Cells need membranous CDH‑1 to provide intercellular molecular adhesion and attachment to extracellular matrices (27) and a decrease in membranous CDH‑1 in PN‑treated CCA cells facilitates subsequent cell detachment and migration. The level of CK‑19 was also reduced by PN, as previously reported in cells treated with TGF‑β and TNF‑α (22,23). Evidently,

Figure 4. Effect of transient ITGα5 knockdown (siITGα5) on the migration response of CCA cells to PN. (A) Downregulation of ITGα5 in siITGα5‑treated CCA cells detected by immunocytochemistry as determined by confocal microscopy. Original magnification, x63. Scale bar, 50 mm. (B) Significant migration ability of CCA cells after treatment with PN compared to the absence of response in siITGα5‑treated CCA cells. (C) Increased level of TWIST‑2 after PN treatment in ITGα5‑expressing CCA cells and attenuated response in the cells with suppressed expression of ITGα5. Results are presented as mean ± SD of triplicate experiments. *P<0.05, **P<0.01 and ***P<0.001. ITG, integrin; CCA, cholangiocarcinoma; PN, peristin; TWIST‑2, Twist‑related protein 2.
the PN-induced EMT phenomenon in CCA cells displays the characteristic decrease in CK-19 and cytoplasmic translocation of CDH-1, which are important steps allowing cells to detach and become ready to move. The reduction in the CK level is an important characteristic of EMT in CCA (24) and is associated with neural invasion, intrahepatic metastasis, and shortened overall survival time of patients (28). Hence, the lower CK-19 level in PN-treated cells than that noted in the untreated controls represents the induction of EMT in PN-treated CCA cells. To accomplish cell motility, cytoskeletal proteins must increase their expression. In CCA cells activated by PN, the expression levels of ASMA and VIM were found to be increased which is consistent with increased motility through ASMA-mediated cell-generated mechanical tension, cytoskeleton remodeling (29) and VIM-regulated cell adhesion and migration (30). VIM appears to be a consistent characteristic of EMT in CCA as it was observed both using TGF-β1 as the positive control EMT inducer (22), and using PN as presented in this present study. Interestingly, PN-treated CCA cells that have undergone EMT in highly confluent cultures were found to exhibit epithelial cell morphology (cobblestone shape) while spindle-shaped cells were observed in areas with low confluence, which is consistent with previous findings (31). Serum MMP-9 has been reported as being increased in CCA patients (32) and high expression of tissue MMP-9 has been correlated with poor patient prognosis (33). Herein, PN-stimulated CCA cells had increased levels of MMP-9 and this alteration may help CCA cells to invade through the degraded extracellular matrix consequently leading to distant metastases. PN-induced MMP expression was observed to display a dose-dependent trend in the range of 10-100 ng/ml, for which 1,000 ng/ml was an excessive level, and a feedback inhibition effect is expected (3). We previously showed that ITGα5β1 is the major receptor for PN on the CCA cell membrane (10). The involvement of ITGα5β1 in the regulation of the PN-activated EMT pathway was confirmed here by the finding that cells silenced for ITGα5β1 had both less migration ability induced by PN treatment and reduced MMP production.

SNAIL-1 has been postulated to control the EMT phenotype in CCA cells after stimulation with TGF-β1 and TNF-α (22,23). In a similar way to these previous findings, the present study showed that PN-treated CCA cells

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<td>Histological type</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WD (n=21)</td>
<td>1.000</td>
<td></td>
</tr>
<tr>
<td>MD (n=8)</td>
<td>1.336</td>
<td>0.563-3.170</td>
</tr>
<tr>
<td>PD (n=6)</td>
<td>4.670</td>
<td>1.648-13.24</td>
</tr>
<tr>
<td>Pap (n=15)</td>
<td>2.070</td>
<td>0.681-6.292</td>
</tr>
<tr>
<td>Vascular invasion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Presence (n=8)</td>
<td>1.071</td>
<td>0.444-2.582</td>
</tr>
<tr>
<td>Absence (n=42)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stage</td>
<td></td>
<td></td>
</tr>
<tr>
<td>I-II (n=4)</td>
<td>1.1763</td>
<td>0.404-7.668</td>
</tr>
<tr>
<td>III-IV (n=46)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tumor size (cm)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤5 (n=27)</td>
<td>1.147</td>
<td>0.553-2.378</td>
</tr>
<tr>
<td>&gt;5 (n=23)</td>
<td></td>
<td></td>
</tr>
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</table>

*P<0.05 indicates significant difference. CCA, cholangiocarcinoma; TWIST-2, Twist-related protein 2; CI, confidence interval; WD, well-differentiated; MD, moderately differentiated; PD, poorly differentiated; Pap, papillary.
exhibited increments of SNAIL-1 and TWIST-2 mRNA. Increased SNAIL-1 and TWIST-2 proteins were also found and the increase in TWIST-2 was statistically significant. This may imply an involvement of TWIST-2 in PN-induced EMT in CCA cells, particularly as cells with suppressed ITGa5β1 had a decreased capability to activate TWIST-2 and this corresponded to a decreased ability of the cells to migrate. TWIST-2 has been shown to control the membrane translocation of CDH-1 through FOXC-2 activation (34) which mediates several EMT phenotypes including those of detached and motile cells (35). This function of TWIST-2 was supported by a finding in cervical cancer cell lines demonstrating that lower intrinsic expression of cytoplasmic CDH-1 was exhibited in cells with high TWIST-2 levels (36). In addition, SNAIL-1 was found to play an important role in CCA activation by PN, probably through the downregulation of CK-19 and upregulation of ASMA, VIM, and MMPs (22,23).

Herein, it was evident that TWIST-2 may have had a strong impact after activation via the PN/ITGa5β1 pathway in EMT-mediated CCA progression. However, TWIST-2 overexpression should be carried out in ITGa5 knockdown cells to rescue the cell response to PN to further validate TWIST-2 in regards to the effects of PN through ITGa5β1 on CCA cells.

Increased expression of TWIST-2 has been associated with metastases in the progression of breast and cervical cancers, and is also correlated with poor cell differentiation and a short patient survival rate (36,37). This is similar to the findings of the present study; those patients with high levels of TWIST-2 had significantly shorter survival times compared to those with low TWIST-2. Increases in the SNAIL-1 level in CCA tissues were previously reported to significantly correlated with lymph node metastasis and a poor survival rate in CCA patients (22). This is in contrast to the findings with extrahepatic CCA, in
which SNAIL, SLUG, TWIST, ZEB1, ZEB2 were not valuable to predict poor outcomes of the patients, whereas VIM, FN, S100A4, CDH1, and CDH2 were associated with short survival times (38).

In conclusion, PN enrichment found in the CCA microenvironment, mainly produced by cancer-associated fibroblasts, activates CCA cells by binding to ITGα5β1 to trigger an AKT-dependent signaling pathway (10) that activates TWIST-2 (Fig. 5G). Although the limitation of FOXC2 was not identified, this study identified alterations in EMT-related genes underlying changes in the phenotype for EMT and driving CCA progression after the binding of PN to ITGα5β1. The change in TWIST-2 levels can be suggested as a marker of poor prognosis that predicts the aggressiveness of CCA. Moreover, clinically targeting this ITGα5β1-mediated pathway and the activated signaling molecules and/or associated transcription factors may help attenuate PN-induced CCA progression.

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Availability of data and material

The datasets used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors' contributions

JS, SS, KU, ST and CT designed the experiments; JS, SS, ST, TK and KU performed the main research work. AP and SW prepared the samples in preparation for the experiments and facilitated the research work. PT and CT discussed and analyzed the data and results. CT reviewed the paper for research grant application, prepared figures, wrote, and improved the scientific quality of the manuscript, and finally submitted the manuscript. All authors read and approved the manuscript and agree to be accountable for all aspects of the research in ensuring that the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Ethics approval and consent to participate

Fifty cases of CCA tissues were obtained from patients who had undergone hepatectomy using the protocol approved by the Human Research Ethics Committee, Khon Kaen University (HE490143). Each patient was informed and written informed consent was obtained.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that there are no competing interests.

Authors' information

JS and SS were enrolled in the Graduate Program in Immunology, Department of Immunology, Faculty of Medicine, Siriraj Hospital, Mahidol University. ST and TK are research assistants under the Research Unit, Faculty of Medicine, Siriraj Hospital, Mahidol University.

References


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