Loss of βPix Causes Defects in Early Embryonic Development, and Cell Spreading and Platelet-Derived Growth Factor-Induced Chemotaxis in Mouse Embryonic Fibroblasts

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βPix is a guanine nucleotide exchange factor for the Rho family small GTPases, Rac1 and Cdc42. It is known to regulate focal adhesion dynamics and cell migration. However, the in vivo role of βPix is currently not well understood. Here, we report the production and characterization of βPix−/− mice. Loss of βPix results in embryonic lethality accompanied by abnormal developmental features, such as incomplete neural tube closure, impaired axial rotation, and failure of allantois-chorion fusion. We also generated βPix−/− mouse embryonic fibroblasts (MEFs) to examine βPix function in mouse fibroblasts. βPix−/− MEFs exhibit decreased Rac1 activity, and defects in cell spreading and platelet-derived growth factor (PDGF)-induced ruffle formation and chemotaxis. The average size of focal adhesions is increased in βPix−/− MEFs. Interestingly, βPix−/− MEFs showed increased motility in random migration and rapid wound healing with elevated levels of MLC2 phosphorylation. Taken together, our data demonstrate that βPix plays essential roles in early embryonic development, cell spreading, and cell migration in fibroblasts.

Keywords: βPix, cell motility, embryonic lethality, focal adhesion, mouse embryonic fibroblast

INTRODUCTION

Cytoskeletal rearrangement is a fundamental step in the regulation of numerous cellular responses, such as proliferation, differentiation, and movement. Rho-family small GTPases, including Rac1, Cdc42, and Rho, are potent determinants that modulate actin cytoskeleton organization by cycling between an active GTP-bound state and an inactive GDP-bound state (Etienne-Manneville and Hall, 2002). Among regulatory proteins of Rho GTPases activity, guanine nucleotide exchange factors (GEFs) activate the GTPases by enhancing the exchange of bound GDP for GTP (Bos et al., 2007). βPix (beta-PAK [p21-activated kinase] interacting exchange factor) is a member of the Dbl family of GEFs, containing DH and PH domains (Zheng, 2001). βPix was firstly identified as a protein localized in focal adhesions (Oh et al., 1997), and it was later shown to act as a GEF for Rac1 and Cdc42 (Koh et al., 2001). Through its GEF activity, βPix modulates diverse cellular events, such as remodeling of the actin cytoskeleton, platelet-derived growth factor (PDGF)-stimulated response, and cell migration (Campa et al., 2006; Kim et al., 2001; ten Klooster et al., 2006). Therefore, investigating the molecular mechanism of βPix in these processes and examining the effect of disrupting βPix expression in animal models are essen-
tial for understanding the function of βPix in vivo.

Previous studies on the role of βPix were mainly carried out using in vitro cell culture systems while βPix genetic manipulation was carried out in model organisms, including Caenorhabditis elegans (Martin et al., 2016) and zebrafish (Liu et al., 2007; Tay et al., 2010). Here, we report the generation of βPix-knockout (βPix-KO) mice through gene targeting. These mice were used for investigating the physiological roles of βPix in vivo. Additionally, we generated βPix-KO mouse embryonic fibroblasts (MEFs) from the βPix-KO embryos to investigate the role of βPix at the cellular level.

In this study, we found that loss of βPix causes defects in early embryonic development that lead to embryonic lethality at embryonic day 9.5 (E9.5). βPix-KO MEFs have defects in cell spreading, PDGF-induced ruffle formation and chemotaxis, and focal adhesion formation. Taken together, our research suggests that βPix plays essential roles in early embryonic development, cell spreading, and cell migration in fibroblasts.

MATERIALS AND METHODS

Generation of βPix-KO mice

Genomic fragment of mouse ARHGEF7 gene was cloned by screening a CIT8 mouse BAC clone library (Invitrogen, USA). To generate a gene-targeting vector for ARHGEF7, a 3.1 kb and a 3.4 kb fragments flanking the exon 19 of this gene were cloned into Os.Dup/Del vector respectively, which carries a neomycin-resistance (Neo+) cassette that replaces the exon 19 and a thymidine kinase cassettes. J-1 ES cells were maintained, electroporated with the targeting vector and neomycin-selected. Survived clones were analyzed for proper recombination after digestion with either EcoRI or SacI. Targeted ES cells were microinjected into the blastocysts of C57BL/6J mice (Charles River Laboratories, USA). After transfection, transfected cells were cultured into single cell colonies, and finally established as wild-type (WT) and βPix-rescue in the cells was confirmed by Western blot analysis.

Antibodies and reagents

Rabbit polyclonal antibodies against the SH3 domain of βPix were prepared as described previously (Oh et al., 1997). The following antibodies were purchased: monoclonal mouse antibody against Actin (clone AC-40; Sigma, USA), monoclonal mouse antibody against Git1 (clone 13/p95PKL: BD Transduction Laboratories, USA), polyclonal rabbit antibody against PAK1 (Santa Cruz Biotechnology, USA), polyclonal rabbit antibody against phospho-PAK1 (Ser199/204)/PAK2 (Ser192/197) (Cell Signaling, USA), monoclonal mouse antibody against Rac1 (clone 23A8: Millipore, USA), monoclonal mouse antibody against phospho-MLC2 (Ser19) (Cell Signaling), monoclonal rabbit antibody against phospho-MLC2 (Thr18/Ser19) (Cell Signaling), monoclonal mouse antibody against Vinculin (clone hVIN-1: Sigma). The following reagents were purchased: Fibronectin from bovine plasma (Sigma) and PDGF-BB human (Sigma).

Western blot analysis

Tissue and cell lysates were prepared as described previously (Oh et al., 1997; Shin et al., 2019; Shrestha et al., 2018). Protein concentrations in the tissue and cell lysates were measured by Bradford assay and with Pierce BCA Protein Assay Reagent (Thermo Scientific, USA), respectively. Equal amounts of protein were resolved by SDS-PAGE and transferred to a PVDF membrane (Millipore). Blots were blocked with 5% skim milk or 3% bovine serum albumin in 0.1% Triton X-100 in PBS (0.1% PBS-T) for 50 min. The blots were incubated with primary antibodies for 1 h at room temperature. Then, the blots were incubated with horseradish peroxidase-conjugated secondary antibodies (Jackson ImmunoResearch Laboratories, USA) for 50 min and analyzed by enhanced chemiluminescence. α-Actin was used as a loading control.

Whole mount and histological analysis of mouse embryos

To observe the whole mounted embryos, embryos of each genotypes were dissected out. Embryos were fixed 4% paraformaldehyde (PFA) (Sigma) and observed under Zeiss Lumar V12 using a 0.8× (Neolumar S. Zeiss, Germany) objective lens. For histological analysis, the whole mouse decidua was dissected out and fixed in Bouin’s fixative or 4% PFA. Fixed embryos were then dehydrated and embedded in paraffin and 10 μm sections were collected on slides. Slides were stained with hematoxylin.

Generation and culture of βPix-KO and βPix-rescue MEFs

βPix-HET mice were intercrossed, and embryos at E8.5 were isolated and triturated in Cellstripper non-enzymatic cell dissection solution (Mediatech, USA), then attached to culture dish. Genomic DNA from yolk sac was used for genotyping. For immortalization, cells were infected with pBabe-Puro-SV40 large T Antigen lentiviral supernatant overnight. After several days or weeks of infection, immortalized cells were cultured into single cell colonies, and finally established as wild-type (WT) and βPix-KO MEF lines. To establish βPix-rescue MEF lines, pCAG-Cre recombinase was transfected in βPix-KO cell with LipofectAMINEPLUS (Invitrogen) according to the manufacturer’s instructions to splice out Neo+ cassette. After transfection, transfected cells were cultured into single cell colonies, and βPix-rescue in the cells was confirmed by genotyping and western blotting. The MEFs were cultured in DMEM (Gibco, USA) supplemented with 10% certified fetal bovine serum (Gibco), 1% MEM non-essential amino acid (Gibco), 1% L-glutamine (Welgene, Korea), 0.1% β-mercaptoethanol (Gibco) and 1% antibiotics/antimycotics mixture (Gibco) in 5% CO2 incubator at 37°C. Coverslips or dishes was coated with 10 μg/ml fibronectin.

Immunocytochemistry

Cultured cells on coverslips were fixed with 3.7% PFA for 10 min, permeabilized with 0.1% Triton X-100 for 10 min and blocked with blocking solution (10% normal goat serum
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[Vector Laboratories, USA], 0.3% bovine serum albumin, and 0.1% PBS-T) for 1 h. Then the coverslips were incubated with primary antibodies diluted in blocking solution for 1 h at room temperature and stained with FITC-conjugated anti-mouse IgG (Jackson ImmunoResearch Laboratories) for 1 h. For actin staining, Rhodamine-phalloidin (Molecular Probes, USA) was used. Following incubation, coverslips were mounted with Vectashield (Vector Laboratories) and observed with the Axiovert 200M microscope (Zeiss) equipped with a Zeiss Axioicam HRm CCD camera using a 100×, 1.40 Plan-Apochromat objective lens or with a LSM700 confocal microscope (Zeiss) using a 10×, 1.20 Plan-Apochromat objective lens.

Cell spreading assay
MEFs were transfected with pEGFP-C1 with Metafectene Pro (Biontex Laboratories, Germany) for 24 h, and re-plated on fibronectin-coated coverslip. After 1 h incubation, cells were fixed with 3.7% PFA, and mounted. Green fluorescent protein (GFP) signals were analyzed for calculating cell spreading area with ImageJ software.

Analysis of ruffle formation
MEFs were cultured on fibronectin-coated coverslip and serum starved overnight. Then treated with PDGF-BB (10 ng/ml) for 7 min, cells were fixed with 3.7% PFA and stained for actin cytoskeleton, and analyzed manually for PDGF-induced ruffle formation. The cells were categorized as peripheral ruffle-, dorsal ruffle-, and peripheral and dorsal ruffle-bearing cells (Abercrombie et al., 1970; Suetsugu et al., 2003).

Transwell assay
For the chemotaxis assay, Costar Transwell Permeable Support (8.0 μm pore size; Corning, USA) was used according to manufacturer’s protocol. In brief, 6000 serum starved MEFs in upper chamber, and filled serum-free media or PDGF-BB (20 μg/ml) containing media in lower chamber. After 4 h incubation, transwell membranes were fixed with methanol, stained with eosin, mounted with Richard-Allan Scientific Mounting Medium (Thermo Scientific) and observed with Axiomager M1 (Zeiss). Migrated cells were manually counted and analyzed.

Analysis of focal adhesion morphology
MEFs were cultured on fibronectin-coated coverslip for 1 h, fixed with 3.7% PFA and stained focal adhesions with Vinculin antibody. The Vinculin-staining puncta were analyzed to quantitate size and number of focal adhesions. The mean fluorescence intensity was measured in ImageJ software in a blinded manner.

Single cell migration assay
MEFs were cultured on fibronectin-coated 35 mm dish for 4 h, and observed in a heated chamber at 37°C. Images were taken every 10 min over 4 h using the Axiovert 200M microscope, and cell migration was analyzed using ImageJ mTrackJ plugin.

Wound healing assay
Confluent MEFs were cultured on fibronectin-coated 35 mm dish for 4 h, then scratched with a pipette tip. Immediately after wounding, cultures were fed with fresh medium supplemented with 0.1% fetal bovine serum and observed in a heated chamber at 37°C. Images were taken every 30 min over 12 h using the Axiovert 200M microscope and the wounded area was measured using ImageJ software.

GST-PBD pull-down assays
For GST-PBD pull-down assay, MEFs were processed as detailed elsewhere (Benard and Bokoch, 2002). In brief, cells were plated on fibronectin-coated 100 mm dish for 24 h. Then, the cells were lysed in pull-down buffer (50 mM HEPES, pH 7.4, 150 mM NaCl, 15 mM NaF, 1 mM NaVO₃, 0.5% NP-40, 1 μg/ml leupeptin, 1 μg/ml aprotonin, and 1 μg/ml pepstatin), and centrifuged for 15 min at 22,250g in 4°C. The supernatant was incubated with 5 μg of purified GST-PBD proteins pre-bound to Glutathione Sepharose 4B for 1 h at 4°C, and washed with pull-down buffer three times to remove unbound proteins. Then, the bound proteins were subjected to SDS-PAGE and Western blotting.

Statistics
All data were expressed as mean ± standard error of the mean. Statistical significances for measurements were calculated using Student’s t-tests and defined as *P < 0.05, **P < 0.01, and ***P < 0.001.

RESULTS
Loss of βPix results in embryonic lethality at E9.5
βPix-HET mice were established (Fig. 1A), and intercrossed to generate homozygous βPix-KO mice. Genotyping of the pups revealed that no homozygous βPix-KO mice were born, and only WT and βPix-HET mice were present in the littermates at an approximately 1:2 ratio. This result indicated that total deletion of βPix resulted in embryonic lethality. To identify when embryonic lethality occurs, the pregnant female mice at different gestation days were sacrificed and the embryos were examined. We found growth-retarded βPix-KO embryos between E8.5 and E9.5, and resorption of these embryos at E10. βPix-KO embryos at E8.5 showed complete lack of βPix expression (Fig. 1B). The expression levels of Rac1 and PAK1, which are binding partners of βPix (Koh et al., 2001), were not affected, but the levels of active PAK1, detected by p-PAK1 antibody (Chong et al., 2001), and Git1 were reduced in βPix-KO embryos (Fig. 1B), indicating that βPix deficiency can affect the activity of PAK1 and the expression level of Git1.

To investigate the morphological phenotype of βPix-KO embryos, E9.5 embryos were dissected from the decidua and their gross appearances were observed under a stereomicroscope. βPix-KO embryos were smaller in size than WT embryos, indicating growth retardation with developmental delays (Fig. 1C). Most βPix-KO embryos exhibited impaired axial rotation and incomplete closure of the neural tube from the hindbrain to the forebrain (Figs. 1C and 1E). Another striking feature was the failure of allantois-chorion fusion (Figs. 1D

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and 1E). The allantois of the βPix-KO embryo was unattached to the chorion and resembled balloon-shaped debris (Fig. 1D). Taken together, this demonstrates that βPix plays essential roles in early embryonic development.

βPix-KO MEFs show defects in cell spreading, and PDGF-induced ruffle formation and chemotaxis

βPix-KO MEFs were generated from E8.5 embryos prior to embryonic lethality. βPix-KO MEFs show no expression of βPix, while βPix-rescue MEFs show recovery of βPix expression (Fig. 2A). Given that βPix is a well-known activator of Rac1, we measured the levels of activated Rac1 in the MEFs using GST-PBD pull-down assay. βPix-KO MEFs exhibited decreased levels of Rac1 activation compared to WT MEFs, while βPix-rescue MEFs showed recovery of Rac1 activation (Fig. 2B).

Previously, it was shown that cell spreading on fibronectin is decreased upon knockdown of βPix expression (ten Klooster et al., 2006). Therefore, we investigated the cell spreading of βPix-KO MEFs on fibronectin (Figs. 2C and 2D). The spread area of βPix-KO MEFs decreased by 34% to 37% compared with that of WT MEFs, and βPix-rescue MEFs effectively recovered cell spreading ability (Fig. 2D). These results demonstrate that βPix is important for cell spreading on fibronectin.

βPix and Rac1 play pivotal roles in PDGF responses, including ruffle formation and chemotaxis (Campa et al., 2006). To investigate the effects of βPix deficiency on PDGF-induced ruffle formation, we examined peripheral and dorsal ruffles in the MEFs upon PDGF treatment. Peripheral and dorsal ruffles were found in 40.2% and 13.6% of the WT MEFs, respectively (Fig. 2E). However, in βPix-KO MEFs, the peripheral ruffle-forming cell population decreased by 25% to 54% and the dorsal ruffle-forming cell population decreased by 79% to 93%. Dorsal ruffle formation was more drastically affected by βPix deficiency than peripheral ruffle formation. Additionally, cells bearing both peripheral and dorsal ruffles were also dramatically decreased in βPix-KO MEFs. βPix-rescue MEFs effectively recovered the ruffle-forming abilities. Next, we examined PDGF-induced chemotaxis of βPix-KO MEFs using a transwell assay. Migration of WT MEFs was increased 1.42 fold in response to PDGF (Fig. 2F). However, this increase was not found in βPix-KO MEFs. PDGF-induced chemotaxis was recovered to WT levels in βPix-rescue MEFs. These data indicate that βPix plays critical roles in ruffle formation and chemotaxis in response to PDGF in fibroblasts.

βPix-KO MEFs have larger focal adhesions and increased random cell motility

βPix is known to regulate focal adhesion (Rosenberger and Kutsche, 2006). To examine the effects of βPix deficiency on focal adhesion, focal adhesions in the MEFs were visualized with Vinculin staining, and the size of the focal adhesion was measured (Fig. 3A). In βPix-KO MEFs, the average size of focal adhesion was approximately 30% larger than that in

Fig. 1. Loss of βPix results in embryonic lethality at E9.5. (A) Gene targeting strategy for generation of βPix-KO mice. The exon 19 is replaced by Neo<sup>R</sup> cassette between LoxP sites. (B) Representative blots for βPix-related proteins. Equal amounts of 5 μg homogenates were loaded per lane. Antibodies used for the blots are shown at the left. (C) Representative images for whole mount analysis of E9.5 embryos. H, head; A, allantois. (D) Representative images for histological analysis of E9.5 embryos in deciduae. Black arrows indicate allantois and white arrowheads indicate chorion. (E) Quantification of the defects in axial rotation, neural tube closure and allantois-chorion fusion. Embryos (9-11) were analyzed in each group. **P < 0.01, ***P < 0.001.
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WT MEFs, and this increase was reduced in βPix-rescue MEFs (Fig. 3B). These results showed that βPix plays a critical role in the regulation of focal adhesion in fibroblasts. The turnover of focal adhesions is highly coordinated and important for adhesion-dependent processes, such as cell migration (Raftopoulou and Hall, 2004). It was reported that reduced expression of βPix by RNA interference resulted in defects in cell migration (Hua et al., 2011; Kuo et al., 2011; Sero and Bakal, 2017). Interestingly, in a single cell motility assay, the total migrated distance of βPix-KO MEFs was increased when compared with that of WT MEFs and this increase was reduced in βPix-rescue MEFs (Fig. 3C). Likewise, in a wound healing assay (Fig. 3D), the rate of wound closure was increased in βPix-KO MEFs compared with that of WT MEFs and this increase was reduced in βPix-rescue MEFs (Fig. 3E). These data indicate that a deficiency of βPix induces an increase in cell motility in the MEFs.

βPix-KO MEFs showed enhanced Ser19 phosphorylation of MLC2

A recent study showed that βPix knockdown promotes keratinocyte motility via myosin light chain activation (Hiroyasu et al., 2017). Given that βPix-KO MEFs have increased motility (Figs. 3C-3E), we investigated the level of MLC2 activation by measuring the Ser19 phosphorylation of MLC2 which is commonly used as an index for MLC2 activation (Amano et al., 1996). βPix-KO MEFs showed increased levels of MLC2 phosphorylation, with this increase reduced in βPix-rescue MEFs (Figs. 4A and 4B). Likewise, βPix-KO MEFs showed an increase in p-MLC2 signal by immunocytochemistry (Fig. 4C), and this increase was reduced in βPix-rescue MEFs (Figs. 4C and 4D). These results suggest that βPix deficiency may induce MLC2 activation, thus increasing actomyosin contractility to enhance cell motility.

DISCUSSION

βPix deficiency results in early embryonic lethality

A recent report from Omelchenko et al. (2014) showed that βPix-KO embryos have defects in the collective cell migration of anterior visceral endoderm cells. The defects, such as incomplete closure of the neural tube and impaired axial
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rotation (Figs. 1C and 1E), may be caused by this abnormal collective cell migration. βPix deficiency also shows failure of allantois-chorion fusion (Figs. 1D and 1E). Allantois-chorion fusion largely relies on the interaction between α4 integrin in the chorion and VCAM1 of the allantois (Kwee et al., 1995; Yang et al., 1995). Genomic ablation of α4 integrin or VCAM1 resulted in phenotypes similar to the βPix-KO embryo, in which early placental formation fails. Abi1 deficient mice also show embryonic lethality and failure of placental development (Ring et al., 2011). Abi1 can directly interact with the α4 integrin cytoplasmic tail and is important for integrin α4 cellular signaling. It has also been reported that Abi1 and βPix can directly interact (Campa et al., 2006). Git1, a stable interacting partner of βPix, is also involved in α4 integrin signaling. Git1-paxillin-integrin α4 interaction is required for Rac activation at the leading edge of migrating cells (Nishiya et al., 2005). βPix may play a role in the α4 integrin-VCAM signaling pathway through Abi1 or Git1 to regulate placental development. Thus, βPix deficiency may distort the early placental development resulting in embryonic lethality.

βPix-KO MEF cells show an increased migration distance in cell migration assay
βPix-KO MEFs have defects in cell spreading, and PDGF-induced ruffle formation and chemotaxis (Figs. 2C-2F). These findings suggest that βPix plays essential roles in cell migration. However, unexpectedly, βPix-KO MEF cells showed increased motility in both single cell migration and wound healing assays (Figs. 3C-3E). Although most studies using βPix knockdown cells reported decreased motility in migration assays (Hua et al., 2011; Kuo et al., 2011; Sero and Bakal, 2017), a recent study showed that βPix knockdown promotes keratinocyte motility (Hiroyasu et al., 2017). They observed no difference in activated Rac1 levels and p-MLC protein levels, but p-MLC localization was altered and traction force was increased in βPix knockdown keratinocytes. They suggested that this promotion of cell motility was due to an increase in the traction force generated by enhanced actomyosin contraction, and fine tuning of spatial MLC activity is important. They also showed an increase in the size of focal adhesions in βPix knockdown keratinocytes. Similar phenotypes were observed in βPix-KO MEFs (Figs. 3A and 3B). Unlike βPix knockdown keratinocytes, βPix-KO MEFs exhibited elevated protein levels of p-MLC2 (Fig. 4). PAK family members can directly regulate phosphorylation of MLC through inhibition of the myosin light chain kinase (Sanders et al., 1999). Since βPix is a positive regulator for PAK activity (Koh et al., 2001), βPix-KO MEFs may induce elevated levels of p-MLC2 by repressing the inhibitory role of PAK on MLC2 phosphorylation.

In conclusion, loss of βPix leads to early embryonic lethality in mice, and βPix-KO MEFs exhibit defects in cell spreading, and PDGF-induced ruffle formation and chemotaxis. Interest-

Fig. 3. βPix-KO MEFs have increase in focal adhesion size and cell motility. (A and B) Representative images for focal adhesions visualized by Vinculin-staining (A) and average focal adhesion area (B) in the MEFs. Cells (65-106) from 5 independent experiments were analyzed per group. (C) Total migrated distance of the MEFs in single cell migration assay. At least 53 cells from 3 independent experiments were analyzed in each group. (D) Representative images of wound healing assay for the MEFs. Two dotted lines indicate original boundary lines of scratch. (E) Ratio of wound area for 12 h to 0 h after wound generation is shown. Four experiments were analyzed in each group. *P < 0.05, **P < 0.01, ***P < 0.001.
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ingly, βPix-KO MEFs showed increased cell motility in single cell migration and wound healing assays, probably via inducing MLC2 activation. Detailed studies will further highlight the underlying roles of βPix in embryonic development and cell migration. βPix-KO MEFs will be a valuable tool for dissecting the functions of βPix in diverse cellular processes.

Disclosure
The authors have no potential conflicts of interest to disclose.

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