Prostaglandin Gβγ signaling stimulates gastrulation movements by limiting cell adhesion through Snai1a stabilization

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SUMMARY
Gastrulation movements form the germ layers and shape them into the vertebrate body. Gastrulation entails a variety of cell behaviors, including directed cell migration and cell delamination, which are also involved in other physiological and pathological processes, such as cancer metastasis. Decreased Prostaglandin E₂ (PGE₂) synthesis due to interference with the Cyclooxygenase (Cox) and Prostaglandin E synthase (Ptges) enzymes hampers gastrulation and limits cancer cell invasiveness, but how PGE₂ regulates cell motility remains unclear. Here we show that PGE₂-deficient zebrafish embryos, impaired in the epiboly, internalization, cell movements in the gastrula. Our analyses reveal that PGE₂ promotes cell protrusive activity and limits cell adhesion by converging and extending gastrulation movements, exhibit markedly increased cell-cell adhesion, which contributes to defective cell movements in the gastrula. Our analyses reveal that PGE₂ promotes cell protrusive activity and limits cell adhesion by modulating E-cadherin transcript and protein, in part through stabilization of the Snai1a (also known as Snail1) transcriptional repressor, an evolutionarily conserved regulator of cell delamination and directed migration. We delineate a pathway whereby PGE₂ potentiates interaction between the receptor-coupled G protein βγ subunits and Gsk3β to inhibit proteasomal degradation of Snai1a. However, overexpression of β-catenin cannot stabilize Snai1a in PGE₂-deficient gastrulae. Thus, the Gsk3β-mediated and β-catenin-independent inhibition of cell adhesion by Prostaglandins provides an additional mechanism for the functional interactions between the PGE₂ and Wnt signaling pathways during development and disease. We propose that ubiquitously expressed PGE₂ synthesizing enzymes, by promoting the stability of Snai1a, enable the precise and rapid regulation of cell adhesion that is required for the dynamic cell behaviors that drive various gastrulation movements.

KEY WORDS: Prostaglandin E₂, Gβγ, Gastrulation, Cell adhesion, Snail, Zebrafish

INTRODUCTION
Prostaglandin signaling is important for homeostasis and contributes to digestion, reproduction, pain, immunity, cardiovascular function and stem cell recovery (Wang and Dubois, 2006). In addition, Prostaglandin E₂ (PGE₂) function is associated with increased cancer cell proliferation, anchorage independence (the ability of a cell to survive without anchorage to an extracellular matrix), invasion and angiogenesis in a variety of tumor types, including colorectal, hepatocellular, transitional bladder carcinoma, medullary thyroid, gall bladder and breast cancer (Buchanan and Dubois, 2006; Wang and Dubois, 2006; Backlund et al., 2008). PGE₂ synthesis begins when arachidonic acid, synthesized from membrane phospholipids, forms PGG₂/PGH₂ through the action of Cyclooxygenases (Cox) 1 and 2 [also known as Prostaglandin-endoperoxide synthases (Ptgs) and Prostaglandin G/H synthases (Pghs)]. PGH₂ is converted by Prostaglandin E synthase (Ptges) to PGE₂, which binds and signals via its downstream G-protein-coupled receptors (GPCRs) E-prostanoid (EP) 1-4 [also known as Prostaglandin E receptors (Ptger)] (see Fig. S1A in the supplementary material) (Regan, 2003; Wu, 2006).

Normal fertility and gestation require prostaglandins. Thus, the phenotypes of mouse mutants with inactive components of prostaglandin synthesis or signaling cannot clarify a role for prostaglandins during embryogenesis (Cha et al., 2006b). However, studies in zebrafish, which develop outside the mother, have shown that prostaglandins are required for early vertebrate embryogenesis, specifically gastrulation, a key process of cell signaling and cell movement that structures the body plan (Grosser et al., 2002; Solnica-Krezel, 2005; Cha et al., 2006a). There are four evolutionarily conserved gastrulation movements: epiboly, internalization, convergence and extension (see Fig. S1B,C in the supplementary material). Epiboly, the initial movement of zebrafish gastrulation, thins and spreads embryonic tissues over the yolk cell. During internalization, mesodermal progenitors move underneath the prospective ectoderm. Following internalization, mesodermal cells migrate towards the animal pole, the future head of the embryo. The movements of convergence and extension narrow the germ layers and embryonic body mediolaterally, while extension movements elongate the embryonic tissues head to tail (Solnica-Krezel, 2005). Each of these cell movements is the outcome of individual cell migratory behaviors that, interestingly, require cell signaling pathways, such as Phosphoinositide-3-kinase (Pik3/Pi3K), that are conserved in migrating cancer cells (Montero et al., 2003; Fujino et al., 2002; Fujino and Regan, 2003).
Previously, prostaglandin synthesis in zebrafish embryos was reduced with enzymic inhibitors of Cox1 (Ptgs1–Zebrafish Information Network) or antisense morpholino oligonucleotides (MO) that inhibit the translation of Cox1 (cox1 MO, MO1-pg) or Ptges (ptges MO, MO2-pgtes). These manipulations resulted in an epiboly delay or arrest, largely owing to depletion of PGE2, the predominant prostaglandin in zebrafish gastrulae (Grosser et al., 2002; Cha et al., 2006a). Additionally, lowering PGE2 signaling with a low dose of pgtes MO (2 ng) resulted in a convergence and extension defect due to the decreased speed of dorsally migrating lateral mesodermal cells (Cha et al., 2006a), suggesting that different gastrulation movements require distinct levels of PGE2.

Here, we extend these analyses to reveal that PGE2 influences all of the gastrulation movements in zebrafish. We have characterized the movement defects manifest in PGE2-deficient gastrulae by time-lapse imaging to evaluate cell motility and protrusive activity, and found that both are impaired with decreased PGE2 synthesis. Further analysis revealed that embryos with decreased PGE2 have markedly increased cell-cell adhesion, which might contribute to the observed movement defects and is the first evidence that PGE2 limits cell adhesion during development. We also delineate a signaling mechanism whereby PGE2 stabilizes the Snai1a protein, an inhibitor of E-cadherin (cdh1) transcription (Barrallo-Gimeno and Nieto, 2005), by preventing its proteasomal degradation as promoted by Gsk3β. Furthermore, PGE2 limits the inhibition of Snai1a by Gsk3β by potentiating a novel interaction between the Gβγ effector subunits of PGE2 signaling and Gsk3β. We propose that ubiquitously expressed PGE2 synthesizing enzymes promote the stability of Snai1a to allow precise and rapid regulation of the cell adhesion that is required for the dynamic cell behaviors of gastrulation.

**MATERIALS AND METHODS**

**Zebrafish strains and maintenance**
Embryos were obtained from natural matings and staged according to morphology as described (Kimmel et al., 1995). With the exception of experiments using the MZ oep^57ts5 (Gritsans et al., 1999) and the Tg(gsc:GFP)^tso (Doitsidou et al., 2002), all experiments were performed using wild-type embryos.

**Embryo injection**
Zebrafish embryos were injected at the one-cell stage, 15-45 minutes post-fertilization. The injected antisense MOs included the control MO5/bp-mismatch MO2-pgtes (5'-GTGTATCTGCTGTTAGGTC-3'), pgtes MO/ MO2-pgtes (Cha et al., 2006a), coxl MO/MO1-pgtes (Grosser et al., 2002) and cdh1 MO/MO3-cdh1 (Babb and Marrs, 2004). RNA constructs for synthetic RNA used for injection included megfp, yfp (Yamashita et al., 2004), snai1a-yfp (T. Hirano laboratory, Osaka University, Osaka, Japan) (Yamashita et al., 2004), zsgkβ3β (M. Hibi, Riken Center for Developmental Biology, Kobe, Japan), βδi, βδi, DNa-catenin and ncd2/cyclops (C. V. Wright laboratory, Vanderbilt University, Nashville, TN, USA) (Elder et al., 1998). Embryos were injected with snai1a-Ha RNA at the 8- to 16-cell stage. All RNA constructs were in the pCS2 vector.

**Snai1a-YFP assay**
Injected embryos were chemically treated as described below and incubated at 28°C until the shield stage. All experiments described were performed at least three times with at least 30 injected embryos per sample (per experiment). Live embryos were oriented using the shield as a morphological landmark in 2% Methylcellulose/0.3× Danieau. Representative embryos were imaged on an LSM 510 confocal microscope (Carl Zeiss MicroImaging, Thornwood, NY, USA), using the 10× objective. Experiments were performed in part through the VUMC Cell Imaging Shared Resource. Images were prepared for publication using Velocity software (Improvision, Coventry, UK) and Adobe Photoshop (Adobe, San Jose, CA, USA).

**Embryo treatment**

**PGE2**
Embryos were treated in 1% DMSO/embryo medium with synthetic PGE2 (10 mM; Cayman Chemical, Ann Arbor, MI, USA) at two time points: following injection and at the dome stage (Cha et al., 2006a).

**Proteasomal inhibitor**
Embryos were treated with Z-Leu-Leu-Leu-H/ MG132 (Peptide Institute, Minoh, Osaka, Japan) (50 μM) (Zhou et al., 2004) from the 128- to 256-cell stage in 1% DMSO/embryo medium until the embryos were imaged at the shield stage.

**Gsk3β inhibitor**
LiCl treatment was performed as described (Stachel et al., 1993). LiCl (Sigma-Aldrich, St Louis, MO, USA) (0.3 M) was added to the embryo medium at the 256-cell stage for 10 minutes, the embryos then rinsed three times in 0.3× Danieau and incubated until the shield stage. Gsk3β BIO (Stemgent, Cambridge, MA, USA) (1 μM) was added in the 1000-cell stage and left until the shield stage.

**Pik3 inhibitor**
Embryos were incubated in 30-50 μM LY294002 (Cayman Chemical) in embryo medium from the dome until the shield stage.

**Cell adhesion assays**
Cell adhesion assays were performed essentially as described (Ulrich et al., 2005). The cells from dissociated blastulae were diluted to 50,000 cells/ml and plated in a fibronectin-coated 96-well plate (5000 cells/well). Images were taken on a SteREO Discovery V12 Dissecting Microscope (Carl Zeiss MicroImaging) every hour for 3 hours.

**Quantitative real-time (qRT) PCR**
RNA was extracted from 20 injected embryos per sample at 60% epiboly using 200 μl Trizol (Tingaud-Sequeira et al., 2004). Following RNA extraction, the samples were diluted to 50 ng/μl. Primers included β-actin (Tingaud-Sequeira et al., 2004), cdh1 (forward, 5'-TGAAGGGCTGCA-GATAACGAC-3'; reverse, 5'-GTGTTGAGGAGCGTAGTGA-3') and snai1a (forward, 5'-GAGCTGGAATGTCAGAAG-3'; reverse, 5'-GTGAAGGGAAGGTAGCAAG-3'). Samples were prepared for qRT-PCR using the iScript One-Step RT-PCR Kit with SYBR Green (Bio-Rad, Hercules, CA, USA). For each sample, 100 ng of template DNA was used, and water was used as a negative control for the no reverse transcriptase (-RT) reaction. The qRT-PCR was performed on an iCycler iQ Multicolor machine (Bio-Rad) at the VUMC Molecular Biology Resource Core. The annealing temperature was 60°C without a temperature gradient. The data shown represent three separate experiments with duplicate samples. The data were analyzed with iQ5 Optical System Software, version 2.0 (Bio-Rad).

**Time-lapse imaging**

**Shield time-lapse**
Embryos were injected with membrane egfp RNA. At the shield stage, embryos were dechorionated in 0.3× Danieau and oriented in MatTek PG35G-0-10-C glass-bottom dishes using 0.8% SeaPlaque Low-Melt Agarose (Lonza, Rockland, ME, USA) in 0.3× Danieau. Images were taken of the shield on a Zeiss Axiovert 200 inverted microscope (Carl Zeiss MicroImaging) with an ERS Spinning Disk Confocal system (PerkinElmer, Fremont, CA, USA) using the 40× oil-immersion (N.A.=1.3) objective. Images of 2-sections (0.5 μm) were taken every minute. The resulting data points were orthogonally reconstructed using Velocity software.

**Protrusion movie analysis**
*membrane egfp or membrane rfp* RNA was injected into Tg(gsc:GFP)^tso zebrafish embryos (labeled donor embryos). Fewer than ten cells were transplanted from donor embryos to unlabeled host embryos at the shield stage. Host embryos containing transplanted cells that were gsc:GFP-positive were mounted at 70-80% epiboly in SeaPlaque agarose in 0.3× Danieau in glass-bottom dishes as described above, with the shield facing downward. Images were taken on the Axiovert 200 inverted microscope/ERS Spinning Disk Confocal system using the 40× oil-immersion.
Whole-mount in situ hybridization (ISH)
Embryos were collected at the indicated stage and fixed with 4% paraformaldehyde overnight at 4°C. Whole-mount ISH was performed as described (Thibie and Thibie, 1998). Probes included snail1 (Hammerschmidt and Nusslein-Volhard, 1993), snailb (Blanco et al., 2007), frizzled 8b (Kim et al., 1998) and no tail (Schulte-Merker et al., 1992).

Western blot
Embryos were homogenized with RIPA buffer (10 mM Tris-HCl pH 7.4, 150 mM NaCl, 1 mM EDTA, 0.1% SDS, 1% Triton X-100, 1% sodium deoxycholate), then frozen at −20°C. Embryo homogenates were lysed with 2× lysis buffer (62.5 mM Tris-HCl pH 6.8, 10% glycerol, 2% SDS, 0.5% β-mercaptoethanol, 0.01% Bromophenol Blue) and heated at 100°C for 5 minutes. Extracts were resolved in a 4-15% polyacrylamide gel and transferred to a PVDF membrane using the Criterion System (Bio-Rad). The antibodies used were anti-zebrafish Snail1 (1:500; Hammerschmidt laboratory, Max-Planck-Institute of Immunobiology, Freiburg, Germany) (Hammerschmidt and Nusslein-Volhard, 1993), anti-zebrafish E-cadherin (1:1000; J. A. Marrs laboratory, Indiana University Medical Center, Indianapolis, IN, USA) (Babb and Marrs, 2004), anti-Gsk3β (1:1000; Cell Signaling Technology, Danvers, MA, USA), anti-phospho-Gsk3β (1:100; Cell Signaling Technology) and anti-Gapdh (1:500; RDI Division of Fitzgerald Industries, Concord, MA, USA). For full scans of all western blots depicted, see Fig. S5 in the supplementary material.

Whole-mount immunohistochemistry
Zebrafish embryos were collected at the indicated stage and fixed with a 1:1 solution of 8% paraformaldehyde and 2× fix buffer (8% sucrose, 0.3 mM CaCl2 in PBS pH 7.3). Embryos were rinsed four times with PBS containing 0.1% Tween 20. Antibodies used were anti-zebrafish E-cadherin (as above) and anti-ZO1 (Tjp1) (1:200; Zymed Laboratories, San Francisco, CA, USA). To visualize nuclei, samples were stained with SYTO 59 (Invitrogen) for 30 minutes prior to imaging, then rinsed twice with PBS containing 0.1% Tween 20 and 2% DMSO.

Cell culture, transfection and immunoprecipitation
HEK-293T cells were cultured in DMEM (Cellgro) media supplemented with 100 units/ml penicillin, 100 unit/ml streptomycin (Gibco) and 10% fetal bovine serum (FBS) (Gibco). Lipofectamine 2000 (Invitrogen) was used for transfections following the manufacturer’s protocol. HEK-293T cells were plated in 30 mm plates and transfected with 1 μg Gsk3β, 1 μg HA-Gβγ, 1 μg Gγ and 2 μg β-Ark (all vectors are pCS2), and cultured for 48 hours post-transfection. For the PGE2 treatment, cells were then treated with 0.1, 1 and 10 μM PGE2 for 14 hours. Cells were washed once with cold PBS and then lysed for 30 minutes on ice in non-denaturing lysis buffer (NDLB) comprising 50 mM Tris-HCl pH 7.4, 300 mM NaCl, 5 mM EDTA, 1% (w/v) Triton X-100 and protease inhibitors (1 mg/ml leupeptin, pepstatin and chymostatin). Then, 750 μg of lysate was diluted to 1 mg/ml with NDLB. Rat anti-HA (Roche) was cross-linked to protein G magnetic beads (New England Biologicals) following the manufacturer’s protocol. Anti-HA protein G beads were then added to the lysate and incubated for 2 hours with rotation at 4°C. Beads were then washed three times with NDLB and once or twice with PBS. Protein was eluted from the beads with sample buffer. Samples, as well as 10 μg of total protein lysate, were then run on SDS-PAGE gels and transferred to a nitrocellulose membrane. Membranes were incubated in 5% milk solution (1 hour at room temperature) and probed with mouse anti-HA (1:1000; 12CA5, Santa Cruz, CA, USA), mouse anti-α-tubulin (1:3000; DM1a, Sigma) and mouse anti-Gsk3 (1:1000; BD Biosciences) for the immunoprecipitation without PGE2 treatment, and with rabbit anti-Gβγ (1:500; T-20, Santa Cruz), mouse anti-α-tubulin (as above), and rabbit anti-Gsk3 (1:1000; 27C10, Cell Signaling) for the immunoprecipitation after PGE2 treatment. HR-conjugated anti-mouse and anti-rabbit antibodies were used to detect primary antibody by chemiluminescence.

RESULTS

Decreased PGE2 synthesis results in global cell movement defects during gastrulation
To detail the mechanisms by which PGE2 regulates gastrulation, we focused on higher dose (4 ng) MO2-ptges-injected embryos that displayed strong and global gastrulation defects (Fig. 1A-C), paralleling those reported for coxl MO or enzymatic inhibitors (Grosser et al., 2002; Cha et al., 2006a). These defects in ptges morphants were suppressed by supplementing the embryo medium with PGE2 and were not observed when a control 5-bp-mismatch MO (control MO) was injected. Ptges- and Coxl-deficient gastrulae showed arrested epiboly and an uneven surface associated with cell clumping (Fig. 1A; see Fig. S2A in the supplementary material). Whereas chordamesodermal cells expressing no tail (ntf; also known as brachury) were in the deeper, mesendodermal layer of control morphant gastrulae, Ptges-deficient gastrulae had ntf-expressing cells in the superficial layer, indicating that internalization was defective in the most strongly affected embryos (Fig. 1B). In less severely affected Ptges-deficient gastrulae, internalization did occur (Fig. 1C). However, migration towards the animal pole, representing the subsequent movement of mesodermal cells, was impaired relative to that in control gastrulae, as indicated by the posteriorly shifted position of frizzled homolog 8b (fzd8b)-expressing prechordal mesoderm cells (Fig. 1C). All these movement defects were partially suppressed with synthetic PGE2 treatment, confirming the specific and essential role of PGE2 in gastrulation movements. Defective gastrulation movements, cell paths and net speed were also revealed by time-lapse analysis of the dorsal shield, which is equivalent to the Spemann-Mangold organizer of amphibians (Fig. 1D,E; see Fig. S1F and Movies 1, 2 in the supplementary material). In addition, the boundary (Brachet’s cleft) between the internalized mesendodermal cells and the superficial layer was distinct in the control, but unclear in Ptges-deficient, gastrulae (Fig. 1D; see Movies 1, 2 in the supplementary material). Together, these data reveal that PGE2 signaling is essential for epiboly and internalization in addition to convergence and extension movements during gastrulation (Grosser et al., 2002; Cha et al., 2006a).

PGE2 synthesis is required for normal protrusion formation in gastrula cells
Defects in cell movements during gastrulation can occur as a result of impaired protrusive activity, cell adhesion, cytoskeleton dynamics and cell polarity (Montero et al., 2003; Montero and Heisenberg, 2004; Solnica-Krezel, 2005; Ulrich et al., 2005). Thus, we sought to evaluate how cell motility was undermined in Ptges-deficient embryos. First, we analyzed protrusive activity, which, when defective, can hinder the motility of gastrula cells (Montero et al., 2003). Time-lapse imaging of membrane EGFPP–labeled cells demonstrated that at the interface of the blastoderm and the yolk cell during early gastrulation, cells exhibited protrusions with few blebs in the control morphants (see Fig. S1D in the supplementary material). ptges morphant cells, however, showed increased blebbing (see Fig. S1E in the supplementary material). From 60-90% epiboly, the protrusions of the prechordal mesodermal cells migrating towards the animal pole in control gastrulae were active and dynamic, with filopodial characteristics (Montero et al., 2003), Statistical analysis
Statistical analyses of the qRT-PCR and cell behavior experiments were performed with Microsoft Excel software. The raw data were processed to calculate the mean and the s.e.m. (as indicated by error bars in the figures). Statistical significance was evaluated by Student’s t-test.
and all protrusions localized to the leading edge of the cell (Fig. 1F; see Fig. S1G-I and Movie 3 in the supplementary material). By contrast, the protrusions of the prechordal mesodermal cells in gastrulae with reduced Ptges function (2 ng MO) were blunted, with lamellipodial, sometimes bleb-like, characteristics (Montero et al., 2003). Moreover, protrusions localized to all cell edges (Fig. 1F; see Fig. S1H and Movie 4 in the supplementary material). Therefore, impaired gastrulation movements in PGE2-depleted embryos are associated with abnormal protrusive activity.

PGE2 negatively regulates cell-cell adhesion in zebrafish gastrulae

Unbalanced cell adhesion, either in excess or deficit, cripples the movement of primordial germ cells in zebrafish, of border cells in D. melanogaster and gastrulation movements in many animals (Blaser et al., 2005; Pacquelet and Rorth, 2005; Hammerschmidt and Wedlich, 2008). In particular, normal gastrulation in all animals requires appropriate levels of E-cadherin (also known as Cadherin 1 or epithelial Cdh1) (Solnica-Krezel, 2006; Hammerschmidt and Wedlich, 2008), a component of the adherens junction that binds β-catenin (Drees et al., 2005). Moreover, zebrafish cdh1 [also known as half baked (hab)] mutants manifest defects in epiboly, internalization, convergence and extension (Babb and Marrs, 2004; Kane et al., 2005; McFarland et al., 2005; Montero et al., 2005; Shimizu et al., 2005; von der Hardt et al., 2007). The requirement of direct and indirect inhibitors of E-cadherin, namely Gα12/13, Wnt11 and p38 (Mapk14a), for normal gastrulation movements underscores the significance of the precise and diverse regulation of cell adhesion in this process (Lin et al., 2005; Ulrich et al., 2005; Zohn et al., 2006; Lin et al., 2009).

Because of the cell clumping observed in Ptges-deficient gastrulae (Fig. 1A), we hypothesized that increased cell adhesion contributed to the gastrulation movement phenotype. To evaluate cell adhesion in Ptges-deficient embryos, we carried out cell adhesion assays using zebrafish embryonic cells ex vivo (Ulrich et al., 2005). First, we induced mesendodermal fates in embryonic cells by co-injecting zygotes with synthetic nodal-related 2 (ndr2; also known as cyclops and znr1) RNA (see Fig. S2F in the supplementary material) and ptges or control MOs. The resulting embryos were dissociated into single cells at the dome stage, then seeded into fibronectin-coated wells for a 3-hour time-course to determine whether cells had formed aggregates. Cells from control morphants showed clumps of two or three cells at the end
of the 3-hour time-course. Strikingly, the ptges MO injection increased cell clumping in a dose-dependent manner. In fact, at the end of the experimental period, most of the cells from embryos injected with the high MO dose were in one or two large three-dimensional clumps (containing thousands of cells) that dominated the well (Fig. 2). We conclude that cell adhesion is increased in Ptges-deficient gastrulae, indicating that PGE₂ signaling can regulate gastrulation movements by limiting cell adhesion.

The excessive cell adhesion in Ptges-deficient gastrulae might be caused by increased E-cadherin. Accordingly, using qRT-PCR we found that cdh1 transcript levels relative to those of β-actin were twofold higher in Ptges-deficient embryos than controls (Fig. 3A). Immunoblotting analysis showed that the protein levels of E-cadherin were increased twofold relative to those in controls (Fig. 3B). Moreover, whole-mount immunohistochemistry revealed that E-cadherin expression was uniformly increased in the mesendoderm of ptges morphants (Fig. 3C). Thus, PGE₂ functions during gastrulation to limit E-cadherin transcript and protein expression.

To investigate whether disrupting E-cadherin translation could suppress aspects of the gastrulation phenotype in PGE₂-deficient gastrulae, we co-injected cdh1 (MO3-cdh1) and ptges MOs. Downregulation of E-cadherin suppressed the cell clumping phenotype, but did not significantly improve the epiboly defect (Fig. 3D,E). Hence, enhanced cell adhesion contributes in part to the gastrulation defects seen in Ptges-deficient embryos.
Snai1a protein expression is stabilized in the presence of PGE2

The increase of \(\text{cdh1}\) transcripts in the Ptges-deficient embryos suggested that PGE2 signaling positively regulates a transcriptional repressor of the \(\text{cdh1}\) gene. Snai1, the best characterized of these, binds E-boxes in the E-cadherin (\(\text{cdh1}\)) promoter to inhibit its transcription (Barrallo-Gimeno and Nieto, 2005). Snai1 conditional mouse mutants show defective formation of the mesoderm layer because of persistent epithelial morphology, as well as impaired anterior migration of mesodermal cells (Carver et al., 2001). Furthermore, Snai1 is required for internalization movements in Drosophila and sea urchin (Barrallo-Gimeno and Nieto, 2005; Wu and McClay, 2007). There are four zebrafish Snail genes, two of which have arisen through duplication of \(\text{snai1}\) in the teleost lineage. \(\text{snai1a}\) and \(\text{snai1b}\) are expressed in the internalizing mesendoderm during gastrulation and are required for the migration of anterior mesendodermal cells towards the animal pole (Hammerschmidt and Nusslein-Volhard, 1993; Yamashita et al., 2004; Blanco et al., 2007). In addition, \(\text{snai1b}\) morphants display convergence and extension defects (Blanco et al., 2007). Altogether, these data imply that Snai1a and Snai1b are required for the gastrulation movements.

Recent work in cell culture has shown that chemical or antisense interference with the prostaglandin signaling pathway components results in decreased Snail transcription and increased E-cadherin protein levels (Dohadwala et al., 2006; Brouxhon et al., 2007). However, our qRT-PCR and whole-mount ISH analyses showed that \(\text{snai1a}\) and \(\text{snai1b}\) levels were not significantly different in \(\text{ptges}\) versus control morphants (Fig. 4A; see Fig. S3A,B in the supplementary material) until 60% epiboly, when the Ptges-deficient gastrulation phenotype is already apparent (see Fig. S3B in the supplementary material). Therefore, we conclude that the regulation of \(\text{snai1a}\) or \(\text{snai1b}\) transcription by PGE2 could not cause the gastrulation defects in this developmental context. By contrast, western blotting with an anti-Snai1a antibody, raised against the full-length protein (Hammerschmidt and Nusslein-Volhard, 1993), revealed that the Snai1a protein level was significantly reduced in embryos injected with \(\text{ptges}\) MO in a dose-dependent manner (Fig. 4B), demonstrating the post-transcriptional regulation of Snai1a by prostaglandins.

To determine whether the localization or stability of Snai1a was affected in Ptges-deficient gastrulae, we analyzed the expression of the Snai1a-YFP fusion protein (Yamashita et al., 2004) in live embryos. We injected synthetic \(\text{snai1a-yfp}\) (Yamashita et al., 2004),
membrane egfp and yfp RNAs containing identical 5’ and 3’ UTR regions into one-cell stage zebrafish embryos and analyzed the resulting proteins by confocal microscopy in the shield stage gastrulae. The injection of ptges MO resulted in the dramatic and dose-dependent reduction of Snai1a-YFP expression as compared with that in control embryos (Fig. 4C). By contrast, YFP and membrane (m) EGFP expression were comparable between gastrulae injected with ptges and control MOs. Loss of Snai1a-YFP expression in Ptges-deficient gastrulae was suppressed by PGE2 treatment. Furthermore, coxl morphants also showed decreased Snai1a-YFP expression (see Fig. S4B in the supplementary material). Interestingly, injection of RNA encoding Snai1b-YFP showed only a mild decrease in YFP expression in the Ptges-deficient as compared with control gastrulae (see Fig. S3C in the supplementary material). These results demonstrate that PGE2 signaling can regulate the protein expression of Snai1a and Snai1b. However, Snai1a seems more sensitive to changes in PGE2 levels than Snai1b.

To test whether the increased cdh1 transcript and E-cadherin protein levels were an outcome of reduced Snai1a expression, snai1a-hA RNA was injected mosaically at the eight-cell stage into embryos injected with ptges MO at the one-cell stage (Fig. 4D). The quantification of E-cadherin immunostaining showed that Snai1a- HA-expressing cells had significantly lower E-cadherin levels than surrounding HA-negative cells in these Ptges-deficient gastrulae (Fig. 4E). This suggested that the increased E-cadherin expression in Ptges-deficient mesendodermal cells was suppressed by restored Snai1a expression. Therefore, we conclude that PGE2 signaling stabilizes Snai1a protein to limit E-cadherin expression during gastrulation.

PGE2 signaling-associated Gβγ subunits stabilize Snai1a by interaction with Gsk3β

To define the molecular mechanism by which PGE2 promotes Snai1a stability, we evaluated the outcome of manipulating possible targets of PGE2 signaling on misexpressed Snai1a-YFP protein levels. First, we tested whether proteolysis contributed to the loss of Snai1a-YFP expression in Ptges-deficient embryos (Dominguez et al., 2003; Zhou et al., 2004). Accordingly, the treatment of Ptges-deficient embryos with a proteasomal inhibitor, MG132 (Zhou et al., 2004), fully suppressed the loss of Snai1a-YFP expression, indicating that proteasomal degradation could decrease Snai1a protein levels (Fig. 5A). Cell culture studies have shown that PGE2 can regulate Glycogen synthase kinase 3β (Gsk3β) (Fujino et al., 2002; Fujino and Regan, 2003). Given that Gsk3β phosphorylates Snai1, thereby targeting it for proteasomal degradation (Zhou et al., 2004; Yook et al., 2006), we investigated whether this kinase regulates Snai1a expression downstream of PGE2 signaling. Consistent with this notion, the injection of gsk3β RNA significantly decreased the level of misexpressed Snai1a-YFP in a dose-dependent manner in control embryos, and caused the complete loss of Snai1a-YFP in Ptges-deficient gastrulae (Fig. 5B). Conversely, Gsk3β inhibition through either LiCl (Stachel et al., 1993) or β-bromoindirubin-3-oxime (BIO) (Goessling et al., 2009) treatment, suppressed the reduction of Snai1a-YFP levels in Ptges-deficient gastrulae. Blocking the function of both Gsk3β and the proteasome resulted in the full restoration of Snai1a-YFP levels in Ptges-deficient embryos, similar to embryos treated with a proteasomal inhibitor alone (Fig. 5D). Thus, PGE2 signaling functions to stabilize Snai1a by inhibiting its proteasomal degradation, which is promoted by Gsk3β.

Previous studies indicate that PGE2 signaling acts in part through heterotrimeric Guanine nucleotide-binding proteins (G proteins) (Buchanan and Dubois, 2006). Following activation of the EP receptors, G protein α and βγ subunits stimulate distinct downstream effectors. Gβγ protein subunits have been shown in cell culture to regulate Gsk3β downstream of PGE2 signaling by the activation of Pik3 (Fujino et al., 2002; Fujino and Regan, 2003; Cha et al., 2006a). We found that overexpression of the Gβγ2 (Gb1n, Gmg2) subunits elevated the Snai1a-YFP levels in Ptges-deficient gastrulae, suggesting that the Gβγ subunits activated by the EP receptors were responsible for conveying the effects of PGE2 on Snai1a (Fig. 5C). Co-injection of RNAs encoding Gβγ2 and Gsk3β into Ptges-deficient embryos, however, blocked the rescue of Snai1a-YFP seen with the RNA encoding Gβγ2 alone, suggesting that Gsk3β regulates Snai1a downstream of the Gβγ subunits to inhibit Snai1a proteasomal degradation (Fig. 5E; Fig. 6). Accordingly, Gsk3β was detected following the immunoprecipitation of anti-HA in HEK-293T cells co-transfected with zebrafish Gsk3β and HA-human Gβγ2. This interaction was inhibited by co-expressing the C-terminal domain of the β-adrenergic receptor kinase (c-βark), which competes for Gβγ2 binding (Fig. 5F). These results identify Gsk3β as a potential new Gβγ effector protein downstream of PGE2 (Fig. 5F). In addition, PGE2 treatment of transfected HEK-293T cells increased, in a dose-dependent manner, the level of Gsk3β following immunoprecipitation of anti-HA (Fig. 5G), suggesting that PGE2 can promote the interaction of Gsk3β with Gβγ. This interaction between the Gβγ subunits and Gsk3β provides a novel mechanism for the regulation of the Snai1a protein downstream of PGE2 signaling.

β-catenin does not promote Snai1 stability

The regulation of Gsk3β by Gβγ signaling downstream of PGE2 presents a possible junction between prostaglandin and Wnt signaling, lending insight into the finding that Cox inhibitors (NSAIDs) decrease the intestinal tumor load of Apcmin mice (Montero et al., 2003). We also employed a Pik3 inhibitor, LY294002, which has been shown to impair zebrafish gastrulation (Montero et al., 2003), to investigate the effects of Pik3 downregulation on Snai1a-YFP levels. Overexpression of Pik3γ or inhibition of Pik3 resulted in gastrulation defects as previously observed (Montero et al., 2003) (F.L. and L.S.-K., unpublished observations), confirming that Pik3 activity was being effectively manipulated (data not shown). However, neither treatment significantly altered Snai1a-YFP expression in control or Ptges-deficient embryos (see Fig. S4A in the supplementary material). Thus, the interaction between Gβγ and Gsk3β to regulate Snai1a is unlikely to involve the regulation of Gsk3β by Pik3.
to enhance β-catenin levels and contribute to hematopoietic stem cell recovery (Goessling et al., 2009). However, we found that the overexpression of ΔNβ-catenin, a constitutively active form, at doses sufficient to induce ectopic body axes in injected embryos (Kelly et al., 1995a) (data not shown) had a minimal effect on Snai1a-YFP expression and could not suppress the decrease in Snai1a-YFP when co-injected with the ptges MO (Fig. 5F). Hence, the effect of PGE2 in stabilizing Snai1a through inhibiting its Gsk3β-mediated degradation does not occur downstream of β-catenin activation.

**DISCUSSION**

Here, we demonstrate that strong reduction of PGE2 synthesis by interference with Cox1 and Ptges enzymes impairs all epiboly and internalization gastrulation movements, adding to the previously documented defects in convergence and extension reported at intermediate PGE2 deficiency (Grosser et al., 2002; Solnica-Krezel, 2005; Cha et al., 2006a). Our data show that these widespread gastrulation defects in PGE2-deficient embryos are in part due to increased cell-cell adhesion. PGE2 signaling limits cell adhesion during gastrulation by modulating E-cadherin transcript and protein, in part through stabilization of Snai1. Our results suggest that PGE2 can regulate the expression of both Snai1a and Snai1b, although Snai1a is much more sensitive to changes in PGE2 synthesis. Snai1a and Snai1b are expressed in different domains of the gastrula (Blanco et al., 2007). Both proteins function non-redundantly in anterior migration of the prechordal plate through repression of E-cadherin. In addition, loss of Snai1b by MO targeting results in a convergence and extension defect. Therefore, it is possible that PGE2 inhibits E-cadherin via distinct Snai1a- and Snai1b-dependent mechanisms, as Snai1b does not contain complete Gsk3β phosphorylation domains. Determining whether PGE2 stabilizes Snai1a and Snai1b by distinct mechanisms is an important future direction because each genetic interaction might lead to the regulation of different cell movements. Although their complementary expression patterns suggest that Snai1a and Snai1b...
deficient gastrulae, but was unable to restore normal gastrulation movements of gastrulation. In this manner, PGE2 impinges on cell adhesion.

We provide evidence that PGE2 negatively regulates E-cadherin expression in part through the stabilization of Snai1a by preventing its Gsk3β-mediated proteasomal degradation (Fig. 6). PGE2 signaling inhibits Gsk3β via its downstream G protein βγ subunits, a novel molecular mechanism by which PGE2 can promote Snai1a function and limit cell adhesion to influence motility. We have shown that the inhibition of Gsk3β by Gβγ is not dependent on Pik3. Moreover, Gsk3β and Gβγ proteins can be co-immunoprecipitated when expressed in mammalian tissue culture (Fig. 5F,G). Therefore, we present an alternative pathway by which PGE2 can regulate Gsk3β without the need for second messengers. This interaction between Gsk3β and Gβγ proteins has also been observed in human cell culture [Gβγ promotes LRP6-mediated β-catenin/TCF signaling by stimulating plasma membrane localization and activation of GSK3 (K.K.J., C. S. Cselényi, C. Thorne, N. Hajicek, W. Oldham, L. A. Lee, H. E. Hamm, J. R. Hepler, T. Kozasa, M. E. Linder and E.L., unpublished)], suggesting that this mechanism occurs in multiple cellular contexts and vertebrate species. Although the molecular mechanism by which Gβγ inhibits Gsk3β during gastrulation remains to be elucidated, the work by Jernigian et al. suggests that Gβγ binds Gsk3β, sequestering it to the membrane to activate its kinase activity on the co-receptor LRP6, leading to the inhibition of β-catenin degradation and the potentiation of β-catenin/TCF-mediated transcription. In addition, in breast cancer cells, the presence of Gsk3β in the nucleus is essential for its silencing of SNAI1 activity. When AXIN2 acts as a nucleocytoplasmic chaperone for GSK3β, exporting it from the nucleus, SNAI1 remains active (Yook et al., 2006). Therefore, we speculate that following activation by PGE2, Gβγ binds Gsk3β at the membrane to prevent it from inhibiting SNAI1 activity in the nucleus.

Although our data suggest that PGE2 does not regulate SNAI1 stability via β-catenin, it is still possible that PGE2 activates β-catenin by the inhibition of Gsk3β. Interestingly, a previous study has shown that β-catenin can repress Cdh1 transcription through noggin-activated LEF1 in mouse hair follicles (Jamora et al., 2003). If this mechanism also operates in gastrulae, PGE2 might employ β-catenin-dependent and -independent mechanisms to repress Cdh1 expression. However, as we have previously demonstrated, PGE2-deficient gastrulae show relatively normal axis specification (Cha et al., 2006a), suggesting that PGE2 and Wnt/β-catenin might interact differently depending on the cellular context.

Sna1a expression is increased in multiple tumor types (Barrallo-Gimeno and Nieto, 2005; Yook et al., 2006) and promotes the recurrence of breast cancer in vivo (Moody et al., 2005). In addition, the activation of EGF signaling, which plays a major role in many cancers, increases the expression of Sna1, emphasizing its role during tumorigenesis (Lu et al., 2003; Mann et al., 2006; Backlund et al., 2008). PGE2 signaling has been correlated with increased cancer cell invasiveness, angiogenesis and anchorage independence (Wang and Dubois, 2006), properties that allow cancer cells to exit the primary tumor and migrate to secondary sites (metastasis). Increased expression of the EP4 receptor has also been reported in colon and breast cancer cells (Chell et al., 2006), indicating that cancer cells can utilize native regulation of the cell motility machinery by PGE2. Therefore, our discovery of the Sna1a-dependent repression of cell adhesion by PGE2 might lend insight into the mechanism by which prostaglandins promote tumor cell motility and metastasis. Because Wnt signaling also inhibits Gsk3β, Wnt and PGE2 pathways can converge, via β-catenin, to promote hematopoietic stem cell survival (North et al., 2007; Goesseling et al., 2009). We suggest that the Wnt and PGE2 pathways...
might also promote cell motility by inhibiting cell adhesion. We also speculate that the repression of E-cadherin by PGE2 might apply to other roles of prostaglandins/PGE2, for example in inflammation and hematopoietic stem cell recovery.

Acknowledgements
We thank L.S.-K. laboratory members for helpful discussions and comments; Heiki Beck and the SC Facility Research Assistants for fish care; Drs M. Hammerschmidt (Snai1 antibody), J. A. Marrs (zCd1 antibody), T. Hirano (Snai1a-YFP construct), M. Hibi (zGsk3b construct), A. Nieto (Snail1b construct), C. Wright (zNdr2 construct), and C. Hong and K. Friedman for providing experimental reagents. Confocal experiments were performed in the VUMC Cell Imaging Core Facility (supported by NIH grant 1S10RR015682). This work was supported in part by NIH grant GM77770 to L.S.-K. Deposited in PMC for release after 12 months.

Competing interests statement
The authors declare no competing financial interests.

Supplementary material
Supplementary material for this article is available at http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.045971/-/DC1

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