The effect of prostaglandin E2 (PGE2) on bone mass has been well-established in vivo. Previous studies have showed that PGE2 increases differentiation, proliferation, and regulates cell morphology through F-actin stress fiber in statically cultured osteoblasts. However, the effect of PGE2 on osteoblasts in the presence of fluid shear stress (FSS), which could better uncover the anabolic effect of PGE2 in vivo, has yet to be examined. Here, we hypothesized that PGE2 modulates F-actin stress fiber in FSS-stimulated MC3T3-E1 osteoblastic cells through protein kinase A (PKA) pathway. Furthermore, this PGE2-induced F-actin remodeling was associated with the recovery of cellular mechanosensitivity. Our data showed that treatment with 10 nM dmPGE2 for 15 min significantly suppressed the F-actin stress fiber intensity in FSS-stimulated cells in a PKA-dependent manner. In addition, dmPGE2 treatment enhanced the cells’ calcium peak magnitude and the percentage of responding cells in the second FSS stimulation, though these effects were abolished and attenuated by co-treatment with phalloidin. Our results demonstrated that 10 nM dmPGE2 was able to accelerate the ‘reset’ process of F-actin stress fiber to its pre-stimulated level partially through PKA pathway, and thus promoted the recovery of cellular mechanosensitivity. Our finding provided a novel cellular mechanism by which PGE2 increased bone formation as shown in vivo, suggesting that PGE2 could be a potential target for treatments of bone formation-related diseases.

Keywords prostaglandin E2; intracellular calcium; cytoskeleton; PKA pathway; fluid shear stress (FSS)

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Introduction

Prostaglandin E2 (PGE2), a 20-carbon proinflammatory prostanoid, has various physiological effects on tissue. In particular, the anabolic effect of PGE2 on bone has been established in vivo [1,2]. In vitro studies have showed that PGE2 stimulates differentiation by elevating the intracellular cAMP level at low concentrations (\(\sim 10^{-8} - 10^{-7} \text{ M}\)), while stimulates proliferation through phosphatidyl inositol turning over at high concentrations (\(\sim 10^{-6} - 10^{-5} \text{ M}\)) in statically cultured MC3T3-E1 osteoblastic cells [3–6]. However, during daily activities, bone cells are constantly stimulated by mechanical force, and the presence of mechanical force alters the release of biochemical factors [7,8] and the involvements of signaling pathways [9,10]. Thus, the cellular effect of PGE2 in the presence of mechanical loading could be different from that in the static condition, which worth to be studied.

It is well accepted that mechanical loading in bone is anabolic in tissue and at cellular levels. Fluid shear stress (FSS), as one kind of common mechanical stimulation in bone [11], deformed the cell body, thus introducing rapid increase in intracellular calcium concentration ([Ca\(^{2+}\)] \(\text{i}\)) [12] and the release of adenosine triphosphate (ATP) and PGE2 [8] in osteoblasts. However, the mechanosensitivity of osteoblasts was found to be decreased during continuous FSS stimulation, but recovered after inserting a rest period [13]. The cytoskeleton, a mechanotransductor during FSS stimulation, was believed to play a central role in the regulation of cellular mechanosensitivity through the modulation of mechanosensitive calcium channel (MSCC) activity [14,15]. While PGE2 has been shown to be capable of affecting cell morphology through modulating F-actin stress fiber in statically cultured osteoblasts [16] and also capable of promoting osteogenic differentiation in rat tendon stem cells [17], the PGE2-regulated F-actin stress fiber remodeling in the presence of mechanical stimulation and its effect on cellular mechanosensitivity have yet to be addressed.

Protein kinase A (PKA) pathway is involved in mechanosensitivity regulation in bone cells. Activation of PKA has been shown to enhance the intracellular calcium response...
Effect of PGE₂ on F-actin remodeling in osteoblasts

Materials and Methods

Materials

16,16-Dimethyl prostaglandin-E₂ (dmPGE₂; Cayman, Ann Arbor, USA) was used in this study due to its prolonged half life and bioactivities similar to PGE₂ [22]. Based on the finding that PGE₂ inhibits DNA synthesis by stimulating differentiation at low concentrations (∼10⁻⁸–10⁻⁷ M) in MC3T3-E1 cells [3,4], dmPGE₂ was used at a final concentration of 10 nM. Phalloidin (Sigma, St Louis, USA), an actin stabilizer, was used at a final concentration of 1 μM. 8-Bromoadenosine 3',5'-cyclic monophosphate sodium salt (8-Br-cAMP, Sigma), an agonist of PKA pathway, was dissolved in distilled water at a stock concentration of 15 mg/ml and used at a final concentration of 100 μM. The cAMP-dependent protein kinase peptide inhibitor (PKI; Promega, Madison, USA), an inhibitor of PKA pathway, was used at a final concentration of 10 μM. All the final solutions were prepared with the flow medium that was used in F-actin staining and calcium imaging studies. The concentration of agonist or inhibitor was chosen based on our previous studies [14,23].

Cell culture

MC3T3-E1 cells were purchased from American Type Culture Collection (ATCC; Manassas, USA). Cells were grown in α-MEM (Sigma) containing 10% fetal bovine serum (FBS; Gibco, Carlsbad, USA) and 1% penicillin G and streptomycin (Sigma) and maintained in a humidified incubator at 37°C with 5% CO₂. Cells were seeded onto type-I collagen coated glass slides at an initial density of 4 × 10⁴ cells/cm² and used for calcium imaging and cytoskeleton staining studies once reaching 80% of confluence. Cells were serum starved overnight prior to each experiment. Cells of 3–10 passages were used in our study.

F-actin staining

The slide containing MC3T3-E1 cells was mounted to a custom designed parallel plate flow chamber [24] and subject to 20 min of FSS at 12 dyn/cm² by using aerated (5% CO₂/95% air) flow medium (α-MEM supplemented with 0.1% FBS, v/v) in a cell incubator (37°C). After FSS loading, cells were treated with dmPGE₂, dmPGE₂ plus phalloidin, dmPGE₂ plus PKI, 8-Br-cAMP, 8-Br-cAMP plus PKI, and vehicle (α-MEM supplemented with 0.1% FBS, v/v) for 15 min, respectively (Fig. 1A). After all treatments, cells were immediately washed with ice cold phosphate buffered saline (PBS), fixed with 2% paraformaldehyde and 0.1% Triton X-100, and stained with Alexa Fluor 488 phalloidin (1:40 dilution; Invitrogen, Grand Island, USA) and DRAQ5™ (1:2000 dilution; Biotatus Ltd., Leicestershire, UK). Stained cells were imaged on an inverted confocal laser scanning microscope (Zesis LSM 510; Heidenheim, Germany) using 488 and 631 nm excitations with a 40 × objective. Since the F-actins network is a 3D structure, to minimize the variation among different batches, we thus recorded our images by focusing the z-level of most nucleuses in a random view. Each experiment was repeated thrice and at least 9 image fields of 40 × magnitude were obtained in each trial.

Image quantification

Velocity software (version4.3; Improvision, Waltham, USA) was used to process the areal fraction of continuous F-actin stress fibers in individual cells from well-documented confocal images (Supplementary Fig. S1). To account for the variations of staining process and laser intensity, which were unavoidably present among different batches of tests, we obtained a correction factor based on the nuclei staining intensity, since the nucleus was supposed to have a relatively constant intensity. This correction factor was used to linearly adjust the actin staining in each image. The continuous F-actin stress fibers were identified by setting two thresholds: objective intensity larger than 60, and an objective size larger than 500 pixels. The total area was obtained by setting a much lower threshold (larger than 7) and using the ‘fill holes in objects’ option. The areal fraction of the continuous F-actin stress fibers was calculated, with the total areal of continuous F-actin stress fibers being divided by the total area of cell. For each experimental group, three slides and 16–36 cells were processed.
Calcium imaging

Cells were washed with PBS, stained with 12 μg/ml Fluo-4 AM (Invitrogen) in α-MEM for 45 min at 37°C, and then washed with PBS again to remove the dye. The slide was mounted to a custom-designed parallel plate flow chamber [24], and fixed to the stage of a fluorescent microscope (Leica Microsystems, Wetzlar, Germany). Flow medium (α-MEM supplemented with 2% FBS, v/v), preserved at 37°C, was slowly driven into the chamber by a peristaltic pump (Longer Peristaltic Pump). The higher concentration of FBS (2%) in calcium imaging than that in F-actin staining study (0.1%) was used to ensure attachment and growth of MC3T3-E1 during the whole experimental process. After resting for 30 min, dynamic fluorescence intensity inside a randomly selected field (1431 × 1059 μm in dimensions) was recorded by real-time [Ca2+]i imaging with a 20× objective at room temperature. The recording time course included a 60-s baseline, followed by two sequential FSS sessions (3 min each at 12 dyn/cm²) with a 15 min rest period inserted in between, during which dmPGE₂, dmPGE₂ plus phalloidin or vehicle (α-MEM supplemented with 2% FBS, v/v) was added to flow medium to test the effect of dmPGE₂ on the calcium response recovery and the involvement of F-actin intensity in second FSS (Fig. 1B). Image J (version 1.44p, http://rsb.info.nih.gov/ij/) was used to analyze the fluorescence intensity in individual cells, which was normalized with the mean background intensity obtained in three randomly chosen blank areas. The calcium peak magnitude was reported as the fold increment of the peak intensity over the mean baseline fluorescence intensity (1). The percentage of responding cells was determined for each test (the number of cells whose calcium peak magnitude was over 1.25 fold of baseline was divided by the number of total cells). The 1.25-fold baseline threshold was chosen based on our previous study [12] using Fluo-2 under 12 dyn/cm², in which MC3T3-E1 cells remained the same, with regard to the ability of reacting with FSS (percentage of responding cells). Each experiment was repeated thrice and 47–102 cells per group were processed.

Statistical analysis

Data were presented as mean ± SD. For the cytoskeleton image quantification data, statistical significance was determined by using one way analysis of variance (ANOVA) with Bonferroni’s post hoc test. A P value of <0.05 indicated statistical significance in all analyses. For calcium imaging data, the statistical significance of the calcium peak magnitude and the percentage of responding cells in all treatments were determined by a one-way ANOVA with Bonferroni’s post hoc test and χ² test, respectively.

Results

Effects of dmPGE₂ and PKA pathway on F-actin intensity

When compared with FSS plus vehicle treatment group (Fig. 2A, b), dmPGE₂ treatment after FSS stimulation significantly decreased the FSS-induced F-actin stress fiber intensity (Fig. 2A, c) to nearly pre-FSS status (Fig. 2A, a). Co-treated cells with dmPGE₂ plus phalloidin (Fig. 2A, d) or dmPGE₂ plus PKI (Fig. 2C, a) caused an impaired inhibitory effect of dmPGE₂. Similar to dmPGE₂ post treatment, 8-Br-cAMP suppressed the F-actin stress fiber intensity in FSS-stimulated cells (Fig. 2C, b), which was again abolished by co-treatment with PKI (Fig. 2C, c).
Quantification of the areal fractions of F-actin stress fibers

Under the present imaging and threshold settings, the areal fraction of continuous F-actin stress fiber in pre-FSS group was $0.28 \pm 0.05$, and was significantly increased to $0.79 \pm 0.03$ by FSS plus vehicle treatment. dmPGE$_2$ significantly suppressed the FSS-induced increment in F-actin stress fiber intensity ($0.32 \pm 0.05$), which was comparable to the static level. Co-incubation of cells with phalloidin significantly attenuated the inhibition of dmPGE$_2$ ($0.56 \pm 0.03$). Similarly, co-treatment with PKI significantly decreased the inhibition of dmPGE$_2$ ($0.6 \pm 0.03$). The 8-Br-cAMP produced similar suppression as dmPGE$_2$ ($0.45 \pm 0.05$) and was completely inhibited by PKI ($0.78 \pm 0.04$) (Fig. 2B,D).

Figure 2. Involvement of PKA in dmPGE$_2$-modulated F-actin remodeling  
(A) Representative F-actin and nuclei staining showing effect of dmPGE$_2$ on F-actin stress fiber intensity in MC3T3-E1 cells (a–d). (B) Quantification results of (A). Compared with Pre-FSS, FSS + vehicle treatment significantly increased the F-actin stress fiber intensity, which was greatly decreased by FSS + dmPGE$_2$. Phalloidin (Phall) significantly attenuated the effect of dmPGE$_2$. (C) Representative F-actin and nuclei staining showing the involvement of PKA in dmPGE$_2$-modulated F-actin stress fiber intensity (a–c). (D) Quantification results of (C). The effect of dmPGE$_2$ was significantly attenuated by PKI. 8-Br-cAMP produced similar effect as dmPGE$_2$ which was blocked by co-treating with PKI. $^aP < 0.05$ vs. FSS + vehicle group, $^bP < 0.05$ vs. Pre-FSS group, $^cP < 0.05$ vs. FSS + dmPGE$_2$ group, $^dP < 0.05$ vs. FSS + 8-Br-cAMP group.
Effects of dmPGE2 on the recovery of mechanosensitivity in MC3T3-E1 cells

Cells responded to the first FSS stimulation with a rapid increase in \([\text{Ca}^{2+}]_i\) in all groups (Fig. 3, first FSS). In detail, the average \([\text{Ca}^{2+}]_i\) peak magnitude and percentage of responding cells were 1.68 ± 0.11 and 77.7 ± 11.6% (52 of 67 cells) in the vehicle group, 1.63 ± 0.07 and 71.2 ± 19.9% (73 of 102 cells) in the dmPGE2-treated group, and 1.62 ± 0.06 and 68.3 ± 7.5% (36 of 51 cells) in the dmPGE2 plus phalloidin group; no statistical difference was found among these groups. However, it was found that dmPGE2 treatment, post the first FSS, significantly increased the cells’ calcium peak magnitude and the percentage of responding cells in the second FSS stimulation by 1.25- and 5.31-fold, when compared with the vehicle treatment, which was respectively abolished and significantly attenuated by co-treating with phalloidin (Fig. 3, 2nd FSS). In detail, the average calcium peak magnitude and percentage of responding cells were 1.26 ± 0.08 and 16.3 ± 4.8% (8 of 47 cells) in the vehicle group, 1.57 ± 0.37 and 86.5 ± 9.3% (58 of 67 cells) in the dmPGE2-treated group, and 1.28 ± 0.02 and 45.7 ± 12.9% (32 of 69 cells) in the dmPGE2 plus phalloidin group.

Discussion

Mechanical forces and the downstream biochemical factors are important regulators for the maintenance of skeletal structure and mass. Osteoblasts respond to FSS with a \([\text{Ca}^{2+}]_i\) transient and a quick burst of ATP release (between 1–5 min post FSS), followed by a delayed, but sustained, PGE2 release [8]. Previous studies have demonstrated a significant role of PGE2 in osteoblasts’ differentiation and proliferation [3–6]. The results from this study suggest a novel biological significance.
of PGE$_2$, which might counteract with early released adaptation-promoting factors, such as ATP. The initially released ATP has been shown to stiffen cells by increasing the polymerization of actin filaments, and thus, promote the adaptation of osteoblasts to their mechanical environment [8]. However, this increased cellular stiffness also caused the loss of mechanosensitivity by decreasing the FSS-induced cellular deformation and opening the possibility of iron channels [14,25]. One major effect of exogenous PGE$_2$, as derived from this study, is the resultant restoration of mechanosensitivity in osteoblasts, by way of softening cells through modulating F-actin stress fiber intensity post FSS stimulation. However, without the additional exogenous PGE$_2$, the effect of released ATP overrode the effect of PGE$_2$ on F-actin stress fiber due to the insufficient endogenous PGE$_2$ release (around $10^{-10}$–$10^{-5}$ M, induced by 20 min of 30 dyn/cm$^2$ FSS loading in MC3T3-E1 cells) [26], which was evidenced by fact that actin polymerization was still observed in vehicle group (Fig. 2A, b).

A reciprocal regulation mechanism existed between FSS-induced [Ca$^{2+}$], transient and cytoskeleton remodeling. On one hand, as the earliest respondent upon receiving mechanical loading, [Ca$^{2+}$], transient was thought to be responsible for the subsequent release of ATP and PGE$_2$, an alteration in gene expression [27] and cytoskeleton remodeling [28] in osteoblasts due to Ca$^{2+}$ influx through the MSCC and voltage-sensitive Ca$^{2+}$ channels (L-VSCC), as well as the intracellular Ca$^{2+}$ release from endoplasmic reticulum [29]. On the other hand, treatments of cytochalasin D or parathyroid hormone disrupted/decreased F-actin stress fiber [14] increased the calcium response induced by FSS by decreasing the cellular stiffness, and modulating the open possibility of MSCC. Both sides suggested a direct connection between F-actin stress fiber and FSS-induced calcium response. In our study, we found that dmPGE$_2$ treatment increased the [Ca$^{2+}$]$_i$ transient during sequential FSS loading, which was abolished or attenuated by phalloidin. Our imaging quantification results showed that dmPGE$_2$ decreased the F-actin stress fiber intensity, while vehicle treatment increased it. These data together showed the participation of F-actin stress fiber in dmPGE$_2$ regulated mechanosensitivity in FSS-stimulated MC3T3-E1 cells. In addition, some cells in dmPGE$_2$ and phalloidin co-treated group showed delayed calcium peak (Fig. 3A, 2$^{nd}$ FSS), which suggested that phalloidin treatment caused the alteration of F-actin dynamics which might also be involved in mechanosensitivity in MC3T3-E1 cells.

Based on the observations that c-AMP/PKI respectively mimicked/attenuated the effect of PGE$_2$ on F-actin remodeling, we conclude that the PKA pathway is involved in PGE$_2$-induced F-actin remodeling. PGE$_2$ acts on osteoblasts through E-prostanoid (EP) receptors, in which EP1, EP2, and EP4 receptors are expressed in MC3T3-E1 cells [30]. EP2 and EP4 receptors are linked to G$\alpha$ alpha subunit (G$\alpha$) to increase intracellular c-AMP level and activate PKA pathway [31]. Our preliminary data showed that PKA activation was able to augment the hypotonic stress induced [Ca$^{2+}$]$_i$ transient in MC3T3-E1 cells (Supplementary Fig. S2A,B), suggesting a potential role of PKA in PGE$_2$ induced mechanosensitivity recovery in osteoblasts. We speculated that when osteoblasts are stimulated by FSS, the released and exogenous PGE$_2$ might act through either or both EP2 and EP4 to activate PKA pathway and thus down-regulate F-actin intensity, eventually leading to the recovery of [Ca$^{2+}$]$_i$ transient (Fig. 4).

Combined with previous reports showing that PKA pathway is involved in low-dose PGE$_2$ induced cell differentiation in osteoblasts [3,4], our and others’ data suggested that PGE$_2$ might facilitate osteoblasts to sense the FSS stimulation, thus amplifying the anabolic effect of FSS by promoting osteogenic differentiation. Our data also suggested that other signaling pathways might synergistically work with the PKA pathway, since PKA inhibition did not entirely blocked the effect of dmPGE$_2$ and c-AMP treatment was found not as effective as dmPGE$_2$ treatment.

In this study, a novel quantification method was used to measure the difference among different treatment groups in F-actin stress fiber. This quantification method has several strengths. First, the fluorescent intensity fluctuation among different batches of tests was unavoidable and exhibited a great impact on the accuracy of quantification results. We thus obtained a correction factor based on the nuclei staining intensity to limit the influence. Second, FSS-induced actin polymerization increased F-actin stress fiber thickness and length [8], causing an elongated cell morphology [32]. By using double thresholds defining system in our method, we were able to detect the area fraction of continuous F-actin stress fiber changes cell by cell between different treatment groups, which provided a better description about the F-actin stress fiber intensity than the total intensity ratio.
measurement used in previous studies [9,10]. However, there is still limitation in our study. The subtype(s) of EP receptor involved in dmPGE2-modulated F-actin remodeling was not clear, since both EP2 and EP4 subtypes have been found to activate PKA pathway in different cell types [31,33,34]. In addition, to better describe the mechanism of PGE2 mediated mechanosensitivity recovery, the involvement of the PKA pathway in PGE2 modulated calcium response and RhoA activation, as well as the long-term effects of PGE2 on mechanical loaded cells, needs to be further studied.

In conclusion, our study demonstrated that 10 nM dmPGE2 was able to accelerate the ‘reset’ process of F-actin stress fiber to its pre-stimulated level partially through PKA pathway, thus promoting the recovery of cellular mechanosensitivity. Our finding provided a cellular mechanism by which PGE2 increased bone formation in vivo, suggesting that PGE2 could be a potential target for treatments of bone formation related diseases.

Supplementary Data

Supplementary data is available at ABBS online.

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References