Prostacyclin primes pregnant human myometrium for an enhanced contractile response in parturition

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An incomplete understanding of the molecular events that regulate the myometrial transition from the quiescent pregnant state to the active contractile state during labor has hindered the development of improved therapies for preterm labor. During myometrial activation, proteins that prime the smooth muscle for contraction are upregulated, allowing maximal responsiveness to contractile agonists and thereby producing strong phasic contractions. Upregulation of one such protein, COX-2, generates PGs that induce contractions. Intriguingly, the predominant myometrial PG produced just prior to labor is prostacyclin (PGI₂), a smooth muscle relaxant. However, here we have shown that activation of PGI₂ receptor (IP) upregulated the expression of several contractile proteins and the gap junction protein connexin 43 through cAMP/PKA signaling in human myometrial tissue in organ and cell culture. Functionally, these IP-dependent changes in gene expression promoted an enhanced contractile response to oxytocin in pregnant human myometrial tissue strips, which was inhibited by the IP antagonist RO3244794. Furthermore, contractile protein induction was dependent on the concentration and time of exposure to the PGI₂ analog iloprost and was blocked by both RO3244794 and PKA knockdown. We therefore propose that PGI₂-mediated upregulation of contractile proteins and connexin 43 is a critical step in myometrial activation, allowing for a maximal contractile response. Our observations have important implications regarding activation of the myometrium prior to the onset of labor. […]

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Prostaglandins (PGs) are a family of lipid mediators that are produced in various tissues in response to cellular stimuli and play a crucial role in regulating various physiological processes. In human pregnancy, PGs are produced by the amnion and chorion to induce labor when necessary. PGs can also be produced locally in the myometrium, where they act to modulate the myometrial contractile response.

The myometrium is the smooth muscle layer of the uterus that undergoes significant structural and functional changes during pregnancy. The myometrial segment of labor is a complex process that involves a series of coordinated myometrial contractions that lead to the delivery of the fetus. The precise mechanisms that regulate the myometrial transition from an inactive, quiescent state to one capable of exhibiting strong phasic contractions are not fully understood.

Several studies have shown that activation of prostacyclin (PGI2) receptor (IP) upregulated the expression of several contraction-associated proteins and the gap junction protein connexin 43 through cAMP/PKA signaling in human myometrial tissue in organ and cell culture. Functionally, these IP-dependent changes in gene expression promoted an enhanced contractile response to oxytocin in pregnant human myometrial tissue strips, which was inhibited by the IP antagonist RO3244794. Furthermore, contractile protein induction was dependent on the concentration and time of exposure to the PGI2 analog iloprost and was blocked by both RO3244794 and PKA knockdown. We therefore propose that PGI2-mediated upregulation of contractile proteins and connexin 43 is a critical step in myometrial activation, allowing for a maximal contractile response. Our observations have important implications regarding activation of the myometrium prior to the onset of labor.

Introduction
Prior to initiation of term labor, the myometrium must become activated (1). This term refers to the conversion of the uterus from an inactive, quiescent physiological state to one capable of exhibiting strong phasic contractions. Early signals, such as expression of surfactant protein A (2) and corticotropin-releasing hormone (3–5), originate with the fetus and placenta, respectively, and, after many intermediate steps, ultimately lead to myometrial activation. The mechanisms and signaling pathways, however, remain poorly understood. Critically absent is knowledge about the signals and cellular mechanisms that directly lead to myometrial activation.

Activation enables the myometrium to be sensitive to contractile stimuli, such as oxytocin and PGs, and to generate synchronous and frequent contractions during labor (1). For smooth muscle cells to become responsive to these agonists, the coordinated expression of several contraction-associated proteins must occur (1, 6, 7). Contraction-associated proteins — which include the oxytocin receptor, PG receptors, COX-2, ion channels, and gap junction proteins, in particular connexin 43 — promote myometrial cell contractility and excitability as well as intracellular connectivity. Gap junctions are present at very low levels during pregnancy, but gap junction proteins, connexin 43 in particular, increase in myometrial cells during labor and delivery (8–10).

An incomplete understanding of the molecular events that regulate the myometrial transition from the quiescent pregnant state to the active contractile state during labor has hindered the development of improved therapies for preterm labor. During myometrial activation, proteins that prime the smooth muscle for contraction are upregulated, allowing maximal responsiveness to contractile agonists and thereby producing strong phasic contractions. Uprogulation of one such protein, COX-2, generates PGs that induce contractions. Intriguingly, the predominant myometrial PG produced just prior to labor is prostacyclin (PGI2), a smooth muscle relaxant. However, here we have shown that activation of PGI2 receptor (IP) upregulated the expression of several contraction proteins and the gap junction protein connexin 43 through cAMP/PKA signaling in human myometrial tissue in organ and cell culture. Functionally, these IP-dependent changes in gene expression promoted an enhanced contractile response to oxytocin in pregnant human myometrial tissue strips, which was inhibited by the IP antagonist RO3244794. Furthermore, contractile protein induction was dependent on the concentration and time of exposure to the PGI2 analog iloprost and was blocked by both RO3244794 and PKA knockdown. We therefore propose that PGI2-mediated upregulation of contractile proteins and connexin 43 is a critical step in myometrial activation, allowing for a maximal contractile response. Our observations have important implications regarding activation of the myometrium prior to the onset of labor.
to determine whether PGI₂ at the time of labor (36) or may play a role in cervical soften
ing. Because the physiologic role of PGI₂ because while its synthesis is increased prior to and during labor
is important for preventing intrapartum thromboembolic compli-
cations. We have previously demonstrated that myometrial tissue strips in organ culture under tension maintain spontaneous contractility as well as contractile response to oxytocin for up to 7 days (45). It is worth noting that tissue obtained from the lower uterine segment is known to vary in contractility, likely because of differences in the site of excision from the uterus. To minimize this effect, we performed experiments only on paired tissue strips that were obtained from the same specimen. Tissue strips were cultured in parallel in the presence or absence of the PGI₂ analog iloprost for 48 h, after which oxytocin-induced contractions of the tissue strips were measured. Usually, spontaneous contractions were irregular and infrequent when the tissue was initially stab-
ilized in the isometric apparatus. Oxytocin was added to initiate rhythmic contractions and test tissue responsiveness. In each case, tissue responded to oxytocin exposure with increased contract-
straction strength. In tissue samples from each of 7 consecutive consenting patients, iloprost pretreatment resulted in greater oxytocin-induced contractile response compared with its control, but the magnitude of the increase varied from 1.1- to 9-fold (Figure 1). Using the Mann-Whitney U test, we determined that this increase achieved a statistical significance of P = 0.0253474. These data reveal that long-term pretreatment of myometrial tissue with iloprost promoted an enhanced contractile response to the agonist oxytocin.

Iloprost upregulates contractile apparatus proteins and connexin 43 in uterine tissue. We investigated possible mechanisms underlying the enhanced contractile response to oxytocin induced by iloprost. Prior to labor, a group of proteins referred to as contraction-associated proteins are expressed. These proteins act to initiate pow-
erful synchronous contractions during labor. Also important are con-
tractile proteins such as α-SMA and SM-MHC, which deter-
mine myocyte contractility. As we have previously shown that PGI₂ can induce contractile protein expression in cultured VSMCs (44), we sought to determine whether iloprost increased expression of these proteins in uterine tissue. We subjected some of the tissue strips from the experiments in Figure 1 to Western blot analysis immediately after the contraction assay and observed an increased expression of SM-MHC isoform 2 (SM2-MHC) and calponin in tissue cultured with iloprost relative to PBS vehicle treatment (Figure 2A). This upregulation also occurred after 6 h of iloprost pretreatment (Figure 2B). To eliminate any potentially confounding effects of oxytocin exposure, we treated myometrial tissue with vehicle or iloprost for 6 or 48 h and then immediately performed Western analysis. Expression of the contractile proteins α-SMA, h-caldesmon, calponin, and SM2-MHC increased in tissue treated with iloprost (Figure 2, C and D).
To begin to address the mechanisms underlying the enhanced contractile protein expression, we next assessed the effects of iloprost exposure on mRNA levels of calponin and SM-MHC in uterine tissue strips. RT-PCR experiments were performed after 48 h of exposure to iloprost or vehicle. Calponin and SM-MHC mRNA levels were increased, which suggests that iloprost may regulate contractile protein expression at the level of transcription (Figure 2E).

Given the important role of gap junctions and connexin 43 in myometrial contractility and labor progression (8–11), we wanted to determine whether iloprost also influences connexin 43 expression in myometrial tissue. Treatment of myometrial tissue with iloprost for 48 h resulted in a significant increase in connexin 43 expression compared with control (Figure 3). Based on previous studies showing the importance of connexin 43 in regulating parturition (11), these data suggest that iloprost may also enhance oxytocin-induced contractility by increasing connexin 43 expression.

Because iloprost at high concentrations can also activate EP1 receptors (46, 47), we used a recently developed potent and selective IP antagonist, RO3244794 (48), as a tool to determine the contribution of the hIP in modulating the iloprost-induced enhanced contractile response to oxytocin. As in Figure 1, we found that myometrial tissue treated with iloprost for 48 h demonstrated increased contractile activity in response to oxytocin relative to control-pretreated tissue (Figure 4A). However, pretreatment with iloprost and RO3244794 blocked the iloprost-enhanced uterine activity, which suggests that the increased activity is specific to activation of the hIP. Interestingly, we observed that RO3244794 alone also reduced the response to oxytocin (Figure 4A), which suggests that endogenous PGI2 produced by the tissue while in organ culture contributes to uterine contractility. Sample traces showing basal and iloprost-induced responses to oxytocin in the presence or absence of RO3244794 are shown in Figure 4B. Western blot analysis of the tissue revealed that RO3244794 also blocked the iloprost-induced increase in h-caldesmon, connexin 43, and SM2-MHC expression (Figure 4, C and D).

In Figure 2, we showed that prolonged iloprost treatment induced contractile protein expression in myometrial tissue whether or not the tissue had been subjected to contraction with oxytocin. In Figure 4D, we further assessed the effects of oxytocin, with and without iloprost. Interestingly, treating tissue with oxytocin alone for 90 min promoted a modest, but measurable, increase in SM2-MHC protein expression relative to vehicle control. However, treatment with iloprost alone for 48 h induced
a greater SM2-MHC expression than did oxytocin alone, and the effects appeared to be additive (Figure 4D). Collectively, our data suggest that long-term pretreatment of myometrial tissue with iloprost, acting through the hIP, modifies the properties of the tissue, upregulating contractile protein and gap junction protein expression and resulting in an enhanced contractile response to the agonist oxytocin.

Iloprost induces SMC-specific contractile apparatus proteins and connexin 43 in monolayer cultures of uterine SMCs. To further investigate the molecular mechanisms underlying iloprost-induced contractile protein expression, we used a monolayer primary culture model of human uterine SMCs (uSMCs) in which cells were cultured from patient specimens obtained as described above. First, dose dependence and kinetics of the response were evaluated. Treatment for 6 h with increasing concentrations of iloprost increased mRNA levels of SM-MHC (Figure 5A). This increase was maximal at 2.5 nmol/l iloprost and decreased slightly at 100 nmol/l, the highest concentration tested. GPCR desensitization is known to occur after exposure to high concentrations of ligand (49). For this reason, 2.5 nmol/l iloprost was used in subsequent cell culture studies.

Iloprost induced an accumulation of SM-MHC message as early as 30 min after treatment, which persisted for up to 18 h (Figure 5, B and C). To determine whether this induction occurs at the level of transcription, we used the RNA synthesis inhibitor actinomycin D (Figure 5D). After 4 h of treatment with iloprost, there was a 1.5-fold increase in SM-MHC mRNA, which did not occur in the presence of actinomycin D. These data indicate that the increase in SM-MHC mRNA with iloprost is not the result of enhanced mRNA stability, and it is therefore likely that iloprost regulates SM-MHC message at the level of transcription.

As in the organ culture model, iloprost induced expression of α-SMA, calponin, h-caldesmon, and SM2-MHC at the protein level (Figure 6A). This induction showed a concentration dependence similar to that seen at the mRNA level (Figure 5A). This upregulation was also time dependent: protein levels of calponin, h-caldesmon, and SM2-MHC increased as early as 2 h, and were sustained for up to 24 h, after treatment with 2.5 nmol/l iloprost (Figure 6, B–D). Notably, the increase in SM2-MHC protein closely followed the kinetics of the mRNA induction (Figure 5, B and C, and Figure 6D). Treatment with 2.5 nmol/l iloprost also increased connexin 43 message and protein levels in a time-dependent manner (Figure 7, A and B).

The cAMP/PKA pathway is necessary for iloprost-induced contractile protein and connexin 43 expression. In direct analogy with vascular smooth muscle (44), we hypothesized that activation of the cAMP/PKA pathway mediates iloprost-induced contractile apparatus protein expression in uSMCs. Indeed, iloprost induced cAMP accumulation in uSMCs in a concentration-dependent manner, with an EC50 of 12.5 nmol/l iloprost (Figure 8A). The main effector of cAMP, PKA, was activated after 20 min in a concentration-dependent manner by iloprost, with an EC50 of 18 nmol/l (Figure 8B). The specific PKA inhibitor peptide PKI inhibited PKA activity induced by 100 nmol/l iloprost, confirming the specificity of the kinase.

Figure 4
An IP antagonist opposes iloprost-enhanced oxytocin-induced contractions and iloprost-induced contractile protein expression. (A) Human myometrial tissue in organ culture was pretreated with 90 μmol/l RO3244794 (RO) for 30 min and then stimulated with or without 25 nmol/l iloprost for 48 h, after which tissue was transferred to an isometric muscle bath and treated with 5 nmol/l oxytocin as described in Methods. Average peak contraction data are represented as fold change relative to vehicle control. Arithmetic mean values are represented by bars, and individual measurements are plotted for the 3 patients. (B) Representative tracings for 1 patient in A. (C) Homogenates of uterine tissue isolated from A were subjected to Western blot analysis with antibodies to h-caldesmon, connexin 43, GAPDH, or β-tubulin. (D) Human myometrial tissue in organ culture was pretreated with 90 μmol/l RO3244794 for 30 min and then treated with vehicle or 25 nmol/l iloprost for 48 h, followed by treatment with 5 nmol/l oxytocin for 8 h, as indicated. Protein was isolated from the tissue and subjected to Western blot analysis with antibodies to SM2-MHC or GAPDH. Lanes were run on the same gel but were noncontiguous (lines).
PKA activity was transient, peaking at 20 min and returning to baseline by 90 min (Figure 8C).

To verify the role of the cAMP in iloprost-induced expression of contractile proteins, we next determined whether the membrane-permeable analog 8-Br-cAMP could mimic the effect of iloprost in uSMCs. Treatment with 8-Br-cAMP resulted in concentration-dependent increases in PKA activity (data not shown). We determined that 0.5 μmol/l 8-Br-cAMP yielded a level of PKA activity similar to that yielded by 2.5 nM iloprost (2.2- and 1.7-fold increase over control, respectively; Figure 9A) and induced expression of calponin and SM-MHC mRNA as early as 1 h (Figure 9B). 8-Br-cAMP increased α-SMA, calponin, h-caldesmon, and SM2-MHC protein in a concentration-dependent (Figure 9C) and time-dependent (Figure 9D) manner, confirming that the second messenger cAMP is sufficient to induce contractile proteins.

To determine whether PKA activation is necessary for the iloprost-induced contractile protein expression, we used siRNA targeted to PKA catalytic subunits to specifically inhibit PKA function. Two highly homologous PKA catalytic subunit isoforms are expressed in uSMCs, PKA catalytic α (PKA Cα) and PKA Cβ. To avoid functional redundancy, we used siRNAs targeting each subunit in combination. Transfection of uSMCs with PKA Cα- and PKA Cβ-specific siRNA reduced subunit expression by 53% and 71%, respectively (Figure 10A), and significantly reduced iloprost-induced PKA activity (Figure 10B). PKA Cα and PKA Cβ knockdown also significantly inhibited the iloprost-induced upregulation of calponin, h-caldesmon, SM2-MHC, and connexin 43 (Figure 10, C–F) proteins to near-control levels. Thus, we conclude that the cAMP/PKA pathway is not only sufficient, but also necessary, for the regulation of contractile protein and connexin 43 expression induced by iloprost.

While these data strongly support a mechanism in which the G protein–coupled hIP induces contractile protein expression by coupling to Gαs, cAMP, and PKA, we tested 2 alternate possibilities. Iloprost has the potential to activate other prostanoid GPCRs, in particular EP1 (46, 47). We found contractile protein induction at low doses of iloprost (2.5 nM, below the EC50 for cAMP production), but we also tested another PGI2 analog, cicaprost, which is more specific for hIP (47). Cicaprost activated PKA in a concentration-dependent manner (Figure 11A), and, similar to iloprost, the enhanced activity was transient, peaking at 20 min and declining by 60 min (Figure 11B). Cicaprost induced dose-dependent increases in h-caldesmon, α-SMA, and SM2-MHC (Figure 11C) and upregulated SM2-MHC, α-SMA, calponin, and connexin 43 protein expression in a time-dependent manner (Figure 11, D–F).

Finally, there is also the possibility that PGI2 and its analogs, at higher concentrations, can activate the intracellular receptor PPARδ, a transcription factor (50, 51). Because we observed effects on contractile protein expression at the level of transcription, we tested whether contractile protein induction was dependent entirely on the hIP cell surface GPCR using RO3244794. RO3244794 inhibited iloprost-induced PKA to baseline levels (Figure 12A) and completely blocked the iloprost induction of h-caldesmon and SM2-MHC (Figure 12, B and C). As a control for receptor specificity, we demonstrated that RO3244794 had no effect on 8-Br-cAMP–induced contractile protein expression (Figure 12D). As demonstrated above, RO3244794 also inhibited contractile protein and connexin 43 expression induced by iloprost in human uterine tissue strips (Figure 4, C and D). Together, these data confirm that iloprost induces contractile protein and connexin 43 expression in uSMCs via the cell surface hIP GPCR and signaling through the cAMP/PKA pathway.

Figure 5
Iloprost induces SM-MHC and calponin message in uSMC culture. (A) Human uSMCs were treated with the indicated concentrations of iloprost for 6 h. Total RNA was isolated and subjected to RT-PCR using primers to SM-MHC and PDH. (B and C) Total RNA was isolated from cells treated with vehicle (Veh) or 2.5 nM iloprost for the indicated times, and RT-PCR was performed as in A. (D) Human uSMCs were treated with vehicle or 2.5 nM iloprost for 4 h, followed by treatment with 1 μg/ml actinomycin D for an additional 4 h. Total RNA was isolated and subjected to RT-PCR using primers to SM-MHC or PDH. Representative gels are shown, as well as densitometric quantitation of 2 separate experiments below. Arithmetic mean values represent fold induction corrected to PDH. Data in A–D were from cell cultures derived from 3 different patients. Lanes in C and D were run on the same gel but were noncontiguous (lines).
PGI₂ and iloprost exert similar effects in vitro. Iloprost and cicaprost are useful analogs to model the action of PGI₂. However, because they are more stable and can potentially activate other prostaglandin receptors depending on the concentration, it is possible that native PGI₂ may have different receptor binding properties than these analogs (47). For this reason, we tested the effects of PGI₂ itself on human uSMCs. Because PGI₂ breaks down more rapidly than do iloprost and cicaprost, we also tested single versus repeated administration of PGI₂. We determined that, similar to iloprost, a single administration of PGI₂ elicited dose-dependent activation of PKA in uSMCs in culture, assayed 20 min after ligand addition (Figure 13A). Repeated administration of PGI₂ every 1.5 h for 6 h increased SM2-MHC and calponin expression in a dose-dependent manner (Figure 13B). Administration of 0.6 μmol/l PGI₂ and 2.5 nmol/l iloprost exhibited similar levels of PKA activation (Figure 13A) and of contractile protein upregulation (Figures 13, B and C). Interestingly, a single treatment of PGI₂ was sufficient to increase contractile protein and connexin 43 expression in a time-dependent manner, up to 24 h (Figure 13C). Taken together, our data with PGI₂, iloprost, cicaprost, and RO3244794 provide evidence that activation of the hIP modulates the expression of contractile apparatus proteins and the gap junction connexin 43 in cultured uSMCs.

**Discussion**

The results of the present study provide evidence that PGI₂ plays an important role in priming the uterus for contraction prior to labor, both by enhancing oxytocin-induced contractions by increasing expression of contractile proteins, which are critical in generating the contractile force of labor, and by increasing the expression of the gap junction protein connexin 43, which is critical in allowing myocytes to act synchronously. Results were consistent across all 21 consecutive consenting (nonselected) patients. We believe that these results provide substantial new
insights into the role of PGI2 during human parturition and suggest that further clinical trials of PGI2-based therapeutics may be warranted.

While we noted a consistent response to PGI2 in all patient samples, the magnitude of the response varied. The range of responses is possibly due to the extent of myometrial activation that had occurred in each specimen in vivo prior to organ culture. We speculate that iloprost would exert a lesser effect in tissue that was partially or fully activated prior to organ culture, whereas tissue that had not yet undergone activation in vivo would demonstrate the greatest increase in contractility after iloprost treatment.

As there is currently no way to accurately predict proximity to labor in patients, we cannot easily test this hypothesis. The degree of endogenous PGI2 produced by the tissue itself is also another critical variable. If a particular specimen produced large amounts of PGI2, then added PGI2 would have only minor effects on contractility. Thus interindividual variability in tissue samples could account for the range of effects that were observed in the contractility experiments.

Our study raises the intriguing question as to how PGI2 can play different roles at different times during pregnancy. We show here that PGI2 upregulation of contractile proteins and connexin 43 depends on a PKA signal. PGI2-dependent smooth muscle relaxation is also thought to be PKA dependent (1, 6). It is likely that multiple factors affect the ultimate outcome of the PKA signal. The ERK/MAPK pathway is the best-characterized example of how the biological outcome of signaling pathways depends upon context and timing. ERK activation can promote opposing outcomes such as proliferation or differentiation, depending on factors such as signal strength and duration, receptor density, extracellular matrix composition, expression of scaffolding proteins and cell type–specific transcription factors, and the presence of other incoming signals (cross-talk; reviewed in ref. S2).

We hypothesize that, because of hormonal changes and other signals, the context for the PGI2 signal at the time of myometrial activation differs from its context earlier in pregnancy. It is known that steroid hormones play a major role in myometrial regulation (6) and that myometrial estrogen receptor levels and estrogen sensitivity are enhanced at term (53). Estrogen receptor, in turn, increases COX-2 expression (53, 54), which has also been implicated in stimulating myometrial activation (53). This elevated COX-2 may, at least in part, explain the increase in PGI2 levels that occurs at term (27). Elevated PGI2 levels could thus alter signal strength and duration, which could generate different effects of PGI2 at term. The number of hIPs is not known to change during the course of pregnancy (55), so the differential signaling may be mediated primarily at the level of ligand expression and/or potential cross-talk mechanisms described below.

The effects of estrogen and other hormonal changes near term may also induce other changes in cellular context, in which cross-talk caused by concomitant activation of other receptors alters the ultimate downstream effectors of the PGI2 signal. Our data suggest that PGI2 and oxytocin signaling may cooperatively induce contractile protein expression. We hypothesize that the different environment during the course of myometrial activation may also result in expression of transcription factors and cofac-

Figure 7
Iloprost induces expression of connexin 43 in human uSMCs. (A) Human uSMCs were treated with vehicle or 2.5 nmol/l iloprost for the indicated times. Total RNA was isolated and subjected to RT-PCR using primers against connexin 43 and PDH. Lanes were run on the same gel but were noncontiguous (lines). (B) Cell lysates from human uSMCs treated with 2.5 nmol/l iloprost for the indicated times were subjected to Western blot analysis with antibodies to connexin 43 and GAPDH.

Figure 8
Iloprost induces cAMP/PKA activity in human uSMCs. (A) Concentration-response curves were determined by treating uSMCs with concentrations of iloprost ranging from 10 pmol/l to 1 μmol/l. After 20 min, cells were harvested, and cAMP was measured as described in Methods. Results are mean ± SEM. EC50 was determined from best-fit curve with nonlinear regression. (B) Human uSMCs were treated with vehicle or the indicated concentrations of iloprost for 20 min, and cell lysates were analyzed for PKA activity as described in Methods. A representative gel is shown. Fluorescence units of phosphorylated kemptide (p-Kemptide) were quantified by densitometry and expressed as fold induction. n = 5 separate experiments. (C) Human uSMCs were treated with 2.5 nmol/l iloprost for the indicated times, and cells were harvested and analyzed for PKA activity.
tors not present earlier in pregnancy. Our ongoing preliminary studies provide evidence that signaling through hIP activates and upregulates multiple transcription factors known to regulate contractile protein expression in smooth muscle, including some that are known to be regulated, at least in part, by calcium signaling (our unpublished observations). These findings suggest an avenue for cooperation between calcium-stimulating agonists such as oxytocin or PGF2α, with the PKA-activating PG1 near term.

Effects on downstream transcription factors also likely explain the sustained effect elicited by single doses of either PGI2 or its analogs, which are all short-lived agonists. We showed that PKA activation by iloprost or cicaprost was rapid and transient, peaking by 20 min. Upregulation of contractile proteins and connexin 43 at the mRNA level was first detected at early time points (as early as 30 min for SM-MHC). The kinetics of contractile protein and connexin 43 protein expression followed those of the mRNA levels, and maximal increases were observed after 24–48 h. Thus, while an initial acute response to cAMP/PKA may in fact be muscle relaxation, our data suggest that the transcriptional changes are also initiated early but result in longer-lived effects on protein expression. Changes in expression of contractile proteins and connexin 43 may be seen as a

Figure 9

cAMP is sufficient to induce contractile protein expression. (A) Human uSMCs were treated with vehicle, 0.25 nmol/l iloprost, or 0.5 μmol/l 8-Br-cAMP for 20 min, and cells were harvested and analyzed for PKA activity. Shown is a representative gel showing phosphorylated kemptide and densitometric quantitation of 2 independent experiments expressed as fold induction. Arithmetic mean values are shown. (B) Total RNA was isolated from human uSMCs at time point 0, treated with vehicle for 7.5 h, or treated with 5 μmol/l 8-Br-cAMP for the indicated times. RNA was subjected to RT-PCR using primers to the basic calponin, SM-MHC, or PDH genes. (C) Human uSMCs were treated with the indicated concentrations of 8-Br-cAMP for 8 h, and cell lysates were subjected to Western blot analysis with antibodies to α-SMA, calponin, h-caldesmon, SM2-MHC, and GAPDH. (D) Human uSMCs treated with vehicle or 0.5 μmol/l 8-Br-cAMP for the indicated times were harvested and subjected to Western blot analysis with antibodies to calponin, h-caldesmon, SM2-MHC, and GAPDH.
PKA is required for iloprost-induced contractile protein and connexin 43 expression. (A) Human uSMCs were transiently transfected with siRNA to PKA Cα and PKA Cβ (siPKA) or negative control siRNA, siControl (siCont). At 40 h after transfection, cells were treated with vehicle or 2.5 nmol/l iloprost for 8 h, harvested, and subjected to Western blot analysis for PKA Cα and PKA Cβ. Representative blots are shown. Amounts of PKA Cα and PKA Cβ proteins relative to GAPDH were quantified by densitometry (n = 7, including cell cultures derived from 2 different patient samples). **P < 0.001 versus vehicle- and iloprost-treated siControl. (B) Human uSMCs were transfected with PKA Cα and PKA Cβ (0.5 μg each) or siControl for 48 h followed by treatment with vehicle or 2.5 nmol/l iloprost (ilo) for 20 min. Cell lysates were analyzed for PKA activity as described in Methods. A representative gel is shown. Densitometric quantitation of fluorescence units of phosphorylated kemptide is expressed as fold induction relative to siControl treated with vehicle (n = 3). (C–F) Human uSMCs were transfected as in A and treated with vehicle or 2.5 nmol/l iloprost for 8 h. Western blots for (C) calponin, (D) h-caldesmon, (E) SM2-MHC, and (F) connexin 43 are shown with β-tubulin or GAPDH loading controls. Lanes in C were run on the same gel but were noncontiguous (lines). Densitometric quantitation of at least 3 independent experiments, including cell cultures derived from 2 different patient samples, are expressed as fold induction corrected to the loading control. All data are mean ± SEM. Significance of differences were determined using 1-way ANOVA with Newman-Keuls post-hoc test.
The priming event, whereby the myometrium is modified such that it becomes sensitized to respond to contractile stimuli — not only oxytocin (our present results), but possibly other PGs, such as PGF$_{2\alpha}$ — during labor.

Previous studies have demonstrated that at higher concentrations, PGI$_2$ and some of its analogs, including iloprost and carbaprostacyclin, can regulate transcription by directly activating the nuclear receptors PPAR$_\alpha$ or PPAR$_\delta$ in a PKA-independent manner (50, 51). Our data do not support a major role for PPARs in upregulating contractile proteins or connexin 43, because RO3244794 or PKA inhibition opposed these effects, and the PGI$_2$ analog cicaprost, which does not activate PPARs (50), mimicked the effects of iloprost.

The local concentrations of PGI$_2$ achieved at term in human myometrium are unknown, because the very short half-life of this agonist precludes accurate measurements in real time. One study measured PGI$_2$ produced by human myometrial tissue isolated at different time points (15–42 weeks) during pregnancy (27). The findings from this study suggest that within a 30-min assay period, the isolated myometrial tissue can produce PGI$_2$ concentrations of 0.4 μmol/l at 38 weeks and 1.6 μmol/l at 40 weeks, a 4-fold increase. Given the short half-life (~3 min), the actual concentrations may be even higher locally in vivo. Notably, these concentrations are entirely consistent with those that we found to activate PKA and to induce contractile protein expression in our present studies (0.6–6.0 μmol/l). Furthermore, the doses of iloprost used in our study that potentiated contractility were fairly low, and concentrations below the EC$_{50}$ for cAMP/ PKA activation exerted potent effects on contractility and protein expression in vitro. It is likely that iloprost was more potent than PGI$_2$ (600 nM PGI$_2$ was comparable to 2.5 nM iloprost) because it is more stable (half-life of ~30 min) and therefore able to amplify signal over a longer period of time.

While relaxation induced by iloprost has been attributed to activation of cAMP/PKA signaling, exact mechanisms have yet to be elucidated. It is thought that PKA inhibits myosin light...
Transiently effective because of eventual PKA-dependent regulation to rapid receptor desensitization and internalization (57). Based on pregnant human subjects, present challenges to advancing our understanding of human parturition and ability to develop betan tocolytic therapies. Therefore, the use of pregnant human myometrial tissue near term, in organ and cell culture, is a particular strength of our study, allowing us to make further inroads into the molecular mechanisms underlying this crucial process.

In conclusion, we have shown that PGI₂ signaling enhanced the contractile response of human pregnant myometrium to oxytocin and upregulated contractile apparatus protein expression and the gap junction protein connexin 43. Our results have important implications for a role of endogenous myometrial PGI₂ as an important factor in regulating activation of the myometrium, a critical step in the initiation and progression of parturition.

**Methods**

**Tissue procurement and culture.** Tissue procurement was performed in accordance with a Dartmouth College Institutional Review Board–approved protocol (IRB protocol no. 15820). Myometrium was obtained as previously described (45), after obtaining informed consent, from women not in labor undergoing Cesarean section delivery between 37.5 and 41 weeks gestation for clinical indications. Indications include repeat Cesarean section and breech presentation. Specifically excluded were patients who were treated with magnesium sulfate, other tocolytic agents, or intravenous oxytocin or who exhibited signs of intrauterine infection. An excisional biopsy was obtained from the upper margin of the uterine incision after removal of the infant and placenta and control of bleeding. Tissue was stored and transported in DMEM supplemented with 10% FBS. Decidua and serosa were dissected from the tissue and discarded. Myometrial tissue strips, approximately 1 × 1 × 15 mm, were cut along the lines of the fascicula using sterile technique. Each end was secured with a 000 vicryl suture. Tissue strips were organ cultured as previously described (45). In brief, tissue strips were cultured by mounting each vertically in its own tube and maintained in culture at 37°C, 5% CO₂ in DMEM. A 400-mg weight was suspended from each strip to apply constant tension. Using this technique, tissue strip contractility was routinely maintained for more than 7 days.

**Contraction assays.** Tissue strips were placed in 1 of 2 identical isometric contraction devices, and experiments were performed in pairs. Forces produced by each tissue were monitored using a Grass-Telefactor FT-03 force transducer. Output was monitored by an analog-to-digital convertor and stored on a personal computer for later analysis. Using micrometer drives, tissues were stretched to 140% of resting length, and the physical distances were not adjusted again for the duration of the experiment. Initial tensions were usually on the order of 1–2 g, which then fell due to viscoelastic creep to values of ∼500 mg. Spontaneous (e.g., applied tension only) contractions were sometimes observed, and at least 3 contractions were measured. After 30 min of observation, 5 nmol/l oxytocin was added to all tissues. In all cases, contractions were then observed, and at least 5 contractions were measured. Sample traces are shown in Figure 4B. For each contraction, peak forces were measured from the baseline to the peak of each contraction. For each experiment, contraction forces were averaged to yield average peak force production with standard deviations. At the conclusion of some experiments, 1 μM A23187 was added to the bath to determine the maximal tension for each tissue.

**Isolation of mRNA and protein from uterine tissue strips.** After treatment, tissue strips were washed in PBS, snap frozen, and stored at −80°C. To isolate protein and mRNA from the same tissue sample, PBS supplemented with

![Figure 12](http://www.jci.org)
RNase inhibitor (0.75 U/μl; Ambion) and a complete protease inhibitors tablet (Roche) was added. Samples were sonicated on ice and then centrifuged at 4,000 g for 15 min at 4°C. The protein-containing supernatant was collected. The pellet was resuspended in TRIzol reagent (Invitrogen) to extract total RNA according to the manufacturer’s protocol.

Cell culture. Human uSMCs were isolated from myometrial tissue obtained as described above. Cells were isolated by enzymatic digestion. In brief, tissue was minced and washed in HBSS. Tissue was digested in Collagenase XI and Collagenase IA (1 mg/ml each; Sigma-Aldrich) in HBSS at 37°C for 1 h, after which the cells were centrifuged and cultured in monolayer in DMEM supplemented with 10% FBS at 37°C. Passages 2–7 were used for experiments. Cultures derived from 4 different patients were used for experiments. All drug treatments were performed in DMEM containing 2.5% FBS. Vehicle-treated cells were incubated with PBS for the maximum duration of the experimental treatment. Cells were treated with various drugs, as indicated in the figure legends. Iloprost was purchased from Amershams Biosciences. 8-Br-cAMP and oysterin were purchased from Sigma-Aldrich. Cicaprost was a gift from Schering. The IP antagonist RO3244794 was a gift from M.-F. Jett (Roche, Palo Alto, California, USA).

Semiquantitative RT-PCR. Total RNA was isolated from monolayer cells using the Qiagen RNeasy kit with DNase I (Qiagen) and quantitated in a Nanodrop 1000 spectrophotometer (Promega). Primers to specifically amplify the human SM-MHC gene transcript, human basic calponin gene transcript, and the pyruvate dehydrogenase (PDH) transcript (housekeeping control gene) were used as described previously (44). Primers were designed to amplify the human connexin 43 gene transcript: sense, 5′-TCACTTGCGCCTCTGGGTTCTT-3′; antisense, 5′-CCTTACGGCCCTTAGAGAAA-3′. A dilution series of the reverse-transcribed cDNA was used to determine the linear range of the PCR for each primer set. PCR was performed using 20 pmol of each primer, 0.04 mmol/l dNTPs, and HotMaster Taq DNA polymerase (Eppendorf). PCR with calponin or PDH primer sets was performed for 32 cycles of 95°C for 30 s; 62°C (connexin 43), 58.5°C (calponin), or 55°C (PDH) for 30 s; and 70°C for 1 min. PCR with MHC primers was performed for 34 cycles of 95°C for 30 s; 54°C for 30 s; and 70°C for 1 min. PCR products were resolved on 1% agarose gels with GelStar nucleic acid stain (Cambrex), and digital images were obtained using a Typhoon Scanner (Molecular Dynamics).

Cell lysis and immunoblotting. Cells were lysed for Western blot analysis as previously described (44). Bradford assay was used to determine total protein concentration (Pierce). Equal amounts of protein (10–20 μg) from each lysate were subjected to Western blot analysis as described previously (44) using primary antibodies against α-SMA, calponin, or β-tubulin (Sigma-Aldrich); SM2-MHC (Seikagaku America); connexin 43 (Cell Signaling); PKA α, PKA β, or β-tubulin (Santa Cruz Biotechnology Inc.); or GAPDH (AbCam). Signal was detected with enhanced chemiluminescence reagents (Pierce). Densitometry was performed using Scion Image Beta (version 4.0.2) to determine relative quantities of protein, with data then normalized to GAPDH or β-tubulin.

Determination of cAMP levels. At confluence, uSMCs in 12-well plates were incubated with DMEM containing 2.5% FBS overnight. Cells were washed twice with PBS plus 4 mmol/l EDTA and 2 mmol/l isobutylmethylxanthine (pH 7.4) and incubated at 20°C for 10 min. cAMP generated in response to defined concentrations of iloprost (10 pmol/l to 10 μmol/l; 20-min treatment) was measured using a radio-receptor competition assay (Amershams Biosciences) as described previously (44). Results were analyzed using GraphPad Prism version 4.0 software. For the concentration response, a nonlinear, curve-fitting program (GraphPad Prism, version 4.0.1) was used and the EC_{50} was determined.

Nonradioactive in vitro assay for PKA activity. We treated uSMCs as indicated in the figure legends. Cells were lysed as described previously (44), and total protein concentration in each lysate was determined by Bradford assay. Equal amounts of protein from each sample (10–30 μg) were subjected to the PepTag Assay for Non-Radioactive Detection of cAMP Dependent Protein Kinase (Promega) using the fluorescence-labeled peptide substrate kemptide according to the manufacturer’s protocol and as described previously (44). The phosphorylated and nonphosphorylated kemptide were separated on a 0.8% agarose gel, and an image was obtained using a Typhoon Scanner (Molecular Dynamics). Densitometry was used to determine the quantity of phosphorylated substrate using Scion Image Beta (version 4.0.2).

Transient transfection of siRNA. Human uSMCs (1–1.5 × 10^6 cells) were transfected with siRNA using the basic SMC Nucleofector Solution and the Nucleofector Device (Amaxa). uSMCs were transfected with 0.5–0.65 μg (52–68 pmol) PKA α and 0.5–0.65 μg PKA β siRNA or 1.0–1.3 μg (80–104 pmol) siCONTROL for 48 h. Cells were treated with vehicle or iloprost for 20 min or 8 h and harvested for PKA activity or Western blotting as described above. The sequence for PKA α siRNA

Figure 13
PGI2 mimics iloprost-induced contractile apparatus and connexin 43 protein expression. (A) Effect of PGI2 on PKA activity. Human uSMCs were treated for 20 min with PGI2 or iloprost at the concentrations indicated and subjected to in vitro PKA assay. The phosphorylated kemptide gel is shown. (B) Effect of PGI2 on contractile apparatus protein and connexin 43 expression. (B) Human uSMCs were treated with PGI2 every 1.5 h up to 6 h or with a single treatment of iloprost for 6 h at the concentrations indicated. Western blots were performed with antibodies against SM2-MHC, calponin, β-tubulin, and GAPDH. (C) uSMCs were subjected to a single treatment of 0.6 μmol/l PGI2 for the indicated times. Western blot analysis was performed with antibodies against SM2-MHC, h-caldesmon, connexin 43, and β-tubulin as a loading control.
was 5′-AAGGUUGUUUGCACAGUCG3′ and the siRNA sequence for PKA Cβ was 5′-AAGAGUUUCCGCAAGGCAAC-3′ (Dharmacon). Microarray-validated nontargeting duplex siRNA (siCONTROL; Dharmacon) was used as a negative control.

**Statistics**

Unless otherwise indicated, data are mean ± SEM. Densitometry results were analyzed using Prism software (version 4.0; GraphPad), and significance of differences was determined using 1-way ANOVA with post-hoc Newman-Keuls test for multiple comparisons or 1-tailed Student’s t test. A P value less than 0.05 was considered significant.

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