Cyclic nucleotide phosphodiesterases (PDE) are the enzymes catalyzing the hydrolysis and inactivation of the second messengers, cAMP and cGMP. Eleven PDE families are described to date, and selective inhibitors of some PDEs families are currently used in clinic for treating cardiovascular disorders, erectile dysfunction, and pulmonary hypertension. Isoforms of the PDE4 family are involved in smooth muscle contraction and inflammation. PDE4 selective inhibitors are currently in clinical trials for the treatment of diseases related to inflammatory disorders. Because of their myorelaxant properties, we first examined their expression in human myometrium and uncover an increased expression of one specific isoform, PDE4B2, in the near-term myometrium as compared to myometrium in the nonpregnant state. Using human myometrial cells in culture, we demonstrated that PDE4B2 can be induced by its own substrate, under the control of one of the major utero-contracrive agonists, PGE2, itself upregulated by the proinflammatory cytokine IL-1β. Functionally, augmentation of global PDE4 activity decreases the ability of β-adrenergic agonists (the most commonly used tocolytic drugs) to inhibit myometrial contraction at the end of pregnancy and during pathophysiological situations, such as persistent intrauterine inflammation which is a major cause of very preterm delivery. Currently exploring the anti-inflammatory properties of PDE4 inhibitors in gestational tissues, we recently demonstrated the ability of these drugs to block a persistent inflammatory response of the foetal membranes in Humans and to prevent inflammation-driven preterm delivery and foetal demise in mice. These data open up a new therapeutical strategy to prevent inflammation-induced preterm delivery and its sequelae in very preterm infants.
Background

Tea leaf has been used for 2,000 years in traditional Chinese medicine for its therapeutical virtues before becoming a traditional drink. One of its main active ingredients, theophylline, is a selective inhibitor of cyclic nucleotide phosphodiesterases (PDE), which hydrolyses the second messengers, cAMP and cGMP. Eleven families of PDE are described to date with different substrate preference, kinetics and modulators [1]. Selective PDE inhibitors of the type 3 and type 5 families are currently used in clinic, respectively for cardiac disease, erectile dysfunction, and pulmonary hypertension. PDE4 specifically hydrolyses cAMP with high affinity and PDE4 inhibitors are highly scrutinized because of their myorelaxant and anti-inflammatory properties.

PDE4 is the largest PDE family known to date, encoded by four distinct genes, PDE4A-D, and generating more than 20 different isoforms in mammals by means of differential promoters and splicing. The PDE4s are further divided in two main classes, the long and short forms that differ with the presence or not of regulatory domains at their amino-terminus [2,3] (Figure 1). In particular, the presence of a PKA-phosphorylation site well-conserved over the PDE4 long forms activates these isoforms in a short-term feedback loop. Conversely, cAMP- and hormonal-sensitive promoters control the transcription of some of the PDE4 short-forms, leading to a decrease in cAMP levels after a sustained elevation of the cyclic nucleotide content. These regulations, as we describe below, are critical to lessen the cAMP signalling in the utero-placental interface at the end of pregnancy.

Results

We examined in human myometrium the expression of the four genes of PDE4 in the nonpregnant state and near term. A concurrent expression of the mRNAs of the four genes with a predominance of the PDE4D and PDE4B mRNAs and a weaker expression of PDE4A and PDE4C genes was observed. Using semi-quantitative PCR and Western blot, we observed, in near term myometrium, an increase of the steady-state levels of PDE4B2 mRNA [4] and protein [5], a short form product of the PDE4B gene. We also explored functionally whether PDE4s are involved in the control of the myometrial contractility. Selective PDE4 inhibitors of the first (rolipram) and of the most recent developed generation achieve a complete relaxation of the myometrial strips, either in nonpregnant or pregnant state [4-6]. Although the β-adrenergic agonists, the most commonly used tocolytic agents, are able to completely relax the nonpregnant myometrium, they are unable to fully inhibit the pregnant uterine contractions. This phenomenon reflects the desensitization of the β-adrenergic system at the end of the pregnancy [7]. Application of a PDE4 inhibitor potentiates the effect of β-mimetics, which then achieves a complete relaxation of the pregnant strip [8] (Figure 2). It is worth noting that PDE4 inhibitors do not potentiate the effect of or act in synergy with β-adrenergic agonists on nonpregnant strips, indicating an increased role of the PDE4s in the contractility of the pregnant state.

We explored in parallel the molecular mechanism of the hormonal regulation of the PDE4 expression in primary culture of myometrial cells. Like in the tissue of origin, PDE4 is the major cAMP-hydrolyzing activity in these cells and the four PDE4 genes are expressed. An increase in intracellular cAMP concentration led to a significant increase in cAMP PDE activity through the selective induction of the short forms of the PDE4B and PDE4D genes, namely PDE4B2, PDE4D1 and PDE4D2 [9]. Hence, PDE4 upregulation provides a feedback mechanism by which cells might adapt from sustained stimulation of the cAMP pathway. We then shown that prostaglandin E2, one of the major effectors of labour, can induce PDE4B2 and PDE4D short forms, through a cAMP-dependent pathway, in the myometrial smooth muscle cells [8]. This regulation may occur during labour, when PGE2 concentration rises dramatically, but also in certain pathophysiological situations, with intrauterine inflammation, because we observed that the proinflammatory cytokine IL1β, by increasing PGE2 concentration, is also able to induce PDE4B2 [10]. Furthermore, impregnation of myometrial strips with PGE2 leads to a heterologous desensitization of the β-adrenergic system. This is translated by a decrease of the relaxant strength of β-adrenergic agonists and is reversed in the presence of PDE4 inhibitors [8]. Therefore, a co-application of β-mimetics and PDE4 inhibitors may greatly improve the efficiency of the tocolysis in case of preterm labour.

Besides, because PDE4 activity is high in the amnion and the chorion, we established that not only the myometrium is a target for PDE4 inhibition but also the fetal membranes. These data are of importance because amniochorionic membranes are major sources of pro-inflammatory mediators and uterotonic agents. Indeed, treatment of amniochorionic explants with the membrane product of E. coli, the lipopolysaccharide (LPS), induced synthesis of a pro-and anti-inflammatory cytokine, TNFα and IL-10 respectively but also induced synthesis of prostaglandins (PGs) and metalloproteases (MMPs) [11]. In addition, PDE4 activity was independently increased by LPS treatment. On this model of explants, we demonstrated that PDE4 selective inhibitors diminished greatly the production of TNFα whereas they enhanced the release of IL-10. They then decreased the production of PGE2 and MMP9. Recently, we derived primary cells from the different cell types present in amniochorionic membranes: amniotic epithelial cells,
Potentiation of the relaxant effect of β-adrenergic agonist in the presence of a PDE4 selective inhibitor in pregnant myometrium

Figure 2
Potentiation of the relaxant effect of β-adrenergic agonist in the presence of a PDE4 selective inhibitor in pregnant myometrium. Strips were mounted for isometric recording under 900 mg tension and after 2 h equilibration, contractile activity was assessed. Myometrial strip of either pregnant (A) or nonpregnant women (B) were preincubated with 3 × 10⁻⁹ M RP73401, a competitive selective PDE4 inhibitor (open circles) or saline (closed circles), 30 min prior the construction of the concentration-response to salbutamol. Results are expressed in percentage of maximal contraction at the end of the equilibration period. Strips from the same patients were subjected in parallel to treatments. Data are expressed as the mean ± SEM for strips isolated from myometrium of five different pregnant women and three different nonpregnant women. (Adapted from [8]).
fibroblasts and chorionic epithelial cells, the choriotrophoblasts, which serve as models to identify signaling pathways involved in a local inflammation.

Very recently, we demonstrated that PDE4 inhibition prevents in vivo preterm delivery and fetal demise in mice [12]. We adapted the inflammation-induced preterm delivery model in mice established by Elovitz et al. [13]. On day 15 of gestation (term 20 days), LPS or saline was injected in the uterus. Two hours after surgery, vehicle or the PDE4 selective inhibitor, rolipram, was injected in the peritoneal cavity and the frequency of preterm delivery and the fetal demises were recorded 48 h post-surgery. In the presence of rolipram, we observed that LPS-induced preterm delivery is prevented as well as fetal demise, the number of live pups in rolipram-LPS injected mice being not statistically different from the control group (Figure 3). Moreover, we showed that the increased cytokine concentrations in the amniotic fluid of LPS-treated dams were significantly reduced by the treatment with rolipram. We evaluated by immunohistochemistry whether the transcription factor of inflammation, NFκB, was translocated in the nucleus in gestational tissues as an index of activation. Interestingly we found that, six hours after LPS injection, NFκB was located in the nucleus of a specific placental cell, at the feto-maternal interface, the glycogen trophoblast whereas this translocation was absent in trophoblasts of dams treated with rolipram. No nuclear translocation of NFκB was observed in myometrial cells.

**Conclusion**

These original studies uncover a new potential target, the myometrial PDE4s, for tocolysis. In addition, PDE4 inhibitors are able to block the production of inflammatory cytokines produced by myometrial and fetal explants that may induce preterm delivery (Figure 4). Moreover the in utero exposure to inflammation is a high risk factor of neurological and respiratory disorders of the preterm infant. Therefore, the combination of the myorelaxant and anti-inflammatory properties of the PDE4 inhibitors make these molecules particularly attractive for the prevention of inflammation-induced preterm delivery and its consequences on the premature infant.

**Competing interests**

The authors declare that they have no competing interests.

**Acknowledgements**

We wish to acknowledge the sponsorship of Ferring, PerkinElmer and Serono, who helped to cover publishing charges of the article. We are also indebted to the financial support and organization by the EU project SAFE (The Special Non-Invasive Advances in Fetal and Neonatal Evaluation Network, LSHB-CT-2004-503243) of the 2005 and 2006 PTL workshops, where data of this article were presented as oral communications.
References
1. Mehats C, Andersen CB, Filopanti M, Jin SL, Conti M: Cyclic nucleotide phosphodiesterases and their role in endocrine prostaglandin signaling. Trends Endocrinol Metab 2002, 13:29-35.
2. Conti M, Richter W, Mehats C, Livera G, Park JY, Jin C: Cyclic AMP-specific PDE4 phosphodiesterases as critical components of cyclic AMP signaling. J Biol Chem 2003, 278:5493-5496.
3. Houslay MD. PDE4 cAMP-specific phosphodiesterases. Prog Nucleic Acid Res Mol Biol 2001, 69:249-315.
4. Leroy MJ, Cedrin I, Breuiller M, Giovagrandi Y, Ferre F: Correlation between selective inhibition of the cyclic nucleotide phosphodiesterases and the contractile activity in human pregnant myometrium near term. Biochem Pharmacol 1989, 38:9-15.
5. Mehats C, Tanguy G, Paris B, Robert B, Perpin N, Ferre F, Leroy MJ: Pregnancy induces a modulation of the cAMP phosphodiesterase 4-conformers ratio in human myometrium: consequences for the utero-relaxant effect of PDE4-selective inhibitors. J Pharmacol Exp Ther 2000, 292:817-23.
6. Oger S, Mehats C, Barnette MS, Ferre F, Cabrol D, Leroy MJ: Anti-inflammatory and utero-relaxant effects in human myometrium of new generation phosphodiesterase 4 inhibitors. Biol Reprod 2004, 70:458-64.
7. Berg G, Andersson RG, Ryden G: Beta-adrenergic receptors in human myometrium during pregnancy: changes in the number of receptors after beta-mimetic treatment. Am J Obstet Gynecol 1985, 151:392-396.
8. Mehats C, Tanguy G, Dallot E, Cabrol D, Ferre F, Leroy MJ: Is up-regulation of phosphodiesterase 4 activity by PGE2 involved in the desensitization of beta-mimetics in late pregnancy human myometrium? J Clin Endocrinol Metab 2001, 86:5358-65.
9. Mehats C, Tanguy G, Dallot E, Robert B, Rebourcec R, Ferre F, Leroy MJ: Selective up-regulation of phosphodiesterase-4 cyclic adenosine 3',5'-monophosphate (cAMP)-specific phosphodiesterase variants by elevated cAMP content in human myometrial cells in culture. Endocrinology 1999, 140:3228-37.
10. Oger S, Mehats C, Dallot E, Ferre F, Leroy MJ: Interleukin-1beta induces phosphodiesterase 4B2 expression in human myometrial cells through a prostaglandin E2- and cyclic adenosine 3',5'-monophosphate-dependent pathway. J Clin Endocrinol Metab 2002, 87:5524-31.
11. Oger S, Mehats C, Dallot E, Cabrol D, Leroy MJ: Evidence for a role of phosphodiesterase 4 in lipopolysaccharide-stimulated prostaglandin E2 production and matrix metalloproteinase-9 activity in human amniochorionic membranes. J Immunol 2005, 174:8082-9.
12. Schmitz T, Souil E, Herve R, Nicco C, Batteux F, Germain G, Cabrol D, Evain-Brisson D, Leroy MJ, Mehats C: PDE4 inhibition prevents preterm delivery induced by an intraterine inflammation. J Immunol 2007, 178:1115-21.
13. Elovitz MA, Wang Z, Chien EK, Rychlik DF, Philippe M: A new model for inflammation-induced preterm birth: the role of platelet-activating factor and Toll-like receptor-4. Am J Pathol 2003, 163:2103-11.