Expression of platelet activating factors and its receptor in cat uterus during early pregnancy

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Summary: Platelet activating factors (PAF1β, PAF1γ, PAF:αh, PAF-R) have a significant role during mammalian pregnancy process. This study aimed to investigate expression profiles of PAF gene family in cat uterus during pregnancy. Tissue samples were collected via ovariohysterectomy at days 7 (Gr1; n=3), 18-20 (Gr2; n=7), 24-25 (Gr3; n=5) after mating from the different regions of the uterus. Relative expression levels of genes were quantified using real-time qRT-PCR. A mixed model was fitted on the normalized data and Least Significant Difference (LSD) test was employed to determine significant differences between groups. Expressions of PAF genes were assessed in the cat uterus during pregnancy around implantation stage; however, no significant differences were detected between the preimplantation and the implantation stage. Similarly, no significant difference concerning mRNA expression was assessed between different parts of the uterus during pregnancy. In conclusion, PAF genes are expressed in the cat uterus during early pregnancy. However, this expression is not associated with changes in the uterus during early pregnancy.

Key words: PAF, cat, pregnancy, qRT-PCR.

Kedi uterusunda, platelet aktive edici faktörler ve reseptörünün erken gebelik döneminde ekspresyonu

Özet: Platelet aktive edici faktörler (PAF1β, PAF1γ, PAF:αh, PAF-R), memeli hayvanların gebelik sürecinde önemli bir rol sahiptir. Bu çalışmamızın amacı, kedi uterusunda PAF gen ailesi ekspresyon profilerini araştırmaktır. Doku örnekleri çiftleşme sonrası 7 (Gr1; n=3), 18-20 (Gr2; n=7), 24-25 (Gr3; n=5) günlerde yapılan ovariohisterektomi operasyonları ile uterusun farklı bölgelerinden toplandı. Nispi mRNA ekspresyon seviyeleri real-time PZR kullanılarak tespit edildi. Normalize edilmiş veriler karsılık model kullanılarak analiz edildi ve istatistik olarak farklı olan gruplar Asgari Önemli Fark (AÖF) testi ile tespit edildi. Kedi uterusunda implantasyon sürecinde PAF genlerinin ekspresre olduğu gözlemlemiştir ancak; istatistik olarak çalışma grupları ve uterus bölgeleri arasında herhangi bir fark bulunmamıştır. Sonuç olarak, kedi uterusunda erken gebelik sürecinde PAF genlerin ekspresre olmaktadır. Ancak; bu ekspresyonların uterusa göre değişikliklerle ilişkiyi yoktur.

Anahtar sözcüklar: PAF, kedi, gebelik, qRT-PZR

Introduction
Platelet activating factor (PAF) has a potential for biological and pathophysiological events (37) and is a mediator synthesized by platelets, neutrophils and macrophages, leading to platelet aggregation (14). However, it is also produced by a number of nonimmune cells such as endothelial (9), vascular, muscle (43), glomerular (13) and endometrial cells (4, 5, 7, 45). PAF plays a role in many reproductive events (16), including ovulation (1), fertilization (27), implantation and early development of embryo (28). PAF has been determined in the uterus of rats (45), rabbits (5), dogs (35) and humans (4). It plays a significant role in mammalian reproduction (29, 32). In peri-implantation period, PAF concentration increases significantly in the human uterus (4) and stimulates prostaglandin E2 (PGE2) production in fetal membranes of humans (8, 24). The PAF and PAF-R (receptor) synthesis is affected by the presence of steroids in the endometrium in humans (42). Endometrial PAF-R mRNA expression was significantly induced by estradiol and progesterone in sheep (11). After ovariectomy, concentration of PAF in the rat uterus was declined (25). PAF could regulate its own receptor in the endometrial cells. Addition of exogenous PAF in
endometrial cell culture increased PAF-R expression in sheep (10). Stimulation by progesterone increased the concentration of PAF in the endometrium of ewe on days 10-16 of the cycle (11).

Tiemann et al. (41) observed that PAF plays an important role in the control of the uterine neovascularization and modulation of the cell growth of the endometrium in preparation of implantation. The concentration of PAF is controlled by synthesis and degradation of PAF in the uterus. Degradation of PAF is governed by PAF acetylhydrolase (PAF:ah), which is the metabolic enzyme for PAF (38). PAF:ah activity is minimum in the uterus during the preimplantation phase of pregnancy in the mouse uterus (30). Implantation is a process between embryo and the endometrium and PAF plays an important role in cell interactions in the uterus during this process (44).

Biological effects of PAF are thought to be receptor (PAF-R) mediated (17). This receptor is present on various cell types, such as smooth muscles, neutrophils, monocytes (12) and also on endometrial cells (20). PAF-R is mainly placed in the endometrial luminal and glandular epithelium in humans (3), rabbits (21), and heifers (40). Investigations on local immunological interactions between fetus and endometrium could be helpful to develop nonsurgical contraceptive methods. A physiological effect of endometrial PAF has not been fully explained in felines. The hypothesis of the study is that the PAF gene family (PAF1β, PAF1γ, PAF:ah, PAF-R) has an important role in feline early and mid-gestation. The objective of this study was to investigate the expression profile of members of the PAF family in the pregnant feline uterus throughout gestation.

Materials and Methods

Animal Material and Experimental Procedure: Cats were ovariohysterectomized at days 7 (Gr1; n=3), 18-20 (Gr2; n=7), 24-25 (Gr3; n=5) after mating, respectively. The preimplantation group (day 7) was proven pregnant (Gr2; n=7), 24-25 (Gr3; n=5) after mating, respectively.

The preimplantation group (day 7) was proven pregnant (Gr2; n=7), 24-25 (Gr3; n=5) after mating, respectively. Briefly, total RNA isolation was performed by using TRIzol Reagent (Invitrogen, Carlsbad, CA, USA) and approximately 50 mg of uterine tissue. The RNA pellets in each tube were dissolved in 100 μl of DEPC-treated sterile water. RNA quality was determined by loading 10 μl of each RNA sample on 2% agarose gel and by measuring 260/280 ratio in NanoDrop ND-100 (Thermo Scientific, Wilmington, DE, USA). Two μg of RNA samples were first cleaned for possible genomic DNA contamination by DNase-I digestion and then subjected to reverse transcription reaction for first strand complementary DNA (cDNA) synthesis using RevertAidTM FirstStandart cDNA Synthesis Kit (Fermentas, Glen Burnie, MD, USA) according to the manufacturer’s instructions.

Primer Design and Real Time PCR: Primers for PAF gene family were derived from multispecies sequences by using the Primer3 program from NCBI (http://www.ncbi.nlm.nih.gov/) database. The primer pair sequences and product sizes are shown on Table 1.

qRT-PCR was used to evaluate the expression profiles of PAF gene families. The reaction was set up as follows: 12.5 μl SYBR Green Master Mix (2X), 5 pMol of each primer (Table 1), 1μl cDNA and ddH2O up to 25 μl of final volume. Thermal cyclic conditions were initial denaturation at 95°C for 10 min followed by 40 cycles of denaturation, annealing and extension (95°C 30 sec, 60°C 30 sec, 72°C 30 sec) on an Applied Biosystems 7500 Real-Time PCR System (Applied Biosystems Foster City, CA, USA) at the Selcuk University Advanced Technology Research and Application Center (ILTEK). From the RNA extraction to the qRT-PCR, the whole procedure was performed twice as technical replicate. As housekeeping gene, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was selected according to Kayis et al. (18).

Statistical Analysis: Before statistical analysis, the efficiencies of amplification of target genes and internal

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Table 1. Primers used in qRT-PCR analyses

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer Sequence (5'-3')</th>
<th>PCR Product (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forward</td>
<td>Reverse</td>
<td></td>
</tr>
<tr>
<td>GAPDH</td>
<td>ATCACCATCTTCCAGGAGCGAGA</td>
<td></td>
</tr>
<tr>
<td></td>
<td>GTCTTCTGGGTGGCAGTATGG</td>
<td>341</td>
</tr>
<tr>
<td>PAF1β</td>
<td>GCCACAGGGCCAAGTATGTGGT</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AACCTTGAGGAGCTGTACCTT</td>
<td>100</td>
</tr>
<tr>
<td>PAF1γ</td>
<td>GTCGTCGACCCACAATGGT</td>
<td></td>
</tr>
<tr>
<td></td>
<td>GTCGTACCGACCAATGGT</td>
<td>142</td>
</tr>
<tr>
<td>PAF:ah</td>
<td>TTTCTGCAAGCAGATCCCCAGAGGA</td>
<td></td>
</tr>
<tr>
<td></td>
<td>GATGCACCTGGCAATTTGCAACAT</td>
<td>125</td>
</tr>
<tr>
<td>PAF-R</td>
<td>TGGCCTCTCCTTTGCTCTGTGCAT</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TGCATGAGTAGCGTGCGAGTG</td>
<td>249</td>
</tr>
</tbody>
</table>

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control (GAPDH) were examined using real-time PCR amplification of serial dilutions (over a 256-fold range) of cDNA. Based on confirmation that the amplification efficiencies of the target and reference genes are nearly the same, data normalization process was performed according to Livak and Schmittengen (23) via $2^{-\Delta C_{i,j}}$ method, where $\Delta C_{i,j} = C_{T, \text{target}} - C_{T, \text{reference}}$ (where $C_{T, \text{target}}$ and $C_{T, \text{reference}}$ are threshold cycles for target and reference gene amplifications, respectively). Following normalization process, Anderson-Darling normality test was performed to obtain distributional properties of data. We performed log transformation on PAF1β, PAF1γ, and PAF:ah and square root transformation on PAF-R to stabilise variance. The following model was fit on the transformed data

$$y_{ijk} = \mu + G_i + L_j + GL_{ij} + e_{ijk}$$

with the effects defined as follows:

- $\mu$ = effect of the overall mean;
- $G_i$ = fixed effect of the $i^{th}$ group;
- $L_j$ = fixed effect of the $j^{th}$ region;
- $GL_{ij}$ = group – region interaction;
- $e_{ijk}$ = residual term for the tissue sample taken from $j^{th}$ region of $k^{th}$ individual that take place in $i^{th}$ group ($i=$ day 7, day 18-20, day 24-25);

Least Significant Difference (LSD) test was applied to detect statistically significant group(s). All analyses were carried out using R 2.14.1 (31).

Results

Resulting qRT-PCR products were separated by electrophoresis on 2% agarose gels and visualized after ethidium bromide staining (Fig. 1). Expressions of PAF1β, PAF1γ, PAF-R and PAF:ah were detected in the cat uterus. Expression profiles with 95% confidence interval (95% CI) of PAF genes during the preimplantation and postimplantation stage were illustrated in Fig. 2. The PAF1γ expression decreased in Gr3 but did not change significantly among the placental, interplacental sites and corpus uteri. Expression of PAF genes did not significantly differ between experimental days and among regions.

Discussion and Conclusion

Successful implantation of blastocysts and gestation requires synergistic interactions between embryo and the uterus. These events are controlled by variety of mediators and factors such as growth factors, cytokines, hormones and enzymes (6, 34, 35). Many of these factors are related to inflammation process. Implantation and inflammation process have several physiological events in common, including increased vascular permeability and some inflammatory mediator release. One of these factors is PAF. Spinks et al. (39), demonstrated that PAF inhibitor has preventive effect for implantation in mouse and rat (2). PAF induces contraction of the smooth muscle of the uterus. PAF was detected in amniotic fluid and PAF:ah decreased during late stage of pregnancy suggesting that PAF plays an important role in progress of gestation (19). Several parts of the reproductive tract and reproductive functions are affected by PAF. PAF affects the release of prostaglandin from endometrial tissue in ewes (33), humans (36), guinea pigs (26) and heifers (15). In the present study, PAF expression in the uterus of pregnant cats was detected between days 7 to 25 of pregnancy. PAF is released by ovine endometrial cells at days 10 and 16 of the sexual cycle. Stimulation of the endometrium by steroid hormones may induce both PAF production and PAF-R expression. Expression of PAF-R increased in the preimplantation period in heifers. These facts support the idea that PAF plays an important role in the implantation process (41).

The result of the current study showed the expression of PAF, PAF:ah and PAF-R within the endometrium during early pregnancy in the cat. PAF gene expression was observed in cat endometrium at days 7, 20 and 25. In contrast to other species, PAF gene expression did not increase significantly in the course of pregnancy. Additionally, the expression of genes was not different between groups and regions (d7, d20, d25). And the PAF1γ expression decreased in Gr3 but did not change significantly among the placental, interplacental sites and corpus uteri. It is known from previous studies in other species that PAF expression is under the control of progesterone. Since pregnant cats are expected to have high concentrations of circulating serum progesterone, a visible influence was expected. The expression of PAF may also be associated with sexual cycle stage and other steroid hormone levels. Further research is needed to evaluate expression profiles of these genes in non-pregnant cats together with the pregnant ones as in this study. Furthermore, the protein expression of fetal and uterine cells and the changes in the course of pregnancy should be investigated.

Fig 1. Expression of PAF genes at mRNA level
Şekil 1. PAF genlerinin mRNA düzeyinde ekspresyonları
Figure 2. Expression and 95% confidence interval of PAF genes in feline uterus during the preimplantation and postimplantation periods.

Şekil 2. Kedi uterusunda PAF genlerinin, preimplantasyon ve postimplantasyon dönemlerindeki ekspresyonları ve %95 güven aralıkları.
References


30. O’Neill C (1998): Autocrine mediators are required to act on the embryo by the 2-cell stage to promote normal...


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