Title:

Uterine and placental distribution of selected extracellular matrix (ECM) components in the dog.

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Running title: ECM modification of canine pregnant uterus

Summary sentence: The canine uterine ECM is only moderately modified in early pregnancy, but undergoes vigorous reorganization processes in the uterus and placenta following implantation.

Key words: dog (*Canis lupus familiaris*), extracellular matrix (ECM), collagen, connexins, TIMPs, pregnancy, remodeling, uterus.

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ABSTRACT

For many years, modifications of the uterine extracellular matrix (ECM) during gestation have not been considered as critical for successful canine (*Canis lupus familiaris*) pregnancy. However previous reports indicated an effect of free-floating blastocysts on the composition

- 40 of the uterine ECM. Here, the expression of selected genes involved in structural functions, cell-to-cell communication and inhibition of matrix metalloproteinases were targeted utilizing qPCR and immunohistochemistry. We found that canine free-floating embryos affect gene expression of *FN1*, *ECM1* and *TIMP4*. This seems to be associated with modulation of trophoblast invasion, and proliferative and adhesive functions of the uterus. Although not
- 45 modulated at the beginning of pregnancy, the decrease of structural ECM components (*i.e.*, *COL1*, *-3*, *-4* and *LAMA*) from pre-implantation towards post-implantation at placentation sites appears to be associated with softening of the tissue in preparation for trophoblast invasion. The further decrease of these components at placentation sites at the time of prepartum luteolysis seems to be associated with preparation for the release of fetal
- 50 membranes. Reflecting a high degree of communication, intercellular cell adhesion molecules are induced following placentation (Cx26), or increase gradually towards prepartum luteolysis (Cx43). The spatio-temporal expression of TIMPs suggests their active involvement in modulating of fetal invasiveness, and together with *ECM1* they appear to protect deeper endometrial structures from trophoblast invasion. With this, the dog appears to be an
- 55 interesting model for investigating placental functions in other species, *e.g.*, in humans in which *Placenta accreta* appears to share several similarities with canine subinvolution of placental sites (SIPS).

INTRODUCTION

60 Interactions between the canine (*Canis lupus familiaris*) uterus and the corpora lutea (CL), and the interchange between the uterus and fetal trophoblast, represent remarkably unique mechanisms among mammalian species.

The bitch is a monoestric, non-seasonal breeder in which ovulation occurs spontaneously (<u>Concannon 1993</u>). As in other domestic animals CL are formed following ovulation, but in

- 65 contrast to other species, in canine pregnancy no placental steroid hormone synthesis arises (<u>Hoffmann *et al.* 1994</u>). Consequently, the increasing circulating steroid levels in both non-pregnant and pregnant bitches fully originate from the CL with luteal progesterone (P4) being responsible for establishment and maintenance of pregnancy until term (<u>Concannon *et al.* 1989</u>). Due to the lack of luteolysis in the absence of pregnancy, peripheral P4 levels are
- 70 almost identical in non-pregnant and pregnant bitches until shortly before parturition when prepartum luteolysis occurs signaling the onset of parturition (Concannon *et al.* 1989). Consequently, in contrast to livestock, in the dog successful pregnancy does not depend on the inhibition of luteolysis to maintain the luteal life span and thereby embryo survival (Kowalewski *et al.* 2009, Gram *et al.* 2013, Kowalewski 2014). In other domestic animal
- 75 species pre-attachment embryos transmit signals needed to extend CL function (Niswender et al. 2000, Bazer et al. 2010, Bazer 2015), referred to as "maternal recognition of pregnancy" (Short 1969). This embryo-maternal communication synchronizes blastocyst development and uterine receptivity in a complex series of molecular and cellular mechanisms (Seshagiri et al. 2009, Paidas et al. 2010, Geisert 2015). In some species, e.g., in rodents or primates, the
- 80 uterine changes result in decidua formation and are therefore very pronounced. In primates, decidualization takes place even in the absence of embryos, while in rodents it is induced by their presence (reviewed in (Geisert 2015)). In early pregnancy, the uterine extracellular matrix (ECM) influences trophoblast invasion (Johnson *et al.* 2003) and remodelling of the decidua (Damsky *et al.* 1993, Lala & Nandi 2016, Smith *et al.* 2016). The formation of
- 85 decidua, *e.g.*, in mice, is associated with clearing of hyaluronic acid from the uterine stroma (Brown & Papaioannou 1992). In addition, *in vitro* studies demonstrate that decidualizing stromal cells of mouse and human origin produce ECM products such as basal lamina-like materials (Wewer *et al.* 1985, Wewer *et al.* 1986, Kisalus *et al.* 1987).
- Despite the absence of an antiluteolytic signal at the beginning of pregnancy, in the dog synchronization between blastocyst development and uterine preparation for implantation is also essential for healthy pregnancy. Thus, in dogs and other carnivores decidualization also takes place and is associated with implantation and placenta formation (<u>Wislocki & Dempsey</u>)

1946, Dempsey & Wislocki 1956, Kautz et al. 2014, Graubner et al. 2017b). In accordance with the aforementioned *in vitro* decidualization studies with mouse and human cells, canine 95 uterine cells can also be decidualized in vitro. Their response to a decidual stimulus (i.e., increased secretory and proliferative activity) provides evidence that also in dogs uterine ECM modulation during decidualization occurs (Kautz et al. 2015). Furthermore, functional genomics studies imply that *in vivo* canine free-floating embryos transmit signals to the uterus and discretely, *i.e.*, without clearly visible morphological changes, affect uterine ECM 100 composition at the early stage of pregnancy (Graubner et al. 2017a). Indicating the beginning of canine decidualization, these embryo-derived signals control the expression of uterine biochemical markers of decidualization (Kautz et al. 2014). The first morphological signs of decidualization are found later, commencing with attachment of the trophoblast (Graubner et al. 2017b). Ultimately, the uterine modifications become advanced and maternal-derived 105 decidual cells are formed as an indispensable component of the endotheliochorial placenta in the dog. Decidual cells are needed for successful pregnancy since they are the only cells in the canine placenta expressing the nuclear P4 receptor (PGR) (Vermeirsch et al. 2000, Kowalewski et al. 2010). Additionally, these cells and maternal endothelium can resist trophoblast digestion. Adequate embryo-maternal contacts between the trophoblast and 110 maternal decidual cells, all embedded in the ECM components, are important for maintenance and termination of pregnancy. Thus, withdrawal of P4 from decidual cells will interfere with

and termination of pregnancy. Thus, withdrawal of P4 from decidual cells will interfere with their physiological function and lead to abortion or preterm parturition (Kowalewski *et al.* 2010). ECM components appear to be involved here in feto-maternal communication leading to the luteolytic cascade, expulsion of fetuses and fetal membranes. Disturbances in embryo-

115 maternal communication may result in clinical conditions like SIPS (subinvolution of placental sites), resembling placenta accreta in humans characterized by excessive trophoblast invasion.

Formation of the endotheliochorial placenta in the dog and other carnivores, indeed represents a unique type of placentation, with strong, but yet restricted, invasion. So far, little attention

120 has been paid to uterine function in dogs during pregnancy, resulting in limited knowledge about its possible involvement in the establishment, maintenance and termination of pregnancy.

Because of this lack of information, driven by the hypothesis that ECM compounds are strongly modified throughout canine gestation, this study focused on the localization (utilizing

125 immunohistochemistry (IHC)) and expression (using semi-quantitative TaqMan PCR) of several ECM proteins in the canine uterus. Selected stages of the reproductive cycle were

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considered, starting with morpho-functional modifications induced by the presence of preattachment embryos, through implantation, early and late placentation, until prepartum luteolysis. A variety of different factors collectively represent the term ECM. Here, the uterine localization and expression of ECM components with structural and adhesive functions were investigated (*e.g.*, collagen (COL) 1, -3 and -4, alpha smooth muscle actin (α SMA), and fibronectin (FN) 1). Additionally, proteins of the basal lamina, proteins involved in cell-to-cell communication, and proteins involved in inhibition of matrix metalloproteinases were targeted (*e.g.*, connexin (Cx) 26 and -43, laminin alpha (LAMA) 2, extracellular matrix protein 1 (ECM1), and tissue inhibitor of metalloproteinases (TIMP) 2

MATERIALS AND METHODS

Sample collection

and -4).

- 140 For the present study, tissue material from 31 crossbred healthy bitches aged 2-8 years was used. Animals were randomly assigned to the following groups: (1) early pregnant (pre-attachment days 10-12; n=10); (2) corresponding non-pregnant controls (n=8); (3) early post-implantation (days 18-25 of pregnancy; n=5); (4) mid-gestation (days 35-40 of pregnancy; n=5); and (5) prepartum luteolysis (n=3).
- 145 Estrus detection was performed by cytological evaluation of vaginal smears and regular measurements of P4 concentrations every 2-3 days starting with the proestrous bleeding. The day when P4 blood levels increased > 5ng/ml for the first time was determined as the day of ovulation. Considering the peculiar need for canine oocyte maturation within the oviduct (on average 2-3 days after ovulation), bitches were mated 2 days after ovulation (defined as day 0
- 150 of gestation). Tissue collection was performed through routine ovariohysterectomy (OHE). In the early pregnant group (group 1), OHE was performed between days 10-12 of gestation. Pregnancy was confirmed by flushing free-floating (*i.e.*, pre-attachment) embryos. Dogs negative for embryo flushing were allotted to group 2 (corresponding non-pregnant control at day 10-12 of the luteal phase). The prepartum luteolysis takes place in the dog 12-24 h before
- 155 any visible physiological signs of parturition. Therefore, for group 5, blood samples were collected every 6 h starting on day 58 of pregnancy. When circulating P4 levels continued to decrease below 2-3 ng/ml in 3 consecutive measurements, indicating the prepartum luteolysis, samples were collected. Tissue collection in groups 1 and 2 consisted of uterine cross-sections (the whole thickness of the uterine wall). For groups 3-5, sampling included utero-placental
- 160 cross-sections (Ut-Pl; the whole thickness of the uterine wall, *i.e.*, uterus with adjacent

placenta). Additionally, for groups 3 and 4 inter-placental uterine cross-sections (inter-Pl; full thickness like groups 1 and 2) were collected. No inter-Pl samples were available from the natural luteolysis group of dogs (group 5).

For semi-quantitative real-time PCR, cellular RNA was stabilized by immersing the samples

- 165 for 24h at 4°C in RNAlater® (Ambion Biotechnologie GmbH, Wiesbaden, Germany); prolonged storage was at -80°C. For immunohistochemistry (IHC) studies, tissues were fixed for 24h in 10% neutral phosphate-buffered formalin at 4°C, and subsequently washed for 1 week with PBS (phosphate buffered saline) and embedded in paraffin wax. For additional information about tissue sampling and processing procedures, see previous reports
- 170 (<u>Kowalewski *et al.* 2010</u>, <u>Gram *et al.* 2013</u>, <u>Kautz *et al.* 2014</u>). All uterine tissue samples were used for semi-quantitative real-time PCR, while 3 randomly chosen samples from each group were used for IHC.

All experimental procedures were carried out in accordance with animal welfare legislation and approved by the respective authorities of the University of Ankara (permits no. 2006/06

 and 2008-25-124) Ankara, Turkey, and Justus-Liebig University Giessen (permit no. II 25.3-19c20-15c GI 18/14 and VIG3-19c-20/15c GI 18,14) Giessen, Germany.

RNA isolation, reverse transcription (RT), semi-quantitative (TaqMan) PCR and evaluation of data

- Semi-quantitative TaqMan PCR was performed on total RNA isolated from all dogs. For this, TRIZOL[®] reagent (Invitrogen, Carlsbad, CA, USA) was used according to the manufacturer's instructions. From each sample 10ng total RNA were used for RQ1 RNAse-free DNAse treatment (Promega, Dübendorf, CH). The RT reaction was performed according to the manufacturer's instructions with the High Capacity cDNA Reverse Transcription Kit including RNase Inhibitor (Applied Biosystems from Thermo Fisher Scientific, Foster City, CA, USA). Following this, amplification of cDNA was performed with the TaqMan[®]PreAmp Master Mix Kit (Applied Biosystems) according to the supplier's protocol. Detailed information about the TaqMan PCR procedure has been previously published (Kowalewski *et al.* 2006b, Kowalewski *et al.* 2011, Kautz *et al.* 2015). In short, reactions were run in
- 190 duplicates with the Fast Start Universal Probe Master (ROX®) (Roche Diagnostics AG, Rotkreutz, CH) in an automated fluorometer (ABI PRISM® 7500 Sequence Detection System, Applied Biosystems). For negative controls, the so-called RT-minus control was used to check for potential genomic DNA contamination. Additional controls consisted of running experiments with autoclaved water instead of cDNA (no-template control) (Kowalewski *et al.*

- 195 2006b, Kowalewski et al. 2011). If canine-specific TaqMan systems were not available commercially, Primer Express Software v 2.0 (Applied Biosystems) was used to select primers and probes which were purchased from Microsynth, Balgach, CH. TaqMan probes were designed with 6-FAM- and TAMRA-labels. For a list of self-designed primer-sequences and TaqMan systems see <u>Table 1</u>. The self-designed expression assays were validated by the
- 200 CT slope method, which tests the TaqMan efficiency at different quantities of the targeted RNA as previously described (Kowalewski et al. 2011). The reaction efficiency for all selfdesigned assays was approximately 100%. Three reference genes, GAPDH, Cyclophilin A (PPIA) and ACTIN-B, were used for normalization to ensure homogeneity of variances. The following canine-specific TaqMan Gene Expression Assays were ordered from Applied
- Biosystems: *PPIA* (Prod. No. Cf03986523-gH); *ACTIN-B* (Prod. No. Cf03023880_g1), collagen type 1, alpha 1 (*COL1A1*) (Prod. No. Cf02741575_mH); COL type 3, alpha 1 (*COL3A1*) (Prod. No. Cf02631366_m1); COL type 4, alpha 1 (*COL4A1*) (Prod. No. Cf02696157_mH). For relative quantification, the comparative CT method (ΔΔCT method) was applied as previously described (Kowalewski *et al.* 2010, Kowalewski *et al.* 2011). The
- 210 sample with the lowest expression was used as a calibrator.
 Statistical differences between the non-pregnant and early pregnant group (groups 1 and 2) were determined by an unpaired, two-tailed Student's t-test; a p-value <0.05 was considered as statistically significant. When target gene expression was compared in more than 2 observational groups (time-dependent expression at selected time points during pregnancy), a</p>
- 215 parametric one-way ANOVA was applied. In the case of p<0.05, the Tukey-Kramer multiple comparisons post-test was performed. Statistical assumptions, such as normality and equality of variances were tested prior to applying the t-test or ANOVA. All analyses were performed with GraphPad 3.06 software (GraphPad Software, San Diego, CA, USA). Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).
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Immunohistochemistry

Standard immunohistochemistry (IHC) was performed to identify the localization of several ECM-proteins. Detailed information about our indirect immunoperoxidase method has been published previously (Kowalewski *et al.* 2006a, Kowalewski *et al.* 2006b). In short, following embedding in paraffin, tissues were sectioned into 2-3µm thickness, mounted onto SuperFrost

microscope slides (Menzel-Glaser, Braunschweig, DE). Following de-paraffinization using xylene and rehydratation in an ethanol series, antigen retrieval was performed using either heat-induced epitope retrieval (HIER) or proteolytic-induced epitope retrieval (PIER). HIER

was performed in a microwave oven at 560 W for 15 min in 10mM citrate buffer with pH 6.0.

- PIER was performed using 0.25% pepsin (P7000-25G Sigma-Aldrich Chemie GmbH, Buchs, CH) in 10mM HCl at 37°C. The duration of PIER varied from 5 min to 90 min depending on the antibody (for detailed information see <u>Table 2</u>). Following antigen retrieval, sections were treated with 0.3% hydrogen peroxide in methanol for 30 min to block endogenous peroxidases. After that, blocking was performed with 10% normal serum from the same species in which the secondary antibody was produced, and overlaid with primary antibodies
- overnight at 4°C. For negative controls, non-immune IgGs of the same species instead of the primary antibody at the same protein concentration as for the primary antibody were used (isotype controls).

Following incubation with the primary antibody, sections were washed with IHC buffer/0.3%

- 240 Triton X pH 7.2-7.4 (0.8 mM Na₂HPO₄, 1.74 mM KH₂PO₄, 2.68 mM KCl, 137 mM NaCl), and incubated with biotin-labeled secondary antibodies at 1:100 dilution. Next, the signals were enhanced using the Vectastain ABC Kit (Vector Laboratories, Inc., Burlingame, CA, USA) for 30 min, and immune reactions were visualized with the Liquid DAB+ substrate kit (Dako Schweiz AG, Baar, CH). Counter-staining was performed with hematoxylin. Next,
- 245 dehydration was performed in an ethanol series and xylene, and mounted using Histokit® (Assistant, Osterode, DE). Detailed information about commercially available primary and secondary antibodies, antigen retrieval and concentrations is given in <u>Table 2</u>. All groups within one experiment were simultaneously stained with Liquid DAB+ substrate kit (Dako) for the same target using the same development time.
- All slides were qualitatively evaluated for localization of the respective target protein and representative pictures were taken. Two researchers with good experience in canine uterine and placental histology evaluated the signals of target genes independently, using an ordinal system of none (-), weak (+/-), strong (+), and very strong (++) present (<u>Table 3</u>).
- To identify the specific localization of TIMP2 und TIMP4 in the canine placenta, 4 255 consecutive slides (2-3µm thickness) were prepared from placenta in the prepartum luteolysis group. Each consecutive slide was stained with a different primary antibody (TIMP2, TIMP4, vimentin (VIM), pan-cytokeratin (pan-CYT)). Vimentin was used as a marker of mesenchymal cells in order to differentiate between different placental cellular compartments. It stains positively for decidual cells, endothelial cells and fibroblasts. Cytokeratin served as a marker for epithelial cells, staining positively in fetal trophoblast. The staining and evaluation
 - procedures were as described above.

RESULTS

mRNA expression of ECM-proteins in canine uterus, inter-Pl and Ut-Pl compartments at selected time points during pregnancy and prepartum luteolysis.

In all samples, the mRNA expression of ECM-proteins, cell adhesion molecules and modulators of ECM was detectable by applying semi-quantitative (TaqMan) PCR.

To evaluate embryo-induced changes in the uterus, a pairwise comparison between nonpregnant and early pregnant (pre-attachment) bitches was performed. In this contrast, *FN1* was significantly suppressed (p=0.01) by the presence of embryos during early pregnancy (pre-attachment), in contrast to increased *ECM1* and *TIMP4* expression (p=0.01) (Figure1). *COL1, COL3, COL4, Cx26* and -43 were not affected at the beginning of early pregnancy (p>0.05, not shown). Effects of the presence of embryos on *LAMA2* and *TIMP2* mRNA expression in the pre-implantation uterus were investigated previously (Graubner *et al.* 2017a)

- 275 and were, therefore, excluded from the present study. Next, the uterine pre-implantation stage was compared to inter-Pl sites and to the Ut-Pl compartments from different stages of pregnancy (*i.e.*, post-implantation, mid-gestation and prepartum luteolysis). In inter-Pl sites uterine gene expression of *ECM1* was significantly upregulated at mid-gestation compared with pre- and post-implantation (p<0.001); it did not</p>
- 280 differ at earlier stages of pregnancy (p>0.05), *i.e.*, pre- and post-implantation (Figure 2). The Ut-Pl compartments showed increased expression of the *ECM1* gene in the prepartum luteolysis group compared with pre-implantation uterus and Ut-Pl sites at early post-implantation stage (p<0.001 and p<0.01, respectively (Figure 2).

The uterine expression of all three major collagens, COL1, COL3 and COL4, showed highest

- 285 mRNA abundance at mid-gestation compared with either pre-implantation (p<0.05 for COL1, p<0.01 for COL3 and p<0.001 for COL4) or post-implantation (p<0.05 for COL1 and COL4). Whereas COL1 and COL4 did not differ between pre-implantation and post-implantation, COL3 was higher in the early post-implantation group (p<0.05) (Figure 2). As for Ut-Pl compartments, early implantation was associated with suppression of COL1 and COL3 levels</p>
- 290 (p<0.001 for *COL1* and p<0.05 for *COL3*). Whereas they did not differ from postimplantation to mid-gestation, further suppression of both *COLs* mRNA expression was observed at prepartum luteolysis (p<0.001 and p<0.01 when compared with mid-gestation for *COL1* and *COL3*, respectively) (Figure 2). In contrast, the expression of *COL4* in the Ut-Pl comparison was unaffected at all examined stages of pregnancy (p>0.05).
- *FN1* showed similar distribution patterns at inter-Pl sites as *COL1*, *COL3* and *COL4*. Following a more or less stable expression at inter-Pl at early gestation (*i.e.*, during pre-

implantation and post-implantation), its uterine mRNA levels were highest at mid-gestation compared with previous gestational stages (p<0.05 and p<0.001 for pre-implantation and post-implantation, respectively) (Figure 3). At placentation sites (Ut-Pl compartments), the

300 prepartum luteolysis was associated with downregulation of *FN1* compared with midgestation (p<0.001) (Figure 3).

The uterine mRNA levels of *LAMA2* were not significantly changed (p>0.05) during preimplantation and post-implantation at the inter-Pl sites, but decreased thereafter, from postimplantation to mid-gestation (p<0.001) (<u>Figure 3</u>). *LAMA2* was strongly modulated at

305 implantation sites, decreasing at Ut-Pl following early implantation (p<0.001), but it was not further changed towards prepartum luteolysis (Figure 3).
 The uterine levels of *Cx26* were lowest at pre-implantation. They increased, however,

significantly (p<0.001) at inter-Pl sites, showing the highest mRNA abundance postimplantation (Figure 3). Although apparently decreased at mid-gestation, Cx26 levels were

- 310 still more highly represented at mid-gestation compared to pre-implantation (p<0.001). When compared with Ut-Pl, the expression of Cx26 was also significantly increased at implantation sites (P<0.001) and maintained its high mRNA levels until mid-gestation. The decrease towards prepartum luteolysis was significant compared with Ut-Pl during post-implantation and mid-gestation (p<0.001 and p<0.001, respectively) (Figure 3).
- 315 Uterine expression of the Cx43 gene did not differ significantly at inter-Pl sites (p>0.05) (Figure 3). It was, however, significantly modulated in Ut-Pl compartments. Thus, implantation was associated with suppression of Cx43 expression at placentation sites compared with the pre-implantation uterus (p<0.05). This expression continued, however, to increase at Ut-Pl gradually towards prepartum luteolysis, showing significant differences
- from post-implantation towards mid-gestation (p<0.05) and from mid-gestation to pre-partum luteolysis (p<0.05).

As for other genes, *i.e.*, *FN1*, *COL1*, -3, -4, the uterine mRNA encoding for *TIMP2* was highest at mid-gestation (p<0.01 and p<0.001 compared with pre-implantation and post-implantation, respectively (Figure 4). *TIMP4* mRNA was already strongly induced at inter-Pl

after implantation (p<0.001) and maintained its high levels through mid-gestation (Figure 4). At placentation sites, both *TIMP2* and *TIMP4* were strongly modulated and showed a gradual increase with progression of pregnancy, starting with early gestational stages until prepartum luteolysis when it was highest (p<0.01 and p<0.05 when compared with post-implantation) (Figure 4).

Localization of ECM proteins in the canine uterus and placenta

Localization and distribution patterns of COL1, COL3, COL4, FN1, ECM1, LAMA2, Cx26, Cx43, TIMP2 and TIMP4 were investigated at the protein level by applying IHC. All experimental groups were examined. A summary of the findings is presented descriptively in

335 <u>Table 3</u>.

COL1 and COL3 were detectable in all maternal and fetal stromal compartments and in myometrium (Figure 5, Supplemental File 1, respectively). While at all compared stages of pregnancy (including non-pregnant controls at days 10-12), COL1 staining appeared to be evenly distributed, the intensity of COL3 was generally weaker compared with COL1.

340 COL4 was clearly detectable throughout gestation and was targeted to the basal lamina of blood vessels (<u>Supplemental Files 2 D and G</u>) in all compartments. Additionally, COL4 was detected in the myometrium of all groups with staining intensity varying, however, individually (<u>Supplemental Files 2 B, F, I</u>).

ECM1 was weakly detectable in luminal epithelium in the non-pregnant group (in one sample

- 345 it was below the detection limit) (Figure 6 A and B). In early pregnancy, strong signals were detectable in luminal epithelium and in superficial and deep glands (Figures 6 C and D). During mid-gestation and at prepartum luteolysis (Figure 6 E and H), it was only weakly detectable in the glandular epithelium of the superficial glands. At prepartum luteolysis, ECM1 was abundantly present in the connective tissue layer separating superficial glands
- 350 from deep glands (<u>Figure 6 H</u>). Following implantation (shown at mid-gestation) (<u>Figure 6 F</u>) and at prepartum luteolysis (<u>Figure 6 I</u>), signals were detectable in glandular epithelium of deep glands. Placental localization of ECM1 was in cytotrophoblast (shown at mid-gestation and prepartum luteolysis) (<u>Figure 6 G and J</u>).

Uterine endometrial expression of Cx43 was localized predominantly in epithelial and myometrial compartments (Figure 7). The luminal epithelium (Figure 7 A and C) and the epithelial cells of superficial and deep glands (Figure 7 A-F, H, I) stained strongly for Cx43. Additionally, Cx43 was present in the *tunica intima* of maternal blood vessels in all samples. In placenta (Figure 7 G and J), Cx43 was diffusely present in cyto- and syncytiotrophoblast and decidual cells. The strongest placental signals were found in the *tunica intima* of blood

360 vessels (Figure 7 G and J).

Regarding TIMP2, it was only weakly expressed in the non-pregnant uterus (sometimes below detection limits) (Figure 8 A-B). In contrast, in the early pregnant uterus, at days 10-12 (prior to attachment), its expression was clearly detectable in the luminal epithelium, and superficial and deep glands (Figure 8 C, D). Myometrial signals were weak in both groups

- 365 (Figure 8 B, D). Following implantation (shown at mid-gestation), endometrial TIMP2 was detectable in epithelial compartments of superficial and deep uterine glands (Figure 8 E and E); stromal compartments of the connective tissue layer appeared only weakly stained (Figure 8 E). At prepartum luteolysis, uterine TIMP2 signals were weaker in the epithelium of superficial glands (Figure 8 H). However, strong staining was observed in the connective
- 370 tissue layer separating the superficial from deep glands (<u>Figure 8 H</u>). Myometrial signals were weak (<u>Figure 8 F, I</u>). Placental TIMP2 at mid-gestation was distributed ubiquitously and was found in cyto- and syncytiotrophoblast, in decidual cells and in vascular endothelial cells (*tunica intima*) (<u>Figure 8 G</u>). During prepartum luteolysis, placental expression was predominantly found in endothelial cells of maternal vessels (<u>Figure 8 J</u>). Sporadic nuclear
- 375 staining was detected in the nuclei of myocytes, which did not allow further interpretation. Compared with TIMP2, TIMP4 appeared to stain generally more weakly in all experimental groups and tissue compartments. Although weakly represented, uterine expression was localized in the luminal epithelium (Figure 9 A and C). Somewhat stronger staining was found in superficial and deep glands (Figure 9 A-D, F). Stronger TIMP4 signals were found
- during prepartum luteolysis in the connective tissue layer separating superficial from deep glands (Figure 9 H). It also appeared to be induced in deep uterine glands (Figure 9 I).
 Within Ut-Pl compartments, placental TIMP4 was found in cytotrophoblast (shown at midgestation and during prepartum luteolysis; Figure 9 G, J). For Supplemental File 3, consecutive cross-sections of Ut-Pl compartment were prepared during prepartum luteolysis
- 385 and were stained with either vimentin (VIM; <u>Supplemental File 3C</u>), pan-cytokeratin (pan-CYT; <u>Supplemental File 3D</u>), TIMP2 or TIMP4, allowing better differentiation between cellular tissue compartments stained by particular antibodies. In particular, localization patterns of both factors within canine placenta are clearly presented, with TIMP2 targeting to vascular endothelial cells and cytotrophoblast, presenting only weak signals in
- 390 syncytiotrophoblast and maternal decidual cells (<u>Supplemental File 3A</u>). Clear TIMP4 signals were localized in cytotrophoblast (<u>Supplemental File 3B</u>).
 In all samples, strong signals for aSMA were found in myometrium and the media of blood vessels (<u>Supplemental File 4</u>). Additionally, in the placenta aSMA was present in decidual cells, as shown at mid-gestation and prepartum luteolysis, (<u>Supplemental Files 4 G, J</u>).
- 395 FN1 was detectable throughout pregnancy. In general, stronger signals were found in epithelial than stromal compartments. The luminal epithelium of the non-pregnant group (Supplemental File 5A) was more strongly stained compared with the pre-attachment group (Supplemental File 5C). The myometrium stained weakly in all experimental groups and was

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below detection limits in some samples (<u>Supplemental Files 5 B, F, I</u>). Additionally, diffuse signals for FN1 were observed throughout all cellular placental compartments, *i.e.*, syncytio-and cytotrophoblast, decidual cells and blood vessels (Supplemental Files 5 G and J).

LAMA2 (<u>Supplemental File 6</u>) was clearly present in all stromal compartments and in the media of blood vessels throughout pregnancy. Additionally, it was detectable in glandular epithelial cells in all groups. In the placental part of Ut-Pl compartments, LAMA2 staining
was stronger in the stroma than in blood vessels (Supplemental Files 6 G and J).

- Regarding Cx26, this protein was present in epithelial compartments throughout pregnancy (<u>Supplemental File 7</u>). Generally, signals were weaker in stromal and myometrial compartments. Cx26 was also detectable in endometrial epithelial cells. Following implantation, as shown during mid-gestation and at prepartum luteolysis, this protein was
- 410 strongly locally induced in the epithelial cells of glandular chambers immediately above the connective tissue layer (Supplemental Files 7 E, H). Placental Cx26 was present diffusely in cyto- and syncytiotrophoblast and in decidual cells (indicated in Supplemental File 6 J).

DISCUSSION

- 415 The effects of modifications of uterine ECM on canine pregnancy have not been considered for many years. Because different tissues consist of dissimilar mixed combinations of cells which evolutionarily developed from different pedigrees, the configuration of ECM varies among tissues (Abedin & King 2010). These ECM components are involved in regulation of cell growth and differentiation, act as extracellular storage of hormones, and are involved in
- activation of signaling cascades (reviewed in: (<u>Hubmacher & Apte 2013</u>, <u>da Anunciacao et al.</u> 2017)). In the uterus, the major ECM components are collagens, proteoglycans, hyaluronan and glycoproteins (<u>Leoni et al. 1990</u>, <u>Oliveira et al. 2015</u>, <u>Franczyk et al. 2017</u>). It is worth noting that in mammals the uterine ECM composition changes dynamically depending on the reproductive status (*i.e.*, non-pregnant vs. pregnant) (<u>Cabrol et al. 1985</u>, <u>Leoni et al. 1990</u>,
- 425 <u>Hjelm et al. 2002</u>, <u>Boos et al. 2003</u>), and in some species, such as humans and cattle, the composition of the uterine ECM is cycle-dependent (<u>Boos 2000</u>, <u>Curry & Osteen 2001</u>). In fact, most details about uterine ECM modifications and functions during pregnancy are known only for humans and rodents. In these species, in early pregnancy the uterine ECM influences trophoblast invasion (Johnson et al. 2003) and remodeling of decidua (<u>Damsky et</u>
- 430 <u>al. 1993</u>, <u>Lala & Nandi 2016</u>, <u>Smith et al. 2016</u>), while the ECM is modulated by matrix metalloproteinases (MMPs) of fetal (trophoblast) origin. As for the canine non-pregnant and pregnant uterus and placenta, the time- and organ-specific expression of MMP2 and -9 have

been investigated (<u>Beceriklisoy et al. 2007</u>, <u>Fellows et al. 2012</u>, <u>Diessler et al. 2017</u>). Their expression patterns indicate pregnancy-associated modifications of ECM.

- The present study explored the spatio-temporal expression of ECM components selected from our preceding microarray analysis (Graubner *et al.* 2017a), and known for their involvement in modulating uterine and placental functions in other animals, in the canine uterus and placenta throughout pregnancy. Quantitative assessment was done by qPCR, and cellular localization was investigated qualitatively by IHC. The antibodies applied herein proved unsuitable for western blot analysis. It appears plausible that linearization of proteins by
- reducing agents during western blot preparation affects the structure of target epitopes, thereby preventing antibody binding.

Structural and adhesive proteins of the ECM

- Expression of the following proteins was assessed: COL1, COL3, COL4, ECM1, FN1, LAMA2 and aSMA. Free-floating embryos were previously reported to increase the uterine expression of *LAMA2* (Graubner et al. 2017a). Here, the uterine adjustments during early decidualization were additionally characterized by increased expression of *ECM1*, whereas *FN1* was suppressed. None of the major COLs was affected by the presence of embryos.
- 450 Showing distinctly different localization patterns and both being affected by presence of embryos, ECM1 and LAMA2 appear to be involved in the establishment of canine pregnancy. As a component of the lamina basalis, increased expression of LAMA is a marker of ongoing stromal decidualization in humans (Church *et al.* 1997). Similarly in the dog, even if devoid of apparent morphological differentiation of the uterus, pre-implantation embryos stimulated
- 455 the expression of LAMA2, predominantly in the uterine stroma. The increased ECM1 expression was abundantly localized in endometrial epithelium. In the human endometrium its expression during implantation is further stimulated by some of the chemokines critically involved in leukocyte migration and trophoblast invasion, *e.g.*, CX3CL1 and CCL14 (Hannan & Salamonsen 2008). Although not yet investigated, similar functional relationships cannot
- 460 be ruled out for the canine species. Further effects of ECM1 in the canine uterus might be associated with its proliferative activities. Thus, by interacting with the epidermal growth factor (EGF) related pathways, ECM1 promotes cellular differentiation and proliferation (Lee et al. 2014). Within the canine uterus, the localization patterns of EGF receptor (EGFR) were described previously (Sagsoz et al. 2014) and reflect the localization of ECM1 presented
- 465 herein. A possible functional relationship between these two entities is certainly worth attention and should be addressed in the future.

500

Regarding the embryo-induced suppression of the cell adhesion molecule FN1, similar to the dog, its decrease during early pregnancy also occurs in rodents (Zollinger & Smith 2017), a phenomenon that has been linked to the early decidualization process around the time of

- 470 implantation (Grinnell et al. 1982). Moreover, reduction of FN1 expression seems to be important to prevent excessive trophoblast invasion into maternal tissues (Kaloglu & Onarlioglu 2010). Despite the less invasive type of canine placentation compared to rodents, the *FN1* decrease observed before trophoblast attachment in dogs could also possibly be linked to similar protective functions in order to control trophoblast invasion. A key
- 475 mechanism in this context seems to be the cooperation of FN1 and integrin molecules in modulating trophoblast proteolytic activities [through/via] the ECM (<u>Sutherland et al. 1993</u>, <u>Fazleabas et al. 1997</u>). Interestingly, however, the canine uterine expression of several integrins (ITG) (*e.g.*, ITGA2B, ITGB2 and ITGB3) increases in early pregnancy (<u>Bukowska et al. 2011</u>), while FN1 expression is diminished by the presence of embryos.
- 480 It needs to be emphasized that, in agreement with our previous findings (Graubner *et al.* 2017a), and in contrast to other species including humans and rodents, the above described functional changes in the canine uterus driven by the presence of embryos, take place without any morphologically visible alterations of uterine structures. The deep structural remodeling processes of the canine uterus start following implantation and placentation and are associated
- 485 with increased expression of COLs (*COL1*, -3, and -4), *ECM1* and *FN1* in the uterine wall (*i.e.*, inter-Pl). The increased expression of structural and basement membrane components, widely localized in stroma and, depending on the type of COL (*e.g.*, COL1 and -3), in myometrium, seems to relate to the uterine growth as a natural phenomenon resulting from the development of conceptuses.
- 490 Conversely, at implantation sites, trophoblast invasion and placentation decreased the expression of two major COLs (COL1 and -3) and LAMA2. This is apparently associated with the remodelling processes and placental development, as all COLs, and in particular the most robust COL1, were clearly detectable in utero-placental stromal compartments.

Similar effects were observed in other animals displaying invasive types of placentation, *e.g.*,

495 in rodents, in which implantation was associated with decreased *COL1* expression at invasion sites, implying the need for *COL1* suppression during remodeling events associated with embryo-induced decidualization and subsequent placentation in these animals (<u>Clark *et al.*</u> <u>1993</u>).

The decrease in *LAMA2* found in our study additionally indicates loss of stability of the basal lamina, and associated with it migration and differentiation of cells (<u>Kleinman *et al.* 1985</u>,

Engvall *et al.* 1990, Aumailley 2013). This differentiates the dog from mammals with more invasive forms of placentation such as rodents and primates. Accordingly, around the time of implantation in the murine uterus, laminin is synthesized by decidual cells (Wewer *et al.* 1985) where it influences trophoblast migration (Zhang *et al.* 2000, Korgun *et al.* 2007). In

- 505 humans also decidual cells produce laminin (Kisalus et al. 1987, Aplin et al. 1988, Haouzi et al. 2011). At later stages of human gestation, several isoforms of laminin and fibronectin are found in the basal lamina of capillaries and in the stroma and trophoblast cells of the villi (Korhonen & Virtanen 2001). As for the dog, LAMA2 seemed to be more strongly represented in the stromal than epithelial compartments, in particular following implantation.
- 510 Besides laminin, the decidualization process in humans is marked by increased expression of aSMA (Geisert 2015). This protein seems to be strongly required during pregnancy since there is a relationship between abnormal actin polymerization and pregnancy failure in humans (Montazeri *et al.* 2015). In dogs, aSMA is only weakly present in uterine stromal cells of non-pregnant animals and at the pre-implantation stage. However, following
- 515 placentation aSMA is strongly expressed in maternal decidual cells and can be used as a cellular marker of decidual cells in the canine species. As a matter of fact, aSMA was also strongly expressed in canine decidualized uterine primary stromal cells *in vitro* (Kautz *et al.* 2015).
- Following attachment and invasion of the trophoblast in dogs the girdle placenta is
 developing. Histologically, at the placentation site a connective tissue layer can be found that separates enlarged superficial glands (so called glandular chambers) from deeper localized parts of the uterus. It functions as a barrier, protecting deep uterine glands and myometrium from proteolytic activity of the trophoblast. When this barrier is breached, an exaggerated invasion of trophoblast may result in SIPS (Al-Bassam *et al.* 1981). Clearly detectable staining of structural collagens (COL1, and -3) and basal lamina components such as LAMA2 was identified throughout pregnancy in the stroma of this connective tissue layer. Their respective mRNA levels were not modulated from post-implantation towards mid-gestation, but the major collagens, COL1 and -3, were strongly suppressed at the time of prepartum luteolysis. This may be seen as an indicator of preparation for parturition and release of the
- 530 fetal membranes (placentolysis). A similar role of physiological degradation of collagens during preparation for the release of fetal membranes has been discussed for cattle (<u>Attupuram et al. 2016</u>). In fact, a retained placenta can be successfully treated by administration of collagenases into the umbilical artery (<u>Eiler & Hopkins 1993</u>, <u>Attupuram et</u> <u>al. 2016</u>).

535 With regards to *FN1*, which was generally more strongly represented in epithelial compartments, its uterine and placental expression followed the expression patterns exhibited by COLs. This leads to the assumption that the main role of *FN1* in the canine uterus is a bridging function between collagens and other ECM components.

The utero-placental *ECM1* expression increased with the progression of pregnancy and was highest at prepartum luteolysis. Within the placenta, ECM1 stained predominantly in cytotrophoblast, which appeared to be the major source of ECM1 in the Ut-Pl compartment. This resembles results observed in humans where the importance of ECM1 during maintenance of pregnancy and fetal development is underlined by its highest expression in the placenta and fetal heart tissue but not in other organs (*e.g.*, brain, lung, liver, kidney, pancreas

- 545 or skeletal muscles) (<u>Smits *et al.* 1997</u>). In addition to the aforementioned function of ECM1 in promoting proliferation and differentiation of cells during the onset of pregnancy, at prepartum luteolysis the co-localization of ECM1 with TIMP2- and -4 in the tissue layer separating superficial from deep uterine glands strongly attracted our attention. This spatiotemporal detection is specific to prepartum luteolysis, but not at earlier stages of pregnancy. A
- 550 possible implication for its function arises from *in vitro* studies showing ECM1-mediated reduction of MMP9 proteolytic activity in a human model (Fujimoto *et al.* 2006). Cumulatively, a protective role of ECM1, possibly interacting with a metalloproteinases activity-balancing system, should be considered.

555 Connexins (Cx26 and -43)

Connexins (Cx) build gap junction proteins, through which two cells can exchange small molecules, electrical charges and second messengers (Bruzzone *et al.* 1996, Kumar & Gilula 1996, Evans & Martin 2002). The uterine expression of Cx26 and -43 has been described during the menstrual cycle in humans as well as during pregnancy in humans, rats and sheep

- 560 (Winterhager et al. 1993, Grummer et al. 1994, Jahn et al. 1995, Johnson et al. 2017). Both hormonal- and embryo implantation-mediated effects were observed (Grummer et al. 2004). Successful implantation of embryos in rodents requires a complete suppression of both connexins prior to implantation (Grummer et al. 1994, Grummer et al. 2004). At the time of implantation connexins are re-induced (Grummer et al. 1996). The importance of Cx during
- 565 establishment of pregnancy has been further strengthened by their abnormal expression patterns reported in pathologies such as recurrent pregnancy loss (<u>Laird 2006</u>, <u>Nair *et al.*</u> 2011).

No such information was so far available regarding the canine uterus and placenta, which prompted us to investigate the expression and distribution patterns of Cx26 and -43 570 throughout pregnancy. The uterine expression of both Cx remained unaffected by the presence of free-floating embryos. The respective proteins were detectable, however, with Cx43 appearing more abundantly expressed than Cx26. Whereas being clearly detectable, Cx43 was more or less constantly present in the uterine wall; its increasing expression in the Ut-Pl compartment followed placental development and reached its highest levels during 575 prepartum luteolysis. It is noteworthy that, although rather ubiquitously expressed, within the placenta Cx43 mostly targeted to the maternal endothelium. This expression pattern differed strongly from that observed for Cx26, which appeared to be in agreement with observations made in other species exhibiting invasive types of placentation. Thus, the uterine and placental expression of Cx26 was strongly induced following implantation. Its expression was 580 predominantly localized in the endometrial tissues, both at the placentation sites and at inter-Pl. The prepartum luteolysis was associated with a strong decrease of the respective mRNA levels.

Regarding the possible functions of connexins within the placenta, in humans Cx43 appears indispensable for proper decidualization and uterine angiogenesis (Laws *et al.* 2008).

- Additionally, it has been shown *in vitro* that Cx43 is involved in regulating differentiation of cytotrophoblast to syncytiotrophoblast and plays roles in feto-maternal exchange (<u>Cronier et al. 2002</u>). Based on the localization pattern of Cx43 described above, in the dog its involvement in uterine and placental angiogenesis also seems likely, since the strongest signals were noticed in the intima and media of placental blood vessels. The involvement of
- 590 Cx43 during canine decidualization cannot, however, be ruled out and merits further investigations.

Spatio-temporal expression of TIMP2 and TIMP4

When discussing the development and function of the canine endotheliochorial placenta, the

- 595 mechanisms controlling the invasive behavior of trophoblast need to be considered. In general, the invasive properties of trophoblast are at least in part provided by MMPs (reviewed in (Goldman-Wohl & Yagel 2002, Cohen *et al.* 2006)). At parturition, MMPs are additionally associated with placental detachment, as shown in humans and cattle (Eiler & Hopkins 1992, Strauss 2013, Menon *et al.* 2016). In dogs, MMPs are present in trophoblast
- 600 cells (<u>Beceriklisoy *et al.* 2007</u>, <u>Fellows *et al.* 2012</u>, <u>Diessler *et al.* 2017</u>). Biologically active inhibitors, *i.e.*, TIMPs, by interacting with MMPs promote cell growth, inhibit angiogenesis

and reveal both anti- and pro-inflammatory effects (reviewed in (Stetler-Stevenson 2008, Brew & Nagase 2010)). Although TIMP2 and TIMP4 are 50% identical in sequence, they still bear differences in specificity to MMPs (Brew & Nagase 2010). *In vitro* studies with human endometrial cells have shown that TIMPs are positively associated with the decidualization process and cell migration (Graham *et al.* 2017). Interestingly, *TIMP2* expression can be induced by P4 (Imada *et al.* 1994, Jo *et al.* 2015). We have previously shown the ability of canine embryos to increase uterine *TIMP2* expression prior to implantation (Graubner *et al.* 2017a). During canine labor also, the placental presence of *TIMP2* has been confirmed at the

- 610 mRNA level (Fellows et al. 2012). It is assumed that interference with TIMPs functions can lead to placental retention and SIPS in dogs, or to spontaneous early pregnancy failure and *Placenta accreta* in humans (Al-Bassam et al. 1981, Ke et al. 2006, Nissi et al. 2013). In addition to the previously reported increase in mRNA expression of *TIMP2* (Graubner et al. 2017a), here also, elevated TIMP4 expression was observed in the pre-implantation canine
- 615 uterus. The stimulatory effect on gene expression was supported by IHC results in which freefloating embryos appeared to increase signals for TIMP2 in both superficial and deep uterine glands. Following implantation, a gradual increase was noted for both TIMPs (*i.e.*, *TIMP2* and -4) in uterine and placental compartments.
- An important finding from our study is the strong presence of TIMP2 and -4 in the endometrial connective tissue layer separating the superficial from deep glands. In a healthy pregnancy, trophoblast invasion is stopped at this bordering protective structure. Additionally, uterine and placental TIMPs were localized around blood vessels. Thus, it is plausible to assume that both proteins are involved in the protection of maternal tissue from trophoblast invasion. Within the placenta, TIMP2 appeared more abundantly represented than TIMP4.
- 625 Both factors were, however, localized in trophoblast cells, which appears to reflect an auto-/paracrine feedback loop acting within these cells, possibly controlling own proteolytic function of trophoblast and protecting the maternal tissues from excessive invasion. Although it is not known for the dog if TIMP2 and TIMP4 interact with active MMP2 and MMP9, our findings (*i.e.*, the gradual increase of *TIMP2*, *TIMP4* and *ECM1*) indicate that MMP-
- 630 mediated-trophoblast invasion might be balanced by these factors. In this regard, future studies should be considered, investigating for example the expression of MMPs and their regulators within excessive trophoblast invasion leading to conditions such as SIPS.

Conclusions

- 635 In this study we investigated the modulation of uterine and placental ECM in defined stages of canine pregnancy. Based on the presented results, it seems that the primary goal of the early embryo maternal communication is the regulation of trophoblast invasion, and the proliferative and adhesive functions of the uterus. Following trophoblast attachment, the ECM is strongly modulated reflecting dynamic feto-maternal interactions during establishment of the canine endotheliochorial placenta.
 - The role of the endometrial connective tissue layer as an active, and not only physical but also a biochemical, barrier protecting maternal tissues from unrestrained trophoblast invasion, is strongly implied. This is supported by the abundant local expression of ECM components potentially actively involved in modulating the invasiveness of fetal cells.
- 645 Regulatory mechanisms involved in modulation of ECM components during establishment and maintenance of gestation modes characteristic of invasive placentation types, appear to be similar to what has been described for other mammalian species exhibiting similar, or even more intense invasion by trophoblast (haemochorial placentation type). In particular, TIMPs and ECM1 appear to be plausible candidates involved in regulating the establishment and
- 650 termination of pregnancy in the dog. With this, the dog could provide an interesting model for investigating placental functions in other species, *e.g.*, in humans in which *Placenta accreta* appears to share several similarities with canine SIPS.

Abbreviations:

655 aSMA: alpha-smooth muscle actin COL: collagen CL: corpus luteum Cx: connexin ECM: extracellular matrix 660 ECM1: extracellular matrix protein 1 FN1: fibronectin1 GAPDH: glyceraldehyde 3-phosphate dehydrogenase HIER: heat initiated epitope retrieval IHC: immunohistochemistry 665 inter-Pl: inter-placenta LAMA2: laminin alpha 2 min: minutes MMP: matrix metalloproteinase

OHE: ovariohysterectomy

670 P4: progesterone pan-CYT: pan-cytokeratin
PGR: nuclear progesterone receptor
PIER: protease initiated epitope retrieval
PPIA: *Cyclophilin A*675 qPCR: quantitative real-time polymerase chain reaction
SD: standard deviation
SIPS: subinvolution of placental sites
TIMP: tissue inhibitor of matrix metalloproteinase
Ut-Pl compartment: utero-placental compartment
680 VIM: vimentin

Declaration of interest:

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported. All authors read and approved the final version of the manuscript.

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Author contribution statement:

FRG: involved in developing the concept of the study, experimental design, generating data, analysis and interpretation of data and writing of the manuscript. AB: knowledge transfer, critical discussion of data, editing of the manuscript. SA and IK: knowledge transfer, critical

695 discussion of data, collection of tissue material. MPK: designed and supervised the project, involved in interpretation of the data, drafting and revising the manuscript. All authors read and approved the final manuscript.

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Figures, Tables and Supplemental Files: (submitted as separated file)

	References:
710	Abedin M & King N 2010 Diverse evolutionary paths to cell adhesion. <i>Trends Cell Biol</i> 20 734-742.
	Al-Bassam MA, Thomson RG & O'Donnell L 1981 Involution abnormalities in the
	postpartum uterus of the bitch. <i>Vet Pathol</i> 18 208-218. Aplin JD, Charlton AK & Ayad S 1988 An immunohistochemical study of human
	endometrial extracellular matrix during the menstrual cycle and first trimester of
715	pregnancy. Cell Tissue Res 253 231-240.
, 10	Attupuram NM, Kumaresan A, Narayanan K & Kumar H 2016 Cellular and molecular mechanisms involved in placental separation in the bovine: A review. <i>Mol Reprod Dev</i> 83 287-297.
	Aumailley M 2013 The laminin family. <i>Cell Adh Migr</i> 7 48-55.
720	Bazer FW 2015 History of Maternal Recognition of Pregnancy. <i>Adv Anat Embryol Cell</i> <i>Biol</i> 216 5-25.
	Bazer FW, Wu G, Spencer TE, Johnson GA, Burghardt RC & Bayless K 2010 Novel
	pathways for implantation and establishment and maintenance of pregnancy in mammals. <i>Mol Hum Reprod</i> 16 135-152.
725	Beceriklisoy HB, Walter I, Schafer-Somi S, Miller I, Kanca H, Izgur H & Aslan S 2007 Matrix metalloproteinase (MMP)-2 and MMP-9 activity in the canine uterus
	before and during placentation. <i>Reprod Domest Anim</i> 42 654-659. Boos A 2000 Immunohistochemical assessment of collagen types I, III, IV and VI in
	biopsy samples of the bovine uterine wall collected during the oestrous cycle.
730	Cells Tissues Organs 167 225-238.
	Boos A, Stelljes A & Kohtes J 2003 Collagen types I, III and IV in the placentome and
	interplacentomal maternal and fetal tissues in normal cows and in cattle with
	retention of fetal membranes. <i>Cells Tissues Organs</i> 174 170-183.
705	Brew K & Nagase H 2010 The tissue inhibitors of metalloproteinases (TIMPs): an
735	ancient family with structural and functional diversity. <i>Biochim Biophys Acta</i> 1803 55-71.
	Brown JJ & Papaioannou VE 1992 Distribution of hyaluronan in the mouse
	endometrium during the periimplantation period of pregnancy. <i>Differentiation</i> 52 61-68.
740	Bruzzone R, White TW & Paul DL 1996 Connections with connexins: the molecular
, 10	basis of direct intercellular signaling. <i>Eur J Biochem</i> 238 1-27.
	Bukowska D, Kempisty B, Jackowska M, Wozna M, Antosik P, Piotrowska H &
	Jaskowski JM 2011 Analysis of integrins and vascular endothelial growth factor
	isoforms mRNA expression in the canine uterus during perimplantation period.
745	Pol J Vet Sci 14 253-258.
	Cabrol D, Dallot E, Cedard L & Sureau C 1985 Pregnancy-related changes in the distribution of glycosaminoglycans in the cervix and corpus of the human uterus.
	<i>Eur J Obstet Gynecol Reprod Biol</i> 20 289-295.
	Church HJ, Richards AJ & Aplin JD 1997 Laminins in decidua, placenta and
750	choriocarcinoma cells. <i>Placenta</i> 18 143-162.
	Clark DE, Hurst PR, McLennan IS & Myers DB 1993 Immunolocalization of collagen
	type I and laminin in the uterus on days 5 to 8 of embryo implantation in the rat.
	Anat Rec 237 8-20.
	Cohen M, Meisser A & Bischof P 2006 Metalloproteinases and human placental
755	invasiveness. <i>Placenta</i> 27 783-793.
	Concannon PW 1993 Biology of gonadotrophin secretion in adult and prepubertal female dogs. <i>J Reprod Fertil Suppl</i> 47 3-27.

	Concannon PW, McCann JP & Temple M 1989 Biology and endocrinology of ovulation,
	pregnancy and parturition in the dog. J Reprod Fertil Suppl 39 3-25.
760	Cronier L, Defamie N, Dupays L, Theveniau-Ruissy M, Goffin F, Pointis G &
	Malassine A 2002 Connexin expression and gap junctional intercellular
	communication in human first trimester trophoblast. <i>Mol Hum Reprod</i> 8 1005-
	1013.
	Curry TE, Jr. & Osteen KG 2001 Cyclic changes in the matrix metalloproteinase system
765	in the ovary and uterus. <i>Biol Reprod</i> 64 1285-1296.
	da Anunciacao A, Mess AM, Orechio D, Aguiar BA, Favaron PO & Miglino MA 2017
	Extracellular matrix in epitheliochorial, endotheliochorial and haemochorial
	placentation and its potential application for regenerative medicine. <i>Reprod</i>
	Domest Anim 52 3-15.
770	Damsky C, Sutherland A & Fisher S 1993 Extracellular matrix 5: adhesive interactions
	in early mammalian embryogenesis, implantation, and placentation. <i>Faseb j</i> 7
	1320-1329.
	Dempsey EW & Wislocki GB 1956 Electron microscopic observations on the placenta
	of the cat. J Biophys Biochem Cytol 2 743-754.
775	Diessler M, Ventureira M, Hernandez R, Sobarzo C, Casas L, Barbeito C & Cebral E
	2017 Differential expression and activity of matrix metalloproteinases 2 and 9 in
	canine early placenta. <i>Reprod Domest Anim</i> 52 35-43.
	Eiler H & Hopkins FM 1992 Bovine retained placenta: effects of collagenase and
	hyaluronidase on detachment of placenta. <i>Biol Reprod</i> 46 580-585.
780	Eiler H & Hopkins FM 1993 Successful treatment of retained placenta with umbilical
	cord injections of collagenase in cows. <i>J Am Vet Med Assoc</i> 203 436-443.
	Engvall E, Earwicker D, Haaparanta T, Ruoslahti E & Sanes JR 1990 Distribution and
	isolation of four laminin variants; tissue restricted distribution of heterotrimers
705	assembled from five different subunits. <i>Cell Regul</i> 1 731-740.
785	Evans WH & Martin PE 2002 Gap junctions: structure and function (Review). <i>Mol</i> <i>Membr Biol</i> 19 121-136.
	Fazleabas AT, Bell SC, Fleming S, Sun J & Lessey BA 1997 Distribution of integrins and the extracellular matrix proteins in the baboon endometrium during the
	menstrual cycle and early pregnancy. <i>Biol Reprod</i> 56 348-356.
790	Fellows EJ, Hazzard TM & Kutzler MA 2012 Gene expression in pre-term, pre-labour
/)0	and parturient canine placenta. <i>Reprod Domest Anim</i> 47 Suppl 6 182-185.
	Franczyk M, Lopucki M, Stachowicz N, Morawska D & Kankofer M 2017
	Extracellular matrix proteins in healthy and retained placentas, comparing
	hemochorial and synepitheliochorial placentas. <i>Placenta</i> 50 19-24.
795	Fujimoto N, Terlizzi J, Aho S, Brittingham R, Fertala A, Oyama N, McGrath JA & Uitto
	J 2006 Extracellular matrix protein 1 inhibits the activity of matrix
	metalloproteinase 9 through high-affinity protein/protein interactions. <i>Exp</i>
	Dermatol 15 300-307.
	Geisert RD 2015 Regulation of Implantation and Establishment of Pregnancy in
800	Mammals Tribute to 45 Year Anniversary of Roger V. Short's "Maternal
	Recognition of Pregnancy" Introduction. Regulation of Implantation and
	Establishment of Pregnancy in Mammals: Tribute to 45 Year Anniversary of Roger
	V. Short's Maternal Recognition of Pregnancy 216 1-4.
	Goldman-Wohl D & Yagel S 2002 Regulation of trophoblast invasion: from normal
805	implantation to pre-eclampsia. Mol Cell Endocrinol 187 233-238.
	Graham A, Holbert J & Nothnick WB 2017 miR-181b-5p Modulates Cell Migratory
	Proteins, Tissue Inhibitor of Metalloproteinase 3, and Annexin A2 During In Vitro

	Decidualization in a Human Endometrial Stromal Cell Line. <i>Reprod Sci</i> 24 1264-
810	1274. Cram A. Buchler II. Boos A. Hoffmann P. & Kowalowski MD 2012 Biosynthesis and
010	Gram A, Buchler U, Boos A, Hoffmann B & Kowalewski MP 2013 Biosynthesis and degradation of canine placental prostaglandins: prepartum changes in expression and function of prostaglandin F2alpha-synthase (PGFS, AKR1C3) and 15- hydroxyprostaglandin dehydrogenase (HPGD). <i>Biol Reprod</i> 89 2.
	Graubner FR, Gram A, Kautz E, Bauersachs S, Aslan S, Agaoglu AR, Boos A &
815	Kowalewski MP 2017a Uterine responses to early pre-attachment embryos in the domestic dog and comparisons with other domestic animal species. <i>Biol</i>
	Reprod. Crawba an ED, Deichlan IM, Dahman NA, Davan, Camping D, Dava A, & Kawalawalii
	Graubner FR, Reichler IM, Rahman NA, Payan-Carreira R, Boos A & Kowalewski MP 2017b Decidualization of the canine uterus: From early until late gestational
820	in vivo morphological observations, and functional characterization of
820	immortalized canine uterine stromal cell lines. <i>Reprod Domest Anim</i> 52 Suppl 2 137-147.
	Grinnell F, Head JR & Hoffpauir J 1982 Fibronectin and cell shape in vivo: studies on
	the endometrium during pregnancy. J Cell Biol 94 597-606.
825	Grummer R, Chwalisz K, Mulholland J, Traub O & Winterhager E 1994 Regulation of
	connexin26 and connexin43 expression in rat endometrium by ovarian steroid hormones. <i>Biol Reprod</i> 51 1109-1116.
	Grummer R, Hewitt SW, Traub O, Korach KS & Winterhager E 2004 Different
	regulatory pathways of endometrial connexin expression: preimplantation
830	hormonal-mediated pathway versus embryo implantation-initiated pathway. <i>Biol Reprod</i> 71 273-281.
	Grummer R, Reuss B & Winterhager E 1996 Expression pattern of different gap
	junction connexins is related to embryo implantation. <i>Int J Dev Biol</i> 40 361-367.
025	Hannan NJ & Salamonsen LA 2008 CX3CL1 and CCL14 regulate extracellular matrix
835	and adhesion molecules in the trophoblast: potential roles in human embryo implantation. <i>Biol Reprod</i> 79 58-65.
	Haouzi D, Dechaud H, Assou S, Monzo C, de Vos J & Hamamah S 2011 Transcriptome
	analysis reveals dialogues between human trophectoderm and endometrial cells
	during the implantation period. <i>Hum Reprod</i> 26 1440-1449.
840	Hjelm AM, Barchan K, Malmstrom A & Ekman-Ordeberg GE 2002 Changes of the
010	uterine proteoglycan distribution at term pregnancy and during labour. <i>Eur J</i>
	Obstet Gynecol Reprod Biol 100 146-151.
	Hoffmann B, Hoveler R, Nohr B & Hasan SH 1994 Investigations on hormonal changes
	around parturition in the dog and the occurrence of pregnancy-specific non
845	conjugated oestrogens. <i>Exp Clin Endocrinol</i> 102 185-189.
	Hubmacher D & Apte SS 2013 The biology of the extracellular matrix: novel insights.
	Curr Opin Rheumatol 25 65-70.
	Imada K, Ito A, Itoh Y, Nagase H & Mori Y 1994 Progesterone increases the production
	of tissue inhibitor of metalloproteinases-2 in rabbit uterine cervical fibroblasts.
850	<i>FEBS Lett</i> 341 109-112.
	Jahn E, Classen-Linke I, Kusche M, Beier HM, Traub O, Grummer R & Winterhager E
	1995 Expression of gap junction connexins in the human endometrium
	throughout the menstrual cycle. <i>Hum Reprod</i> 10 2666-2670.
055	Jo YS, Lee GS, Nam SY & Kim SJ 2015 Progesterone Inhibits Leptin-Induced
855	Invasiveness of BeWo Cells. Int J Med Sci 12 773-779.
	Johnson GA, Burghardt RC, Bazer FW & Spencer TE 2003 Osteopontin: roles in implantation and placentation. <i>Biol Panrod</i> 60 1459, 1471
	implantation and placentation. <i>Biol Reprod</i> 69 1458-1471.

	Johnson ML, Redmer DA, Reynolds LP & Grazul-Bilska AT 2017 Gap junctional
	connexin messenger RNA expression in the ovine uterus and placenta: effects of
860	estradiol-17beta-treatment, early pregnancy stages, and embryo origin. Domest
	Anim Endocrinol 58 104-112.
	Kaloglu C & Onarlioglu B 2010 Extracellular matrix remodelling in rat endometrium
	during early pregnancy: The role of fibronectin and laminin. <i>Tissue and Cell</i> 42
0.65	301-306.
865	Kautz E, de Carvalho Papa P, Reichler IM, Gram A, Boos A & Kowalewski MP 2015 In vitro decidualisation of canine uterine stromal cells. <i>Reprod Biol Endocrinol</i> 13
	85.
	Kautz E, Gram A, Aslan S, Ay SS, Selcuk M, Kanca H, Koldas E, Akal E, Karakas K,
	Findik M, Boos A & Kowalewski MP 2014 Expression of genes involved in the
870	embryo-maternal interaction in the early-pregnant canine uterus. <i>Reproduction</i>
	147 703-717.
	Ke Y, Lu JH, Yang BL, Guo HQ, Ma QY, Zhu H, Shu HM & Li DJ 2006 [Involvement of
	matrix metalloproteinase-2, -9, and tissue inhibitors of metalloproteinase-1, 2 in
~ 	occurrence of the accrete placenta]. <i>Zhonghua Fu Chan Ke Za Zhi</i> 41 311-314.
875	Kisalus LL, Herr JC & Little CD 1987 Immunolocalization of extracellular matrix
	proteins and collagen synthesis in first-trimester human decidua. <i>Anat Rec</i> 218 402-415.
	402-415. Kleinman HK, Cannon FB, Laurie GW, Hassell JR, Aumailley M, Terranova VP,
	Martin GR & DuBois-Dalcq M 1985 Biological activities of laminin. J Cell
880	Biochem 27 317-325.
	Korgun ET, Cayli S, Asar M & Demir R 2007 Distribution of laminin, vimentin and
	desmin in the rat uterus during initial stages of implantation. J Mol Histol 38 253-
	260.
005	Korhonen M & Virtanen I 2001 Immunohistochemical localization of laminin and
885	fibronectin isoforms in human placental villi. <i>J Histochem Cytochem</i> 49 313-322. Kowalewski MP 2014 Luteal regression vs. prepartum luteolysis: regulatory
	mechanisms governing canine corpus luteum function. <i>Reprod Biol</i> 14 89-102.
	Kowalewski MP, Beceriklisoy HB, Aslan S, Agaoglu AR & Hoffmann B 2009 Time
	related changes in luteal prostaglandin synthesis and steroidogenic capacity
890	during pregnancy, normal and antiprogestin induced luteolysis in the bitch. Anim
	<i>Reprod Sci</i> 116 129-138.
	Kowalewski MP, Beceriklisoy HB, Pfarrer C, Aslan S, Kindahl H, Kucukaslan I &
	Hoffmann B 2010 Canine placenta: a source of prepartal prostaglandins during
00 г	normal and antiprogestin-induced parturition. <i>Reproduction</i> 139 655-664.
895	Kowalewski MP, Mason JI, Howie AF, Morley SD, Schuler G & Hoffmann B 2006a Characterization of the canine 3beta-hydroxysteroid dehydrogenase and its
	expression in the corpus luteum during diestrus. J Steroid Biochem Mol Biol 101
	254-262.
	Kowalewski MP, Meyer A, Hoffmann B, Aslan S & Boos A 2011 Expression and
900	functional implications of peroxisome proliferator-activated receptor gamma
	(PPARgamma) in canine reproductive tissues during normal pregnancy and
	parturition and at antiprogestin induced abortion. <i>Theriogenology</i> 75 877-886.
	Kowalewski MP, Schuler G, Taubert A, Engel E & Hoffmann B 2006b Expression of
905	cyclooxygenase 1 and 2 in the canine corpus luteum during diestrus.
202	<i>Theriogenology</i> 66 1423-1430. Kumar NM & Gilula NB 1996 The gap junction communication channel. <i>Cell</i> 84 381-
	388.

910	 Laird DW 2006 Life cycle of connexins in health and disease. <i>Biochem J</i> 394 527-543. Lala PK & Nandi P 2016 Mechanisms of trophoblast migration, endometrial angiogenesis in preeclampsia: The role of decorin. <i>Cell Adh Migr</i> 10 111-125. Laws MJ, Taylor RN, Sidell N, DeMayo FJ, Lydon JP, Gutstein DE, Bagchi MK & Bagchi IC 2008 Gap junction communication between uterine stromal cells plays a critical role in pregnancy-associated neovascularization and embryo survival. <i>Development</i> 135 2659-2668.
915	Lee K-m, Nam K, Oh S, Lim J, Kim Y-P, Lee JW, Yu J-H, Ahn S-H, Kim S-B, Noh D-Y, Lee T & Shin I 2014 Extracellular matrix protein 1 regulates cell proliferation and trastuzumab resistance through activation of epidermal growth factor signaling. <i>Breast Cancer Research : BCR</i> 16 479.
920	 Leoni P, Carli F & Halliday D 1990 Collagen in the Pregnant and Non Pregnant Uterus. <i>Clinical Science</i> 79 10P-11P. Menon R, Bonney EA, Condon J, Mesiano S & Taylor RN 2016 Novel concepts on pregnancy clocks and alarms: redundancy and synergy in human parturition. <i>Hum Reprod Update</i> 22 535-560.
925	Montazeri M, Sanchez-Lopez JA, Caballero I, Maslehat Lay N, Elliott S, Lopez-Martin S, Yanez-Mo M & Fazeli A 2015 Activation of Toll-like receptor 3 reduces actin polymerization and adhesion molecule expression in endometrial cells, a potential mechanism for viral-induced implantation failure. <i>Hum Reprod</i> 30 893- 905.
930	 Nair RR, Jain M & Singh K 2011 Reduced expression of gap junction gene connexin 43 in recurrent early pregnancy loss patients. <i>Placenta</i> 32 619-621. Nissi R, Talvensaari-Mattila A, Kotila V, Niinimaki M, Jarvela I & Turpeenniemi-Hujanen T 2013 Circulating matrix metalloproteinase MMP-9 and MMP-2/TIMP-2 complex are associated with spontaneous early pregnancy failure. <i>Reprod Biol</i>
935	 Endocrinol 11 2. Niswender GD, Juengel JL, Silva PJ, Rollyson MK & McIntush EW 2000 Mechanisms controlling the function and life span of the corpus luteum. <i>Physiol Rev</i> 80 1-29. Oliveira GBd, Vale AMd, Santos ACd, Moura CEBd, Rocha HAdO & Oliveira MFd 2015 Composition and significance of glycosaminoglycans in the uterus and
940	 placenta of mammals. <i>Brazilian Archives of Biology and Technology</i> 58 512-520. Paidas MJ, Krikun G, Huang SJ, Jones R, Romano M, Annunziato J & Barnea ER 2010 A genomic and proteomic investigation of the impact of preimplantation factor on human decidual cells. <i>Am J Obstet Gynecol</i> 202 459.e451-458. Sagsoz H, Liman N, Saruhan BG & Kucukaslan I 2014 The expression of epidermal
945	growth factor receptors and their ligands (epidermal growth factor, neuregulin, amphiregulin) in the bitch uterus during the estrus cycle. <i>Anim Reprod Sci</i> 147 161-179. Seshagiri PB, Sen Roy S, Sireesha G & Rao RP 2009 Cellular and molecular regulation
950	of mammalian blastocyst hatching. <i>J Reprod Immunol</i> 83 79-84. Short RV 1969. Implantation and the Maternal Recognition of Pregnancy. In <i>Ciba</i> <i>Foundation Symposium - Foetal Autonomy</i> , pp. 2-31. Smith SD, Choudhury RH, Matos P, Horn JA, Lye SJ, Dunk CE, Aplin JD, Jones RL & Harris LK 2016 Changes in vascular extracellular matrix composition during decidual spiral arteriole remodeling in early human pregnancy. <i>Histol Histopathol</i> 24 557 574
955	 31 557-571. Smits P, Ni J, Feng P, Wauters J, Van Hul W, Boutaibi ME, Dillon PJ & Merregaert J 1997 The human extracellular matrix gene 1 (ECM1): genomic structure, cDNA cloning, expression pattern, and chromosomal localization. <i>Genomics</i> 45 487-495.

Stetler-Stevenson WG 2008 Tissue inhibitors of metalloproteinases in cell signaling:
metalloproteinase-independent biological activities. <i>Sci Signal</i> 1 re6.

960 **Strauss JF, 3rd** 2013 Extracellular matrix dynamics and fetal membrane rupture. *Reprod Sci* **20** 140-153.

Sutherland AE, Calarco PG & Damsky CH 1993 Developmental regulation of integrin expression at the time of implantation in the mouse embryo. *Development* **119** 1175-1186.

- 965 **Vermeirsch H, Simoens P & Lauwers H** 2000 Immunohistochemical detection of the estrogen receptor-alpha and progesterone receptor in the canine pregnant uterus and placental labyrinth. *Anat Rec* **260** 42-50.
 - Wewer UM, Damjanov A, Weiss J, Liotta LA & Damjanov I 1986 Mouse endometrial stromal cells produce basement-membrane components. *Differentiation* **32** 49-58.
- 970

980

- Wewer UM, Faber M, Liotta LA & Albrechtsen R 1985 Immunochemical and ultrastructural assessment of the nature of the pericellular basement membrane of human decidual cells. *Lab Invest* **53** 624-633.
- Winterhager E, Grummer R, Jahn E, Willecke K & Traub O 1993 Spatial and temporal expression of connexin26 and connexin43 in rat endometrium during trophoblast invasion. *Dev Biol* 157 399-409.

Wislocki GB & Dempsey EW 1946 Histochemical reactions in the placenta of the cat. *Am J Anat* **78** 1-45.

Zhang C, Duan E, Cao Y, Jiang G & Zeng G 2000 Effect of 32/67 kDa laminin-binding protein antibody on mouse embryo implantation. *J Reprod Fertil* **119** 137-142.

Zollinger AJ & Smith ML 2017 Fibronectin, the extracellular glue. *Matrix Biol* **60-61** 27-37.

Figures, Tables and Supplemental Files:

Tables:

Table 1:

List of primers used for real-time (TaqMan) PCR.

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Table 2:

List of primary and secondary antibodies used for indirect immunohistochemistry (IHC). PIER = protease initiated epitope retrieval; HIER heat initiated epitope retrieval

10 **Table 3:**

Tabular, descriptive presentation of localization patterns of extracellular matrix proteins (ECM) as determined by immunohistochemistry (IHC) in canine uterine and placental compartments. (-) = not present; (+/-) = weakly present; (+) = strongly present; (++) = very strongly present. COL = collagen, FN1 = fibronectin 1, ECM1 = extracellular matrix protein

15 1, LAMA2 = laminin alpha 2, Cx = connexin, TIMP = tissue inhibitor of metalloproteinase, aSMA = alpha-smooth muscle actin.

Figures:

Figure 1:

Expression of selected extracellular matrix (ECM) genes in the canine pre-implantation uterus (days 10-12 of pregnancy) affected by the presence of free-floating embryos, compared with gene expression in their non-pregnant counterparts as determined by Real Time (TaqMan) PCR. *FN1*= fibronectin 1; *ECM1*= extracellular matrix protein 1, *TIMP4* = tissue inhibitor of metalloproteinases 4. An unpaired, two-tailed Student's t-test was applied. P<0.05 was defined as significant. Bars with different asterisks differ at P=0.01. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).

Figure 2:

Expression of extracellular matrix protein 1 (*ECM1*), collagen (*COL*) 1, -3 and -4 as
determined by Real Time (TaqMan) PCR in the canine uterus during different times of pregnancy. Two comparisons are presented for each gene: First, the pre-implantation (Pre-imp.) stage was compared to the inter-placental sites during post-implantation (Post-imp.) and mid-gestation (Mid-gest.), and next to the utero-placental compartments (placentation sites) of post-implantation (Post-imp.), mid-gestation (Mid-gest.) and prepartum luteolysis. Asterisks

35 indicate $\star = p < 0.05$, $\star \star = p < 0.01$, $\star \star \star = p < 0.001$. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).

Figure 3:

Expression of fibronectin 1 (*FN1*), laminin alpha 2 (*LAMA2*), connexin (*Cx*) 26 and -43 as
determined by Real Time (TaqMan) PCR in canine uterus during different times of pregnancy. Two comparisons are presented for each gene: First, the pre-implantation (Pre-imp.) stage was compared to inter-placental sites during post-implantation (Post-imp.) and mid-gestation (Mid-gest.), and next to the utero-placental compartments (placentation sites) of post-implantation (Post-imp.), mid-gestation (Mid-gest.) and prepartum luteolysis. Asterisks

45 differ at $\star = p < 0.05$, $\star \star = p < 0.01$, $\star \star \star = p < 0.001$. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).

Figure 4:

Expression of tissue inhibitor of metalloproteinase (TIMP) 2 and -4 as determined by Real

- 50 Time (TaqMan) PCR in canine uterus during different times of gestation. Two comparisons are presented for each gene: First, the pre-implantation (Pre-imp.) stage was compared to the inter-placental sites during post-implantation (Post-imp.) and mid-gestation (Mid-gest.), and next to the utero-placental compartments (placentation sites) of post-implantation (Post-imp.), mid-gestation (Mid-gest.) and prepartum luteolysis. Asterisks differ at ★= p<0.05, ★★=</p>
- 55 p<0.01, ★★★= p<0.001. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).</p>

Figure 5:

compartments.

Representative pictures of immunohistochemical detection of collagen 1 (COL1) in the canine
uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: the
non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine
uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at
prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with surface area; (B, D, F
and I): deep glands area at the border with the myometrium; (E and H): area above the
connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental

Page 2 of 9

<u>Non-pregnant (A-B)</u>, and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium;

70 <u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel:

Collagen 1 was clearly detectable throughout pregnancy in all stromal compartments. Before implantation and placentation, strong signals were detected in stroma and myometrium of

both non-pregnant (A, B) and pregnant dogs (C, D). Following implantation and placentation, strong signals were localized in the connective tissue layer (stromal area) separating superficial from deep glands (E, H) as well as in myometrial (F, I) and placental stromal (G, J) compartments. Inlet in picture B shows a negative (isotype) control.

80 **Figure 6**:

Representative pictures of immunohistochemical detection of extracellular matrix protein 1 (ECM1) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-

Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J) : placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface

- 90 (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;
 <u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel;
- 95 ECM1 was weakly detectable in luminal epithelium in the non-pregnant group (A, B). In early pregnancy, strong signals were detectable in luminal epithelium and in superficial and deep glands (C, D). During mid-gestation (E) and at prepartum luteolysis (H), ECM1 was weakly detectable in glandular epithelium of the superficial glands. At prepartum luteolysis, ECM was strongly present in the connective tissue layer separating superficial glands from
- 100 deep glands (H). In mid-gestation (F) and at pre-partum luteolysis (I), ECM1 was detectable

in the glandular epithelium of deep glands. Placental localization of ECM1 was in cytotrophoblast, shown at mid-gestation (G) and prepartum luteolysis (J). Inlet in picture B shows respective negative (isotype) control.

105 **Figure 7:**

Representative pictures of immunohistochemical detection of cell adhesion molecule connexin 43 (Cx43) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F,

G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J) : placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface

115 (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium;

120 cytotrophoblast

125

Cx43 was present in epithelial compartments and myometrial compartments. The luminal epithelium (A, C) and the epithelial cells of superficial and deep glands (A-F, H, I) stained strongly for Cx43. Additionally, the protein was present in tunica intima of blood vessels in all samples. Placental Cx43 localization was diffusely detected in cyto- and syncytiotophoblast and decidual cells (G, J), additionally strong signals for Cx43 were detected in the intima and media of placental blood vessels. Inlet in picture B shows respective negative (isotype) control.

Figure 8:

130 Representative pictures of immunohistochemical detection of tissue inhibitor of metalloproteinases 2 (TIMP2) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); A and C:

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135 luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments.

<u>Non-pregnant (A-B), and before implantation and placentation (C-D)</u>: open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium;

<u>After implantation and placentation (E-J):</u> open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; D = decidual cell; S = syncytiotrophoblast; C = cytotrophoblast</u>

- 145 In the non-pregnant group, TIMP2 was weakly detectable in the deep glands (B). In early pregnancy, TIMP2 was clearly detectable in luminal (C) and superficial (C) epithelium, deep glands (D) and myometrial compartments (D). At mid-gestation, TIMP2 was detectable in the glandular epithelium of the superficial glands (E), but not in the stroma of the connective tissue layer separating superficial from deep glands (E). At prepartum luteolysis, TIMP2
- 150 signals appeared weaker (compared to mid-gestation) in the luminal epithelium of the superficial glands; additionally, strong signals were detected in the connective tissue layer (H). At mid-gestation (F) TIMP2 appeared in the *tunica media* of blood vessels in the myometrium. At midgestation (F) and at prepartum luteolysis (I), TIMP2 was present in the myometrium and in deep glands. In placenta at midgestation (G) and at pre-partum luteolysis
- 155 (J), TIMP2 was present in cyto- and syncytiotrophoblast, decidual cells and the intima of blood vessels. In placenta the endothelium appeared to stain strongest for TIMP2. Inlet in picture B shows the respective negative (isotype) control.

Figure 9:

- 160 Representative pictures of immunohistochemical detection of tissue inhibitor of metalloproteinases 4 (TIMP4) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during midgestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C):
 165 luminal surface area; (B, D, F and I): deep glands area at the border of the myometrium; (E
- and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments.

<u>Non-pregnant (A-B)</u>, and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;

- glandular epithelium; asterisk = blood vessel;
 <u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel;
- Before trophoblast attachment (*i.e.* at early pregnancy), and in non-pregnant controls, TIMP4
 appeared weakly present in the surface epithelium (A, C). Also superficial epithelium stained weakly throughout pregnancy. Stronger signals were seen in the epithelium of deep glands, in particular in non-pregnant dogs (B) and at mid-gestation and in the prepartum luteolysis group (F, I). TIMP4 was strongly detected at prepartum luteolysis in the connective tissue layer separating superficial from deep glands (H). In the placenta, shown at mid-gestation and
- 180 prepartum luteolysis, TIMP4 was present in the cytotrophoblast (G, J). Inlet in picture B shows respective negative (isotype) control.

Supplemental Files

Supplemental File 1

- 185 Immunohistochemical detection of collagen 3 (COL3) in the canine uterus and utero-placental (Ut-Pl) compartments at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands
- 190 area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments. Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;
- 195 <u>After implantation and placentation (E-J):</u> open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; open arrowhead = fetal stroma. Inlet in picture B shows the respective negative (isotype) control.

200 **Supplemental File 2:**

Immunohistochemical detection of collagen 4 (COL4) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I,

- J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments.
 <u>Non-pregnant (A-B), and before implantation and placentation (C-D)</u>: open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead =
- 210 glandular epithelium; asterisk = blood vessel; <u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands (*i.e.*, glands above connective layer); solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel. Inlet in picture B shows the respective negative (isotype) control.
- 215

Supplemental File 3

Immunohistochemical detection of TIMP2 and TIMP4 on consecutive slides. Four consecutive slides were prepared from placental compartments of the prepartum luteolysis group. Of these, IHC staining was performed against TIMP2 (A), TIMP4 (B), Vimentin (C)

220 and pan-Cytokeratin (D). (A-D): open arrows = decidual cells; solid arrows = cytotrophoblast; solid arrowhead = intima of maternal blood vessels; open arrowhead = syncytiotrophoblast.

Supplemental File 4

- Immunohistochemical detection of alpha smooth muscle actin (aSMA) in the canine uterus
 and utero-placental compartments (Ut-Pl) at selected time points during pregnancy; in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer; (G and J): placental compartments.
- <u>Non-pregnant (A-B), and before implantation and placentation (C-D)</u>: open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;

After implantation and placentation (E-J): open arrows = glandular epithelium of superficial

235 glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel. Inlet in picture B shows the respective negative (isotype) control.

Supplemental File 5

- 240 Immunohistochemical detection of fibronectin 1 (FN1) in the canine uterus and uteroplacental (Ut-Pl) compartments at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep
- 245 glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments. Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium;
- 250 <u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; D = decidual cell; S = syncytiotrophoblast; C = cytotrophoblast. Inlet in picture B shows the respective negative (isotype) control.

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Supplemental File 6

Immunohistochemical detection of laminin alpha 2 (LAMA2) in the canine uterus and uteroplacental (Ut-Pl) (the whole thickness of the uterine wall, *i.e.*, uterus with adjacent placenta) compartments at selected time points during pregnancy: in the non-pregnant canine uterus at

- 260 early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective layer (superficial glands, so-called glandular chambers); (G and J) : placental compartments.
- 265 <u>Non-pregnant (A-B), and before implantation and placentation (C-D)</u>: open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;

<u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium

270 of deep glands; asterisk = blood vessel. Inlet in picture B shows the respective negative (isotype) control.

Supplemental File 7

- 275 Immunohistochemical detection of cell adhesion molecule connexin 26 (Cx26) in the canine uterus and utero-placental (Ut-Pl) compartments at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B,
- D, F and I): deep glands area at the border to the myometrium; (E and H): area above the connective tissue layer; (G and J): placental compartments.
 <u>Non-pregnant (A-B), and before implantation and placentation (C-D)</u>: open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium;
- 285 <u>After implantation and placentation (E-J)</u>: open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (cicular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; D = decidual cell; S = syncytiotrophoblast; C = cytotrophoblast. Inlet in picture B shows the respective negative (isotype) control.

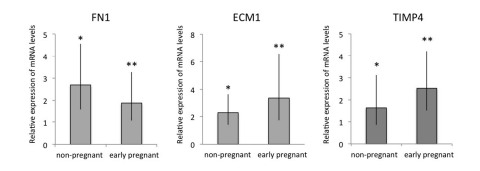
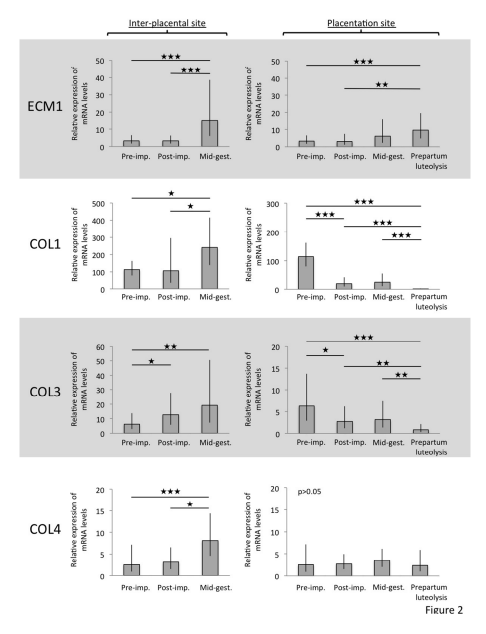
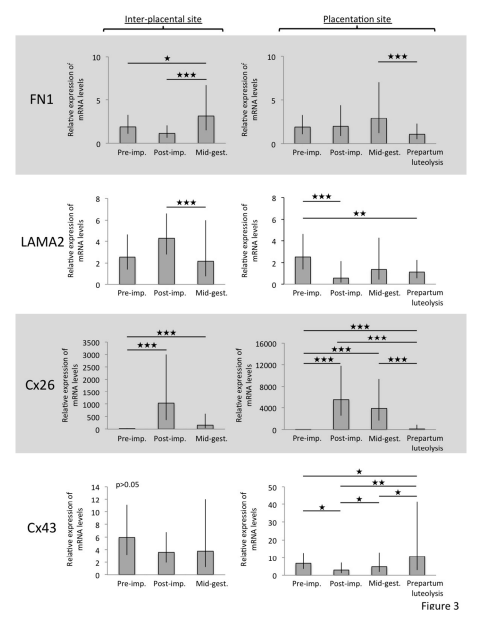


Figure 1

Expression of selected extracellular matrix (ECM) genes in the canine pre-implantation uterus (days 10-12 of pregnancy) affected by the presence of free-floating embryos, compared with gene expression in their non-pregnant counterparts as determined by Real Time (TaqMan) PCR. FN1= fibronectin 1; ECM1= extracellular matrix protein 1, TIMP4 = tissue inhibitor of metalloproteinases 4. An unpaired, two-tailed Student's t-test was applied. P<0.05 was defined as significant. Bars with different asterisks differ at P=0.01. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).



Expression of extracellular matrix protein 1 (ECM1), collagen (COL) 1, -3 and -4 as determined by Real Time (TaqMan) PCR in the canine uterus during different times of pregnancy. Two comparisons are presented for each gene: First, the pre-implantation (Pre-imp.) stage was compared to the inter-placental sites during post-implantation (Post-imp.) and mid-gestation (Mid-gest.), and next to the utero-placental compartments (placentation sites) of post-implantation (Post-imp.), mid-gestation (Mid-gest.) and prepartum luteolysis. Asterisks indicate $\star = p < 0.05$, $\star \star = p < 0.01$, $\star \star \star = p < 0.001$. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).



Expression of fibronectin 1 (FN1), laminin alpha 2 (LAMA2), connexin (Cx) 26 and -43 as determined by Real Time (TaqMan) PCR in canine uterus during different times of pregnancy. Two comparisons are presented for each gene: First, the pre-implantation (Pre-imp.) stage was compared to inter-placental sites during post-implantation (Post-imp.) and mid-gestation (Mid-gest.), and next to the utero-placental compartments (placentation sites) of post-implantation (Post-imp.), mid-gestation (Mid-gest.) and prepartum luteolysis. Asterisks differ at $\star = p < 0.05$, $\star \star = p < 0.01$, $\star \star \star = p < 0.001$. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).

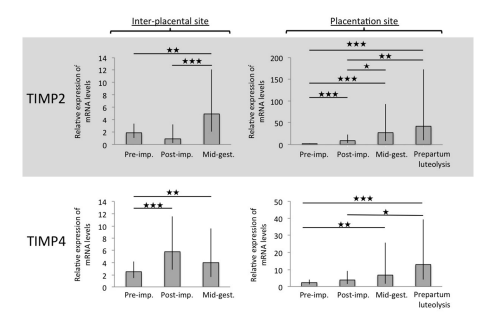
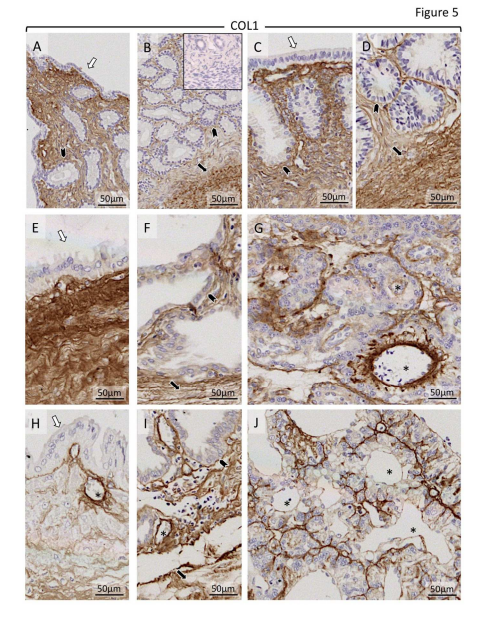


Figure 4

Expression of tissue inhibitor of metalloproteinase (TIMP) 2 and -4 as determined by Real Time (TaqMan) PCR in canine uterus during different times of gestation. Two comparisons are presented for each gene: First, the pre-implantation (Pre-imp.) stage was compared to the inter-placental sites during postimplantation (Post-imp.) and mid-gestation (Mid-gest.), and next to the utero-placental compartments (placentation sites) of post-implantation (Post-imp.), mid-gestation (Mid-gest.) and prepartum luteolysis. Asterisks differ at $\star = p < 0.05$, $\star \star = p < 0.01$, $\star \star \star = p < 0.001$. Numerical data are presented as geometric means Xg ± geometric standard deviation (SD).

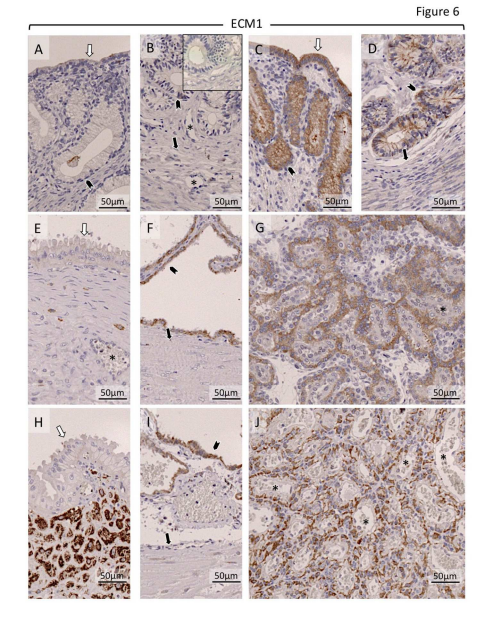


Representative pictures of immunohistochemical detection of collagen 1 (COL1) in the canine uterus and utero-placental compartments (Ut-PI) at selected time points during pregnancy: the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-PI units during mid-gestation (E, F, G), and in Ut-PI compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; After implantation and placentation (E-J): open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel:

Collagen 1 was clearly detectable throughout pregnancy in all stromal compartments. Before implantation

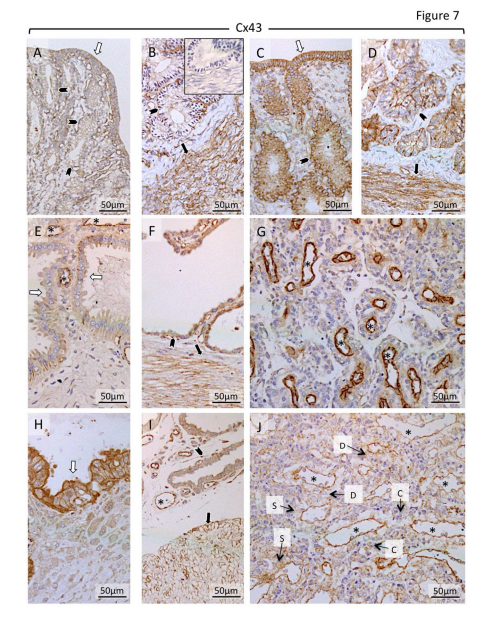
and placentation, strong signals were detected in stroma and myometrium of both non-pregnant (A, B) and pregnant dogs (C, D). Following implantation and placentation, strong signals were localized in the connective tissue layer (stromal area) separating superficial from deep glands (E, H) as well as in myometrial (F, I) and placental stromal (G, J) compartments. Inlet in picture B shows a negative (isotype) control.



Representative pictures of immunohistochemical detection of extracellular matrix protein 1 (ECM1) in the canine uterus and utero-placental compartments (Ut-PI) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-PI units during mid-gestation (E, F, G), and in Ut-PI compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J) : placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;

After implantation and placentation (E-J): open arrows = glandular epithelium of superficial glands solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; ECM1 was weakly detectable in luminal epithelium in the non-pregnant group (A, B). In early pregnancy, strong signals were detectable in luminal epithelium and in superficial and deep glands (C, D). During midgestation (E) and at prepartum luteolysis (H), ECM1 was weakly detectable in glandular epithelium of the superficial glands. At prepartum luteolysis, ECM was strongly present in the connective tissue layer separating superficial glands from deep glands (H). In mid-gestation (F) and at pre-partum luteolysis (I), ECM1 was detectable in the glandular epithelium of deep glands. Placental localization of ECM1 was in cytotrophoblast, shown at mid-gestation (G) and prepartum luteolysis (J). Inlet in picture B shows respective negative (isotype) control.

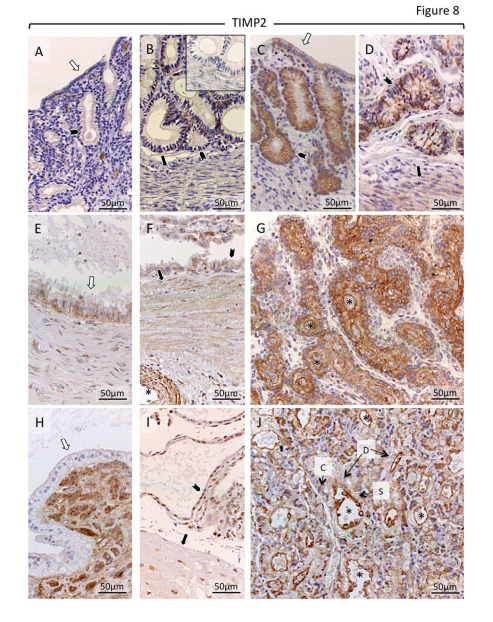


Representative pictures of immunohistochemical detection of cell adhesion molecule connexin 43 (Cx43) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J) : placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; After implantation and placentation (E-J): open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; D = decidual cell; S = syncytiotrophoblast; C = cytotrophoblast

Cx43 was present in epithelial compartments and myometrial compartments. The luminal epithelium (A, C)

and the epithelial cells of superficial and deep glands (A-F, H, I) stained strongly for Cx43. Additionally, the protein was present in tunica intima of blood vessels in all samples. Placental Cx43 localization was diffusely detected in cyto- and syncytiotophoblast and decidual cells (G, J), additionally strong signals for Cx43 were detected in the intima and media of placental blood vessels. Inlet in picture B shows respective negative (isotype) control.

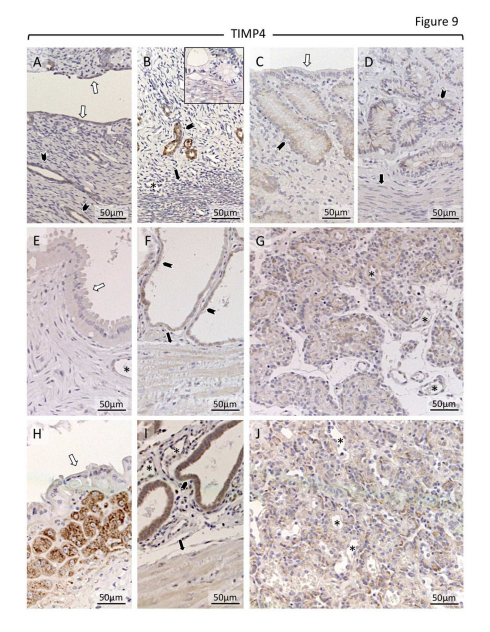


Representative pictures of immunohistochemical detection of tissue inhibitor of metalloproteinases 2 (TIMP2) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); A and C: luminal part of uterus with uterine surface area; (B, D, F and I): deep glands area at the border with the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; After implantation and placentation (E-J): open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; D = decidual cell; S = syncytiotrophoblast; C = cytotrophoblast

In the non-pregnant group, TIMP2 was weakly detectable in the deep glands (B). In early pregnancy, TIMP2

was clearly detectable in luminal (C) and superficial (C) epithelium, deep glands (D) and myometrial compartments (D). At mid-gestation, TIMP2 was detectable in the glandular epithelium of the superficial glands (E), but not in the stroma of the connective tissue layer separating superficial from deep glands (E). At prepartum luteolysis, TIMP2 signals appeared weaker (compared to mid-gestation) in the luminal epithelium of the superficial glands; additionally, strong signals were detected in the connective tissue layer (H). At mid-gestation (F) TIMP2 appeared in the tunica media of blood vessels in the myometrium. At midgestation (F) and at prepartum luteolysis (I), TIMP2 was present in the myometrium and in deep glands. In placenta at midgestation (G) and at pre-partum luteolysis (J), TIMP2 was present in cyto- and syncytiotrophoblast, decidual cells and the intima of blood vessels. In placenta the endothelium appeared to stain strongest for TIMP2. Inlet in picture B shows the respective negative (isotype) control.



Representative pictures of immunohistochemical detection of tissue inhibitor of metalloproteinases 4 (TIMP4) in the canine uterus and utero-placental compartments (Ut-Pl) at selected time points during pregnancy: in the non-pregnant canine uterus at early diestrus (A, B), early-pregnant (pre-implantation) canine uterus (C, D), Ut-Pl units during mid-gestation (E, F, G), and in Ut-Pl compartments at prepartum luteolysis (H, I, J); (A and C): luminal surface area; (B, D, F and I): deep glands area at the border of the myometrium; (E and H): area above the connective tissue layer (superficial glands, so-called glandular chambers); (G and J): placental compartments.

Non-pregnant (A-B), and before implantation and placentation (C-D): open arrows = surface (luminal) epithelium; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium; asterisk = blood vessel;

After implantation and placentation (E-J): open arrows = glandular epithelium of superficial glands; solid arrows = myometrium (circular layer); solid arrowhead = glandular epithelium of deep glands; asterisk = blood vessel; Before trophoblast attachment (i.e. at early pregnancy), and in non-pregnant controls, TIMP4 appeared weakly present in the surface epithelium (A, C). Also superficial epithelium stained weakly throughout pregnancy. Stronger signals were seen in the epithelium of deep glands, in particular in non-pregnant dogs (B) and at mid-gestation and in the prepartum luteolysis group (F, I). TIMP4 was strongly detected at prepartum luteolysis in the connective tissue layer separating superficial from deep glands (H). In the placenta, shown at mid-gestation and prepartum luteolysis, TIMP4 was present in the cytotrophoblast (G, J). Inlet in picture B shows respective negative (isotype) control.

Primer	Accession number		Product length (bp)	
extracellular matrix protein 1 (ECM1)	XM_845921.4	Forward	5'-CAG TCT GGC TTC TCC CAC CTT A-3'	
		Reverse	5'-GCG GTT TGT GTG GCT GTG A-3'	99
		TaqMan Probe	5'-AGA CTA GAT ATT CCC GCT GCT GCC GCT-3'	
connexin 26 (<i>Cx26</i>)	AJ439693.1	Forward	5'-CCA CTA CTT CCC CAT CTC TCA CA -3'	
		Reverse	5'-TCC GGT AGG CGA CAT GCA T -3'	98
		TaqMan Probe	5'-CCG ACT CTG GGC TCT GCA GCT GAT C-3'	
connexin 43 (<i>Cx43</i>)	AY462223	Forward	5'-AAA AGA GAA CCC TGC CCT CAT C-3'	
		Reverse	5'-AGG ACA CGA CCA GCA TGA AGA-3'	91
		TaqMan Probe	5'-ACT GCT TCC TCT CTC GCC CCA CG-3'	
fibronectin 1 (FN1)	XM_014110981	Forward	5'-CAC GCC GAA CTA CGA TGC-3'	
		Reverse	5'-TGC GAT ACA TGA CCC CTT-3'	95
		TaqMan Probe	5'-AAG TTT GGA TTT TGC CCC ATG GCC-3'	
laminin alpha 2 (LAMA2)	XM_014113700.1	Forward	5′-AAA CCG GCT CAC GAT TGA G-3′	
		Reverse	5'-AGT TGA ACG GTG GCG AAG T-3'	99
		TaqMan Probe	5'-CCT GCT CTT CTA CAT GGC TCG GAT CAA-3'	
glyceraldehyd 3-phosphate dehydrogenase (GAPDH)	AB028142	Forward:	5'-GCT GCC AAA TAT GAC GAC ATC A-3'	
		Reverse:	5'-GTA GCC CAG GAT GCC TTT GAG-3'	75
		TaqMan Probe	5'-TCC CTC CGA TGC CTG CTT CAC TAC CTT-3'	
tissue inhibitor of matrix metalloproteinase-2 (<i>TIMP2</i>)	AF188489	Forward	5'-CAT AGG TAC CAG ATG GGC TGT GA-3'	
		Reverse	5'-CAG TCC ATC CAG AGG CAC TCA-3'	95
		TaqMan Probe	5'-TGA TCC CGT GCT ATA TCT CGT CTC CGG-3'	
tissue inhibitor of matrix metalloproteinase-2 (<i>TIMP4</i>)	NM_001314106	Forward	5'-CTG TGG CTG CCA AAT TAC CA-3'	
		Reverse	5'-CCC ATA GAG CTT CCG TTC CA-3'	103
		TaqMan Probe	5'-ACC ATC TCA GCC CCT AAC GAG TGC CTC-3'	

Table 1

Antibody	Company	Reference Number	Antigen-Retrieval	Dilution	
collagen I (COL1)	Rockland	31680	PIER 60min	1:800	
collagen III (COL3)	Quartett	292302	PIER 45min	1:200	
collagen IV (COL4)	Quartett	250424	PIER 60min	1:500	
laminin 2 alpha (LAMA2)	Bioss Antibodies	bs-8561R	PIER 90min	1:300	
fibronectin 1 (FN1)	Novus Biologicals	NBP1-91258	PIER 30min	1:300	
alpha-smooth-muscle-actin (aSMA)	Dako	M0851	HIER	1:100	
connexin 26 (Cx26)	Thermo-fisher	QC215178	PIER 5min	1:200	
connexin 43 (Cx43)	Abcam	AB11370	PIER 15min	1:750	
tissue inhibitor of matrix metalloproteinase-2 (TIMP2)	Merck Millipore	AB2965	HIER	1:1500	
tissue inhibitor of matrix metalloproteinase-2 (TIMP4)	LifeSpan Biosciences	LS-C116809	HIER	1:750	
anti-vimentin clone vim3B4 (VIM)	Dako	M7020	HIER	1:100	
anti-cytokeratin (pan-CYT)	Dako	Z0622	HIER	1:700	
extracellular matrix protein 1 (ECM1)	Proteintech	11521-1-AP	HIER	1:200	
biotinylated goat anti-rabbit IgG (H+L)	Vector Laboratories	BA-1000	-	1:100	
biotinylated horse anti-mouse IgG (H+L)	Vector Laboratories	BA-2000	-	1:100	

Table 2

	COL1	COL3	COL4	FN1	ECM1	LAMA2	Cx26	Cx43	TIMP2	TIMP4	aSMA
		L	NO	N-PREGI	NANT		<u> </u>	I		I	
			1			1.	. /	<u> </u>	. /	. /	
Luminal epithelium	-	-	-	+	+	+	+/-	+	+/-	+/-	-
Stroma	+	+	-	+	-	+	-	-	-	-	+
Glandular epithelium	-	-	-	+	-	+	+/-	+	+/-	+	-
Blood vessels	-	-	+	-	-	+	-	+	+	-	++
MYOMETRIUM	1		r	r	1			1	r		
Smooth muscle cells	-	-	-	-	-	-	+	+	+/-	-	++
Stroma	+	+	+	+	-	+	-	-	+	-	-
Blood vessels	-	-	+	-	-	+	-	+	+	-	++
PRE-ATTACHMENT ENDOMETRIUM											
Luminal epithelium	-	-	-	+	++	+	+	+	++	+/-	-
Stroma	+	+	-	+	-	+	-	-	-	-	-
Glandular epithelium	-	_	_	+	+	+	+	+	++	+	-
Blood vessels	-	_	+	-	_	+	-	+	+	-	+
MYOMETRIUM									•		•
Smooth muscle cells - - - - + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + + +											
Stroma	+	+	+/-	+	_	+	-	-	+	_	-
Blood vessels			+	-	_	+	_	+	+	_	++
						<u> </u>		<u> </u>		I	
PLACENTA			IVIII	D-GESTA	TION			•			
Cytotrophoblast cells	-	-	-	+	+	-	+	+/-	+	+	-
Syncytiotrophoblast	-	-	-	+	-	-	+	+/-	+	-	-
Decidual cells	-	-	-	+	-	-	+	+/-	+	-	++
Blood vessels	-	-	+	+	-	+	+	++	+	-	+
Stroma	+	+	-	+	-	+	-	-	-	-	-
ENDOMETRIUM											
Epithelium of superficial glands	-	-	-	+	+	+	++	+	+	+/-	-
Connective tissue layer separating superficial from deep glands	+	+	-	+	-	++	-	+	+/-	-	-
Epithelium of deep glands	-	-	-	+	+	+	+	+	+	+	-
Blood vessels	-	-	+	+	-	+	+/-	++	+	-	+
MYOMETRIUM								•		•	
Smooth muscel cells	-	-	-	-	-	+	+	+	+	-	+
Blood vessels	-	-	+	-	-	+	-	+	+	-	++
Stroma	+	+	+	+	-	+	-	-	+	+	-
PREPARTUM LUTEOLYSIS PLACENTA											
Cytotrophoblast cells	-	-	-	+	+	-	++	+	+	+	-
Syncytiotrophoblast	-	-	-	+	-	-	+	+	+	-	_
Decidual cells	-	_	_	+	_	_	+	+	+	_	++
Blood vessels	-	-	+	+	_	+	+/-	++	++	-	++
Stroma	+		-	+		+	- +/-	-	-	-	1.4
	+	+	-	+	-	+	-	-	-	-	-
ENDOMETRIUM			l							,	
Epithelium of superficial glands Connective tissue layer	-	-	-	+	+	+	++	+	+/-	+/-	-
separating superficial from deep glands	+	+	-	+	++	+	+	+	++	++	-

Epithelium of deep glands	-	-	-	+	+	+	+	+	+	++	-
Blood vessels	-	-	+	-	-	+	+	++	+	-	++
MYOMETRIUM											
Smooth muscel cells	-	-	-	-	-	+	+	+	+	-	+
Blood vessels	-	-	+	-	-	+	+	+	+	-	+
Stroma	+	+	+/-	+	-	+	-	-	+	+	-

Table3