1	EXPRESSION OF UTERINE OXYTOCIN RECEPTORS AND BLOOD PROGESTERONE,
2	13,14-DIHYDRO-15-KETO-PROSTAGLANDIN F2A, AND IONIZED CALCIUM LEVELS
3	IN DYSTOCIC BITCHES
4	
5	Tuire Tamminen <sup>a</sup> , Lena Sahlin <sup>b</sup> , Britt Masironi-Malm <sup>b</sup> , Merja Dahlbom <sup>c</sup> , Terttu Katila <sup>a</sup> , Juhani
6	Taponen <sup>a</sup> , Outi Laitinen-Vapaavuori <sup>d</sup>
7	
8	<sup>a</sup> Department of Production Animal Medicine, University of Helsinki, Paroninkuja 20, 04920
9	Saarentaus, Finland
10	<sup>b</sup> Department of Women's and Children's Health, Karolinska Institutet and Karolinska University
11	Hospital; 171 64 Solna, Sweden
12	<sup>c</sup> Veterinary Clinic of Mäntsälä, Mäntymäentie 3, 04600 Mäntsälä, Finland
13	<sup>d</sup> Department of Equine and Small Animal Medicine, University of Helsinki, Viikintie 49, 00790
14	Helsinki, Finland
15	Corresponding author. Tel.: +358 50 4150196. E-mail address: tuire.tamminen@helsinki.fi (T.
16	Tamminen).
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#### 18 Abstract

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20 This study aimed to examine the etiology of canine dystocia by measuring the relative expression of 21 oxytocin receptor (OXTR) mRNA and the concentration of serum progesterone, plasma 22  $PGF_{2\alpha}$  metabolite (PGFM), and blood ionized calcium (iCa) near term and in dystocia. Altogether 58 23 bitches were included in this study, 41 of which underwent cesarean section (CS). The four CS groups 24 were based on history: complete uterine inertia (CUI; n = 7), partial uterine inertia (PUI; n = 13), 25 obstructive dystocia (OD; n = 10), and elective cesarean section (ECS; n = 11). An additional group 26 of medically treated dystocia without CS (MD; n = 8) and a control group (C; n = 9) with normal 27 parturition (without CS and medical treatment) were also formed. Blood samples were taken prior to 28 CS or medical treatment. Progesterone concentrations were highest in the ECS and a significant 29 difference (p<0.05) was observed between the ECS and the OD and between the ECS and the combined dystocia (CUI, PUI, OD, MD) groups (COMB). Highest concentrations of PGFM was 30 31 observed in the C, the difference being significant (p<0.05) between the C and the ECS and between 32 the C and the COMB group. The progesterone: PGFM ratio was significantly (p<0.05) higher in the 33 ECS than in the C and the COMB group. No significant difference (p>0.05) was observed in iCa 34 concentrations between the groups. Relative OXTR mRNA expression was evaluated with real-time 35 PCR from full-thickness uterine samples taken from the incision site during CS. The expression was highest in the ECS and the difference in expression was significant (p<0.05) between the ECS and 36 37 the OD and between ECS and the combined dystocia (CUI, PUI, OD) groups (COMB2). The study 38 supports previous reports of decreasing progesterone and increasing PGFM during prepartum 39 luteolysis. Upregulation of OXTR occurs near term. In obstructive dystocia, a prolonged influence of 40 oxytocin and uterine exhaustion may lead to downregulation of OXTR. Complete primary uterine 41 inertia may have a different etiology as no clear decrease in OXTR was observed in CUI as in OD. It

42 remains unclear if parturition ceases because of uterine inertia or if uterine inertia occurs because of

43 ceased parturition and desensitization of receptors.

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45 Keywords:

- 46 Canine, uterus, birth, progesterone, prostaglandin  $F_{2\alpha}$
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### 49 **1. Introduction**

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51 Parturition is a complex event and includes hormonal and behavioral changes, neural activity, and interaction between the dam and the offspring. Near term, canine plasma  $PGF_{2\alpha}$  levels increase 52 leading to luteolysis, followed by a decrease in peripheral plasma progesterone levels [1,2,3] that 53 54 allow for contractions of the uterus and for parturition to proceed [1]. The secretion of  $PGF_{2\alpha}$  in the 55 bitch is suggested to originate from placental trophoblast cells [4]. During parturition, increase in 56 peripheral plasma cortisol [2,5,6], vasopressin [6], and oxytocin (OT) [6,7] occur. However, changes 57 in cortisol levels vary greatly between individuals during parturition [2,5,6]. While estrogen 58 concentrations are somewhat higher in the last trimester of pregnancy in the bitch, there is no marked 59 prepartum increase as detected in many other species [8,9,10]. Two days prior to parturition, estrogen levels of the bitch decrease suddenly during prepartum luteolysis indicating its luteal source [11]. 60

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Occurrence of dystocia in bitches varies greatly depending on the population studied; the average is estimated to be below 5% [12]. In a group of 200 000 insured bitches (excluding Boston Terrier, English Bulldog, and French Bulldog) in Sweden, dystocia occurred in 16% of parturitions [13]. In the UK, the occurrence varied from 0% to 92% among 151 breeds (22, 005 litters) [14]. While dystocia seems to be more common in miniature and small breeds [13,15,16], several medium- and
large-size breeds also have a higher than average proportion of litters born by cesarean section (CS)
[14]. Approximately 60% of dystocia cases undergo CS [13,15,16,17]. In brachycephalic breeds, the
proportion of CS is very high [14,18]. There may be a risk of bias in statistics of dystocia in these
breeds due to the popularity of elective CS (ECS).

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72 Dystocia is sometimes difficult to diagnose. Therefore, a complete history and physical examination 73 is required. The suggested causes vary slightly according to different authors [12,19,20]. There are 74 several, sometimes simultaneous, causes of dystocia. Maternal factors are more common than fetal 75 factors. The most common maternal cause is primary uterine inertia, which can be complete or partial [12]. In complete primary uterine inertia, the uterus fails to initiate parturition due to absence of 76 77 uterine contractions and thus no puppies are born [12,17]. In partial primary uterine inertia, the bitch 78 may have weak uterine contractions or contractions that cease without any obvious reason (such as 79 obstruction) before all puppies are born [12,17]. Secondary uterine inertia is caused by prolonged 80 parturition due to obstruction in the birth canal [12,17].

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Oxytocin is a nonapeptide hormone produced mainly in the hypothalamus and stored in the posterior 82 83 pituitary gland. Oxytocin is released after suitable stimulus, such as intracervical pressure. As one of 84 the most potent uterotonic hormones, OT enhances the contractility of the uterus. During parturition, 85 plasma OT concentration increases [6,7]; this may not occur in dystocia [21,22]. The effect of OT in 86 the uterus is mediated through specific, class I G-protein-coupled transmembrane receptors known as 87 oxytocin receptors (OXTR) [23]. Near term, during prepartum luteolysis, OXTR are upregulated [24,25,26]. In humans, continuous exposure to OT leads to desensitization of OXTR by reduction of 88 89 OT binding sites in the myometrial cell membrane and by downregulation of OXTR mRNA in 90 myometrial cells [27]. While desensitization may also have a role in canine dystocia due to prolonged

91 influence of OT, there is no published evidence of OXTR desensitization in bitches. The aim of this 92 study was to examine the relative expression of OXTR mRNA in the canine uterus near parturition 93 and in dystocia. Levels of serum progesterone, plasma prostaglandin  $F2_{\alpha}$  metabolite (PGFM), 94 progesterone: PGFM ratio, and blood ionized calcium (iCa) were also analyzed to clarify possible 95 causative factors for dystocia. 96 97 98 2. Materials and methods 99 100 The study was approved by the Ethics Committee of the Viikki Campus, University of Helsinki, Finland. Blood sampling from bitches with normal parturition was authorized by the National Animal 101 Experiment Board (ESAVI, Hämeenlinna, Finland), license number ESAVI/3802/04.10.03/2011. 102 103 104 2.1. Groups 105 106 Client-owned pet bitches that had CS performed either at the Small Animal Clinic of Mäntsälä or the 107 Veterinary Teaching Hospital of the University of Helsinki were enrolled in the study (Table 1). The inclusion criteria were a diagnosis of dystocia resulting in CS or ECS due to small litter size or 108 109 previous dystocia. In addition, one group was established from bitches with medically treated mild 110 dystocia that gave birth without CS. Bitches with normal parturitions served as controls for blood parameters. The owners of the bitches were requested to sign a written consent and complete a 111 112 questionnaire to obtain the history of the bitch including previous and present parturitions. Any systemic disease was an exclusion criterion. 113

115 The following study groups were formed: 1) complete primary uterine inertia (CUI; n = 7, no puppies 116 born, parturition does not proceed, discharge of fetal fluids >3 hours or green discharge, no response to vaginal stimulus), 2) partial primary uterine inertia (PUI; n = 13, at least one puppy born, parturition 117 118 ceases without obstruction), 3) obstructive dystocia (OD; n = 10, fetal oversize/narrow birth canal, malpresentation, malformation), 4) elective caesarean section (ECS; n = 11, 58-66 days from mating, 119 120 previous dystocia, one or two puppies, before the onset of the stage 1 of parturition), 5) medically treated dystocia (MD; n = 8, no CS, medical treatment), 6) control (C; n = 9, no CS, no medical 121 122 treatment, normal parturition). Dystocia groups were also combined (COMB: CUI, PUI, OD, MD 123 and COMB2: CUI, PUI, OD) to compare with ECS and C. The diagnosis and treatment decisions 124 were performed by the veterinarian on call. After blood sampling, the bitches were treated, if necessary, with calcium glubionate (Calcium-Sandoz<sup>®</sup>, Sandoz A/S, Copenhagen, Denmark) and 125 oxytocin (Vetox<sup>®</sup>, Vetcare, Salo, Finland) (Table 1). 126

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The individual and average data of the bitches are presented in Table 1. Altogether 35 different breeds were included in the study. Four bitches in the ECS group had had previous history of dystocia and CS. In the PUI and MD groups, each had one bitch with mild, medically treated dystocia without CS in the previous pregnancy. The previous parturitions of the other multiparous bitches were normal. The gestation length was calculated from ovulation day (at progesterone level 16-32 nmol/l) and from the first and the last mating according the available information (Table 1).

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#### 135 2.2. Blood sampling

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Blood samples were taken prior to CS or medical treatment from the vena cephalica into a syringe
(Radiometer Safe Pico, ref: 956-610, Radiometer Medical, Copenhagen, Denmark), to an EDTA tube
(Vacuette<sup>®</sup>, Mekalasi Oy, Nurmijärvi, Finland) with 5000 KIU aprotinin/ml EDTA blood (Aprotinin,

Roche Diagnostics GmbH, Mannheim, Germany) and a serum tube with clotting activator (Vacuette<sup>®</sup>, Mekalasi Oy, Nurmijärvi, Finland). Blood samples were taken prepartum in the ECS group and peripartum (second stage of parturition) in the other groups. EDTA tubes and syringes were stored in an ice-water bath and serum tubes at room temperature. Blood samples were centrifuged (Eppendorf Centrifuge 5810R, Eppendorf Nordic A/S, Hørsholm, Denmark) as follows: EDTA tubes at 4 °C, 1200 x g, 10 min, and serum tubes at 22 °C, 1700 x g, 10 min. Plasma and serum were divided into aliquots, frozen at -20 °C, and stored at -70 °C until analyzed.

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#### 148 2.3. Progesterone assay

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Serum progesterone concentrations were measured in one run using a commercial RIA kit (Progesterone Coat-A-Count<sup>®</sup> RIA, Siemens Healthcare Diagnostics Oy, Espoo, Finland) according to the manufacturer's instructions. The concentrations were measured in duplicate with a gamma counter (1272 GliniGamma, LKB Wallac Oy, Turku, Finland). The intra-assay coefficient of variation was 3.4% at a serum concentration of 4.4 nmol/L and 2.0% at a concentration of 32.5 nmol/L.

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#### 157 2.4. $PGF_{2a}$ metabolite assay

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159 Concentrations of the major metabolite of PGF2 $\alpha$ , 13,14-dihydro-15-keto-prostaglandin F<sub>2 $\alpha$ </sub> (PGFM), 160 were measured from plasma using a commercial immunoassay kit (DetectX<sup>®</sup> 13,14-dihydro-15-keto-161 PGF<sub>2 $\alpha$ </sub> (PGFM) Enzyme Immunoassay Kit, Arbor Assays, Michigan, USA) according to the 162 manufacturer's instructions. Prior to performing the assay, plasma samples were diluted 1:15 with 163 the assay buffer provided in the kit. The optical density of each well was measured with a Multiscan 164 GO Spectrophotometer with SkanIt software 4.1 (Thermo Fisher Scientific Oy, Vantaa, Finland). The intra-assay coefficient of variation of duplicates was 12.0%. The inter-assay coefficient of variation was 13.0% at a plasma concentration of 15.8 nmol/L and 3.5% at a concentration of 58.5 nmol/L. The linearity of the assay was evaluated by diluting the canine plasma sample (1/10, 1/20 and 1/40) with the assay buffer provided in the kit. Observed to expected ratios were calculated for the dilutions. The mean recovery of the expected PGFM concentrations in different dilutions was 102% and dilutions of the canine plasma sample showed linearity over the studied range (R<sup>2</sup>=0.996). The detection limit was 0.13 nmol/L.

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173 *2.5. iCa* 

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Blood iCa was analyzed instantly with Roche Electrolyte Analyzer (9180, Fisher Scientific Oy,
Vantaa, Finland) from Safe Pico syringes stored in an ice-water bath. The syringes contained 60 IU
of dry electrolyte-balanced heparin. Contact with air was minimized with a specific cap to remove
possible air bubbles.

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180 *2.6. Uterine samples* 

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Uterine samples were obtained only from bitches undergoing a CS. Immediately after the removal of the puppies from the uterus, a full-thickness sample of uterine wall (approximately 5 x 30 mm) was taken from the incision site (interplacental area, uterine body or proximal horn). The sample was immediately frozen in liquid nitrogen and stored at -70 °C for PCR analysis to measure the relative expression of OXTR mRNA.

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188 *2.7. PCR* 

#### 190 2.7.1. RNA preparation and reverse transcription

The full procedure has been described previously [28]. In brief, a 2- $\mu$ g aliquot of total RNA from each canine uterine sample was reverse transcribed at 37°C for 60 min in a final volume of 20  $\mu$ L with a reaction mixture (Qiagen) containing 1× RT buffer, dNTP mix (0.5 mM each dNTP), 600 ng random primers (Invitrogen, Paisley, UK), 2 units of RNase inhibitor (Qiagen), and 4 units of Omniscript<sup>TM</sup> reverse transcriptase (Qiagen).

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#### 197 2.7.2. Real-time PCR analysis

The real-time PCR analysis and the primers used have been described previously [28]. The 198 199 oligonucleotide primer pair for the OXTR was designed with NCBI/Primer-BLAST. To standardize 200 the quantification method, RPL27 and HPRT1 were selected as non-regulated reference genes with 201 primer pairs obtained from Silva et al. [29] and Bhatti et al. [30], respectively. The primers were based on the sequences of the canine genes, and were the following: OTR forward primer: 5'-202 203 TGCTGGCCTTCATCGTGTGCT-3'; OTR primer: 5'reverse GATGAAAGCCGAGGCTTCCTTGGG-3' from NM\_001198659.1 with predicted size 95 bp; 204 205 RPL27 forward primer: 5'-ACAATCACCTCATGCCCACA-3'; RPL27 reverse primer: 5'-CTTGACCTTGGCCTCTCGTC-3' from NM\_001003102.2 with the predicted size 122 bp; HPRT1 206 207 forward primer: 5'-AGCTTGCTGGTGAAAAGGAC-3'; HPRT1 reverse 5'primer: 208 TTATAGTCAAGGGCATATCC-3' from NM\_001003357.1 with predicted size 104 bp. All samples 209 were run in duplicate and the purity of PCR products was confirmed by a melting-curve analysis in 210 all experiments. Each PCR assay included a negative control containing an RNA sample without 211 reverse transcription. The PCR amplification rate and the cycle threshold (C<sub>t</sub>) values were analyzed using iCycler<sup>TM</sup> iQ 3.1 software (Bio-Rad). The OXTR product was normalized against the mean of 212 213 RPL27 and HPRT1 products to yield the relative expression of OXTR mRNA.

# 215 2.8. Statistical analysis

217	Data were analyzed using IBM SPSS Statistics 24 software for Windows. The non-parametric
218	Kruskal-Wallis one-way ANOVA test with Bonferroni correction was used to detect possible
219	differences in serum progesterone levels, plasma PGFM levels, blood iCa, and relative expression of
220	OXTR mRNA between the groups. Differences were considered statistically significant at p<0.05.
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223	3. Results
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225	3.1. Progesterone
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227	Serum progesterone concentrations in the different groups are presented in Fig. 1a. The
228	concentrations were highest in the ECS group; the largest variation in levels was also observed in this
229	group. There was a significant difference (p<0.05) between the ECS and the OD and between the
230	ECS and the COMB groups.
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232	3.2. PGFM
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234	Plasma PGFM concentrations were highest in the C and lowest in the ECS group (Fig. 1b). A
235	significant difference (p<0.05) was detected between the C and the ECS and between the C and the

- 236 COMB groups.

	238	3.3.	Progesterone	e:PGFM
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The progesterone: PGFM ratio was highest in the ECS group (Fig. 1c). A significant difference (p<0.05) was observed between the ECS and the C and between the ECS and the COMB groups. 3.4. iCa Blood iCa concentrations were lowest in the PUI group but no significant difference (p>0.05) was observed between the groups (Fig. 1d). No hypocalcemia was detected (reference interval 1.16-1.40 nmol/l). 3.5. qPCR The mean relative expression of OXTR mRNA was highest in the ECS group (Fig. 1e). The difference was significant (p<0.05) between the ECS and the OD and between the ECS and the COMB2 groups. There was no significant difference between bitches treated or not treated with calcium glubionate and OT. 4. Discussion Our study indicates that in complete primary uterine inertia the etiology may not be the absence or downregulation of OXTR, as there was no difference in OXTR expression in comparison of CUI to bitches near term but before the first stage of parturition (ECS group). Upregulation of OXTR occurs near term, and the prolonged influence of OT and uterine exhaustion in obstructive dystocia may lead
to downregulation of OXTR.

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265 Our results support previous reports [1,2,3] on decreasing progesterone and increasing PGFM levels during prepartum luteolysis in pregnant bitches. As expected, progesterone levels were higher in the 266 267 ECS group than in the other groups, as CS was performed in this group before the onset of parturition 268 (before stage 1). A sudden decrease of progesterone is observed in near term pregnant bitches at the 269 end of the luteal phase [11]. Termination of corpora lutea function in non-pregnant bitches is 270 suggested to be more likely regressive than the active luteolytic process found in pregnant bitches, 271 which indicates a different regulation mechanism [31]. Failure of luteolysis can lead to prolonged 272 gestation [32]. Except in the ECS group, all bitches in this study had undergone luteolysis.

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274 One possibility for the etiology of complete primary uterine inertia could be a problem in parturition 275 initiation. Excessive progesterone and insufficient PGF2a levels could prevent sufficient uterine 276 contractions and thus interfere with parturition. However, our results suggest that this might not be 277 the case, as the progesterone and PGFM levels in CUI group were similar to other dystocia groups. 278 This may indicate that the etiology is more likely at the level of uterine function, such as myometrial 279 distention beyond its capacity to contract or the lack of cervical pressure to stimulate OT release. The 280 progesterone: PGFM ratio was highest in the ECS group, where the highest progesterone and lowest 281 PGFM concentrations were also found. This indicates that luteolysis had not yet occurred in this 282 group. A high progesterone: PGFM ratio has been reported in dystocic bitches with complete primary 283 uterine inertia in comparison to a control group [22]. In our study no such difference was observed.

284

Calcium and OT injections are used as a treatment for uterine inertia to enhance contractions of the uterus [19,20]. Batra [33] reported that OT-induced myometrial contractions in the rat depend on the

influx of extracellular calcium, and this influx is directly increased by OT. The action of OT has also 287 been postulated to occur by inhibiting the  $Ca^{2+}$ -extrusion pump in humans [34]. Hypocalcemia was 288 not diagnosed in any of the bitches in this study. However, there are reports of hypocalcemia in risk 289 290 groups of uterine inertia [35] and in dystocic bitches [36]. In our study, a single treatment with calcium glubionate and OT did not seem to affect the expression of OXTR mRNA or distribution of 291 292 OXTR. In dystocia, the uterus has been under the influence of OT, and exhaustion and desensitization 293 may prevent medical treatment to induce uterine contractions. However, in this study the number of 294 bitches treated or not treated was low and further investigation is necessary.

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296 Veiga et al. [26] reported higher expression of OXTR mRNA in both endometrium and myometrium 297 of late pregnant and parturient bitches than in earlier stages of pregnancy. In our study, full-thickness 298 samples were used for real-time PCR; endometrium and myometrium thus cannot be compared 299 separately. The samples of this study were run together with samples from our earlier report on non-300 pregnant bitches [28], and the relative expression of OXTR mRNA was higher in pregnant bitches 301 than in non-pregnant ones. Expression of OXTR in the canine uterus is probably not regulated only 302 by a decrease of progesterone. In anestrous bitches with basal levels of progesterone, OXTR 303 expression does not differ from diestrous bitches with uteri under the influence of progesterone [28]; 304 the expression is thus likely a part of more complex regulatory pathways. In the OD group, OXTR 305 mRNA expression was significantly decreased. In the PUI group the decrease also approached 306 significance. A large variation of OXTR mRNA expression in the CUI group may be due to the heterogeneity of this group. It is also possible that in the CUI group the mechanism of dystocia is 307 308 different than that of the PUI and OD groups. The uterus does not contract in complete primary uterine 309 inertia, which may be due to the lack of cervical stimulus and insufficient release of OT to systemic 310 circulation. Thus, desensitization might not occur and expression of OXTR mRNA could remain 311 high. Bergström et al. [21] reported lower plasma OT concentrations in primary uterine inertia cases than in bitches with normal parturition. In obstructive dystocia, and possibly in partial uterine inertia,
uterine exhaustion possibly with paracrine or autocrine signaling may result in OXTR
downregulation.

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Although strict criteria were defined to include the bitches in the groups in this study, some 316 heterogeneity probably exists. Breed diversity also increases the heterogeneity of the groups. In this 317 318 study, the bitches with normal parturition were used as controls only for the blood parameters. For 319 OXTR gene expression only ECS samples from prepartum bitches were used. Further studies are 320 necessary to compare OXTR gene expression also with samples from bitches with normal parturition. 321 The number of bitches was quite low (particularly in the CUI group), which may affect the results. A greater number of individuals is necessary to more properly evaluate the effect of calcium and OT 322 323 treatment on OXTR. Furthermore, an uterokinetic study in vitro with myometrial muscle strips, as 324 described by Gogny et al. [37], may provide information on myometrial contractions and 325 desensitization under prolonged influence of OT. Further studies of genetic background with breeds 326 and lines susceptible to complete primary uterine inertia are needed.

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#### 329 **5. Conclusions**

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This study provides evidence of prepartum upregulation of OXTR in the canine uterus. Expression of OXTR was increased near term. A decrease in expression was observed in obstructive dystocia and may also occur in partial primary uterine inertia. However, no clear decrease in expression was observed in the CUI group, which may indicate a different etiology for inertia than in OD. The etiology in complete primary uterine inertia is more likely at the level of uterine function, such as myometrial distention beyond its capacity to contract or the lack of cervical pressure to stimulate OT

337	release. A decrease of OXTR may also occur during normal parturition; the role of desensitization
338	of OXTR in dystocia should to be clarified. It remains unclear if parturition ceases because of uterine
339	inertia or if uterine inertia occurs because of ceased parturition and desensitization of receptors.
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352	References
353	
354	[1] Concannon PW, Isaman L, Frank DA, Michel FJ, Currie WB. Elevated concentrations of 13,14-
355	dihydro-15-keto-prostaglandin F-2 alpha in maternal plasma during prepartum luteolysis and
356	parturition in dogs (Canis familiaris). J Reprod Fertil 1988;84:71-7.
357	[2] Veronesi MC, Battocchio M, Marinelli L, Faustini M, Kindahl H, Cairoli F. Correlations among
358	body temperature, plasma progesterone, cortisol and prostaglandin F2 $\alpha$ of the periparturient bitch. J
359	Vet Med A 2002;49:264-8.
360	[3] Nohr B, Hoffmann B, Steinetz BE. Investigation of the endocrine control of parturition in the dog
361	by application of an antigestagen. J Reprod Fertil Suppl 1993;47:542–3.

- [4] Kowalewski MP, Beceriklisoy HB, Pfarrer C, Aslan S, Kindahl H, Kücükaslan I et al. Canine
  placenta: a source of prepartal prostaglandins during normal and antiprogestin-induced parturition.
  Reproduction 2010;139:655-64.
- [5] Concannon PW, Butler WR, Hansel W, Knight PJ, Hamilton JM. Parturition and lactation in the
  bitch: serum progesterone, cortisol and prolactin Biol Reprod. 1978;19:1113-8.
- 367 [6] Olsson K, Bergström A, Kindahl H, Lagerstedt A-S. Increased plasma concentrations of
  368 vasopressin, oxytocin, cortisol and the prostaglandin F2α metabolite during labour in the dog. Acta
  369 Physiol Scand 2003;179:281-7.
- [7] Klarenbeek M, Okkens AC, Kooistra HS, Mol JA, Bevers MM, Taverne MAM. Plasma oxytocin
- 371 concentrations during late pregnancy and parturition in the dog. Theriogenology 2007;68:1169-76.
- [8] Yoshinaga K, Hawkins RA, Stocker JF. Estrogen secretion by the rat ovary in vivo during the
  estrous cycle and pregnancy. Endocrinology 1969;85:103-12.
- [9] Robertson HA, Smeaton TC. The concentration of unconjugated oestrone, oestradiol-17 alpha and
  oestradiol-17 beta in the maternal plasma of the pregnant ewe in relation to the initiation of parturition
  and lactation. J Reprod Fertil 1973;35:461-8.
- 377 [10] Robertson HA. Changes in the concentration of unconjugated oestrone, oestradiol-17alpha and
- oestradiol-17beta in the maternal plasma of the pregnant cow in relation to the initiation of parturition
- and lactation. J Reprod Fertil 1974;36:1-7.
- [11] Concannon PW, Hansel W, Visek WJ. The ovarian cycle of the bitch: Plasma estrogen, LH and
  progesterone. Biol Reprod 1975;13:112-21.
- 382 [12] Linde-Forsberg C. Abnormalities in pregnancy, parturition, and the periparturient period. In:
- 383 Ettinger SJ, Feldman EC, editors. Textbook of veterinary internal medicine, 7th ed. St. Louis: Elsevier
- 384 Saunders; 2010, p. 1890–1901).

- [13] Bergström A, Nødtvedt A, Lagerstedt AS, Egenvall A. Incidence and breed predilection for
  dystocia and risk factors for cesarean section in a Swedish population of insured dogs. Vet Surg
  2006;35:786-91.
- [14] Evans KM, Adams VJ. Proportion of litters of purebred dogs born by caesarean section. J Small
  Anim Pract 2010;51:113-8.
- 390 [15] Gaudet DA. Retrospective study of 128 cases of canine dystocia. J Am Anim Hosp Assoc
  391 1985;21:813-8.
- 392 [16] Münnich A, Küchenmeister U. Dystocia in numbers Evidence-based parameters for
  393 intervention in the dog: Causes for dystocia and treatment recommendations. Reprod Dom Anim
  394 2009;44:Suppl 2:141–7.
- 395 [17] Darvelid AW, Linde-Forsberg C. Dystocia in the bitch: A retrospective study of 182 cases. J
  396 Small Anim Pract 1994;35:402-7.
- [18] Linde Forsberg C, Persson G. A survey of dystocia in the Boxer breed. Acta Vet Scand
  2007;49:8.
- 399 [19] Johnston SD, Root Kustritz MV, Olson PNS. Canine and feline theriogenology. 1st ed.
  400 Philadelphia: WB Saunders; 2001.
- 401 [20] Davidson A. Problems during and after parturition. In: England G, von Heimendahl A, editors.
- BSAVA Manual of canine and feline reproduction and neonatology, Gloucester: Brittish Small
  Animal Veterinary Association; 2010, p. 121–34.
- 404 [21] Bergström A, Fransson B, Lagerstedt A-S, Olsson K. Primary uterine inertia in 27 bitches:
  405 aetiology and treatment. J Small Anim Pract 2006;47:456-60.
- 406 [22] Bergström A, Fransson B, Lagerstedt AS, Kindahl H, Olsson U, Olsson K. Hormonal
  407 concentrations in bitches with primary uterine inertia. Theriogenology 2010;73:1068-75.
- 408 [23] Gimpl G, Fahrenholz F. The oxytocin receptor system: structure, function, and regulation.
- 409 Physiol Rev 2001;81:629-83.

[24] Derussi AA, de Souza RW, Volpato R, Guaitolini CR, Ackermann CL, Taffarel MO et al.
Progesterone (PR), oestrogen (ER-alpha and ER-beta) and oxytocin (OTR) gene expression in the
oviduct and uterus of pregnant and non-pregnant bitches. Reprod Domest Anim 2012;47:Suppl
6:197-9.

414 [25] Gram A, Boos A, Kowalewski MP. Uterine and placental expression of canine oxytocin receptor

415 during pregnancy and normal and induced parturition. Reprod Domest Anim 2014;49:Suppl 2:41-9.

416 [26] Veiga GA, Milazzotto MP, Nichi M, Lúcio CF, Silva LC, Angrimani DS, et al. Gene expression

of estrogen and oxytocin receptors in the uterus of pregnant and parturient bitches. Braz J Med Biol
Res 2015;48:339-43.

[27] Phaneuf S, Asbóth G, Carrasco MP, Europe-Finner GN, Saji F, Kimura T, et al. The
desensitization of oxytocin receptors in human myometrial cells is accompanied by down-regulation
of oxytocin receptor messenger RNA. J Endocrinol 1997;154:7-18.

[28] Tamminen TM, Sahlin L, Masironi B, Taponen J, Laitinen-Vapaavuori O, Katila T. Oxytocin
receptors in dioestrous and anoestrous canine uteri. Reprod Domest Anim. 2017;52:153-9.

424 [29] Silva E, Leitão S, Ferreira-Dias G, Lopes da Costa L, Mateus L. Prostaglandin synthesis genes

425 are differentially transcripted in normal and pyometra endometria of bitches.

426 Reprod Domest Anim 2009;44:Suppl 2:200-3.

427 [30] Bhatti SF, Rao NA, Okkens AC, Mol JA, Duchateau L, Ducatelle R et al. Role of progestin-

428 induced mammary-derived growth hormone in the pathogenesis of cystic endometrial hyperplasia in

the bitch. Domest Anim Endocrinol 2007;33:294-312.

[31] Kowalewski MP. Luteal regression vs. prepartum luteolysis: regulatory mechanisms governing
canine corpus luteum function. Reprod Biol. 2014;14:89-102.

432 [32] Irons PC, Nöthling JO, Volkmann DH. Failure of luteolysis leads to prolonged gestation in a

433 bitch: a case report. Theriogenology. 1997;48:353-9.

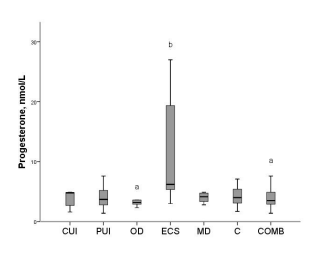
- [33] Batra S. Effect of oxytocin on calcium influx and efflux in the rat myometrium. Eur J Pharmacol.
  1986 Jan 14;120:57-61.
- [34] Popescu LM, Nutu O, Panoiu C. Oxytocin contracts the human uterus at term by inhibiting the
  myometrial Ca2+-extrusion pump. Biosci Rep. 1985;5:21-8.
- 438 [35] Hollinshead FK, Hanlon DW, Gilbert RO, Verstegen JP, Krekeler N, Volkmann DH. Calcium,
- parathyroid hormone, oxytocin and pH profiles in the whelping bitch. Theriogenology.
  2010;73:1276-83.
- 441 [36] Frehner BL, Reichler IM, Keller S, Goericke-Pesch S, Balogh O. Blood calcium, glucose and
- 442 haematology profiles of parturient bitches diagnosed with uterine inertia or obstructive dystocia.
- 443 Reprod Domest Anim 2018;53:680-7.
- 444 [37] Gogny A, Mallem Y, Destrumelle S, Thorin C, Desfontis JC, Gogny M et al. In vitro comparison
- 445 of myometrial contractility induced by aglepristone-oxytocin and aglepristone-PGF2alpha
- 446 combinations at different stages of the estrus cycle in the bitch. Theriogenology 2010;74:1531-8.
- 447
- 448

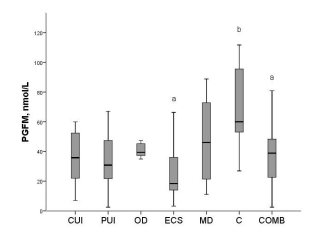
#### 449 Author contributions

- 450 TMT: design of the study, RNA isolation, real-time PCR, data and statistical evaluation, manuscript
- 451 writing. OV, MD, JT, TK: design of the study, data and statistical evaluation, manuscript editing. LS,
- 452 BM: RNA isolation, real-time PCR, manuscript editing.
- 453
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#### 455 **Conflicts of interest**

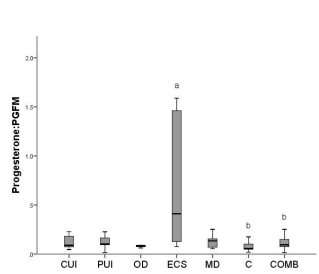
- 456 The authors have no conflicts of interest to declare.
- 457

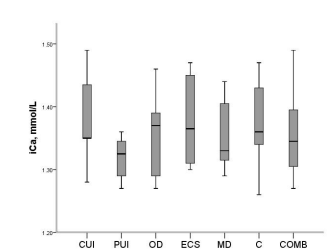




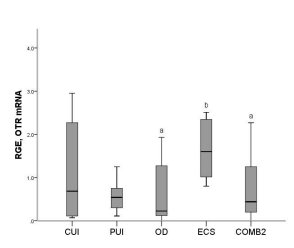


a)









d)

## Table 1. The individual and average data of 58 bitches divided in six groups. Parity includes the current parturition. Medication: 0, no medication; 1, calcium and oxytocin after blood sample. n/a: not available.

Group	Mean age, min-max (years)	Mean weight, min-max (kg)	Mean litter size, min- max	Breed	Parity	Gestation length (days from the last and first mating)	Gestation length (days from ovulation)	Duration of the first stage of parturition (hours)	Duration of the second stage of parturition before intervention (hours), discharge	Litter size	Number of puppies born before dystocia/by cesarean section	Medication
1.	4.1,	23.7,	6.4,	Border Collie	1	56-61	n/a	26	3, fetal fluids	4	0/4	1
Complete primary	2.3-6.2	4.7-35.0	1.0-14.0	Airedale Terrier	1	60-62	63	45	6, fetal fluids	7	0/7	0
uterine inertia				Karelian Bear Dog	3	68	n/a	n/a	n/a, green fluids	1	0/1	0
(CUI)				Dogo Argentino	1	65	n/a	29	6, green fluids	2	0/2	0
n = 7				Chinese Crested Dog	1	62	n/a	16	4, fetal fluids	6	0/6	1
				English Springer Sp.	2	60	n/a	48	6, fetal fluids	14	0/14	1
				Labrador Retriever	1	59	60	65	6, fetal fluids	11	0/11	0
2.	5.1,	29.2,	7.0,	Mixed	1	64-65	n/a	n/a	4	5	1/4	1
2. Partial primary	2.6-8.0	6.5-65.0	5.0-15.0	Labrador Retriever	2	60-62	62	5	4	8	5/3	1
uterine inertia	2.0-0.0	0.3-03.0	5.0-15.0	Dalmatian	3	63	63	6	4	6	1/5	0
(PUI)				Poodle, Standard	1	61-62	62-63	7	9.5	5	3/2	0
n = 13									5			1
11-10				Labrador Retriever	2	63	n/a	10	-	8	6/2	-
				Miniature Schnauzer	1	60-61	n/a	24	3.5	7	1/6	1
				Tibetan mastiff	2	59-61	62	24	4	5	4/1	1
				Mixed	1	62	n/a	27	n/a	1	1/6	1
				Border Collie	1	57-61	n/a	22	5.5	6	4/2	1
				Bullmastiff	2	56-57	57	65	n/a, fetal fluids	15	1/14	0
				Great Dane	1	62	n/a	26	6	6	2/4	1
				Bearded Collie	1	54-56	58	14	5.5	8	4/4	1
				Giant Schnauzer	2	58	60	12	10	5	2/3	1
<ol><li>Obstructive</li></ol>	5.1,	23.7,	5.0,	Cairn Terrier	3	64	66	n/a	4	1	0/1	0
dystocia	2.3-8.3	7.0-65.0	1.0-9.0	Border Collie	3	59-63	63	n/a	n/a, green fluids	7	0/7	1
(OD)				Saluki	2	61	64	18	4	3	0/3	0
n = 10				Cavalier King Ch. Sp.	1	59-60	62	24	4	7	0/7	0
				French Bulldog	1	61-63	n/a	9	6	3	0/3	0
				Dog de Bordeaux	2	65-67	n/a	11	4	9	0/9	1
				Smooth Collie	3	60-61	n/a	24	8	6	3/3	1
				Cavalier King Ch. Sp.	1	56-58	59	n/a	3	4	1/3	1
				Spanish Mastiff	1	62	62	11	n/a, fetal fluids	3	0/3	1
				Border Collie	3	60-61	61	24	3	7	0/7	1
4.	A 6	22.0	3.8,	Cavalier King Ch. Sp.	3	n/a	60	0	0	4	0/4	0
4. Elective cesarean			3.8, 1.0-10.0	Border Collie	3	64-65	64	0	0	2	0/2	0
section			1.0-10.0	St. Bernard	2	n/a	61	0	0	2	0/2	0
(ECS)												-
n = 11				Cavalier King Ch. Sp.	3	58-60	60	0	0	5	0/5	0
				Great Dane	2	62-63	62-63	0	0	1	0/1	0
				Shetland Sheepdog	4	60	n/a	0	0	1	0/1	0
				Boston Terrier	2	63-63	62	0	0	2	0/2	0
				Newfoundlander	1	58-64	n/a	0	0	10	0/10	0
				Cavalier King Ch. Sp.	3	59-60	59	0	0	8	0/8	0
				Boston Terrier	1	63-65	n/a	0	0	3	0/3	0
				Golden Retriever	1	64-66	65	0	0	4	0/4	0
5.	4.7,	13.3,	5.1,	Dachshund	1	58-60	n/a	1	2	4	0/-	1
Medically treated	2.6-6.5 4.5-20.0	4.5-20.0	4.5-20.0 2.0-8.0	Dachshund, miniature	1	63	n/a	15	5	6	0/-	1
dystocia, no cesarean section (MD) n = 8				Finnish Lapphund	1	n/a	n/a	n/a	7	5	4/-	1
				Jack Russell Terrier	2	58-61	n/a	n/a	4	2	0/-	1
				Australian Kelpie	1	60	n/a	15	7	8	6/-	1
				English Springer Sp.	3	n/a	n/a	12	10	4	2/-	1
				Finnish Lapphund	2	n/a	n/a	n/a	5	6	3/-	1
				Lapponian Herder	2	n/a	63	7	5	6	2/-	1
5	4.20,	10.9,	5.4,	Cavalier King Ch. Sp.	1	n/a	60	n/a	n/a	5	-	0
6. Control, normal parturition	4.20, 1.9-6.5	8.0-30.0	5.4, 4.0-10.0	Labrador Retriever	3	n/a	60	n/a	n/a	10	-	0
					4					4		0
(C)				Cavalier King Ch. Sp.		n/a	59	n/a	n/a		-	
1 = 9				Pug	2	n/a	61-62	n/a	n/a	5	-	0
				Pug	1	n/a	61	n/a	n/a	5	-	0
				Cavalier King Ch. Sp.	4	61	n/a	n/a	n/a	5	-	0
				Cavalier King Ch. Sp.	1	62	n/a	n/a	n/a	4	-	0
				Cavalier King Ch. Sp.	2	n/a	58-59	n/a	n/a	5	-	0
	1	1	1	Cavalier King Ch. Sp.	2	n/a	60	n/a	n/a	6	-	0