INVITATION

COMMITTEES

• Local Organising Committee
• Scientific Programme Committee
• International Advisory Board

ORAL COMMUNICATIONS

Monday, 27th June 2022

• SYMPOSIUM 1 O01 - O04
  The nonconformist: conceptus-maternal communication in the dog
• SYMPOSIUM 2 O05 - O06
  Devising new gonadotropins for the future... which strategies, which uses?
• SYMPOSIUM 3 O07 - O08
  Cloning and genome editing
• SYMPOSIUM 4 O09 - O10
  New aspects of corpus luteum regulation toward successful pregnancy
• WORKSHOP 5 O11 - O14
  Improving livestock production: beyond genetic again

Tuesday, 28th June 2022

• SYMPOSIUM 5 O15 - O16
  Reproduction in non-domestic and endangered species
• SYMPOSIUM 6 O17 - O20
  Immune regulation of oviduct/uterine function
• SYMPOSIUM 7 O21 - O23
  Effects of heat stress on reproduction: from conception to lactation
• SYMPOSIUM 8 O24 - O27
  Cryopreservation: freezing, vitrification, freezing drying

Wednesday, 29th June 2022

• SYMPOSIUM 9 O28 - O29
  New approaches in buffalo reproductive management
• SYMPOSIUM 11 O30 - O31
  New imaging systems for assessing gamete and embryo quality
• SYMPOSIUM 12 O32 - O33
  Novel insights on uterine immunology during pregnancy and disease
POSTERS

Monday, 27th June 2022

- Avian species reproduction        M01 - M05
- Bovine reproduction              M06 - M107
- Canine and feline reproduction   M109 - M127
- Control of estrous cycle         M128 - M133
- Cryobiology of gametes and embryos M134 - M156
- Reproductive pathologies         M157 - M165

Tuesday, 28th June 2022

- Buffalo reproduction              T01 - T09
- Male reproductive physiology     T10 - T18
- Metabolism and reproduction       T19 - T28
- Ovary and oocyte                 T29 - T64
- Pregnancy, placental function and parturition T66 - T76 and T168
- Reproduction exotic anim and wild species T77 - T99
- Reproduction in fish              T100 - T101
- Reproduction in sheep and goats   T102 - T139
- Testis                           T141
- Uterus                           T143 - T146
- Oviduct                          T147 - T153
- Embryogenesis in vitro           T154 - T167

Wednesday, 29th June 2022

- Camalid reproduction             W01 - W10
- Embryo development and differentiation W11 - W28
- Female reproductive physiology   W29 - W34 and W171
- Horse reproduction               W35 - W81
- Imaging methods in reproduction   W82 - W90
- Neuroendocrine control of reproduction W91 - W93
- Nutrition and reproduction       W94 - W102 and W170
- Other                            W106 - W111
- Pig reproduction                 W112 - W135
- Reproductive system diseases     W136 - W141
- Spermatology and sperm quality   W142 - W169

ABSTRACTS WORKSHOPS

25

SPONSORS & PARTNERS

250

287
GENE EXPRESSION PATTERNS IN UTERUS AND OVIDUCT DURING THE PREOVULATORY PERIOD IN Ewes

F. Cuadro 1, V. De Brun 1, C. Brochado 2, M. Souza 2, C. García Pintos 1, R. Nuñez-Olivera 1, C. Menezes 1, G. Gastal 1, A. Menchaca 1,
1Facultad de Veterinaria, Universidad de la Republica, Uruguay
2Fundación IRAUy, Uruguay
3Instituto Nacional de Investigación Agropecuaria -INIA, Uruguay
4Instituto Nacional de Investigación Agropecuaria -INIA, Uruguay; Fundación IRAUy, Uruguay

BACKGROUND-AIM
The uterine and oviduct environment soon after ovulation plays a major role on fertilization and embryo development. However, the knowledge about the uterine/oviductal environment during the preovulatory period is not fully understood. This study investigated the gene expression oviduct (ampulla and isthmus) and uterus (horns) during the preovulatory period in multiparous ewes.

METHODS
Multiparous Corriedale ewes (n=10) were synchronized during breeding season with intravaginal sponges and equine chorionic gonadotrophin (Menchaca and Rubianes 2004). Estrus was detected after sponge removal by using an androgenized ram. The ewes were euthanized (thiopental sodium Tiobarbital and T-61) from 36 to 48 h after sponge removal (i.e., before ovulation), the reproductive tract was dissected and the oviducts (ampulla and isthmus) and a portion of the upper third of the uterine horns were sampled to be stored at -80°C. Gene expression of PGR (nuclear progesterone receptor), ESR1 (estriadiol receptor alpha), IGF2 (Insulin-like Growth Factor 2), PTGS2 (Prostaglandin-Endoperoxide Synthase 2), Serpina14 (Uterine milk protein), SOD1 and SOD2 (Superoxide Dismutases 1 and 2) as housekeeping genes (BACT (Beta-Actin), and HPRT (Hypoxanthine-Guanine Phosphoribosyltransferase) were determined by real time PCR. All variables were analyzed using a mixed model procedure in SAS.

RESULTS
The expression of most of the genes (PGR, ESR1, IGF2, PTGS2, SOD1, SOD2) was greater (P<0.001) in the oviduct compared to the uterus tissue. Only the Serpina14 gene had higher (P<0.001) RNA expression in the uterus (62.3±9.0) compared to both ampulla (1.9±9.7) and isthmus (18.1±9.0). Regarding the oviduct tissue, the genes (IGF2, PTGS2, SOD1 and SOD2 had greater (P<0.001) RNA expression in the isthmus (2.1±0.2; 8.6±1.4; 2.0±0.2; 2.1±0.1) compared to the ampulla (0.1±0.2; 1.3±1.2; 0.9±0.2; 0.7±0.1), respectively.

CONCLUSIONS
During the preovulatory period, the isthmus segment seems to have a greater gene expression activity of certain genes compared to the other regions, likely improving the environment for the cumulus oocyte complex transport, sperm migration for fertilization, and early embryo passage to the uterus.

BISPHENOL S ADMINISTERED TO PREGNANT EWES WITH CONTRASTED METABOLIC STATUS IMPAIRED THE OVARIAN FOLLICULAR DEVELOPMENT OF FETUSES AND LAMBS
M. Lebachelor; De La Rivière 2, O. Téteau 3, P. Jarrier-Gaillard 2, O. Lasserre 1, P. Papillier 2, A. Binet 3, A. Desmarchais 2, S. Ellis 2, V. Maillard 2, N. Picard-Hagen 5, C. Vignault 4
1 UEPAO, INRAE, 37380, Nouzilly, France
2CNRS, IFCE, INRAE, Université de Tours, PRC, F-37380, Nouzilly, France
3CNRS, IFCE, INRAE, Université de Tours, PRC, F-37380, Nouzilly, France; Service de Chirurgie pédiatrique viscérale, urologique, plastique et brûlés, CHRU de Tours, 37000 Tours, France
4CNRS, IFCE, INRAE, Université de Tours, PRC, F-37380, Nouzilly, France; Service de Médecine et Biologie de la Reproduction, CHRU de Tours, 37000 Tours, France
5ToxAlim (Research Centre in Food Toxicology), Université de Toulouse, INRA, ENV7, INP-Purpan, UPS, Toulouse, France

BACKGROUND-AIM
The size of the ovarian reserve, which is established by the number of primordial follicles, is defined before birth in ovine. The prenatal period is a critical window of sensitivity to environmental factors. Maternal metabolic status during gestation can influence fetal programming, and affect the ovarian reserve. Bisphenol A (BPA), a plasticizer used in food packaging, has deleterious effects on fetal folliculogenesis. It has been banned from the food industry and mainly replaced by bisphenol S (BPS). The objective of this study was to determine the effects of BPS 50 µg/kg/d in utero exposure on female fetuses and lambs from ewes with contrasted metabolic status on follicular population by ovarian histology.

METHODS
This study was divided into two experiments performed on pregnant ewes exhibiting a contrasted metabolic status (lean versus well-fed). First, pregnant ewes were exposed to BPS by daily subcutaneous injection for 3 months, to analyze the ovarian development of the fetuses at 130 days of gestation. Second, pregnant ewes were exposed daily to BPS through food for 3 months and the lambs were then monitored up to 4 month-old (pre-puberty). Follicular population was characterized through follicle classification and counting on ovarian sections.

RESULTS
In utero BPS exposure led to an increase in the number of pre-antral (p = 0.005) and antral follicles (p < 0.001) in fetuses and conversely to a decrease in 4 month-old lambs (p = 0.007). Besides, a significant interaction between maternal metabolic status and BPS exposure during gestation was reported for the number of primordial (p = 0.019), pre-antral (p = 0.003) and antral (p <0.001) follicles of fetuses. In offspring of fat mothers, the plasma anti-Mullerian hormone decreased in 1 and 2 month-old lambs (p < 0.001). In offspring of well-fed mothers, the body weight of the female fetuses increased (p < 0.001). Moreover, the body weight of female lambs that were in utero exposed to BPS increased compared to control (p = 0.046).

CONCLUSIONS
In conclusion, bisphenol exposure during gestation had deleterious consequences on the body and folliculogenesis of the offspring. BPS effects also vary according to the maternal metabolic status. Further research will be needed to determine if these alterations will have deleterious effects on the ovarian reserve and on the reproductive function in adulthood.
W31
BISPHENOL S IMPAIRED HUMAN AND OVINE GRANULOSA CELL STEROIDOGENESIS

M. Lebachelier De La Rivière 1, O. Téteau 1, S. Amar 1, A. Binet 1, A. Desmarchais 1, S. Elvis 1, F. Guérif 1, M. Jaubert 1, V. Maillard 2, P. Papillier 3, C. Vignault 1

1 CHRU de Tours, 37000 Tours, France
2 CNRS, IFCE, INRAE, Université de Tours, PRC, F-37380, Nouzilly, France

BACKGROUND-AIM
Human being is exposed to bisphenol A (BPA), a plasticizer used in food containers, through diet. Granulosa cell (GC) play a fundamental role for oocyte growth and maturation. BPA impaired GC steroidogenesis and has been classified as an endocrine disruptor (ED). BPA was banned from the food industry and replaced by structural analogues, particularly bisphenol S [BPS]. Given that the ewe has similar follicle kinetic compared to women, the evaluation of the effects of BPS on the ovine would allow to investigate whether the ewe is a suitable model to study ED effects in human. Our objective was therefore to assess BPS effects, and its mechanisms of action, on both human and ovine GC.

METHODS
After puncture of follicles from women undergoing in vitro fertilization or ewe ovaries (slaughterhouses), GC were collected, purified and treated in complemented serum-free Mc Coy Medium, with BPS (1 µM, 10 µM or 50 µM) for 48-h. We analysed GC viability (adenylate kinase activity assay) and proliferation (incorporation of BrDU), secretion of oestradiol and progesterone (ELISA essay), expression of steroidogenic enzymes (Western Blot), hormonal receptor gene expression (quantitative RT-PCR) and MAPK3/1 signalling pathway, as it is involved in both survival and MAPK3/1 phosphorylation.

RESULTS
BPS 10 µM significantly decreased progesterone secretion of both human (16 %; p = 0.0059) and ovine (22 %; p = 0.0402) GC compared to control. BPS 50 µM significantly decreased oestradiol secretion in human GC (46%; p < 0.0001), while it was significantly increased with BPS 10 µM in ovine GC (198%; p = 0.0082). In both ovine and human GC, BPS did not affect cell viability, proliferation, protein expression of steroidogenic enzymes, PR gene expression, or MAPK3/1 phosphorylation.

CONCLUSIONS
Thus, the effects of BPS on GC appear harmful and similar between human and ovine; except for oestradiol secretion, likely due to the ovulatory stage of GC collection for women. Our data confirmed that the ewe is a relevant model, in term of sensitivity, for the human female reproduction. Ewe GC will allow us to further study the detailed mechanisms of action of BPS on female reproduction.

W33
INTERRELATIONSHIP BETWEEN INTRAFOLLICULAR DOPAMINE AND ESTRADIOL CONCENTRATIONS IN CYCLING MARES

K. Satué Ambrojo 1, M. Marcilla Corzano 2, E. Fazio 1, P. Medica 1

1 Departament of Veterinary Medicine-University of Messina
2 Faculty of Veterinary-Cardenal Herrera CEU University

BACKGROUND-AIM
Follicular hormonal changes associated with oocyte maturation can be modulated locally by monoamines (Bódis et al., 1993). Animal models, ovariectomy decreases the release of dopamine (DA), that can be reversed with subsequent treatment of estradiol (E2) (Thompson and Moss, 1994). This is consistent with the elevation of DA presumably in relation to high levels of E2 during the estrous cycle. This response suggests that DA in FF could play an important role in the mechanism of ovulation in humans (Bódis et al., 1994). In mares, the effects of DA on follicular growth may be mediated by FSH secretion (King et al., 2008). However, the relationship between DA and E2 concentrations has not yet been investigated. The objective of this study was to investigate the interrelationship between DA and E2 in follicular fluid (FF) in different categories of follicular sizes in cycling mares.

METHODS
A total of 60 samples of FF by aspiration of slaughterhouse ovarian follicles of 30 clinically healthy mares aged 6.6 ± 1.3 years are evaluated. FF samples were classified according to diameter as: small (20-30 mm), medium (31-40 mm) and large (≥ 41 mm) size. Intrafollicular DA (pg/mL) concentrations were measured with a radioimmunoassay technique (Labor Diagnostika Nord GmbH, Nordhorn, Germany). The concentrations of E2 (ng/mL) in FF are determined by a competitive enzyme-linked immunosorbent assay (Estradiol sensitive ELISA Demeditec DE4399) validated specifically for FF in the equine species.

RESULTS
The intrafollicular concentrations of E2 in small follicle were significantly lower (652.9 ± 241.3 pg/mL) than in medium (1,498.9±205.8 pg/mL) and in large (1,692.7±146.8 pg/mL) follicle sizes (P < 0.05). In FF, the concentrations of DA in medium size follicles were significantly higher (707.9±360.7 pg/mL) than in small (396.2±113.8 ng/mL) and large (351.2±132.5 ng/mL) follicle sizes (P < 0.05). DA and E2 concentrations are positively correlated (r=0.66; P < 0.05).

CONCLUSIONS
These results suggest that, as is the case at the circulating level, ovarian estrogens are involved in dopaminergic activity in follicular development, or that catecholamines modulate the stimulating effect of estradiol in follicular fluid.