Embryonic diapause modulates ovarian follicular dynamics and oocyte quality in the European roe deer (*Capreolus capreolus*)

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Abstract

The phenomenon of embryonic diapause has been reported to possibly affect ovarian follicular development. The aim of the current study was to identify the impact of the embryonic diapause period on follicle number and oocyte quality in the European roe deer (*Capreolus capreolus*). In this species, the embryo undergoes five months of developmental delay. We have determined the antral follicle count (AFC), oocyte morphological quality, oocyte meiotic resumption capacity and the transcriptome profile in immature (GV) and *in vitro* matured (MII) oocytes obtained during embryonic diapause and after embryo reactivation. A drop in AFC, number of oocytes suitable for *in vitro* maturation and oocytes reaching the MII stage was observed after embryonic reactivation. A large impact of the diapause period was evident when MII oocytes were compared against GV oocytes using an RNA-seq approach. A total of 1028 genes significantly differed between MII and GV oocytes during diapause and 1795 genes after reactivation. These genes included *PDE3A* and *SOD2*, known for being involved in oocyte meiotic arrest and oxidative stress. In addition, tumour-related genes included *NDRG3*, *NCSTN* and *UGDH*, which have been associated with cell proliferation, migration and metabolism, were found. Comparing diapause versus reactivation, differential transcript abundance in both GV and MII oocytes was observed (44 and 32 differentially expressed genes, respectively). Our results suggest that the embryonic diapause and reactivation periods modulate follicular dynamics, influencing the oocyte towards lower quality after reactivation. This is may be related to increased apoptosis, abnormal mitochondria function/structure and metabolic dysfunction in the maturing oocyte.

Introduction

Embryonic developmental delay, termed embryonic diapause, occurs in more than 130 mammalian species. However, this process is still enigmatic and not fully understood to date [1, 2]. In the European Roe Deer (*Capreolus capreolus*), the only known ungulate that exhibits this phenomenon, the rut starts at the end of July/beginning of August. The female roe deer is considered a seasonally monoestrus breeder [3], with an estrus duration of approximately 36h [4]. The start of the breeding season and further ovulation are modulated by melatonin. In animals receiving melatonin implants in mid-April, a sharp decreased in prolactin and an advancement of the ovulatory period by 72 days, indicated by the increment of blood progesterone (P4) concentrations, has been reported [5].
Furthermore, the exposure to artificial long days from the summer solstice delayed ovulation by 46 days [5]. In the absence of a male, peripheral P4 concentrations rise after spontaneous ovulation and corpus luteum (CL) formation occurs. Thereupon, P4 levels are maintain until March [6, 7]. Furthermore, oxytocin (OXT) has been found in the CL during the breeding season and the CL was able to secrete OXT in response to the administration of the PGF2α analogue cloprostenol [3]. However, administration of OXT did not stimulate uterine secretion of PGF2α [3], although OXT receptors have been found in the uterus during the breeding season and diapause, thus differing from other ruminants [3]. After fertilization, the roe deer embryos develop to the blastocyst stage and hatch from the zona pellucida. Afterwards, blastocysts enter a phase of developmental delay or obligate embryonic diapause [8]. Embryonic reactivation and implantation only take place at the end of December/beginning of January [8]. Peripheral plasma/serum oestradiol concentrations remain low during diapause. A following increase after embryo elongation has been reported [9]. Furthermore, basal prolactin (PRL) concentrations during diapause displaying a substantial increase after implantation have been observed. In contrast, peripheral and luteal P4 concentrations remain constant throughout diapause, elongation and implantation [9]. Additional reports have suggested a rise in P4 levels between January and March prior to a reported decrease in March in pregnant does [3]. Furthermore, no differences in CL mass in does carrying a blastocyst and non-pregnant does have been reported [3]. Up to date, most efforts on the field of embryonic diapause in mammals have focused on studying the maternal endocrinology, embryonic morphology, the uterine environment as well as their interaction during the developmental delay and reactivation processes. Little is known about the female gonads and oocyte characteristics in roe deer.

Interestingly, some of the potential diapause regulatory factors described in mice, mink and invertebrates (reviewed by [2]) are also involved in follicle/oocyte mitotic/meiotic arrest maintenance and activation. These comprise the mammalian target of rapamycin (mTOR) [10, 11], the MYC proto-oncogene, bHLH transcription factor (MYC) [12] and the forkhead box class O family member 3 (FOXO3) [13, 14]. Additionally, the same factors have also been related to the aging process (reviewed by [15]).

Recently, a decrease in ovarian follicle number following embryonic reactivation was reported in the roe deer [16]. A potential impact of diapause on follicular physiology was likewise suggested [16]. We hypothesise that diapause could play a role in modifying the ovarian activity and therefore oocyte quality. Hence, we studied the ovarian characteristics and oocyte quality during the embryonic diapause and reactivation period in the roe deer by combining morphological observations, cellular and transcriptomics analyses to identify whether these developmental stages influence the follicle population, the oocyte mRNA abundance profiles and the oocyte capacity to resume meiosis.

Materials and methods

All reagents of analytical grade were supplied by Sigma–Aldrich Co. (St. Louis, USA) unless otherwise stated. Sample collection did not require ethical approval, since it was performed during regular hunts and none of the animals was culled for scientific purposes. In the present study, the status of the retrieved ovaries and their respective oocytes was allocated to one of two different periods, according to the corresponding embryo stage. The periods were determined as “Diapause (D)” (embryo in embryonic diapause characterized by a round morphology) and “Reactivation (R)” (embryo elongated or implanted). To determine the reproductive status of the females and the embryonic developmental stage, the uterus horns were either flushed or opened by cutting (when implantation was observed) for embryo retrieval. Since non-pregnant does from hunted animals cannot be confirmed, females in which an embryo was not found were excluded. Therefore, only does with a known embryonic developmental stage were included in the analyses.
**Roe deer ovarian characteristics**

Roe deer ovaries were retrieved in northern Switzerland and southern Germany between October 2016 - January 2017 and October 2017 - January 2018. A total of 141 hunted animals were considered for the analyses. The length and width of each ovary was determined to estimate the ovarian surface area (mm$^2$) using the formula for an ellipse (length $\times$ width $\times \pi/4$) [17]. Follicles visible on the ovary surface were counted and classified according to their size in three different groups: small <1 mm, medium 1.1-4.9 mm and big >5 mm. Corpora lutea (CL) were documented and included for ovary classification, namely no presence of a CL (0 CL), one CL present (1 CL), or two or more CL present (≥2 CL), respectively.

**Roe deer oocyte retrieval**

After ovarian morphological features were documented, both ovaries from the same female were sliced in BO-OPU medium (IVF Bioscience, IVF Limited T/A IVF Bioscience, Bickland Industrial Park, Falmouth, Cornwall, United Kingdom) at 35°C. Oocytes were collected and classified in BO-Wash medium (IVF Bioscience) according to six (I-VI) morphological categories [18]: Category I: Oocytes with more than four layers of compact cumulus cells and a regular cytoplasm. Category II: Oocytes with three to four layers of compact cumulus cells, regular cytoplasm or lightly granulated. Category III: Oocytes with one to two layers of cumulus of corona radiata, cytoplasm with irregular appearance, such as dark areas. Category IV: Denuded oocytes. Category V: Oocytes with expanded cumulus. Category VI: Degenerated oocytes. Oocytes classification was performed randomly for 15 animals for each embryo developmental group/phase.

**In vitro oocyte maturation (IVM)**

After classification, roe deer oocytes were washed in 3 ml maturation medium (BO-HEPES IVM, IVF Bioscience). Thereafter, oocytes were transported to the lab at 38.5°C in 1 ml glass tubes containing 500 µl of the maturation medium and a maximum of 20 oocytes per tube. Oocytes were cultured for 22 - 26 h at 38.5°C and only oocytes from categories I, II and III were used for IVM.

**Processing of immature and in vitro mature oocytes for analyses**

Immature (GV, germinal vesicle stage) and in vitro matured (MII, metaphase II stage) roe deer oocytes were denuded by incubation in Dulbecco’s phosphate buffer saline (PBS, AppliChem GmbH, Darmstadt, Germany), supplemented with 0.1 % hyaluronidase and 1mg/ml BSA (Fraction V) for 3 min at 38.5° and after vortexed for 1 min at 1400 rpm. The remaining cumulus cells were removed by gentle pipetting. Maturation was confirmed by the presence of the first polar body in the perivitelline space (MII). Single denuded immature and matured oocytes were snap frozen at -80°C, in PBS enriched with 0.1% polyvinyl alcohol (PVA) (PBS-PVA solution) for further analysis.

**Evaluation of oocyte maturation**

After maturation, oocytes were denuded as described above, fixed in PBS-PVA solution containing 10% paraformaldehyde, and maintained at 4°C until evaluation. Prior to analysis, oocytes were washed twice in PBS-PVA and stained with a PBS-PVA solution containing 0.2 mg/ml propidium iodide and 0.4% triton for five minutes. For nuclear evaluation, oocytes were mounted on glass slides and covered with a cover slip. Oocyte evaluation was immediately performed using a fluorescence microscope (Axioplan 2, Carl Zeiss AG, Oberkochen, Germany) at 400x magnification. The number of oocytes at germinal vesicle (GV), germinal vesicle breakdown (GVBD), metaphase I (MI), anaphase I (AI), telophase I (TI), metaphase II (MII) and non-evaluable stages (NE) was determined. Oocyte nuclear stage was evaluated in 11 and 8 randomly selected animals for the diapause and reactivation periods, respectively.
Histology of the ovary
Ovaries from five randomly selected animals in each group (D and R) were cut at middle of the sagittal plane in two halves and fixed using 10% formalin for 24 hours. Thereafter, ovaries were washed in PBS for seven days (PBS was changed daily). Subsequently, samples were dehydrated in graded ethanol series (30%, 50% and 70%, 24 h each) and finally embedded in paraffin. For histological analyses, ovarian slices were cut at a thickness of 2-3 μm on a rotary microtome in the sagittal plane, and processed in routine hematoxylin and eosin staining.

Oocyte mRNA library preparation
Pools of oocytes were processed using the Smart-seq2 single cell protocol for full-length cDNA and library preparation as previously described [19]. Single oocytes were grouped into two pools of 10 immature and mature oocytes for both, diapause and reactivation groups, for the analyses (at least four donors/pool, eight libraries in total). Lysis buffer containing free dNTPs and tailed oligo-dT oligonucleotides (30-nt-poly-dT stretch and 25-nt universal 5’ anchor sequence) was added to each pool of oocytes. Afterwards, reverse transcription (RT) was performed at 42°C for 90 min, followed by 50°C for 2 min. Thereafter, temperature was lowered to 42°C for 2 min. This cycle was repeated ten times. Template switching oligos were added to the RT. After RT, 18 cycles preamplification were performed using KAPA HiFi HotStart Ready mix (KAPA Biosystems, Wilmington, Massachusetts, USA) and IS PCR primers. High Sensitivity Bioanalyzer chip (Agilent, Santa Clara, California, USA) was used to assess the quality of the cDNA samples. Tagmentation was carried out using the Illumina Nextera XT DNA sample preparation kit (Illumina Inc, San Diego, California, USA) at 55°C for 5 min. Afterwards, 12 cycles PCR amplification were performed, followed by clean up and concentration measurement. Libraries were pooled and sequenced using a HiSeq 4000 platform (Illumina).

Bioinformatics and data analyses
The obtained sequence reads (FastQ files) were analysed with a locally installed version of Galaxy (https://doi.org/10.1093/nar/gky379). The adapter sequence was clipped with Trim Galore (visited on 2019/04/24) [20] and quality control was done using FastQC. Sequences were mapped with Salmon [21] against a roe deer transcriptome containing 63,935 transcripts (own unpublished data) and annotated against human, bovine and a number of other RefSeq transcripts obtained from NCBI. The read count table was filtered by counts per million (CPM) cut-off to remove transcripts with neglectable read counts and used for statistical analysis with ABSSeq to identify differentially expressed genes (DEG) [22] (parameters: afold = TRUE and normalization = “qtotal”). Because of some large differences in excounts numbers (normalized count data) between biological replicates from the same oocyte stage and period of time (in both the D and R group), an additional filtering using <60 percentage of change and a standard deviation < 13 was performed. Four different comparisons were performed: i) GV oocytes from the embryo developmental diapause period vs. GV oocytes from the embryo reactivation period (GV_D_vs_GV_R); ii) MIi oocytes from the embryo developmental diapause period vs. MIi oocytes from the embryo developmental reactivation period (MIi_D_vs_MII_R); iii) GV vs. MIi oocytes from the embryo developmental diapause period (GV_D_vs_MIi_D); iv) GV vs. MIi oocytes from the embryo developmental reactivation period (GV_R_vs_MIi_R). To determine differentially expressed genes (DEG), the adjusted p-value for the analyses was set to <1% and a log2 fold change (FC) > 1 / < 1 was applied for all comparisons as a cut-off. Hierarchical cluster analysis was performed for DEGs using MultiExperiment Viewer (MeV). Venn diagrams were performed using jvenn interactive Venn diagram viewer [23]. The obtained DEG were uploaded to the Ingenuity Pathway Analysis software (IPA: Qiagen, Hilden, Germany) to identify enrichment for Canonical pathways (CP) and Disease and Functions (DF). The level of significance for enrichment in the IPA output is assigned by a p-value in the uploaded data set compared with the known molecules in the database for a particular CP and DF entry. Only entries...
with a p-value < 0.01 were included for the analyses. Furthermore, IPA calculates a z-score to infer the activation state: activation/increase or inhibition/decrease. A z-score > 2 was considered statistically significant.

Statistical analysis of oocyte and ovary data
Data from ovary size and total and suitable oocyte numbers were evaluated using an analysis of variance model. Follicle numbers and oocyte maturation rates were compared by Kruskal-Wallis test. These statistical tests were performed in R, version 3.6.1 [24] and the significance was set at P < 0.05.

Results and discussion
Here, we report for the first time, morphological and molecular oocyte characteristics during the embryonic diapause and reactivation periods in the roe deer. Additionally, we present ovarian characteristics including the follicular growth and ovary size. We have applied next generation transcriptome sequencing to characterize the effects of the embryo developmental stage on the transcriptome relating to GV and MII-staged oocytes during the embryonic diapause and after the reactivation phase.

Ovarian characteristics
The average surface area of the 282 collected ovaries between October and January did not change between the diapause (D) and embryo reactivation (R) group (p>0.05) (see Table 1). As expected, the presence of 1 or 2 CL increased the surface area. The total number of follicles and the number of small follicles decreased after reactivation (p<0.05) (see Figure 1a-b). The number of large and medium sized follicles did not differ between D and R (p>0.05) (see Figure 1b). The ovarian histology confirmed the observed findings regarding different populations of follicles, including intact antral and atretic follicles in both D and R. Furthermore, in both D and R, there were CL not fully and fully integrated in the ovarian structures and covered by the surface epithelium (see Figure 2). Although the number of antral follicles appeared higher in ovaries displaying not fully integrated CL, the number of atretic follicles seemed higher in those ovaries with fully integrated CL, regardless of the experimental group (see Figure 2).

It has been reported that during early pregnancy (between days 6 - 34 post oestrus), bovine ovaries bearing a CL show decreased follicular development as a local effect, meanwhile follicular growth is sustained in the contralateral ovary [25]. However, differences in follicle number in non-pregnant vs. pregnant bovine donors used for ovum pick up have not been reported [26]. In the ewe, a seasonal species, a suppression of antral follicle growth during early gestation remained also restricted to CL-bearing ovaries (days 10 – 30 post oestrus [27]) as in the cow. In the red deer, using ultrasound examination, seasonality (anoestrous) has negatively affected the follicle size and longevity. Additionally, little or no luteal activity was observed [28]. In the roe deer using ultrasonography, antral follicular development during diapause in both pregnant and non-pregnant females has been reported [29]. However, the authors did not study the differences in follicular populations. Previous studies from ovarian histological cuts in the roe deer (from August to January hunting season) reported the presence of follicles at all stages of development and atretic follicles present in the majority of the ovaries [30]. Interestingly, mitosis was observed in some follicles, meaning active development [30]. However, the authors did not distinguish diapause from reactivation. In our study, we have observed a drop in antral follicle number in the roe deer potentially related to embryonic reactivation. Interestingly, a comparable pattern in CL number distribution and ovary size in both developmental periods (diapause vs. reactivation) was also evident. Furthermore, previous studies have revealed no differences in the CL mass in non-pregnant and pregnant roe deer females [3], suggesting that the CL presence influenced the
follicular growth in a similar manner and that the observed reduction in follicle count might be due to the reactivation period. Although, differences in estradiol-17β (E2) and progesterone (P4) levels have not been found in the roe deer between embryonic diapause and reactivation [16], the possible effects of the CL ovarian integration and its functionality deserves further research.

Table 1. Effects of the embryonic diapause and reactivation on the ovary area in the roe deer.

| Number of CL per ovary | Diapause | Reactivation |
|------------------------|----------|--------------|
|                        | Number of ovaries | Area (mm²) | Number of ovaries | Area (mm²) |
| 0 CL                   | 48       | 67.6 ± 3.7   | 15               | 59.2 ± 4.5 |
| 1 CL                   | 121      | 76.0 ± 2.2   | 37               | 72.8 ± 4.0 |
| ≥2 CL                  | 45       | 95.5 ± 5.2   | 16               | 89.3 ± 4.9 |

Data are the mean ± SEM (p>0.05).

**Oocyte quality and maturation rates**

Since oocyte quality depends on the follicular environment, we have further analysed the oocyte morphological quality and its ability to resume meiosis. Oocyte quality was evaluated in 15 does at D and R. The total number of oocytes did not differ between developmental stages (784 and 711 oocytes from the embryonic diapause and reactivation, respectively, p>0.05). The total number of oocytes per animal was 52.2 ± 5.0 vs. 47.4 ± 8.0 for D and R, respectively (p>0.05). When ovary slicing is performed, oocytes are retrieved from follicles differing in size, which can lead to the similar observed oocyte numbers. The total average number of oocytes suitable for IVM (Qualities I-III) per animal was 27.7 ± 3.0 for D and 12.8 ± 2.0 for R (p<0.05). Maturation rates were randomly analysed in oocytes from 11 and 8 animals for the diapause and reactivation group, respectively. A total of 154 and 118 oocytes were fixed and stained for nuclear status after IVM for the two evaluated periods, respectively. The proportion of oocytes in GVBD and MI stages was significantly higher for the reactivation group (see Figure 3, p<0.05). No differences were observed in the proportion of GV, AI, TI, MII and degenerated oocytes (p>0.05) (see Figure 3). In cattle, no differences have been reported regarding the number of follicle number, of collected oocytes, of the cleavage rates and of blastocyst formation, when ovum pick up has been performed in pregnant and non-pregnant donors [26]. Thus, the lower number in oocytes suitable for IVM and reaching the MII stage might suggest an increased follicular atresia after embryonic reactivation.
Figure 1. Impact of the embryonic diapause period on the ovarian follicle number in the roe deer. Ovaries were retrieved between October - January. a Total number of antral follicles present during embryonic diapause and reactivation. b Antral follicle number for big: ≥5 mm, medium: 1.1-4.9 mm and small: ≤ 1 mm follicle sizes. Each dot corresponds to one animal, including both ovaries. Data were compared by Kruskal-Wallis test (p<0.05).
Figure 2. Histological sections of roe deer ovaries during embryonic diapause and after embryonic reactivation. Samples were stained with hematoxylin and eosin. Solid arrows indicate atretic follicles. CL, corpus luteum.
Figure 3. Effects of the embryonic developmental stage (determined after embryo retrieval by uterine flushing) on the meiotic resumption capacity of roe deer oocytes. The developmental state was assessed 22-26 hours post IVM. Bars represent the percentages calculated using the total number of oocytes per treatment. *Represents statistical significance for the respective meiotic status between treatments (p<0.05). NE, non-evaluable; GV, germinal vesicle stage; GVBD, germinal vesicle breakdown; MI, metaphase I; AI, anaphase I; TI, Telophase I; MII, metaphase II; Deg, Degenerated.

Transcriptome analysis of roe deer oocytes
We have performed RNA-sequencing on freshly collected GV and in vitro matured oocytes (MII). A total of 220.2 million raw reads were obtained from the eight libraries. Low-quality reads and PCR duplicates were removed and a total of 72.8 million clean reads (20.6 million for GV diapause, 19.4 million for GV reactivation, 16.0 million for MII diapause and 16.7 million for MII reactivation) were further processed for the analyses. A total of 7995 genes were identified and annotated. Additionally, 4997 and 4868 transcripts for GV and MII oocytes were identified but could not be annotated and were not included for further analyses, limiting the current results. Most of the identified genes (6239, 78 %) were detected in both GV and MII oocytes. Moreover, some genes were only present in GV (778, 9.7 %) or MII (397, 4.9 %) oocytes (see Figure 4). The results from MII oocytes might not completely resemble the portrait of in vivo matured oocytes, because it has been demonstrated that the expression pattern in these two types of oocytes can differ [31].

The unsupervised clustering of the samples by multiple dimension scaling (MDS) analysis showed a tight grouping according to the oocyte stage and the embryo developmental period (Figure 5). For the dimension 1, an evident separation of the GV oocyte from MII oocytes was observed. The dimension 2 displayed the separation of the oocyte samples according to the diapause and reactivation period (see Figure 5). Additionally, a hierarchical clustering analysis of the differentially expressed genes (DEG) was performed for the four different comparisons (see Figure 6). For oocytes in the GV stage, 18 genes were found to be up-regulated (40.9 % of DEG) and 26 down-regulated (59.1 %) during diapause (see Figure 6a). For MII oocytes, 14 genes were up-regulated (43.7 %) and 18 down-regulated (56.2 %) during diapause (see Figure 6b). When metaphase II oocytes were compared to GV oocytes, 81 (7.9 %) and 220 (12.2 %) genes were up-regulated and 947 (92.1%) and 1’575 (87.7%) were down-regulated during the diapause and reactivation periods,
respectively (Figure 6c-d, and Bernal Supplementary Tables 1 and 2
https://www.diapause2019.ethz.ch/abstract-submission/).

Figure 4. Venn diagram displaying the overlaps of detectable genes. GV, Germinal vesicle; MII, metaphase II; D, diapause; R, reactivation

Figure 5. Multidimensional scaling plot for all identified transcripts based on pairwise fold changes between samples. GV, germinal vesicle; MII, metaphase II; D, diapause; R, reactivation; 1, sample 1; 2, sample 2.
Differentially expressed transcripts in immature and mature oocytes

The potential effects of the embryonic stage (D vs R) on the oocyte transcriptome were evaluated. Only few genes were found to be differentially expressed in GV and MII oocytes between the two evaluated embryonic periods (GV: 44, MII: 32) (see Figure 7, Table 2-3). From the total number of DEG, more than 700 genes were present in more than one of the evaluated comparisons (overlapping sector of the Venn diagram) (see Figure 7). However, an important number of DEG were found to be only present in the non-overlapping sector of the Venn diagram for each comparison (see Figure 7). When the oocyte profiles after maturation were evaluated against GV oocytes, a lower number of DEG in MII oocytes from the diapause group (GV_D_vs_MII_D) was observed (1’028 and 1’795 for D and R, respectively). As the culture conditions were identical for both groups of oocytes, the observed changes are most likely a result of intrinsic anomalies of the oocytes, probably due to the embryonic developmental stage and not due to the maturation system. However, the effects of hormonal profiles, seasonality or time of conception cannot be excluded. Oocytes at the germinal vesicle stage are transcriptionally active and can replace degraded transcripts [32]. Nevertheless, during maturation, oocytes undergo selective degradation of transcripts [33] and MII oocytes become quiescent, while they remain able to use the stored transcripts for protein synthesis. This transcriptional activity pattern could cover-up early or increased consumption of some transcripts, which might be only detectable after the oocyte reaches the MII stage. An increased number of DEG in MII relative to GV oocytes has likewise been reported in oocytes from advanced-age women related to younger-age women, which was a result of a stress-induced usage of transcripts in the aged oocytes [34]. This might indicate that roe deer oocytes after embryonic reactivation experience additional cellular stress, comparable to aging-related processes. This hypothesis needs further analyses for validation.
Some of the identified DEG in GV and MII oocytes during diapause were related to cell cycle, oxidative stress, metabolism and apoptosis. A selection of up-regulated DEG in GV and/or MII included important signalling-related genes such as the N-myc downstream-regulated gene 3 (NDRG3), involved in cell proliferation, apoptosis and differentiation and up-regulated in malignant tumours [35]. Furthermore, Nicastrin (NCSTN) as a crucial component of the γ-secretase (GS) enzyme, plays a role in developmental signalling via the transcription factor Notch. Its expression has been involved in propagation and invasion of tumours [36]. SOD2 [37] and TXNIP [38] are modulators of cellular redox signalling protecting the cells from oxidative stress. The adiponectin receptor 2 (ADIPORE2) has been found to be down-regulated in GV oocytes of 32-week-old mice (aged) [39]. The activation of PDE3A, an oocyte specific phosphodiesterase, which remains inactive during the oocyte arrest at the prophase I, leads to a drop in intra-oocyte cAMP levels and meiotic resumption [40] (see Table 2-3).

Among the down-regulated DEG were UDP-glucose 6-dehydrogenase (UGDH), Morf4 family associated protein 1 (MRFAP1), protein phosphatase Mg2+/Mn2+ dependent 1B (PPM1B), transmembrane protein 59 (TMEM59) and cyclin dependent kinase 12 (CDK12). UGDH is involved in signal transduction, cell migration, and cancer growth and metastasis and has been reported to be down-regulated in slowed tumour cell metabolism [41]. Additionally, overexpression of MRFAP1 has resulted in decreased cell growth and increased mitotic apoptosis in HeLa cells [42]. The PPM1B
catalyses the dephosphorylation of a selection of proteins. Its overexpression causes cell-growth arrest or cell death [43]. The overexpression of TMEM59, has been found to stimulate apoptosis in glioma cells [44]. Downregulation of CDK12 has been observed in diapause like-stage with slowed aging in Drosophila melanogaster [45].

Altogether, the expression patterns of the DEG in both GV and MII oocytes match a profile of decreased apoptosis, reduced oxidative stress, higher oocyte survival and slowed aging during embryonic diapause. This suggests a role of this developmental stage on oocyte quality (see Table 2-3).

![Venn diagram displaying the overlaps of differentially expressed genes. GV, germinal vesicle; MII, metaphase II; D, diapause; R, reactivation](image)

**Figure 7.** Venn diagram displaying the overlaps of differentially expressed genes. GV, germinal vesicle; MII, metaphase II; D, diapause; R, reactivation

**Table 2.** List of differentially expressed genes (DEG) in GV roe deer oocytes during embryonic diapause related to after embryo reactivation

| Gene symbol | Gene description                                                                 | Log2 fold change* | adj.pvalue  |
|-------------|-----------------------------------------------------------------------------------|-------------------|-------------|
| LOC110743349 | Uncharacterized                                                                   | -10.33            | 1.8E-19     |
| PNRC1       | proline rich nuclear receptor coactivator 1                                       | -7.19             | 2.8E-16     |
| PLPP1       | phospholipid phosphatase 1                                                        | -7.15             | 1.5E-11     |
| PYURF       | PIGY upstream reading frame                                                        | -6.89             | 2.5E-15     |
| TMEM123     | transmembrane protein 123                                                         | -6.20             | 1.3E-06     |
| GPSM2       | G protein signalling modulator 2                                                   | -6.04             | 6.6E-05     |
| RBMS1       | RNA binding motif single stranded interacting protein 1                           | -4.09             | 4.6E-04     |
| C1orf115    | chromosome 1 open reading frame 115                                               | -3.46             | 1.5E-04     |
| SOD2        | superoxide dismutase 2                                                            | -2.71             | 5.9E-10     |
| OGT         | O-linked N-acetylglucosamine (GlcNAc) transferase                                 | -2.24             | 6.4E-03     |
| CNOT1       | CCR4-NOT transcription complex subunit 1                                           | -2.01             | 4.0E-04     |
| Gene Name   | Description                                           | Log2 Fold Change | P-Value |
|------------|-------------------------------------------------------|-----------------|---------|
| ELP3       | elongator acetyltransferase complex subunit 3         | -2.01           | 4.0E-03 |
| BUD31      | BUD31 homolog                                         | -1.83           | 1.1E-04 |
| MORF4L2    | mortality factor 4 like 2                             | -1.79           | 2.3E-06 |
| GPATCH11   | G-patch domain containing 11                         | -1.75           | 1.1E-05 |
| CLCC1      | chloride channel CLIC like 1                          | -1.75           | 2.4E-07 |
| MAPRE2     | microtubule associated protein RP/EB family member 2  | -1.67           | 1.8E-03 |
| GCH1       | GTP cyclohydrolase 1                                  | -1.61           | 5.6E-05 |
| DDIT3      | DNA damage inducible transcript 3                     | -1.56           | 6.8E-08 |
| NMRK1      | nicotinamide riboside kinase 1                        | -1.39           | 1.4E-03 |
| LOC109552164 | Uncharacterized                                    | -1.38           | 6.7E-08 |
| OSBPL9     | oxysterol binding protein like 9                      | -1.25           | 6.5E-03 |
| CDC40      | cell division cycle 40                                | -1.21           | 5.1E-03 |
| LRC1H      | leucine rich repeats and calponin homology domain containing 1 | -1.17 | 3.6E-03 |
| SRP9       | signal recognition particle 9                         | -1.05           | 2.4E-03 |
| LOC113609157 | Uncharacterized                              | -1.01           | 5.0E-08 |
| ATIC       | 5-aminooimidazole-4-carboxamide ribonucleotide formyltransferase/IMP cyclohydrolase | 1.25 | 3.6E-08 |
| TARS       | threonyl-tRNA synthetase                               | 1.27            | 1.3E-03 |
| KMT5B      | lysine methyltransferase 5B                           | 1.41            | 2.0E-07 |
| HSPA1A     | heat shock protein family A (Hsp70) member 1A         | 1.64            | 4.9E-04 |
| LOC113615135 | Uncharacterized                              | 1.73            | 3.5E-16 |
| FH         | fumarate hydratase                                    | 1.85            | 1.8E-03 |
| TAC1       | tachykinin precursor 1                                | 2.19            | 1.6E-08 |
| PDE3A      | phosphodiesterase 3A                                  | 2.21            | 9.8E-11 |
| RTRAF      | RNA transcription, translation and transport factor    | 3.68            | 1.0E-08 |
| FBXO38     | F-box protein 38                                      | 5.00            | 1.6E-04 |
| COG1       | component of oligomeric golgi complex 1               | 5.12            | 2.4E-04 |
| LOC109488965 | Uncharacterized                              | 5.52            | 9.8E-06 |
| SH3GL1     | SH3 domain containing GRB2 like 1, endophilin A2      | 5.54            | 7.1E-03 |
| N4BP2L2    | NEDD4 binding protein 2 like 2                        | 5.66            | 2.3E-06 |
| CCNG1      | cyclin G1                                            | 6.00            | 1.4E-05 |
| STAG1      | stromal antigen 1                                     | 6.07            | 1.1E-07 |
| TXNIP      | thioredoxin interacting protein                       | 7.05            | 9.8E-06 |
| NDRG3      | NDRG family member 3                                  | 9.29            | 1.6E-40 |

* A negative log2 fold change indicates higher expression at reactivation, while a positive log2 fold change indicates higher expression at diapause.
Table 3. List of differentially expressed genes in MII roe deer oocytes during embryonic diapause related to reactivation

| Gene symbol | Gene description                                         | Log2 fold change * | adj.pvalue   |
|------------|----------------------------------------------------------|--------------------|--------------|
| RSL24D1    | ribosomal L24 domain containing 1                        | -7.58              | 3.6E-15      |
| UGDH       | UDP-glucose 6-dehydrogenase                              | -7.06              | 5.9E-14      |
| MRFAP1     | Morf4 family associated protein 1                        | -4.88              | 1.3E-03      |
| AP3B1      | adaptor-related protein complex 3, beta 1 subunit        | -4.70              | 5.5E-03      |
| PPM1B      | protein phosphatase, Mg2+/Mn2+ dependent 1B              | -4.50              | 5.4E-04      |
| CACNA2D4   | calcium voltage-gated channel auxiliary subunit alpha2delta 4 | -4.25              | 1.3E-03      |
| FAM91A1    | family with sequence similarity 91 member A1             | -3.25              | 8.5E-03      |
| MCC        | MCC regulator of WNT signaling pathway                   | -2.38              | 1.4E-04      |
| GPSM2      | G protein signalling modulator 2                         | -2.15              | 6.9E-04      |
| LOC107401058 | Uncharacterized                                      | -1.95              | 4.8E-03      |
| COPZ1      | coatomer protein complex subunit zeta 1                  | -1.74              | 1.6E-03      |
| CLIP1      | CAP-Gly domain containing linker protein 1               | -1.62              | 4.6E-08      |
| CDK12      | cyclin dependent kinase 12                               | -1.46              | 8.5E-03      |
| TOR1A      | torsin family 1 member A                                 | -1.25              | 7.9E-03      |
| TMEM59     | transmembrane protein 59                                 | -1.21              | 5.1E-04      |
| LOC109552164 | Uncharacterized                                      | -1.21              | 1.2E-06      |
| ACSL3      | acyl-CoA synthetase long chain family member 3           | -1.14              | 3.2E-04      |
| G3BP1      | G3BP stress granule assembly factor 1                    | -1.10              | 3.4E-04      |
| NIFK       | nucleolar protein interacting with the FHA domain of MKI67 | 1.03               | 8.5E-03      |
| ZKSCAN8    | zinc finger with KRAB and SCAN domains 8                 | 1.06               | 2.1E-03      |
| SNX10      | sorting nexin 10                                        | 1.51               | 1.2E-03      |
| UBE2R2     | ubiquitin conjugating enzyme E2 R2                       | 1.55               | 1.3E-03      |
| ELOVL7     | ELOVL fatty acid elongase 7                              | 1.63               | 7.9E-03      |
| ECHDC1     | ethylmalonyl-CoA decarboxylase 1                         | 1.81               | 2.8E-05      |
| SYNJ2      | synaptojanin 2                                          | 2.42               | 2.6E-03      |
| TAC1       | tachykinin precursor 1                                  | 2.56               | 4.4E-09      |
| H310_01334 | hypothetical protein                                    | 3.94               | 6.9E-04      |
| HSP90AB1   | heat shock protein 90 alpha family class B member 1      | 4.48               | 4.4E-04      |
| ADIPORE2   | adiponectin receptor 2                                   | 4.99               | 2.1E-03      |
| SOD2       | superoxide dismutase 2                                   | 5.63               | 5.7E-06      |
| RBMS1      | RNA binding motif single stranded interacting protein 1  | 5.63               | 2.4E-03      |
| NCSTN      | nicastrin                                               | 5.71               | 2.1E-04      |

* A negative log2 fold change indicates higher expression at reactivation, while a positive log2 fold change indicates higher expression at diapause
Functional analysis of DEG in GV and MII oocytes and GV versus MII oocytes

For the GV_D_vs_MII_D and GV_R_vs_MII_R comparisons, an IPA analysis was performed including canonical pathways, functions and upstream regulators. Due to the low number of differentially expressed genes (DEG) for the GV_D_vs_GV_R and MII_D_vs_MII_R comparisons, the IPA analysis did not provide statistically significant results.

A function enrichment analysis of DEG in MII oocytes relative to GV oocytes was performed and several categories were overrepresented for the DEG of both developmental stages, such as, “Cellular development, Cellular growth and proliferation”, “Cell death and survival”, “Cancer”, “Cancer, Organismal injury and abnormalities”, “DNA replication, recombination, repair”, “Protein synthesis”, and “Nucleic Acid Metabolism, Small Molecule Biochemistry” (see Table 4). Additionally, “Amino acid metabolism, Post-translational modification, Small molecule biochemistry” and “Molecular transport” were only identified in the diapause group. “Cellular function and maintenance”, “Cellular assembly and Organization”, “Cell cycle, DNA replication, recombination, and repair”, “Carbohydrate Metabolism”, “Cancer, Organismal injury and abnormalities”, “DNA replication, recombination, repair”, “Protein synthesis”, and “Nucleic Acid Metabolism, Small Molecule Biochemistry” were only enriched at reactivation (see Table 4).

We further analysed the pathway enrichment in MII oocytes compared to GV oocytes. From a total of 39 enriched canonical pathways that were identified (see Table 5), 26 showed a similar inhibition pattern between the diapause and reactivation groups, including “Oxidative phosphorylation”, known as one the major pathways for ATP production and also related to the transcript selective degradation process during oocyte maturation in other species [33]. A total of 11 additional canonical pathways that appeared as inhibited were only enriched in the reactivation period and two were only enriched in MII oocyte from the diapause period. The observed increase in number of DEG in the GV_R_vs_MII_R comparison was also reflected in the enrichment of additional functions and pathways. It is known that mitochondria play a crucial role to provide the ATP required for oocyte maturation. In addition, they contain anti-apoptotic and apoptogenic components, which have been related to abnormal activation or fragmentation in aged oocytes (reviewed by [46]). “Function of mitochondria”, “Formation of cytoskeleton”, “Synthesis of ATP”, and “Development of mitochondria” were some of the functions additionally found inhibited or decreased in the oocytes from the reactivation group, suggesting a possible dysfunction of mitochondria in these oocytes. Furthermore, “Homologous recombination” and “Repair of DNA” functions were also found in these oocytes, which might implicate altered meiosis. Two additional depleted pathways in the GV_R_vs_MII_R group (“Cysteine biosynthesis” and “Methionine degradation I”) are necessary for “Protein synthesis”, another dysregulated function. These modifications might be related to the also found “Eif2 Signalling”, since they are involved in global protein synthesis and translation of specific stress-associated mRNA transcripts [47]. Although “Cell survival” was decreased in both groups of oocytes (GV_D_vs_MII_D and GV_R_vs_MII_R), “Apoptosis” was increased after embryonic reactivation, leading to speculate that apoptosis could have already started in the follicle, thereby decreasing the oocyte quality and meiotic resumption capacity, as also observed in our study.

Selective degradation of maternal RNA in oocytes is necessary for oocyte to zygote activation and oocyte meiosis to blastomere mitosis transition [48]. The abnormal degradation of determined transcripts might affect oocyte quality and further embryo development [49], which could be the consequence of the observed expression pattern in the oocytes from the reactivation group.
| Function categories                                      | Diseases and Bio Functions                                | Z-score MII vs. GV reactivation | Z-score MII vs. GV diapause | Number of genes MII vs. GV reactivation | Number of genes MII vs. GV diapause |
|---------------------------------------------------------|----------------------------------------------------------|---------------------------------|----------------------------|----------------------------------------|----------------------------------|
| Cell Death and Survival                                 | Apoptosis                                                | 3.6                             | -                          | 353                                    | -                                |
| Cellular Development, Cellular Growth and Proliferation | Cell proliferation of tumour cell lines                  | -3.6                            | -3.4                       | 222                                    | 140                              |
| Cell Death and Survival                                 | Cell survival                                            | -7.6                            | -4.4                       | 200                                    | 114                              |
| Cancer                                                  | Cell transformation                                      | -3.0                            | -2.7                       | 63                                     | 47                               |
| Cellular Function and Maintenance                       | Cellular homeostasis                                     | -2.6                            | -                          | 203                                    | -                                |
| Cancer, Organismal Injury and Abnormalities             | Development of malignant tumour                          | 2.6                             | 2.2                        | 785                                    | 483                              |
| Cellular Assembly and Organization                      | Development of mitochondria                              | -2.1                            | -                          | 10                                     | -                                |
| DNA Replication, Recombination, and Repair              | Excision repair                                          | -2.8                            | -2.2                       | 22                                     | 14                               |
| Cellular Assembly and Organization                      | Formation of cytoskeleton                                | -4.0                            | -                          | 55                                     | -                                |
| Cellular Assembly and Organization, Cellular Function and Maintenance | Function of mitochondria                                | -2.0                            | -                          | 20                                     | -                                |
| Cell Cycle, DNA Replication, Recombination, and Repair  | Homologous recombination                                 | -2.6                            | -                          | 19                                     | -                                |
| Carbohydrate Metabolism                                | Metabolism of carbohydrate                              | -2.3                            | -                          | 90                                     | -                                |
| Nucleic Acid Metabolism                                | Metabolism of nucleic acid component or derivative       | -4.0                            | -                          | 66                                     | -                                |
| Protein Synthesis                                       | Metabolism of peptide                                   | -2.0                            | -2.2                       | 32                                     | 24                               |
| Amino Acid Metabolism, Post-Translational Modification, Small Molecule Biochemistry | Phosphorylation of L-serine                            | -                               | -2.2                       | -                                      | 5                                |
| Post-Translational Modification                        | Phosphorylation of protein                               | -2.8                            | -                          | 88                                     | -                                |
| Cellular Growth and Proliferation, Connective Tissue Development and Function, Tissue Development | Proliferation of connective tissue cells                 | -2.3                            | -                          | 79                                     | -                                |
| DNA Replication, Recombination, and Repair              | Repair of DNA                                            | -3.6                            | -                          | 50                                     | -                                |
| Energy Production, Nucleic Acid Metabolism, Small Molecule Biochemistry | Synthesis of ATP                                        | -3.1                            | -                          | 25                                     | -                                |
| Nucleic Acid Metabolism, Small Molecule Biochemistry    | Synthesis of purine nucleotide                           | -3.6                            | -2.4                       | 36                                     | 22                               |
| Gene Expression                                         | Transactivation                                          | -3.0                            | -                          | 77                                     | -                                |
| Gene Expression, Protein Synthesis                      | Translation of RNA                                       | 2.5                             | -                          | 35                                     | -                                |
| Molecular Transport                                     | Transport of molecule                                    | -                               | -3.0                       | -                                      | 126                              |

**Table 4.** Biological functions in MII oocytes relative to GV oocytes during embryonic diapause and reactivation.
Table 5. Canonical pathways in MII oocytes relative to GV oocytes during embryonic diapause and reactivation

| Canonical Pathways                                           | Z-score MII vs. GV reactivation | Z-score MII vs. GV diapause |
|-------------------------------------------------------------|---------------------------------|-----------------------------|
| Acute Myeloid Leukaemia Signalling                         | -2.1                            | -                           |
| α-Adrenergic Signalling                                     | -2.3                            | -2.2                        |
| Aldosterone Signalling in Epithelial Cells                  | -                               | -2.4                        |
| BMP signalling pathway                                      | -2.9                            | -2.4                        |
| CD40 Signalling                                             | -2.5                            | -2.5                        |
| Cysteine Biosynthesis III (mammalia)                       | -2.4                            | -                           |
| D-myo-inositol-5-phosphate Metabolism                       | -2.1                            | -                           |
| EIF2 Signalling                                             | -2.6                            | -                           |
| ErbB Signalling                                             | -2.7                            | -2.8                        |
| Gluconeogenesis I                                          | -2.4                            | -                           |
| Glycolysis I                                                | -2.4                            | -                           |
| Fatty Acid β-oxidation I                                   | -2.5                            | -2.0                        |
| Gα12/13 Signalling                                         | -3.0                            | -2.5                        |
| IL-1 Signalling                                             | -3.2                            | -2.8                        |
| Ketolysis                                                   | -2.0                            | -                           |
| IL-6 Signalling                                             | -3.1                            | -3.6                        |
| iNOS Signalling                                             | -2.0                            | -2.4                        |
| LPS-stimulated MAPK Signalling                             | -3.1                            | -3.2                        |
| Methionine Degradation I (to Homocysteine)                  | -2.4                            | -                           |
| Melatonin Signalling                                        | -2.9                            | -2.4                        |
| NGF Signalling                                              | -3.1                            | -2.8                        |
| NRF2-mediated Oxidative Stress Response                     | -3.7                            | -3.2                        |
| Oxidative Phosphorylation                                  | -4.7                            | -3.0                        |
| PFKFB4 Signalling Pathway                                  | -3.3                            | -2.4                        |
| 3-phosphoinositide Degradation                              | -2.5                            | -                           |
| PI3K Signalling in B Lymphocytes                            | -2.8                            | -2.6                        |
| PPAR Signalling                                             | 2.1                             | 2.5                         |
| Pyridoxal 5′-phosphate Salvage Pathway                      | -2.3                            | -2.3                        |
| Pyrimidine Ribonucleotides De Novo Biosynthesis             | -3.0                            | -2.0                        |
| Rac Signalling                                              | -3.5                            | -2.8                        |
Regulation of eIF4 and p70S6K Signalling: -2.1
Pyrimidine Ribonucleotides Interconversion: -2.8
Salvage Pathways of Pyrimidine Ribonucleotides: -3.0
STAT3 Pathway: -2.5
Superpathway of Cholesterol Biosynthesis: -
Superpathway of Methionine Degradation: -2.6
TGF-β Signalling: -2.5
TCA Cycle II (Eukaryotic): -2.4
Tryptophan Degradation III (Eukaryotic): -2.0

Conclusions
The present study provides novel insights into ovarian and oocyte physiology in the roe deer. We have observed that ovarian antral follicle number decreased after embryonic reactivation had occurred. Thus, reactivation might be associated with reduced oocyte quality, reflected by the lower number of oocytes suitable for IVM and oocytes capable to reach the MII stage. A deeper view to the oocyte quality as seen from RNA-Seq data revealed differential gene expression between oocytes retrieved during embryonic diapause and after reactivation. The findings could potentially be related to extended lifespan and diminished senescence of the oocyte during embryonic diapause and alleviation of cellular stress after meiotic resumption (see Figure 8). For the interpretation of the present results caution is needed. Since does exhibit high fertility rates, sampling of confirmed non-pregnant animals is not possible under hunting conditions. Therefore, seasonality effects cannot be excluded. Furthermore, the follicle counts were performed visually, which might lead to underestimation of the exact number and size of present antral follicles. Additionally, the time between fertilization and sample collection cannot be estimated, adding high variability to the sampling process. A validation of the discovered effects together with further analyses might provide novel quality markers.
Figure 8. Potential mechanisms of the embryonic diapause and reactivation periods on the antral follicular count and oocyte quality.

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.

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