Seasonal effects on miRNA and transcriptomic profile of oocytes and follicular cells in buffalo (*Bubalus bubalis*)

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Season clearly influences oocyte competence in buffalo (*Bubalus bubalis*); however, changes in the oocyte molecular status in relation to season are poorly understood. This study characterizes the microRNA (miRNA) and transcriptomic profiles of oocytes (OOs) and corresponding follicular cells (FCs) from buffalo ovaries collected in the breeding (BS) and non-breeding (NBS) seasons. In the BS, cleavage and blastocyst rates are significantly higher compared to NBS. Thirteen miRNAs and two mRNAs showed differential expression (DE) in FCs between BS and NBS. DE-miRNAs target gene analysis uncovered pathways associated with transforming growth factor β (TGFβ) and circadian clock photoperiod. Oocytes cluster in function of season for their miRNA content, showing 13 DE-miRNAs between BS and NBS. Between the two seasons, 22 differentially expressed genes were also observed. Gene Ontology (GO) analysis of miRNA target genes and differentially expressed genes (DEGs) in OOs highlights pathways related to triglyceride and sterol biosynthesis and storage. Co-expression analysis of miRNAs and mRNAs revealed a positive correlation between miR-296-3p and genes related to metabolism and hormone regulation. In conclusion, season significantly affects female fertility in buffalo and impacts on oocyte transcriptomic of genes related to folliculogenesis and acquisition of oocyte competence.

Water buffalo (*Bubalus bubalis*) is an important livestock resource for both developing and developed countries. The major factor affecting buffalo farming profitability is reproductive seasonality, resulting in cycles of calving and milk production. Buffalo is a short-day breeder, with increased fertility in response to decreasing day length.¹,² This photoperiod-dependent seasonality pattern is more pronounced as distance from the equator, together with variations in the light/dark ratio, increases. In Italy, in order to satisfy market demand, out of breeding mating strategy (OBMS), consisting in interrupting sexual promiscuity or the use of artificial insemination (AI) during the breeding season (BS), is commonly utilized.² The OBMS improves the distribution of calving throughout the year, but it reduces fertility.³ Longer post-partum anoestrus periods as well as higher incidence of embryonic mortality are observed in months with increasing daylight length and particularly in mid-winter, which coincides with the transition to seasonal anoestrus at Italian latitudes.⁴ The embryonic mortality is due to inadequate luteal growth and function, resulting in reduced progesterone secretion.¹ This has a negative impact on embryo growth, associated with alterations in transcriptomic and proteomic profiles of the embryos and chorioamnios/caruncles,⁵,⁶ which ultimately impair embryo attachment to the uterine endometrium.

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An additional factor determining reproductive failure in the non-breeding season (NBS) is the oocyte developmental competence. Indian authors reported decreased efficiency of ovum pick-up (OPU) during the NBS, mainly due to the reduced follicular population. A seasonal effect on the number of follicles and oocytes, as well as on oocyte competence, has also been reported in Egyptian buffaloes. In Italian Mediterranean buffaloes, season clearly influences oocyte competence, as indicated by improved blastocyst yields recorded during months with decreasing daylight. In Murrah buffalo heifers, the decreased oocyte quality recorded during long day months was associated to reduced concentration of oestradiol both in plasma and follicular fluid, as well as of intrafollicular IGF-1. Despite the evidence of a seasonal influence in buffalo, the molecular mechanisms affecting oocyte competence in the NBS are poorly understood.

A fine-tuned spatio-temporal expression of multiple genes is known to be essential for follicular development and oocyte maturation, and requires a strict interaction between mRNAs and regulatory miRNAs. In addition, in many tissues a time-controlled gene expression is mediated by miRNAs, which regulate core clock genes coordinating daily rhythms in physiology and behaviour. A relationship between the variation of mRNAs abundance for specific genes related to folliculogenesis in ovaries and changes in photoperiods was previously reported in other non-ruminant species such as the Siberian hamster. Changes in transcriptome and miRNA expression in relation to season were further investigated in sheep, another short-day breeder, where oocyte competence was observed to decline during the NBS, as indicated by impaired in vitro embryo development. Furthermore, transcriptome variations potentially associated with off-season reproduction were reported in sheep ovaries.

Again, differences in miRNA profiling in ovaries of Tan sheep and Small Tail Han (STH) sheep were related to oovine anoestrus and BS. Seasonal differences in the expression of miRNAs involved in hormone regulation, follicular growth and angiogenesis were also observed in Kazakh sheep ovaries during oestrus. Recently, an integrated analysis of mRNA and miRNA expression in European mouflon (Ovis musimon) and sheep (Ovis aries) depicted a miRNA-mRNA regulatory network associated with reproductive traits in Ovis species.

MicroRNAs play a significant role during follicle development in bovine and the role of miRNAs is also demonstrated in the regulation of lactating physiology in the buffalo mammary gland, and miRNA expression changes were observed in buffalo corpus luteum during pregnancy.

Therefore, in this study we investigated, for the first time, if reproductive failure in the NBS is associated to changes in gene expression affecting oocyte developmental competence in buffalo. To evaluate seasonal effects on oocyte competence, the miRNA and transcriptomic profiles of oocytes and corresponding follicular cells were characterized from abattoir-derived ovaries collected in the BS and NBS.

**Results**

**Cleavage and blastocyst rates.** With regard to oocyte competence, cleavage rate was higher in the BS compared to the NBS (69.4%; vs. 60.7; \( P < 0.05 \)). In addition, an improvement in blastocyst yields was recorded in the BS, both in relation to total COCs (26.5 vs. 16.3%; \( P < 0.01 \)) and cleaved oocytes (38.7 vs. 27.4%; \( P < 0.05 \)).

**miRNAs.** Twenty samples, i.e. 10 pools of OOs (sample OO1-OO5 from NBS and OO6-OO10 from BS) and 10 pools of FCs (sample FC1-FC5 from NBS and FC6-FC10 from BS), were characterized for their miRNA content.

About 10 million reads were sequenced for both OOs and FCs (see Supplementary File S1 for statistics). About 1% and 20% of them were assigned to miRNAs in OOs and FCs, respectively.

In total, 769 miRNAs were identified in at least three samples in all conditions (468 *Bos taurus* bta-miRNAs, 279 novels, and 22 novels homologous to related species). Among them, 467 were detected in at least three OO samples and 635 in at least three FC samples. Principal Component Analysis (PCA) clearly separates OOs and FCs according to their miRNA content, with 44% of the variance explained by component 1 (Supplementary file S2).

Few of the most expressed miRNAs showed a similar relative abundance in OOs and FCs (bta-miR-10b, bta-miR-148a and bta-miR-26a); on the contrary, expression rate of most miRNAs differed in the two cellular types (bta-miR-21-5p was highly expressed in FCs, whereas bta-miR-423-3p in OOs). In fact, there was a statistically significant difference in the expression of a high proportion of the miRNAs (n = 413) (False Discovery Rate FDR < 0.05) between OOs and FCs (Supplementary file 4). When the two seasons were considered, the PCA produced a good distinction between oocytes collected in BS and NBS, whereas FCs from BS and NBS could not be clearly distinguished (Fig. 1).

The number of differentially expressed miRNAs (DE-miRNAs, FDR < 0.05) between the two seasons was 13 for both OOs and FCs (Supplementary file 5 and Table 1). A view of the normalized expression of the most representative DE-miRNAs is shown in Fig. 2. Target prediction using human miRNAs homologous to buffalo DE-miRNAs led to the identification of 6,712 and 4,847 genes potentially regulated in OOs and FCs, respectively (\( P < 0.05 \)). GO analysis using a subset of more significant target genes (n = 136) identified pathways related to triglyceride and cholesterol metabolism and transport, and mesoderm and epithelial cell morphogenesis differentiation for OOs, and related to photoperiodism, circadian clock regulation, and transforming growth factor beta signalling for FCs (Table 2).

**RNAseq.** RNA-seq analysis was performed on the same samples used for miRNA profiling to evaluate the gene expression variation between the two cellular types and seasons. Approximately 23.5 ± 4.4 and 54.5 ± 10.5 millions of reads were obtained for OOs and FCs samples with a mapping rate of 93.5% and 92.4%, respectively (Supplementary file 6). A total of 22,013 unique genes present in at least three samples from both cellular types were identified (19,240 counted in at least three OOs and 21,277 counted in at least three FCs samples). PCA
considering the relative expression of these genes showed a clear separation between the two cellular types (Supplementary file 7). There seems to be no seasonal effect in the overall transcript abundance for both OOs and FCs (Fig. 3).

The relative expression of mRNAs in the two cellular types was very different, with 14,680 (66.7%) DEGs between OOs and FCs. When DEGs were calculated between the two seasons, 22 mRNAs were found to differ between NBS and BS for OOs, whereas only two DEGs were present in FCs (Supplementary file 8 and Table 3).

**Figure 1.** Principal component analysis considering (a) the 467 miRNAs expressed at least in triplicate in the oocytes (OOs), (b) the 635 miRNAs expressed at least in triplicate in the Follicular cells (FCs). Samples 1–5 from non-breeding season (NBS), samples 6–10 from breeding season (BS).

**Table 1.** Differentially expressed miRNAs DE-miRNAs (false discovery rate (FDR) < 0.05) between the two seasons (NBS = non breeding season, BS = breeding season) for oocytes (OOs) and follicular cells (FCs).

| miRNAs             | logFC | FDR    | miRNAs             | logFC | FDR    |
|--------------------|-------|--------|--------------------|-------|--------|
| bta-miR-143        | −2.07 | 5.42E−05 | Novel:NC_037550.1_18643 | 1.58 | 7.24E−04 |
| Novel:NC_037550.1_18643 | 1.58 | 7.24E−04 | Novel:chi-miR-184 | −4.24 | 1.54E−04 |
| bta-miR-199a-3p    | −2.10 | 7.24E−04 | Novel:NC_037550.1_18643 | −2.37 | 4.25E−04 |
| bta-miR-1468       | −1.83 | 3.69E−03 | Novel:NC_037550.1_18643 | −2.37 | 4.25E−04 |
| bta-miR-1388-5p    | −4.55 | 3.55E−02 | Novel:NC_037550.1_18643 | −2.37 | 4.25E−04 |
| bta-miR-296-3p     | −1.41 | 3.91E−02 | Novel:NC_037550.1_18643 | −2.37 | 4.25E−04 |
| Novel:NC_037550.1_18643 | −1.36 | 3.91E−02 | Novel:miR-2332 | −1.47 | 4.48E−03 |
| Novel:NC_037569.1_47305 | −1.36 | 3.91E−02 | Novel:miR-141 | −3.73 | 4.48E−03 |
| Novel:NC_037550.1_18643 | −1.36 | 3.91E−02 | Novel:miR-2478 | 1.68 | 4.97E−03 |
| bta-miR-331-5p     | −4.19 | 4.12E−02 | Novel:miR-34b | −4.11 | 8.16E−03 |
| bta-miR-199a-5p    | −2.09 | 4.47E−02 | Novel:miR-34c | −4.02 | 8.16E−03 |
| bta-miR-222        | −1.35 | 4.95E−02 | Novel:miR-486 | 0.96 | 3.65E−02 |
Although a limited number of DEGs was found to differ in OOs between the two seasons, GO analysis revealed that some of them were related to lipid storage and localization and regulation of interleukin-8 (IL8) production (Table 4).

**miRNAs and mRNA interaction.** In order to evaluate whether miRNAs could potentially regulate the expression of specific genes, the list of genes differentially expressed in oocytes between the two seasons was intersected with the list of DE-miRNA target genes observed in the same experimental condition. Six genes (CCL1, FOLR2, IGF2, HSPA1A, IL1B, CTSK) were found to be potentially regulated by specific DE-miRNAs. Interestingly, among the 8 DE-miRNAs (miR-143, miR-1468, miR-199a-3p, miR-199a-5p, miR-222, miR-25, miR-296-3p, miR-331-5p) targeting 6,712 genes, miR-296-3p targets 4 out of the 6 shared DEGs. In addition, all the miRNAs show a positive correlation with gene expression (Fig. 4).

**Discussion**
The present study aims to investigate the causes of the decreased oocyte competence during the NBS in buffalo. Many biological processes are required for developmental competence, with the exchange of information between oocyte and follicular environment promoting oocyte maturation. Therefore, the focus of this study was to evaluate differences in the miRNA and transcriptomic profiles of OOs and corresponding FCs from buffalo.
| GOID      | Associated genes found                  | GO term                                                      | P value*  |
|-----------|----------------------------------------|--------------------------------------------------------------|-----------|
| 0010866   | [DGAT2, FITM2, NR1H3]                  | Regulation of triglyceride biosynthetic process              | 0.004     |
| 0060742   | [NOTCH1, SFTPA1, SFTPA2]               | Epithelial cell differentiation involved in prostate gland development | 0.005     |
| 0015918   | [ABC4G, APOB, NR1H3, OSBPL6, PNLIP]    | Sterol transport                                            | 0.009     |
| 0030850   | [NOTCH1, SFTPA1, SFTPA2, WNT5A]        | Prostate gland development                                   | 0.012     |
| 0048332   | [AXIN1, GNPD1A, WNT5A]                 | Mesoderm morphogenesis                                      | 0.014     |
| 0019915   | [APOB, DGAT2, FITM2, NR1H3]           | Lipid storage                                               | 0.026     |
| 0001707   | [AXIN1, GNPD1A, WNT5A]                 | Mesoderm formation                                          | 0.027     |
| 0006536   | [ATAT1, GGT1, GLU2D]                  | Glutamate metabolic process                                 | 0.028     |
| 0006641   | [APOB, DGAT2, FITM2, INSIG1, NR1H3]   | Triglyceride metabolic process                              | 0.028     |
| 0042116   | [IL13, NR1H3, WNT5A]                  | Macrophage activation                                       | 0.028     |
| 0090207   | [DGAT2, FITM2, NR1H3]                 | Regulation of triglyceride metabolic process                | 0.029     |
| 0008206   | [ACA1A1, CYP27A1, OSBPL6]              | Bile acid metabolic process                                 | 0.030     |
| 0002657   | [EXOSC3, GNPD1A, IL13]                | Regulation of immunoglobulin production                     | 0.033     |
| 0070527   | [FERMT3, MYH9, PRKCCQ]                | Platelet aggregation                                        | 0.033     |
| 0020067   | [IL13, NOTCH1, WNT5A]                 | Glandular epithelial cell differentiation                   | 0.033     |
| 0019217   | [DGAT2, INSIG1, NR1H3, PDHB]          | Regulation of fatty acid metabolic process                   | 0.034     |
| 0097006   | [APOB, DGAT2, PLAGL2]                 | Regulation of plasma lipoprotein particle levels            | 0.035     |
| 0055090   | [DGAT2, FITM2, NR1H3]                 | Acylglycerol homeostasis                                    | 0.035     |
| 0070328   | [DGAT2, FITM2, NR1H3]                 | Triglyceride homeostasis                                    | 0.035     |
| 0030301   | [ABC4G, APOB, NR1H3, PNLIP]           | Cholesterol transport                                       | 0.037     |
| 0050830   | [DROSHA, H1ST1H2BK, PGLYRP1, PGLYRP3]| Defence response to Gram-positive bacterium                 | 0.037     |
| 0033344   | [ABC4G, APOB, NR1H3]                  | Cholesterol efflux                                          | 0.038     |
| 0002702   | [EXOSC3, GNPD1A, IL13, WNT5A]         | Positive regulation of production of molecular mediator of immune response | 0.038     |
| 0030514   | [CHRDL1, NOTCH1, WNT5A]               | Negative regulation of BMP signaling pathway                | 0.038     |
| 0010883   | [APOB, FITM2, NR1H3]                  | Regulation of lipid storage                                 | 0.039     |
| 0019432   | [DGAT2, FITM2, NR1H3]                 | Triglyceride biosynthetic process                            | 0.039     |
| 0043030   | [IL13, NR1H3, WNT5A]                  | Regulation of macrophage activation                         | 0.039     |
| 0045599   | [AXIN1, INSIG1, WNT5A]                | Negative regulation of fat cell differentiation              | 0.040     |
| 0002639   | [EXOSC3, GNPD1A, IL13]                | Positive regulation of immunoglobulin production            | 0.040     |
| 0046460   | [DGAT2, FITM2, NR1H3]                 | Neutral lipid biosynthetic process                          | 0.040     |
| 0046463   | [DGAT2, FITM2, NR1H3]                 | Acylglycerol biosynthetic process                            | 0.040     |
| 0043153   | [BHLHE40, PPL1CR, PPI1CC]             | Entrainment of circadian clock by photoperiod               | 0.004     |
| 1903844   | [ING3, ONECUT2, SKI, STRAP, XBPI]     | Regulation of cellular response to transforming growth factor beta stimulus | 0.008     |
| 0017015   | [ING3, ONECUT2, SKI, STRAP, XBPI]     | Regulation of transforming growth factor beta receptor signaling pathway | 0.008     |
| 0096848   | [BHLHE40, PPL1CR, PPI1CC]             | Photoperiod                                                 | 0.008     |
| 0096849   | [BHLHE40, PPL1CR, PPI1CC]             | Entrainment of circadian clock                              | 0.008     |
| 1903845   | [ONECUT2, SKI, STRAP, XBPI]           | Negative regulation of cellular response to transforming growth factor beta stimulus | 0.012     |
| 0035012   | [ONECUT2, SKI, STRAP, XBPI]           | Negative regulation of transforming growth factor beta receptor signaling pathway | 0.012     |
| 0010923   | [FKBP18, PPL1R1B, TMEM225]            | Negative regulation of phosphatase activity                 | 0.014     |
| 0032755   | [IL1R12, TL16, XBPI]                  | Positive regulation of interleukin-6 production             | 0.025     |
| 0045582   | [IL1R12, TIPK, XBPI]                  | Positive regulation of T cell differentiation               | 0.025     |
| 0035304   | [FKBP18, PPL1R1B, SMPD1]              | Regulation of protein dephosphorylation                     | 0.035     |
| 0071230   | [CASTOR1, PGDFC, XBPI]                | Cellular response to amino acid stimulus                    | 0.035     |
| 1903036   | [FKBP18, SCARF1, XBPI]                | Positive regulation of response to wounding                 | 0.040     |
| 0010257   | [NDUFAF6, NDUFC1, NDUF87]             | NADH dehydrogenase complex assembly                         | 0.042     |
| 0032981   | [NDUFAF6, NDUFC1, NDUF87]             | Mitochondrial respiratory chain complex I assembly          | 0.042     |
| 0097031   | [NDUFAF6, NDUFC1, NDUF87]             | Mitochondrial respiratory chain complex I biogenesis        | 0.042     |
| 0032922   | [BHLHE40, PPL1CR, PPI1CC]             | Circadian regulation of gene expression                     | 0.047     |

Table 2. GO terms identified for the target genes of differentially expressed miRNAs between the two seasons for oocytes (OOs) and Follicular Cells (FCs). Indicated are gene ontology IDs (GO-ID), gene ontology terms (GO-term), associated genes found and corrected P values as determined by ClueGO (https://apps.cytoscape.org/apps/cluego). *Term P value corrected with Bonferroni step down.
ovaries collected in the BS and NBS. To our knowledge, this is the first study reporting together the miRNA and mRNA profiling from pools of low numbers of oocytes and corresponding FCs collected from abattoir-derived ovaries in livestock, as well as the first time that the seasonal effects on miRNA and mRNA profiling of oocytes and FCs are investigated in buffalo. Unfortunately, the amount of RNA obtained from such a limited number of oocytes was not sufficient to perform further experiments to validate our results.

In accordance with previous findings, in the present study we observed a reduced oocyte developmental competence in the NBS, as indicated by decreased cleavage and blastocyst rates after in vitro fertilization (IVF), and this was associated to changes in miRNA and transcriptomic profiles both in OOs and FCs. Being miRNAs only one of the small RNA components, as expected those identified in buffalo represent only a fraction of the total small non-coding RNA present in both OOs and FCs. The overall miRNA expression pattern was different between OOs obtained in the two seasons. However, FCs from the two seasons cluster together. Considering that different phenotypes of FCs can be observed within the follicle, it is likely that miRNA expression in FCs is mainly driven by cell position and function, thus masking seasonal effects. Recently, Zhang et al. reviewed miRNA profile studies related to ovarian development and function in mammals. Although many studies reported miRNA ovary profiling, only a few investigated miRNA variations in granulosa cells and only one in OOs.

Interestingly, many of the differentially expressed miRNAs in the two seasons for both OOs and FCs are involved in follicular maturation and development regulation. In our study seasonal changes modified the expression of miR-143, miR-25, miR-222 and miR-199a in buffalo OOs. In the mouse ovary, miR-143 is highly expressed and related to oestradiol production and steroidogenesis gene expression. MiR-143 and miR-25 were also shown to promote progesterone release in human ovarian granulosa cells. Furthermore, cyclic variations in the expression of miR-222 and miR-199a were reported in cattle during follicle maturation, with expression increasing until the mid-luteal phase, and decreasing in the late follicular phase in the bovine dominant follicle.

Some of the differentially expressed miRNAs identified between the NBS and BS in buffalo FCs were also reported to exhibit expression modulation during the cycle in cattle. In particular, temporal miRNA expression dynamics were observed for miR-184 in FCs between days 3 and 7 of the bovine oestrous cycle and for miR-2411 and miR-34c between subordinate and dominant follicles during the early luteal

Figure 3. Principal component analysis considering (a) the 19,240 mRNAs expressed at least in triplicate in the oocytes (OOs), (b) the 21,277 mRNAs expressed at least in triplicate in the Follicular cells (FCs). Samples 1–5 from non-breeding season (NBS), samples 6–10 from breeding season (BS).
MiR-34c was shown to exert anti-proliferative and pro-apoptotic effects in porcine granulosa cells by targeting Forkhead box O3a (FoxO3a) \(^3^4\). In addition, the expression of some of the DE-miRNAs detected in our study differs in several ovarian disorders. It was reported that mir-141 and miR-199a are respectively up and down regulated in human ovarian cancer \(^3^5\), miR-184 is a potential predictor of recurrence in human ovarian granulosa cell tumours \(^3^6\), and miR-486-5p is downregulated in cumulus cells collected from women affected by polycystic ovary syndrome \(^3^7\).

Interestingly, GO analysis of the predicted target genes for DE-microRNAs uncovered pathways associated with OOs and FCs physiology. Oocytes collected from the BS and NBS showed DE-miRNAs able to regulate genes for triglyceride and sterol biosynthesis essential for lipid metabolism, which provides a potent source of energy during oocyte maturation \(^3^8\). In FCs, the DE-miRNA target genes were related to pathways involved in transformation of growth factor β (TGFβ) and circadian clock photoperiod. TGFβ promotes granulosa cell proliferation regulating the expression of luteinizing hormone receptor (LH-R) \(^3^9–4^1\). Altered photoperiod can affect mRNA expression in ovaries \(^1^5\), in fact, transcriptome changes occurred between BS and NBS samples.

Considering DEGs between seasons, it is interesting to note that although only two DEGs were found in FCs, many of the DEGs in the OOs are known to be related to oocyte competence. In the NBS, decreased oocyte competence in buffalo was associated to change in the expression of secreted phosphoprotein 1 (\(SPP1\)), \(RUNX\) family transcription factor 2 (\(RUNX2\)) and Cathepsin K (\(CTSK\)) in OOs. Both \(SPP1\) and \(RUNX2\) expression was observed to change in oocyte and-granulosa cell complexes at various stages of follicle development in pigs \(^4^2\). In addition, variations in the expression of \(SPP1\) were recorded in cumulus cells derived from

| **GO terms** | **Associated genes found** | **P value**
|-------------|----------------------------|-----------------
| GO:19915    | [APOE, IL1B, MSR1]         | Lipid storage  | 7.17E−05 |
| GO:32370    | [APOE, IL1B, SPP1]         | Positive regulation of lipid transport | 8.94E−05 |
| GO:32677    | [CD14, HSPA1A, IL1B]       | Regulation of interleukin-8 production | 4.09E−05 |
| GO:32757    | [CD14, HSPA1A, IL1B]       | Positive regulation of interleukin-8 production | 6.29E−05 |
| GO:1905954  | [APOE, IL1B, MSR1, SPP1]   | Positive regulation of lipid localization | 5.76E−06 |

Table 3. Differentially expressed gene DEGs (false discovery rate (FDR) < 0.05) calculated between the two seasons (non breeding season NBS, breeding season) for oocytes (OOs) and follicular cells (FCs).

Table 4. GO terms identified for the differentially expressed gene (DEGs) between the two seasons for oocytes (OOs). Indicated are gene ontology IDs (GO-ID), gene ontology terms (GO-term), associated genes found and corrected P values as determined by ClueGO (https://apps.cytoscape.org/apps/cluego). * Term P value corrected with Bonferroni step down.
cumulus-oocyte-complexes (COCs) collected from cows undergoing FSH priming, as a model of high oocyte competence43, and RUNX2 expression was associated with controlled ovarian stimulation outcome in assisted reproductive technology treatment in women44. Furthermore, CTSK in cumulus cells was suggested as a predictive marker for oocyte competence in bovine COC45. In buffalo OOs during the NBS, a decreased expression of heat shock protein family A (Hsp70) member 1A (HSPA1A), known to be related to oocyte survival and apoptosis, was also observed. The HSPA1A plays a critical role through its protective action against apoptosis and its expression is reduced in poor, as compared to competent, ovine COCs46.

Another transcript down-regulated in the NBS OOs is interleukin-1 beta (IL-1β). Although IL-1β deficiency in mice prolongs ovarian lifespan47), IL-1β stimulates the growth and sustains maturation in mare48 and bovine oocytes49. In addition, IL-1β was postulated to be involved in different ovulation-associated events such as prostaglandin production and steroidogenesis49. Modulation of expression levels was observed in this study also for other genes related to gonadotropic hormone synthesis and metabolism, showing a reduced expression in buffalo OOs during NBS. Apolipoprotein E (APOE) is expressed in cultured ovarian granulosa cells, and is present in human follicular fluid where its relative levels are correlated with serum estrogen concentration51. In rats, APOE exerts a role in directing cholesterol during steroidogenesis and regulating follicular estrogenic production52. Insulin like growth factor 2 (IGF2) was observed to be expressed in bovine oocytes53. The expression of IGF2 is modulated by growth hormone (GH) in vitro matured Rhesus macaque oocytes54. Furthermore, it is known that IGF2, in combination with follicle stimulating hormone (FSH), acts directly on oocyte competence in caprine follicles55. Finally, the folate receptor beta (FOLR2), also down-regulated in OOs during the NBS, is a key gene linked to methionine/folate cycles in bovine oocyte56, also involved in folate transport in mice oocytes during follicular development57.

In our study, a positive correlation between DEGs and DE-miRNA target genes was observed. MiRNAs usually mediate repression of their target mRNAs by inhibiting their translation, therefore reducing the abundance of their products58. However, several studies reported a positive miRNA-mRNA regulation with a feed-forward mechanism probably mediated by transcription factors59. Notably, among all DE-miRNAs, miR-296-3p is the most correlated with transcripts changing in OOs between BS and NBS. MicroR-296-3p was previously reported to be expressed in ovaries in mice60 and to repress cell plasticity in different tumour lines61, promoting apoptosis in liver62 and in mammalian pancreatic α cells63. Recently, altered expression of bta-miR-296-3p was detected in muscle, kidney, and liver, in bovine foetuses with large offspring syndrome (LOS)64. In addition, miR-296 was also observed to be epigenetically regulated as a part of the imprinted Gnas/GNAS clusters65.

**Conclusion**

In conclusion, the reduced oocyte developmental competence recorded during the NBS in buffalo is associated with changes in miRNA and mRNA content in OOs and corresponding FCs. The GO analysis showed over-representation of key genes related to lipid and sterol biosynthesis and hormone regulation, crucial for folliculogenesis and acquisition of oocyte competence. These observations might help to explain the seasonal difference in the potential of buffalo oocytes, thus providing the basis for the development of strategies to improve oocyte competence in the NBS. Nevertheless, further efforts are still needed to validate expression modulation of miRNAs and key genes identified in our study and deeply investigate their role in seasonal reproduction in buffalo.
Materials and methods

Collection of oocytes and granulosa cells. The study was carried out in Southern Italy (latitude 40.5°–41.5° N and longitude 13.5–15.5) in October, i.e. autumn (BS) and January, i.e. mid-winter (NBS). Buffalo ovaries were collected at a local slaughterhouse (Real Beef s.r.l., Flumeri (AV), Italy) under national food hygiene regulations, and transported to the laboratory in physiological saline supplemented with 150 mg/L kanamycin at 30–35 °C within 4 h after slaughter. In order to reduce variability, the ovaries were collected from a homogeneous population of buffaloes, i.e. 134 cyclic multiparous Italian Mediterranean Buffalo cows with a mean weight of 552 ± 0.4 kg and age of 5.3 ± 0.4 years, over a total of 10 replicates (5/season). Cyclic ovarian activity was assessed by two clinical examinations carried out 12 days apart before slaughter, to detect the presence of a follicle greater than 1 cm and/or corpus luteum on the ovary.

For each day of collection (n = 10), 2–8 mm follicles were aspirated under controlled pressure to collect both OOs and FCs for molecular analyses, while a group of cumulus oocyte complexes (COCs) were in vitro matured, fertilized and cultured up to the blastocyst stage (n = 238 and 234, respectively in the BS and NBS).

Follicular fluid was aspirated using an 18 G needle under vacuum (40–50 mm Hg) in Falcon tubes and poured into a petri dish for COC recovery. The COCs were evaluated according to morphology and classified according to Di Francesco et al.30. Grade A and B COCs, considered suitable for in vitro embryo production (IVEP), were quickly selected from the dish and washed thoroughly in medium H199.

For each replicate, COCs were denuded of their cumulus cells by gentle pipetting and denuded oocytes were washed in phosphate buffer solution (PBS) + 0.1% polyvinyl alcohol (PVA), pooled (20/pool), snap frozen in liquid nitrogen and stored at −80 °C until RNA isolation.

The follicular fluid was centrifuged at 300×g for 10 min at 4 °C to separate the follicular fluid and the FCs. After centrifugation, the supernatant was centrifuged again at 2000 g for 10 min. Insemination was performed in 50 µL drops of IVF medium, stripped of cumulus cells by gentle pipetting and allocated to 20 µL drops of IVC medium, i.e. in TCM199 buffered with 25 mM sodium bicarbonate and supplemented with 10% FCS, 0.2 mM sodium pyruvate, 0.5 µg/mL FSH, 5 µg/mL LH, 1 µg/mL 17-β-estradiol and 50 µg/mL kanamycin, and incubated at 38.5 °C for 21 h in a controlled gas atmosphere of 5% CO2 in humidified air.

The methods for in vitro fertilization (IVF) and culture (IVC) described below have been reproduced from Di Francesco et al. 201232. Frozen straw from a bull previously tested for IVF were thawed at 37 °C for 40 s and sperm was selected by centrifugation (25 min at 300g) on a Percoll discontinuous gradient (45% and 80%). The sperm pellet was re-suspended to a final concentration of 2 × 10⁶ mL⁻¹ in the IVF medium, consisting of Tyrode albumin lactate pyruvate57 supplemented with 0.2 mM penicillin, 0.1 mM hypotaurine and 0.01 mM heparin. Insemination was performed in 50 µL drops of IVF medium under mineral oil (5 oocytes per drop) at 38.5 °C under humidified 5% CO2 in air. Twenty hours after IVF, putative zygotes were removed from the IVF medium, stripped of cumulus cells by gentle pipetting and allocated to 20 µL drops of IVC medium, i.e. synthetic oviduct fluid (SOF) including essential and non-essential amino acids and 8 mg/mL bovine serum albumin58. Culture was carried out under humidified air with 5% CO2, 7% O2 and 88% N2 at 38.5 °C. On day 5 post-insemination (pi) the cleavage rate was assessed and the embryos transferred into fresh medium for further 2 days of IVC, when blastocyst rates were recorded.

RNA isolation. Samples for RNA isolation were obtained from pools (n = 20) of OOs and FCs for both conditions (BS and NBS). The methods described below have been reproduced in part from Lange-Consiglio et al.46. Total RNA was isolated by NucleoSpin miRNA kit (Macherey–Nagel, Germany), following the protocol at 38.5° C for 21 h in a controlled gas atmosphere of 5% CO2 in humidified air.

The methods for in vitro fertilization (IVF) and culture (IVC) described below have been reproduced from Di Francesco et al. 201232. Frozen straw from a bull previously tested for IVF were thawed at 37 °C for 40 s and sperm was selected by centrifugation (25 min at 300g) on a Percoll discontinuous gradient (45% and 80%). The sperm pellet was re-suspended to a final concentration of 2 × 10⁶ mL⁻¹ in the IVF medium, consisting of Tyrode albumin lactate pyruvate57 supplemented with 0.2 mM penicillin, 0.1 mM hypotaurine and 0.01 mM heparin. Insemination was performed in 50 µL drops of IVF medium under mineral oil (5 oocytes per drop) at 38.5 °C under humidified 5% CO2 in air. Twenty hours after IVF, putative zygotes were removed from the IVF medium, stripped of cumulus cells by gentle pipetting and allocated to 20 µL drops of IVC medium, i.e. synthetic oviduct fluid (SOF) including essential and non-essential amino acids and 8 mg/mL bovine serum albumin58. Culture was carried out under humidified air with 5% CO2, 7% O2 and 88% N2 at 38.5 °C. On day 5 post-insemination (pi) the cleavage rate was assessed and the embryos transferred into fresh medium for further 2 days of IVC, when blastocyst rates were recorded.

In vitro embryo production. Unless otherwise stated, reagents were purchased from Sigma Chemical Company (Milano, Italy). The methods for in vitro maturation (IVM) described below have been reproduced in part from Gasparri et al.46. For each replicate, Grade A and B COCs recovered by follicular aspiration were rinsed in HEPES-buffered TCM199 supplemented with 10% fetal calf serum (FCS) and in vitro matured, fertilized and cultured to the blastocyst stage. Briefly, COCs were allocated to 50 µL drops (10 per drop) of TCM199 medium, i.e. in TCM199 buffered with 25 mM sodium bicarbonate and supplemented with 10% FCS, 0.2 mM sodium pyruvate, 0.5 µg/mL FSH, 5 µg/mL LH, 1 µg/mL 17β-estradiol and 50 µg/mL kanamycin, and incubated at 38.5 °C for 21 h in a controlled gas atmosphere of 5% CO2 in humidified air.

The methods for in vitro fertilization (IVF) and culture (IVC) described below have been reproduced from Di Francesco et al. 201232. Frozen straw from a bull previously tested for IVF were thawed at 37 °C for 40 s and sperm was selected by centrifugation (25 min at 300g) on a Percoll discontinuous gradient (45% and 80%). The sperm pellet was re-suspended to a final concentration of 2 × 10⁶ mL⁻¹ in the IVF medium, consisting of Tyrode albumin lactate pyruvate57 supplemented with 0.2 mM penicillin, 0.1 mM hypotaurine and 0.01 mM heparin. Insemination was performed in 50 µL drops of IVF medium under mineral oil (5 oocytes per drop) at 38.5 °C under humidified 5% CO2 in air. Twenty hours after IVF, putative zygotes were removed from the IVF medium, stripped of cumulus cells by gentle pipetting and allocated to 20 µL drops of IVC medium, i.e. synthetic oviduct fluid (SOF) including essential and non-essential amino acids and 8 mg/mL bovine serum albumin58. Culture was carried out under humidified air with 5% CO2, 7% O2 and 88% N2 at 38.5 °C. On day 5 post-insemination (pi) the cleavage rate was assessed and the embryos transferred into fresh medium for further 2 days of IVC, when blastocyst rates were recorded.

Library preparation and sequencing. In total, 20 libraries of small RNA and 20 libraries of RNA-Seq were obtained from five animals per group (n = 5) of two cellular types (OOs and FCs) in both seasons (BS and NBS). Small RNA libraries were prepared using TruSeq Small RNA Library Preparation kit, according to manufacturer’s instructions (Illumina). Small RNA (sRNAs) libraries were pooled together and purified with Agen- court AMPure XP (Beckman, Coulter, Brea, CA) (1 Vol. sample: 1.8 Vol. beads) twice49. The methods described below have been reproduced in part from Frattini et al. 201770. RNA-Seq libraries were generated using the Illumina TruSeq RNA Sample Preparation v2 Kit but with one-half of the recommended reagent volumes. Concentration and quality of RNA were determined by Agilent 2,100 Bioanalyzer (Santa Clara, CA, USA). The isolated RNAs were stored at −80 °C until use.

Data analysis. **miRNA analysis.** Illumina raw sequences were quality checked with FastQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and trimmed with Trimomatic (version 0.32)31, then miRDeep2 (mirDeep2 (version 2.0.0.5)72 was used for miRNA detection and discovery. Known miRNAs available at MirBase (https://www.mirbase.org/) were used to support miRNA identification. In particular, Bos taurus miRNAs were input to support known miRNA detection and miRNAs from related species (sheep, goat and human) were input to support novel miRNA identification. All the identified miRNAs were quantified using the
miRDeep2 quantifier module. The Bioconductor edgeR package (version 2.4) was used to identify statistically significant differential expression between groups of samples (false discovery rate [FDR] < 0.05)73. Predicted miRNA gene targeting of differentially expressed *Bos taurus* miRNAs (DEmiRNAs) was performed with miRWalk2.074, using homologous human miRNAs as input identifiers.

Target genes were submitted to GO analysis. GO classification of the DEGs was performed according to canonical GO categories, using the Cytoscape (version3.2.1) plug-in ClueGO (version 2.3.5) which integrates GO and enhances biological interpretation of large lists of genes75. MicroRNA cluster analysis was performed with Genesis (version1.8.1)76.

**RNA-seq analysis.** RNA-Seq raw data were trimmed using Trimmomatic (version 0.32)71. Sequences were aligned to the buffalo reference genome version UOA_WB_1 (GCF_003121395.1) using STAR_2.3.076. Subsequently, HTSeq-count (version 0.6.1p1) was used to count sequences aligned to each gene. The software package EdgeR of Bioconductor (version 3.6) was used to estimate differential expression between groups of samples77. RNAseq cluster analysis was performed with Genesis (version1.8.1)76. Differentially expressed genes DEGs were submitted to GO analysis, using the Cytoscape (version3.2.1) plug-in ClueGO (version 2.3.5)75. GO and enhances biological interpretation of large lists of genes75. MicroRNA cluster analysis was performed with Genesis (version1.8.1)76.

**In vitro embryo production.** Differences in cleavage and blastocyst rates between seasons were analyzed by Chi square test. The level of significance was set at \( P < 0.05 \).

**Data availability**

RNA-Seq data are available in the Sequence Reads Archive (SRA), BioProject accession number, PRJNA599337. Novel miRNA precursors and novel miRNA mature sequences are reported in Supplementary files S10 and S11.
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Author contributions
E.C. performed the genomic study and was co-author of the manuscript. B.L. performed the bioinformatic study. A.S., B.L. and E.C. assembled and analyzed bioinformatic data and produced figures and supplementary files. M.R., M.K., G.D.V., V.L. and B.G. collected oocytes and produced embryos. A.L.C. coordinated fieldwork logistic, assembled and analyzed data from embryo culture, and was co-author of the manuscript. B.G. designed the study and was co-author of the manuscript. All co-authors provided useful comment on the manuscript.

Competing interests
The authors declare no competing interests.

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