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# Gene expression of pregnancy-associated glycoproteins-1 (PAG-1), interferon-tau (IFNt) and interferon stimulated genes (ISGs) as diagnostic and prognostic markers of maternal-fetal cellular interaction in buffalo cows

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THERIOGENOLOGY

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# ABSTRACT

The aim of this study was to determine the presence of Pregnancy-associated glycoprotein -1 (PAG-1) mRNA expression in the maternal circulation of pregnant buffaloes during the early stage of pregnancy. Contemporaneously, the mRNA expression levels of Interferon-tau (IFNt) and some Interferon stimulated genes (ISGs) (interferon stimulated gene 15 ubiquitin-like modifier interferon, ISG15; Mixoviruses resistance 1 and 2, MX1 and MX2; 2',5'-oligoadenylate synthase 1,OAS1) were evaluated in order to expand our knowledge of the molecular processes involved in the early stages of pregnancy and to identify potential biomarkers of maternal-fetal cellular interaction in buffalo. The study was conducted on 38 synchronized and artificially inseminated buffalo cows (d 0), divided *ex post* into 3 groups: Pregnant (n = 17), Nonpregnant (n = 15) and Embryo mortality (n = 6). Blood samples were collected on d 14, 19, 28 and 40 after artificial insemination (AI) for peripheral blood mononuclear cells (PBMCs) isolation. Expression levels of mRNA of PAG-1, IFNt, ISG15. MX1, MX2 and OAS1 were measured using RT-qPCR. No significant changes were observed in IFNt and PAG gene expressions between groups, while significant differences (p < 0.001) were found for ISG15, MX1, MX2, and OAS1. Pairwise comparisons revealed that the differences between groups occurred on days 19 and 28 post-AI. ISG15 proved to have the best diagnostic performance for distinguishing between pregnant animals and animals that experienced embryo mortality with the ROC analysis. According to the results of the univariate analyses, day 19 was identified as the most indicative to discriminate between groups while the most reliable genes for this differentiation were ISG15, MX1 and MX2. MX2 proved to be the best gene for discriminating pregnant buffaloes using the discriminant analysis, while MX1 was the gene that best predicted embryo mortality. Our results showed that among PAG-1, IFNt and ISGs expression as diagnostic and prognostic markers of maternalfetal cellular interaction in buffalo cows, ISGs proved to be the best peripheral biomarkers for predicting pregnancy and embryonic mortality during the peri-implantation period. These insights into the mechanisms behind maternal-fetal interaction and the development of a method for the early detection of embryo distress may enable us to implement effective strategies to support embryo survival.

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#### 1. Introduction

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https://doi.org/10.1016/j.theriogenology.2023.06.028 0093-691X/© 2023 Elsevier Inc. All rights reserved. During pregnancy, proteins play crucial roles as hormones, regulatory and signalling molecules from embryo formation, implantation in the uterus and further development to term [1,2].



Over the years, various molecular complexes that promote pregnancy maintenance have been identified. Starting from the 1980s [3,4], in ruminants the molecules called pregnancy-associated glycoproteins (PAGs), which are involved in immunological mechanisms in the maintenance of pregnancy and fetal well-being, have been identified [5-7]. These molecules are synthesized by the mononucleate and binucleated trophoblastic cells of species belonging to the order Cetartiodactyla [8–13]. The proteolytic activity that some PAGs share with the aspartic proteinase family members, suggests their possible involvement in placentogenesis, feto-maternal unit remodeling and implantation [14-16]. Once produced and secreted, these glycoproteins enter the maternal bloodstream and can be used as biomarkers for early pregnancy diagnosis in ruminant species [13] including buffalo [8,10,17,18]. Phylogenetic analyses have divided PAGs into two groups: the "ancient" PAGs group (PAG-2), which is deemed to play an important role in implantation and placentation and the "modern" PAGs group (PAG-1), whose function is not entirely clear [13,19]. Detailed information on the expression of PAG-1 mRNA is lacking in ruminants [20,21], especially when detected in peripheral blood cells. To date, the few studies on buffalo PAG gene expression during pregnancy have focused on PAG-2 in maternal blood leukocytes [22] and on PAG-1 on gravid uteruses [23].

Interferon-tau (IFNt) is one of the first molecules involved in early maternal recognition of pregnancy in ruminants [24]. It is secreted by the trophectodermal cells from blastocysts, which generally occurs between day 16 and day 25 of pregnancy [25]. The role of IFNt is to inhibit oxytocin receptor formation in the endometrium in order to prevent the production of prostaglandin F2alpha (PGF2 $\alpha$ ) [26]. This mechanism allows the inhibition of luteolysis, the maintenance of progesterone production by the corpus luteum increasing successful embryo implantation [27]. A recent study that provides strong evidence for the existence of embryo-uterine crosstalk during pre-implantation in buffalo [28], showing that the developing embryo produces IFNt and regulates an immune response by inducing an anti-inflammatory action in uterine epithelial cells for acceptance of the semi-allogenic embryo in the uterus to sustain the pregnancy. Once secreted into the uterine cavity, some of the IFNt enters the uterine vein and stimulates the expression of some genes called interferon-stimulated genes (ISGs), including the interferon stimulated gene 15 ubiquitinlike modifier interferon (ISG15), Mixoviruses resistance 1 and 2 (MX1 and MX2) and 2',5'-oligoadenylate synthase 1 (OAS1) in various cells, namely endometrial cells, peripheral blood cells and luteal cells [29,30]. In cattle, several studies have demonstrated that ISG expression increases in peripheral blood leukocytes during early gestation [31-34], while to date few data are available for buffalo [35-38].

The aim of this study was to evaluated mRNA expression levels of PAG-1, IFNt and some ISGs (ISG15; MX1; MX2 and OAS1) in order to identify potential biomarkers of maternal-foetal cellular interactions in buffalo cows, in an earlier period of gestation respect that in which the proteins can be detect in the maternal blood. To this aim, peripheral blood cells as PBMCs represent a less invasive, and alternative to caruncular tissue, compatible with the maintenance of pregnancy. Identifying relevant circulating biomarkers reflecting the embryo well-being could represent a powerful tool for investigators involved in studying factors affecting embryo mortality.

## 2. Materials and methods

## 2.1. Animals and experimental design

The study was carried out at the CREA Animal Production and

Aquaculture experimental farm, in Monterotondo, Rome, Italy. The experimental procedures were assessed and approved by the CREA Committee of Ethics in Animal Research (Protocol N.0081676–02/11/2020).

In this study 38 Italian Mediterranean pluriparous buffalo cows (days from calving 107.6  $\pm$  20.2) undergoing a synchronization and artificial insemination (AI) program were enrolled and grouped *ex post* as Pregnant (n = 17), Non-pregnant (n = 15) and Embryo mortality (n = 6), based on diagnostic criteria described below in Section 2.2.

The buffalo cows were synchronized with a progesteronereleasing intravaginal device (PRID®) associated with PMSG, PGF2 $\alpha$  and GnRH as reported by Barbato et al. [22]. The cows were artificially inseminated using frozen-thawed semen 72 h after PRID® removal. Blood samples were collected from the jugular vein in 10 mL EDTA tubes on days 14, 19, 28 and 40 post AI, and immediately processed for PBMCs isolation as described below in Section 2.4.

## 2.2. Pregnancy diagnosis

Pregnancy was diagnosed using both conventional transrectal ultrasonography on day 30 post AI and PAGs determination.

By transrectal ultrasonography, the buffaloes were considered pregnant if an embryonic vesicle and embryo proper with beating heart were detected, while in the absence of these conditions buffaloes were considered as non-pregnant. When a vesicle without an embryo proper and/or an embryonic heart beat were not visible [39]an embryo mortality was supposed to be occurred and validated with the PAGs results.

PAGs determination was adjunct to conventional ultrasonography to diagnose pregnant and not pregnant animals and monitor the pregnant animals throughout the sampling period to highlight embryo mortality. Blood sample were collected from the jugular vein in 10 mL EDTA tubes on days 0, 14, 19, 28 and 40 post AI, and the plasma was separated by centrifugation at  $2700 \times g$  for 10 min and then stored at -20 °C until assayed. Based on PAGs plasma levels (cut-off value: 1 ng/mL) [39], the buffaloes were considered non-pregnant when concentrations remained close to zero throughout sampling and pregnant when the concentrations were >1 ng/mL at d 28 and d 40 post-AI. The animals in which PAGs concentration was found close to the cut-off (0.8–1 ng/mL) between 14 and 28, and then dropped below 0.2 ng/mL on day 40, embryo mortality was deemed to have occurred.

PAGs concentration was used to differentiate *ex post* the groups.

# 2.3. PAGs radioimmunoassay

RIA-860 previously described by Barbato et al. [10] was used to determine PAGs concentration. Pure boPAG<sub>67kDa</sub> preparation was used as the standard and tracer. Iodination (Na–1<sup>125</sup>, Amersham Pharmacia Biotech, Uppsala, Sweden) was carried out according to the Chloramine-T method previously described by Greenwood et al. [40]. The samples were assayed in a preincubated system in which the standard curve ranged from 0.2 to 25 ng/mL. The minimum detection limit (MDL), calculated as the mean concentration minus twice the standard deviation (mean-2 SD) of 20 duplicates of the zero (B0) standard [41], was 0.4 ng/mL. The intra- and inter-assay coefficients were 2.5% and 7.2%, respectively.

## 2.4. PBMCs isolation

The purification of the PBMCs was obtained following a density gradient using Lymphoprep<sup>TM</sup> (1.077 g/mL; Axis-Shield). The blood samples (10 mL) were collected from the external jugular vein in

vacutainer tubes containing anticoagulant EDTA. The protocol used for isolating cells was described by Barbato et al. [22]. In short, 8 mL of whole blood was diluted 1:1 with Hank's Buffered Salt Solution (HBSS) (without  $Ca^{2+}$  and  $Mg^{2+}$ ) into 50 mL Falcon® tubes and then carefully layered over 8 mL of Lymphoprep<sup>™</sup> at room temperature. The tubes were centrifuged at  $800 \times g$  for 30 min at 20 °C. The phase containing erythrocytes settled at the bottom of the tube. The phase containing PBMCs settled on the plasma:Lymphoprep™ interface. The upper plasma layer was removed and discarded without disturbing the plasma:Lymphoprep<sup>™</sup> interface. The PBMCs layer was collected at the plasma:Lymphoprep<sup>™</sup> interface without disturbing the erythrocyte/polymorphonuclear leukocytes pellet. Afterward, the PBMCs were washed twice with HBSS (without  $Ca^{2+}$  and  $Mg^{2+}$ ). Total cell counts and cell viability were determined by LUNA-II<sup>™</sup> automated cell counter (Logos Biosystems, Anyang, South Korea). The purity of the isolated PBMCs fraction was evaluated with a CytoFlex flow cytometer (Beckman Coulter) based on the forward scatter channel (FSC) vs the side scatter channel (SSC). PBMCs purity was on average 98%.  $5 \times 10^6$  cells were lysed in TRIzol<sup>TM</sup> Reagent (Invitrogen, Life Technologies, Carlsbad, CA, USA) and immediately stored at -80 °C until further investigation analysis.

## 2.5. RNA extraction

Total RNA was extracted from the PBMCs lysed in 1.0 mL TRIzol<sup>TM</sup> Reagent according to the manufacturer's protocol. Lastly, the RNA concentration and quality were determined using a spectrophotometer (BioPhotometer, Eppendorf, Hamburg, Germany) with the measurement of absorbance and ratio at 260 nm and 280 nm wavelength. In order to ensure the RNA purity and that it is free from DNA contamination, a further treatment was performed with DNase I, RNase-free (1 U/µL) (Invitrogen, Life Technologies, Carlsbad, CA, USA). The samples were then stored at -80 °C until the next analysis.

#### 2.6. Reverse-transcription and RT-qPCR

Total RNA (1  $\mu$ g) was reverse-transcribed using the High Capacity cDNA Reverse Transcription Kit (Applied Biosystem, Foster City, CA, USA), according to the manufacturer's instructions. The reverse transcription was performed in the following sequence: step 1, incubation at 25 °C for 10 min; step 2 incubation at 37 °C for 2 h; and step 3 incubation at 85 °C for 5 min.

The cDNA obtained from each sample was used as a template for RT-qPCR. In order to optimized the 25  $\mu$ l reaction volume in MicroAmp optical 96-well plates, as previously described by Riva et al. [42], each plate contained duplicates of each sample, 2X Power Sybr Green PCR Master Mix (Applied Biosystem, Foster City, CA, USA) and primers at 500 nM each (0.5  $\mu$ l of 10  $\mu$ M solution).

The species-specific primer pairs were designed by Primer Express Software (Applied Biosystems, Foster City, CA, USA) using the buffalo sequences taken from NCBI database as target, and the primer pairs specificity checked using NCBI Primer-BLAST (https://www.ncbi.nlm.nih.gov/tools/primer-blast/). The primers were purchased from IDT<sup>™</sup> (Integrated DNA Technologies, Coralville, Iowa, USA).

The primer sequences are listed in Table 1. *BuPAG 18* isophorm was selected for PAG-1 group, because among those present in the database is the earliest expressed. Relative quantification of PAG-1, IFNt and ISGs transcript was carried out following the MIQE guidelines [43].

A duplicate no-template control (NTC) was also included on each plate. RT-qPCR was carried out using the 7000 Sequence Detection System (Applied Biosystem, Foster City, CA, USA) at the following thermal cycle conditions: 10 min at 95 °C followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. Quantification was determined after applying an algorithm to the data analyzed by the software of the 7000 Sequence Detection System (Applied Biosystem, Foster City, CA, USA).

Sample amplification specificity was verified by dissociation curves and agarose gel electrophoresis. Prior to sequencing, the PCR products were purified using the QIAquick PCR Purification Kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions. Each PCR-fragment was sequenced and its sequence aligned to the corresponding reference sequence before conducting the qPCR-analyses to assess the amplification of the correct gene.

qPCR amplification efficiency was determined using the slope of the standard curve: Efficiency =  $10^{-1/\text{slope}}$  –1. Only PCR reactions with PCR efficiencies >95% were included in the subsequent analyses. In detail, the amplification efficiency was: for the IFNt E = 112.7%; ISG15 E = 104.62%, for MX1 E = 101.91%, for MX2 E = 95.72%, for OAS1 E = 103.27%, for PAG 18 E = 144.12%, and finally for ACTB the amplification efficiency was 98.80%.

The expression of buffalo target genes was normalized to betaactin (ACTB) reference gene levels (mean) of the same sample and run. The  $2^{-\Delta Ct}$  method was used to calculate the relative expression levels of the target genes [44]. The obtained value was multiplied by 10.000 in order to obtain the test Arbitrary Units (AU).

## 2.7. Statistical analysis

Diagnostic graphics were used to check assumptions and outliers. Gene expressions and PAG plasma concentrations were analyzed by the Generalized Linear Model (GLM) using gamma as probability distributions and log as link function as these variables were positively skewed with a long tail on the right [45]. Animals were included in the models as subjects while Time was included as repeated effects. The models evaluated the main effects of Time (4 levels: 14, 19, 28 and 40 days post-AI), Group (3 levels: Pregnant, Non-pregnant, and Embryo mortality), and their interaction. The last significant difference was used for carrying out multiple comparisons.

Receiver Operating Characteristic (ROC) analyses and the Youden index were used to evaluate the diagnostic performance of each gene and to determine their optimal cut-off value to distinguishing between pregnant and non-pregnant cows and pregnant cows and cows experiencing embryonic mortality [22,46].

Based on the results of the univariate analysis, a genes panel that best discriminates among groups was selected. The discriminant analysis (DA) was then performed to identify the linear combinations of genes (discriminant functions (Dfs)) that best discriminate between Pregnant, Non-pregnant, and Embryo mortality groups. The DA also classified the genes according to their importance in this discrimination. The relative contribution of each gene in classifying the buffalo cows was evaluated using the structure coefficients (correlations between each independent variable and the Df). The centroids indicated the mean discriminant scores of Df for pregnant, non-pregnant, and buffaloes that experienced embryo mortality, which were used to establish the cut-point for classifying samples during cross-validation. The performance of the DA was evaluated by classification-related statistics [47,48].

The statistical analyses were performed with SPSS 25.0 software (SPSS Inc. Chicago, IL, USA). A p value  $\leq$  0.05 was considered to be statistically significant and p < 0.1 indicated a trend.

# 3. Results

A total of 17 out of 38 buffalo cows enrolled in this study became pregnant and 15 remain non-pregnant, while 6 experienced embryo mortality.

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### Table 1

The primer sequences of buffalo assays used in this study.

Name	Sequence $(5' \rightarrow 3')$	NCBI RefSeq.	Amplicon
IFNt Fw Rev	CATCTTCCCCATGGCCTTC CAACCAAGAGATTCTGCCGG	XM_025281435.1	81
ISG15 Fw Rev	AGAACCCACGGCCATGG GATCTTCTGGGCGATGAACTG	NM_001291322.1	121
MX1 Fw Rev	TTGAAAAACGGTATCGTGGCA TCTTGCTGTCACTGTGCGC	XM_025279556.1	131
MX2 Fw Rev	TGCAATCTGCATCAAACAGTCC GGTCTTGACAAAAAACCAGCTTCT	NM_001290849.1	121
OAS1 Fw Rev	ATCGTCCACCCAAGCTGAAG GCAGGTTATTCTCATGCTCCTTACA	XM_025267539.1	81
PAG-1 (BuPAG 18) Fw Rev	TTTGCACCAGTCCAACCTGTT AGGTCTTATTGGTAGGCCGGA	XM_025286464.1	81
ACTB Fw Rev	AACCGTGAAAAAATGACCCAGAT CCGTCACCGGAGTCCATC	XM_025274489.1	131

# 3.1. Univariate analyses

No significant changes were found in IFNt and PAG gene expressions (p value for all the effects >0.1) while all the evaluated effects (i.e., Time, Group, and their interaction) were significant at 0.1% significance level for ISG15, MX1, MX2, and OAS1 (Fig. 1).

Pairwise comparisons pointed out that the differences between groups occurred on days 19 and 28 post-AI. In particular, the ISG15, MX1, MX2, and OAS1 expression levels of pregnant buffalo cows were found to be highest on days 19 and 28 (p < 0.001). The buffalo

cows that experienced embryo mortality tended to have higher values of ISG15, MX1 and MX2 than non-pregnant animals on day 19. Their expressions decreased on day 28 (p < 0.1), and other differences between the embryo mortality and non-pregnant groups were not observed (Fig. 1).

# 3.2. ROC analyses

ROC analyses were performed using the data of day 19 post-AI when previous analyses showed the greatest between-groups



Fig. 1. IFNt (A), PAG (B), MX1 (C), MX2 (D), OAS1 (E) ISG15 (F) and gene expression in PBMCs of pregnant and non–pregnant buffalo cows and buffalo cows that experienced embryonic mortality. Bars not sharing any superscript within each time are significantly different at P < 0.05. When letters are not present, no effect was statistically significant.

differences. Table 2 and Fig. 2 show the area under the ROC curves and the cut-off value of the examined genes. ISG15, MX1, and MX2 were perfectly capable of distinguishing pregnant from nonpregnant buffalo cows (for each gene, AUC = 1.000) and, showed 100% sensitivity and specificity at their optimal cut-off value. OAS1 also proved to have excellent diagnostic performances (AUC = 0.996) while the diagnostic performance of IFNt and PAG performed poor in all classifications.

ISG15 proved to have the best diagnostic performance for distinguishing pregnant cows from cows that experienced embryonic mortality (AUC = 0.980, p < 0.01). The optimal cut-off value showed a sensitivity and a specificity of 88.2% and 100%, respectively.

The performance of genes to distinguish Non-pregnant from Embryo mortality groups was generally lower. The highest accuracy was found for MX2 (AUC = 0.789, p < 0.05) and the optimal cut-off value showed 66.7% of sensitivity and 93.3% of specificity.

#### 3.3. Discriminant analysis

Based on the results of univariate analyses, day 19 proved to be the most indicative for differentiating the groups, while ISG15, MX1 and MX2 were identified as the best genes for this differentiation. Therefore, the DA was performed using the data collected on day 19 and including the above-mentioned genes as independent variables.

Centroids (Table 3) and the plot of Df scores (Fig. 3) indicated that Df1 was able to discriminate the pregnant cows from the others (i.e., non-pregnant cows and those experiencing embryonic mortality). The results of the multivariate analysis showed that MX2 was the gene that best discriminated pregnant cows (followed by MX1 and ISG15). The second Df separated the Embryo mortality group (with a positive coefficient) from the others (i.e., Pregnant and Non-pregnant groups showing negative coefficients). MX1 was the gene that best discriminated embryo mortality (followed by ISG15 and MX2). Overall, the DA correctly classified 76.3% of the buffalo cows.

### 4. Discussion

This study investigated the mRNA expression of two molecules that play key roles in maternal-fetal recognition namely PAG-1 and IFNt, as well as the mRNA expression of several IFN-stimulated genes (ISG15; MX1; MX2 and OAS1) on blood circulating cells in order to identify potential biomarkers reflecting embryo well-being during early pregnancy in buffalo.

To our knowledge no previous research has investigated PAG-1 mRNA expression in circulating PBMCs of buffalo cows.

In this study, the PAG-1 expression levels (BuPAG 18 isoform) in PBMCs did not show statistical differences over all time points and groups analyzed. This lack of difference may be due to the fact that PAGs belong to a very large and complex family of genes [13]. The PAGs family exhibits a spatially and temporally distinct expression patterns during pregnancy: PAG-2 group expression occurs throughout the outer epithelial layer of the placenta (trophectoderm), while PAG-1 group expression is restricted to the binucleate cells [49,50]. Consequently, some PAGs are expressed in certain stages of pregnancy and not in others. In bovine cotyledon tissues, Wiedemann et al. [21] observed high expression levels in middle and late pregnancy for the PAG-1 group and in early pregnancy for the PAG-2 group. Therefore, PAG-1 group is supposed to play an essential role in the maintenance of pregnancy, while the PAG 2 group is supposed to play an important role in placentation and the maintenance of early pregnancy [20,51,52]. However, these findings derived from studies conducted on PAG mRNA expression in placental tissue. Research carried out on PAG-2 mRNA expression in circulating blood cells reported in our previous study [22] highlights early PAG-2 expression on day 14 post AI in pregnant buffalo. This result is in line with those found by other authors on placental tissue: days 17–19 in bovine [53], day 18 in caprine [49] and day 13 in sheep [50], which coincide with the beginning of implantation.

These findings confirm that PAG-1 expression occurs later in pregnancy in buffalo as in other animal species, which is in line with the results obtained by Lotfan et al. [23], who found high levels of *BuPAG 18* isoform expression starting from day 45 of pregnancy in buffalo. Therefore, our results could suggest that PAG-1 is required for the maintenance of pregnancy rather than for pregnancy establishment.

As previously reported for PAG-1 expression, the results obtained for IFNt expression showed no significant differences between the groups and time points considered. In fact, according to our results it is difficult to find INFt gene expression in blood cells. Actually, despite the fact that IFNt transits from the uterus into the

#### Table 2

Results of the Receiver Operating Characteristic (ROC) curve analyses. Sensitivity, specificity and area under the curve (AUC) for the optimal cut-off value for the genes under study.

Discrimination	Gene	Area	95% Confidence Interval	Sig. <sup>a</sup>	Optimal Cut-off	Sensitivity (%)	Specificity (%)
Pregnant vs Non-pregnant <sup>b</sup>	IFNt ISG15 MX1 MX2 OAS1 PAG	0.529 1.000 1.000 1.000 0.996 0.476	0.313-0.745 1.000-1.000 1.000-1.000 1.000-1.000 0.983-1.000 0.256-0.696	0.777 <0.001 <0.001 <0.001 <0.001 0.827	0.20 253.1 995.4 568.7 753.6 0.01	64.7 100 100 94.1 80.0	53.3 100 100 100 100 35.7
Pregnant vs Embryo mortality <sup>b</sup>	IFNt	0.627	0.347-0.908	0.363	0.19	64.7	83.3
	ISG15	0.980	0.931-1.000	0.001	470.39	88.2	100.0
	MX1	0.882	0.698-1.000	0.006	1241.83	88.2	83.3
	MX2	0.961	0.877-1.000	0.001	969.81	76.5	100.0
	OAS1	0.912	0.745-1.000	0.003	666.88	100.0	83.3
	PAG	0.452	0.202-0.703	0.741	0.02	57.1	50.0
Non pregnant vs Embryo mortality <sup>c</sup>	IFNt	0.444	0.199-0.690	0.697	0.12	100.0	26.7
	ISG15	0.700	0.396-1.000	0.161	79.71	66.7	93.3
	MX1	0.744	0.509980	0.087	830.83	66.7	80
	MX2	0.789	0.566-1.000	0.043	326.34	66.7	93.3
	OAS1	0.689	0.443-0.935	0.186	486.59	50.0	80.0
	PAG	0.548	0.297-0.798	0.741	0.01	100.0	35.7

<sup>a</sup> Null hypothesis: true area = 0.5.

<sup>b</sup> The positive actual state is Pregnant.

<sup>c</sup> The positive actual state is Embryo mortality.



Fig. 2. Receiver Operating Characteristic (ROC) curves of ISG15, MX1, and MX2 to discriminate between pregnant and non-pregnant buffalo cows (A), ISG15 to discriminate pregnant buffalo cows from buffalo cows that experienced embryonic mortality (B), and MX2 to discriminate non-pregnant buffalo cows from buffalo cows that experienced embryonic mortality (C).

systemic circulation to exert its effects on maternal physiology, it has been noted that IFNt expression in blood is not easily detectable [54,55]. Therefore, it is preferable to measure the total protein in blood.

However, in our recent study [38] significant differences in INFt expression were observed in PBMCs between pregnant and nonpregnant buffaloes on days 28 and 40 after Al. In our opinion, these dissimilarities may be due to differences between the methodology applied in this study and the methodology used in our previous study such as different primer-sequence (Fw: CCATTCTGACCGTGAAGAAGTA- Rev: TCATCTCCACTCTGATGATTTCC, (RefSeq.: *AY535404.1*) [38] vs Fw: CATCTTCCCATGGCCTTC - Rev: CAACCAAGAGATTCTGCCGG, (RefSeq. *XM\_025281435.1*) respectively) and different fluorescence detection methods in qPCR (Probe system vs Sybr Green, respectively).

Since it is difficult to evaluate the direct expression of IFNt in blood, the response of circulatory leucocytes to IFNt through the Interferon-Stimulated Genes (ISGs) was used as an alternative method for IFNt expression evaluation. In fact, Matsuyama et al., [33] detected a significant increase of ISGs mRNA expressions in PBMCs, especially ISG15 and MX2, following exogenous IFNt administration on days 16–17 after standing oestrus in

### Table 3

Results of the Discriminant analysis: structure coefficients and centroids of discriminant function 1 (Df1, separating the Pregnant from the Non-pregnant and Embryo mortality groups) and function 2 (Df2, separating Embryo mortality from the Pregnant and Non-pregnant groups).

Gene	Structure Matrix		
	Df1	Df2	
MX2 MX1 ISG15	0.986 0.964 0.893	-0.077 0.256 -0.160	
Group	Functions at Group Centroids		
	Df1	Df2	
Pregnant Non-pregnant Embryo mortality	1.348 -1.210 -0.794	-0.050 -0.290 0.866	

synchronized and artificially inseminated cows. This result is in line with the findings of Oliveira et al. [56] and Bazer et al. [57] who observed that ISGs expre during early pregnancy in sheep is induced by the endocrine release of IFNt from the uterine vein. These findings suggest that the quantity of IFNt produced by conceptus could be monitored indirectly.

In our study, all of the ISGs analyzed were more strongly expressed in the Pregnant group than in the Non —pregnant and Embryo mortality groups, and were highly expressed on d 19 and d 28 post AI. These results are in line with the observations by Thakur et al. [36] and Mishra and Sarkar [58], who reported that the gene expression profiles of OAS1, MX2 and ISG15 in buffalo PBMCs increased between days 14 and 20 of pregnancy and declined thereafter. Our findings are also in agreement with the results reported by Nag et al. [37], which showed a significant increase in the expression of ISG15 in buffalo PBMCs on days 18 and 20 and of MX1 on days 14–20 of pregnancy. Recently, Huidrom et al. [28] investigating the effect *in vitro* of pre-implantation embryo on the ISGs



Fig. 3. Plot of discriminant function scores.

expression in buffalo, also found that ISG15, MX2 and OAS1 were upregulated in PBMCs when treated with conditioned medium derived from embryos culture. Still with regard to buffalo, Jain et al. [35] reported that during early pregnancy ISG15 was detected in different reproductive tissues.

A substantial body of literature on other ruminants shows also that the gene expression of MX-1 and OAS-1 genes in peripheral blood is higher in pregnant than in non-pregnant animals during early pregnancy [29,34,59,60]. Studies conducted on cattle in early pregnancy have shown that ISGs are up-regulated in leukocytes and the mean blood levels of ISG15 mRNA of pregnant cows are higher from days 15–32 of gestation compared to non-pregnant cows [32]. In addition to ISG15, the same results were also observed for MX1, MX2 and OAS1 on day 18 of pregnancy [31,61,62]. In sheep, 15–30 days after insemination, higher levels of MX1 mRNA in PBMCs were found in pregnant than in non-pregnant ewes [63].

To our knowledge, no data have been reported in literature regarding a cut-off value for predicting pregnancy or embryonic mortality for the examined genes in buffalo.

Our results showed that all ISGs were perfectly capable of distinguishing pregnant from non-pregnant animals on day 19 post AI, while IFNt and PAG genes performed poorly. The ROC analysis showed that the MX genes are the most reliable markers for distinguishing pregnant animals and those that experienced embryonic mortality, which was also determined using the discriminant analysis. ISG15 proved to have the best diagnostic performance for discriminating pregnant cows and animals that experienced embryo mortality using the ROC analysis.

By measuring ISGs expression in bovine blood to diagnose pregnancy within 18–20 days post-AI, Green et al. [32] reported an optimal cut-off value for d 18 of pregnancy using OAS1 for heifers while there was a lesser response for cows that reduced the sensitivity of the ISG test on that day.

Yoshino et al. [64], suggested that the ISG15 and MX2 values in blood were reliable indicators of pregnancy between 20 and 22 days post-AI in cows, with 80% accuracy.

In our study, buffalo cows that experienced embryo mortality tended to have higher ISG15, MX1 and MX2 values than nonpregnant cows on day 19. The genes that showed the best diagnostic performance (i.e., ISG15, MX1, and MX2) were then selected for the Discriminant analysis (DA). The DA was conducted with the gene expression recorded on day 19 post AI, when between-group differences were greatest. Discriminant analysis is a multivariate technique that identifies the combinations of variables that best discriminate between groups. We hypothesized that a combination of genes could discriminate between pregnant and non-pregnant buffalo cows and cows that experienced embryo mortality. This analysis could also classify the genes according to their importance in this discrimination. Our findings indicated that MX2 is the gene that best distinguishes the pregnant buffalos from the other group (i.e., non-pregnant ones and those experiencing embryonic mortality). This finding is in line with the observations made by Buragohain et al. [65] who showed an increase of MX2 gene expression in the peripheral blood of pregnant buffalo cows on days 14 and 28, with a ten-fold and fourteen-fold increase in plasma protein levels, respectively. This trend of MX2 expression was also observed in our previous study [38] in which mRNA ISG expression was either evaluated in PBMCs or PMNs.

In cattle, Miagawa et al. [66] and Pugliesi et al. [67] reported MX2 expressed at higher levels in PBMC samples starting from day 15 which peaked on day 20 of pregnancy. The high gene expression of MX2 in pregnant animals could due to the fact that blastocyst elongation coincided with these days, which corresponds to the period when the conceptus synthesizes large amounts of IFNt [24].

In our study, the gene that could best predict embryo mortality according to the multivariate analysis proved to be MX1 on day 19 post AI. In cattle, Sheikh et al. [55] showed that MX1, ISG15 and OAS1 showed higher levels of expression in pregnant cows between 14 and 21 days of gestation than in cows with fertilization failure or embryonic mortality. Studies conducted by Kose et al. [68] in sheep showed that MX1 and ISG15 expression could be used as markers for embryo mortality after observing that the expression of these genes was up-regulated on day 15 and remained high until day 23 and only decreased on day 25 in the group that experienced embryonic mortality.

The results of the discriminant analysis showed that MX2 and MX1 were the best genes to discriminate pregnancy or embryonic mortality, respectively, which may be due to the fact that MX gene expression is up-regulated during early pregnancy to block potential viral infections that could lead to pregnancy loss [67,69] or because of their role in regulating endometrial secretion, uterine remodeling and anti-luteolytic activities [70]. It has been suggested that MX proteins possess GTPase activity which might be involved in the calcium metabolism of endometrial cells during early pregnancy [71] and might trigger maternal neutrophil migration towards the developing embryo [62].

## 5. Conclusions

Our results indicated that when PAG-1, IFNt and ISGs expression were used as diagnostic and prognostic markers of maternal-fetal cellular interaction in buffalo cows, ISGs proved to be more reliable peripheral biomarker for the prediction of pregnancy and embryonic mortality in buffalo during the peri-implantation period. These insights into the mechanisms underlying maternalfetal interaction and the identification of a strategy for the early detection of fetal distress could prove useful for embryo survival.

Although a larger sample size of animals is required to provide more accurate values, these preliminary results may help us to identify a biomarker capable of correctly predicting reproductive failure in early pregnancy, which could facilitate the implementation of reparative mechanisms with the aim of improving fetal well-being and reproductive performance in buffalo cows.

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#### **CRediT authorship contribution statement**

**A.B. Casano:** Conceptualization, Methodology, Writing – original draft. **L. Menchetti:** Formal analysis, Visualization. **M. Trabalza-Marinucci:** Writing – review & editing. **F. Riva:** Investigation, and, Writing – review & editing. **G. De Matteis:** Investigation, and, Writing – review & editing. **G. Brecchia:** Writing – review & editing. **A. Inglesi:** Investigation. **E. Rossi:** Investigation. **F. Signorelli:** Investigation. **V.L. Barile:** Conceptualization, Methodology, Writing – original draft. **O. Barbato:** Conceptualization, Methodology, Writing – original draft.

## **Declaration of competing interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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