Comparison of Cleavage Rate of Buffalo Oocytes on Supplementing with Different Concentration of Serum Gonadotropin, Estradiol 17b, Estrus Buffalo Serum and Buffalo Follicular Fluid

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Abstract

Serum gonadotropin @50 and 100IU/ml, estrodiol-17β @0.5 and 1μg/ml, EBS @5 and 10% and BFF @5 and 10% supplementation is done on cleavage rate of buffalo oocytes for the comparison of efficacy of hormones. Ovaries were collected from buffaloes slaughtered at local abattoir. Oocytes were aspirated and good and excellent quality oocytes were matured in basic maturation media TCM199 supplementing above mentioned supplements separately. The recovery rate of culturable oocytes was 85.33%. The highest cleavage rate was documented with supplementation of 10% EBS and the lowest was observed with 10% BFF. The cleavage rate was significantly (p≤0.05) higher in the group supplemented with 5%EBS and 10% EBS as compared to all other groups of the experiment. It can be concluded that the supplementation of EBS to IVM medium has positive effect on cleavage rate.

Keywords
Cleavage rate, Oocytes, Serum gonadotropin, Follicular fluid

Introduction

The buffalo forms the backbone of India’s dairy industry and is considered as the ‘bearer cheque’ of the rural flock and India’s milking machine (Balain, 1999). As per the 19th livestock census India has around 108.7 million of buffalo population and constitutes around 21.23% of total livestock population. Buffalo population of Karnataka is 33.06 lakhs. Buffalo contributes 51% (132430 tonnes) of total nation’s milk production and around 1103.85 tonnes of meat production of total nation’s meat production (Department of Animal Husbandry Dairying and Fisheries, ministry of agriculture, New Delhi, 2014).

The buffalo is the predominant domestic animal for milk and meat production. On average, buffalo is four times as productive as
an average indigenous cow in India. India has world’s best buffalo dairy breeds and provides superior buffalo germplasm to several countries of the world (Kaikini, 1992).

The domestic water buffalo plays a key role in milk production in many Asian countries including India, and they are able to withstand the adverse environmental conditions of the tropics. However the buffalo are traditionally considered to have a low reproductive efficiency (Pankaj, 2015). The low reproductive efficiency in female buffalo can be attributed to delayed puberty, higher age at calving, long postpartum anoestrus period, long calving interval, lack of overt sign of heat, and low conception rate. In addition, female buffaloes have few primordial follicles and a high rate of follicular atresia. These factors are responsible for driving a large number of buffaloes to slaughter house either prior to maturity or after calving, which has resulted in enormous loss of genetic resources and significant decline in their number in recent years. Thus there is a need to improve reproductive performances of buffaloes.

**In vitro** maturation (IVM) of oocytes and **in vitro** fertilization (IVF), **in vitro** culture of embryos and embryo transfer technology appear to be useful techniques for the improvement of reproductive efficiency of buffaloes (Uma, 1997). Embryo transfer technology has made rapid strides in dairy cattle industry and has become viable commercially in some of advanced countries. Therefore, the emphasis has now shifted to **in vitro** embryo production (IVEP) as it can salvage the genetic potential from infertile female and can yield large number of embryos (Kumar and Anand, 2012). Ovaries of the slaughtered animals are the cheapest and the most abundant source of primary oocytes for large scale production of embryos through **in vitro** maturation (IVM) and **in vitro** fertilization (IVF) (Agrawal et al., 1995).

**In vitro** maturation (IVM) is the one of the essential step and starting point for lot of biotechnological applications in animals like **in vitro** fertilization (IVF), cloning, transgenic animal production and embryonic stem cell research. Oocytes maturation is the process of complex changes in the protein phosphorylation which transform the primary oocytes in to mature secondary oocytes. Maturation of the oocytes included two aspects viz., nuclear and cytoplasmatic maturation. The nucleus and the cytoplasm of the oocytes undergo many changes during maturation, making it receptive to fertilization and competent to support embryonic development (Tomek et al., 2002).

Several workers have studied different aspects of IVM in mammalian oocytes. In most of the studies revealed that media designed originally for cell culture, are not capable of supporting high levels of oocytes maturation and, therefore, are commonly supplemented with sera and hormones, growth factors and antioxidants to improve the maturation rates at high level (Kumar and Anand, 2012). The maturation medium with the selection of protein supplements, hormones for IVM play an important role in subsequent IVF and **in vitro** development.

The identification of substances capable of delaying the nuclear maturation time and thus allowing cytoplasmic and nuclear changes to occur synchronously has been the subject of several studies. Follicular fluid (FF), consisting of electrolytes, hormones, amino acids, growth factors, among other components, has been used as a natural substance for blocking the meiosis (Aguilar et al., 2001). Some studies have documented that maturation medium supplemented with FF provides appropriate environment to bovine oocytes development (Romero-Arredondo and Seidel, 1996), since it increases the degree of cumulus cells
expansion (Aguilar et al., 2001) and enhances the embryonic development (Algriany et al., 2004).

Supplementation of serum in media had favourable effect on maturation. The serum contains a number of known growth factors that have an important role in the regulation of oocyte maturation, particularly via cumulus cells, it also prevents the hardening of the zona pellucida; moreover, the beneficial action of serum may be due to its antioxidant properties (Mahmoud and Nawito, 2003). Estrus buffalo serum (EBS), a rich source of hormones and growth factors, could be used to improve the developmental competence of buffalo oocytes culture in vitro (Abid et al., 2008).

An attempt to induce in vitro maturation can only be successful in conditions that are similar to those existing in vivo. Therefore, the biochemical composition of the oocyte culture media has great influence on the success rate of in vitro maturation. Since, under in vivo conditions the oocytes maturation is influenced by various steroid and gonadotropic hormones, it is necessary to supplement the steroid and hormones to the medium in order to sustain the growth and development of oocytes either by directly or through the hormone rich serum medium. Therefore the development of a suitable culture system and components of maturation media for in vitro maturation of oocytes is a major component of the in vitro embryo production procedures.

Perusal of literature indicates less number of works on optimizing the in vitro maturation of buffalo oocytes as compared to the other species.

Considering all these points present study was designed to investigate the effect serum gonadotrophin, estradiol-17β, estrous buffalo serum (EBS) and buffalo follicular fluid (BFF) supplementation on CLEAVAGE rate with the following objective

Materials and Methods

Materials

Media, chemical and plastic wares

All the media and chemicals were procured from Himedia Laboratories, Mumbai unless otherwise stated. Folligon® (Pregnant Mare Serum Gonadotrophin, 1000IU/vial) and Chorulon® (Human Chorionic Gonadotrophin, 1500IU/vial) were purchased from Intervet International B.V., Boxmeer, Netherland. The disposable plastic wares used in this experiment were obtained either from Tarsons Products Private Limited, Kolkata or from Himedia Laboratories, Mumbai.

All the media were prepared by using sterile tissue culture grade water. All the working solutions/media excluding OCM were kept for at least 3-4hr in CO₂ incubator at 38°C, 5%CO₂ and 95 % relative humidity for quenching before use. Prepared OCM was kept in BOD at 37°C. The stocks of media were stored at 4°C and used within one month.

Sterilization procedures

Sterilization of all media was done by filtering through 0.2µm syringe driven filters. The glassware and micropipette tips were sterilized by autoclaving at 121°C for 30min. Fresh sterilized and disposable culture bottles, petridishes, tubes and syringe were used every time. All the equipments were exposed to UV light for 15 minutes before use. In order to avoid bacterial and fungal contamination all the procedure except the aspiration of oocytes, starting from oocyte searching to in vitro culture works and media preparation
were carried out in highly sterile condition under laminar flow cabinet.

**Preparation of buffalo serum and estrous buffalo serum (EBS)**

The blood was collected from jugular vein of normal cycling buffaloes that are not in estrus as well as from the buffaloes that are in estrus for the separation of buffalo serum and estrus buffalo serum respectively. The collected blood was kept in slant position and allowed to clot, later blood was centrifuged at 3000rpm for 15min for serum separation. The separated serum was heat inactivated at 56°C for 30minutes, filtered through 0.45µm filter and stored at -20°C in 2.0ml micro centrifuge tubes as aliquots for future use. Same batch of pooled serum was used for all the trials. The buffalo serum was used in the preparation of working oocyte collection medium (OCM) whereas EBS was used as supplement in T5 and T6 groups.

**Collection of buffalo follicular fluid (BFF)**

Ovaries were collected immediately after slaughter of buffaloes of unknown reproductive status at the local abattoir. The ovaries were maintained in a thermos flask containing warm (35-37°C) normal saline (0.9% NaCl) fortified with 50µg/ml gentamycin sulphate (Gentalab). The ovaries were transported to the laboratory within 2 hrs after slaughter of animals. At the laboratory, the buffalo ovaries obtained from the abattoir were rinsed thoroughly with fresh sterile normal saline supplemented with gentamycin @50µg/ml 5-6 times and final was done with Phosphate buffered saline (PBS). Follicular fluid was aspirated from non atretic, surface follicles (>3mm diameter) of the ovaries using 5ml syringe attached with 18-gauge needle. The pooled follicular fluid was allowed to settle for 10mins and the supernatant was collected. The collected follicular fluid was sterilized by filtering through 0.22µm syringe driven filter and stored in sterile micro centrifuge tubes of 2.0ml capacity at -20°C for subsequent use for IVM as supplement in T7 and T8 groups.

**Collection of ovaries and semen sample**

Buffalo ovaries were collected immediately after slaughter of the animals of unknown reproductive status at the local abattoir. The ovaries were maintained in a thermos flask containing warm (35-37°C) normal saline (0.9% NaCl) fortified with 50µg/ml gentamycin sulphate (Gentalab). The ovaries were transported to the laboratory within 2 hrs after slaughter of animals.

The frozen buffalo bull semen straws supplied by Department of Animal Husbandry and Veterinary Services, Government of Karnataka to the Department of Veterinary Gynaecology and Obstetrics, Veterinary College, Bidar were utilized for *in vitro* fertilization.

**Methods**

**Oocytes collection, processing and grading**

Ovaries were collected immediately after slaughter of adult female buffaloes of unknown reproductive status at the local abattoir. The ovaries were maintained in a thermos flask containing warm (35-37°C) normal saline (0.9% NaCl) fortified with 50µg/ml gentamycin sulphate. The ovaries were transported to the laboratory within 2hrs after slaughter of animals. In the laboratory, ovaries were washed 6-7 times in warm normal saline solution (37°C) fortified with antibiotic and final two washings with PBS. Washed ovaries were submerged in PBS solution in sterile beaker and were used for oocyte collection. Oocytes were aspirated from all visible non atretic follicles (2-8mm in
diameter) by an 18gauge needle attached to 5ml sterile disposable syringe (Dispovan, India) containing 0.5ml oocyte collection media (OCM) (Appendix I). The cumulus oocyte complexes (COC) along with follicular fluid was pooled into 50ml sterile plastic tube containing 1-2ml OCM at 37°C and were allowed to settle for 10minutes.

Finally the sediments were taken in large petridish (90mm) and oocytes were searched under zoom stereo microscope (Motic, Germany). The cumulus oocyte complexes (COC) were isolated, evaluated and graded. Only excellent (>5 layers of cumulus cells and evenly granulated cytoplasm) and good (>3 layers of cumulus cells and evenly granulated cytoplasm) COC were collected and washed several times in OCM followed by maturation media (MM)

In vitro maturation of oocytes

Graded oocytes were washed in respective maturation media for 4-5 times. After washing, 15-20 oocytes were cultured in 50µl droplets of respective maturation media in 35mm sterile petridish. The droplet was covered with warm, non toxic mineral oil and incubated at 38°C, 5% CO₂, 95% relative humidity for 24hrs in CO₂ incubator (Nuaire, USA). The experiment was repeated 6 times for each group as replicates.

The selected oocytes were cultured in TCM 199 medium containing 10%FCS, 0.3% BSA and 10IU/ml of hCG with different supplements as shown in Table No 1.

In vitro fertilization

Sperm preparation for in vitro fertilization

Semen straw was thawed in warm water (35-37°C) for 1min and was emptied in to 15ml tube add 5ml of sperm TALP medium (Appendix IIIa) and centrifuged at 800-1000 rpm for 5min. The supernatant was removed and process was repeated again by adding 5ml sperm TALP. Sperm pellet was formed finally after discarding the supernatant. Sperm pellet was dissolved in 1ml of fertilization TALP (Appendix IIIb), kept in CO₂ incubator for 30min before inseminating the matured oocytes. The sperm concentration was adjusted to 4-5×10⁶ cell/ml.

In vitro fertilization of matured oocytes

The expanded cumulus cells of Invitro matured oocytes were removed by washing (only degree1 and 2 of cumulus cell expansion) several time with fertilization TALP. Denuded oocytes were transferred to 60µl droplet of fertilization TALP medium in 35mm sterile petridish and were inseminated with 40µl of processed spermatozoa. The droplet was covered with sterile mineral oil and kept in CO₂ incubator at 38°C, 5% CO₂, 90-95% RH) for 18h.

In vitro culture of fertilized oocytes and assessment of cleavage rate

After 18h of sperm oocyte co-incubation, oocytes were washed several times with mSOF to remove attached and died spermatozoa from fertilized oocytes and washed oocytes were cultured in modified synthetic Oviductal fluid (mSoF) (appendix IV) supplemented with 0.8% bovine serum albumin (BSA), essential and nonessential amino acids.

Cleavage rate was observed after 96h culture at 38°C, 5% CO₂, 90-95% RH. Cleavage rate was calculated by dividing total number cleaved oocytes (2-4 cell stage embryos) at 96h with number of oocytes utilized for in vitro fertilization and expressed in percent cleavage rate.
Cleavage rate (%) =

\[
\frac{\text{Total number of cleaved oocytes}}{\text{Number of oocytes utilized for } \textit{in vitro} \text{ fertilization}} \times 100
\]

The results of this study describe the cleavage rate achieved by maturing the oocytes in different supplemental groups are presented in this chapter.

**Cleavage rate**

\textit{In vitro} fertilization was carried out for the matured oocytes after removing the expanded cumulus cells completely. Semen sample was processed for capacitating the spermatozoa and finally inseminate the denuded oocytes with capacitated spermatozoa. Cleavage rate was calculated by dividing total number cleaved oocytes (2-4 cell stage embryos) at 96h by number of oocytes utilized for \textit{in vitro} fertilization.

The number fertilized oocytes that reached 2 cell stages were 5/53, 7/55, 4/49, 4/46, 6/55, 8/55, 3/50, and 4/48 respectively for serum gonadotrophin @50IU/ml, serum gonadotrophin @100IU/ml, estradiol-17β @0.5µg/ml, estradiol-17β @1µg/ml, 5% estrus buffalo serum, 10% estrus buffalo serum, 5% buffalo follicular fluid and 10% buffalo follicular fluid supplemented groups respectively. Similarly the number of oocytes that reached 4 cell stage of cleavage was 2/53, 2/55, 1/49, 0/46, 4/55, 3/55, 1/50, and 0/48 respectively for T1, T2, T3, T4, T5, T6, T7 and T8 groups.

The cleavage rate was calculated by considering the oocytes that reached 2-cell and 4-cell stage of cleavage together (Plate 8). The mean percentage values of cleavage rate in serum gonadotrophin @50IU/ml, serum gonadotrophin @100IU/ml, estradiol-17β @0.5µg/ml, estradiol-17β @1µg/ml, 5% estrus buffalo serum, 10% estrus buffalo serum, 5% buffalo follicular fluid and 10% buffalo follicular fluid supplemented group were 12.98±1.07, 16.36±3.27, 8.51±2.76, 9.09±2.97, 18.23±1.98, 20.00±0.87, 8.27±2.70 and 7.98±2.59 respectively (Plate 10).
In the present experiment, highest cleavage rate was shown by the oocytes cultured in IVM medium supplemented with 10% estrus buffalo serum and the lowest cleavage rate was observed in the oocytes supplemented with 10% buffalo follicular fluid. The statistical analysis of the data revealed that cleavage rate was significantly (p≤0.05) higher in the group supplemented with 5% estrus buffalo serum and 10% estrus buffalo serum as compared to all other groups of the experiment. Though the mean cleavage rate in serum gonadotrophin supplemented groups at both the concentrations was more than that in estradiol-17β and buffalo follicular fluid supplemented groups, statistically there was no significant difference between groups supplemented with serum gonadotrophin, estradiol-17β and buffalo follicular fluid.

The comparison of mean values of cleavage rate between two different concentrations of each supplement suggested that there was no significant (p≤0.05) difference between serum gonadotrophin @50IU/ml and @ 100 IU/ml, estradiol-17β @ 0.5µg/ml and @1µg/ml, 5% and 10% estrus buffalo serum, 5% and 10% buffalo follicular fluid.

**Serum gonadotropin**

Supplementation of serum gonadotrophin@100IU/ml to IVM medium showed better cytoplasmic maturation rate and cleavage rate though there was no significant effect in terms of nuclear maturation where as supplementation of the same @50IU/ml did not show much beneficial effects. The beneficial effects of gonadotrophins could be attributed to the fact that they are the primary regulators of oocyte maturation in mammalian oocytes in vivo and one of the functions of its preovulatory surge is to suppress the granulosa cell factor that inhibits meiosis (Umadevi, 1997). Farag et al., (2013) showed that the supplementation of gonadotrophin (PMSG-hCG) to culture media significantly (P <0.05) improved meiotic maturation rate of camel denuded oocytes than that cultured in hormone-free media. Caprine oocytes matured with 20IU/ml PMSG had a good cytoplasmic maturation rate that allows normal embryo development up to blastocyst stage (Kouamo and Kharche, 2015)

**Estrodiol-17β**

The cleavage rate in the oocytes supplemented with estradiol-17β at both the concentration was better than the supplementation of 10% buffalo follicular fluid. This inhibitory effect of Estradiol-17β on oocyte maturation is observed by McGaughey (1977) in pigs, Eppig and Koide (1978) and Dianne and Tenney (1980) in mouse. Estradiol-17β inhibited the cAMP phosphodiesterase activity of mouse oocyte in a concentration-dependent manner (Kaji, 1987). This finding provides an explanation for the inhibitory effect of steroid hormones on germinal vesicle breakdown (GVBD) of mouse oocytes in vitro.

Similar to the present results Tesarik et al., (1995) reported that in humans the addition of E2 to oocyte maturation medium did not produce any apparent effects on either germinal vesicle breakdown or further progression of meiosis, but it did increase the fertilization and cleavage rates of the in vitro matured oocytes. Beket et al., (2002) matured bovine oocytes in TCM199 in the presence of 1µg/ml E2 with or without 0.05 IU/ml recombinant hFSH. They concluded that supplementation of 1µg/ml E2 to a serum free maturation medium negatively affects bovine oocyte nuclear maturation and subsequent embryo development and these negative effects of E2 could be attenuated in the presence of FSH.
**Estrus buffalo serum**

The mean values of cleavage rate were 21.12±1.98 and 23.52±0.87 respectively in the oocytes matured in IVM supplemented with 5% and 10% estrus buffalo serum respectively. Among the eight experimental groups under study the mean values of cleavage rate were highest in the oocytes cultured in IVM medium supplemented with 10% estrus buffalo serum. The beneficial effect of supplementation of estrus buffalo serum to IVM medium on oocyte maturation and subsequent better cleavage rate could be attributed to the fact that estrus buffalo serum contains various hormones like FSH, LH and E2. Biological role of estrus serum is to compensate for whatever essential elements are missing from the medium by serving as a reservoir for many of the beneficial components, such as different energy substrates, steroids, amino acids, fatty acids, vitamins and growth factors. Serum also serves as a protective compound against scavenging ions and small molecules secreted from the developing embryo. Similarly, Totey et al., (1992) observed higher rates of maturation and development to the blastocyst stage in the buffalo oocytes cultured in medium supplemented with EBS and gonadotrophin. Karami et al., (2010) reported high maturation, fertilization and embryo development rate of ovine oocytes in human menopausal serum, estrous sheep serum, and estrous goat serum, than ovine follicular fluid, bovine follicular fluid and control media.

**Buffalo follicular fluid**

The mean values of cleavage rate were 8.27±2.70 and 7.98±2.59 respectively in the oocytes matured in IVM medium supplemented with 5% and 10% buffalo follicular fluid respectively. Among all the experimental groups the lowest cleavage rate was observed in the oocytes supplemented with 10% buffalo follicular fluid. The observation of Stephen et al., (1984) that the follicular fluid transiently suppresses the germinal vesicle breakdown (GVBD) in mouse oocytes by elevating cAMP in the oocyte cumulus cell complex supports the present findings. However these results are in contrast to the findings of Masudul et al., (2012) who concluded that that the GFF has a positive effect on in vitro production of embryos in Black Bengal goats and a 10% level of GFF is recommended based on the improvements observed and the associated economic benefits. Maria et al., (2014) evaluated the influence of follicular fluid added to the maturation medium on the quality of bovine embryos produced in vitro. Even though higher concentration of follicular fluid added to the maturation medium reduced embryonic development rates, in lower concentrations, follicular fluid contributed to increases in inner cell mass number. They concluded that, follicular fluid added to the maturation medium enhances the number of cells in bovine embryos produced in vitro, especially for inner cell mass. The discrepancy in the results of present experiment with others may be because of variations in the size and maturational status of the follicles from which the follicular fluid was collected. Because, Ayoub and Hunter (1993) collected bovine follicular fluid at different stages of the estrous cycle from small, medium, and large follicles. Follicular fluid from both small and medium follicles at estrus had the greatest ability to prevent germinal vesicle breakdown but became less potent at postestrus. Follicular fluid from large follicles at estrus had less ability to inhibit germinal vesicle breakdown than fluid from small and medium follicles. However, follicular fluid from large follicles had less germinal vesicle breakdown inhibiting activity at proestrus than follicular fluid from large follicles at late metestrus, early diestrus, and mid diestrus.
Table 1 Composition and supplements of the maturation media in different groups

| Groups | Supplements | Detailed Composition of maturation medium |
|--------|-------------|------------------------------------------|
| T1     | Serum Gonadotrophin @50 IU/ml | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + serum gonadotrophin @50IU/ml |
| T2     | Serum Gonadotrophin @100 IU/ml | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + serum gonadotrophin @100IU/ml |
| T3     | Estradiol-17β @0.5µg/ml | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + Estradiol-17β @0.5µg/ml |
| T4     | Estradiol-17β @1µg/ml | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + Estradiol-17β @1µg/ml |
| T5     | 5% Estrous buffalo serum | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + 5%EBS |
| T6     | 10% Estrous buffalo serum | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + 10%EBS |
| T7     | 5% Buffalo follicular fluid | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + 5%BFF |
| T8     | 10% Buffalo follicular fluid | TCM 199+10% FCS+ 0.3% BSA+ hCG @10IU/ml + 10%BFF |

Table 2 Cleavage rate (%) of in vitro fertilized buffalo oocytes matured in IVM medium with different supplements (Mean±SE)

| Groups | No. of replicate | Number of matured oocytes utilized | Total number of cleavage | Cleavage rate |
|--------|-----------------|-----------------------------------|--------------------------|--------------|
|        |                 |                                   | 2 cell | 4 cell |                |
| T1     | 6               | 53                                | 5      | 2     | 12.98±1.07abc (7) |
| T2     | 6               | 55                                | 7      | 2     | 16.36±3.27abc (9) |
| T3     | 6               | 49                                | 4      | 1     | 8.51±2.76c (5)   |
| T4     | 6               | 46                                | 4      | 0     | 9.09±2.97c (4)   |
| T5     | 6               | 55                                | 6      | 4     | 18.23±1.98a (10) |
| T6     | 6               | 55                                | 8      | 3     | 20.00±0.87a (11) |
| T7     | 6               | 50                                | 3      | 1     | 8.27±2.70c (4)   |
| T8     | 6               | 48                                | 4      | 0     | 7.98±2.59c (4)   |

a,b,c: Mean values of Cleavage rate with different superscripts differ significantly (p≤0.05)
PLATE No. 7

A. Denudation of Matured Oocyte
B. Capacitation Spermatozoa (Phase Contrast)
C. Ovum Sperm Co-Incubation
D. Attachment of Spermatozoa to Zona

In vitro Fertilization Procedure
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How to cite this article:
Ambadas, K. Shrikant, G. Sarvadnya and Satish, K. B. 2020. Comparison of Cleavage Rate of Buffalo Oocytes on Supplementing with Different Concentration of Serum Gonadotropin, Estradiol 17b, Estrus Buffalo Serum and Buffalo Follicular Fluid. Int.J.Curr.Microbiol.App.Sci. 9(02): 2072-2084. doi: https://doi.org/10.20546/ijcmas.2020.902.235

2084