Observations on Estrus as Monitored by Receptivity to Mating Activity and Pattern of Vaginal Exfoliates In *Trypanosoma brucei* – Infected WAD Goat does Synchronized with Estrumate®

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**SUMMARY:** The effects of experimental *Trypanosoma brucei* infection on receptivity to mating activity and pattern of vaginal exfoliates were monitored using twenty-one adult WAD goats which were synchronized with double injection, seven days apart of Estrumate®. The twenty-one goats consisted of 3 bucks and 18 does. The does were randomly divided into control group ‘A’ having 10 does and test group ‘B’ with 8 does. The goats were fed with Elephant grass in the morning and commercial feed containing 15.23% CP at the rate of 0.25kg/head in the afternoons. Freshwater was provided *ad libitum.* Results showed that while all the control does were observed to stand to be mounted and mated, none of the infected does did. Also, the pattern of the mean percentage vaginal exfoliated cell types encountered between the control and infected doe groups were converse. While parasabal cells changed from 3.05±0.46% during proestrus through 2.42±0.08% at diestrus in the control does, it changed from 22.07±0.56% during expected proestrus through 8.48±0.05% during expected estrus to 28.05±1.09% respectively in the infected does. In like manner, intermediate cell changed from 27.27±0.08% during expected proestrus through 42.37±2.39% during expected estrus to 40.24±1.06 % during expected diestrus in infected does. Similarly, superficial cells changed from 56.25±0.75% during proestrus through 63.70±1.05% during estrus to 7.37±0.01% during diestrus while it changed from 0.00% during expected proestrus through 3.39±0.02% during expected estrus to 63.70±1.05% during estrus to 6.10±0.01% during expected diestrus. The cytology of vaginal exfoliates have been described in the goat (Pretorius, 1997; Perez-Martinez et al., 1999). Its application in the detection of ‘heat’ and perhaps, the optimum breeding time in both natural mating as well as artificial insemination purposes in the WAD goat had also been reported (Ola et al., 2006; Leigh & Ajibade, 2010). There is a paucity of information on the possibility

**KEY WORDS:** Receptivity to mating; Vaginal exfoliates; *Trypanosoma brucei*; WAD does; Estrumate®.

**INTRODUCTION**

The devastating effects of trypanosome infection on the health and productivity of people and livestock in tropical Africa has been well documented (Gasser, 1963; Apter, 1970; Sekoni, 1994). Reproductive symptoms that have been associated with trypanosomosis in the females of most species include irregular oestrus, infertility, sterility, abortion, still birth and neonatal death (Ikede, 1979; Faye et al., 2004).
of association of any reported reproductive disorder in animal trypanosomiasis with exfoliated vaginal cell pattern. The WAD goats are found in areas endemic/encezootic with Trypanosomiasis. This study was therefore designed to investigate the success of estrus monitored by willingness to accept the male (i.e. buck) and vaginal exfoliate pattern in Trypanosoma brucei infected female WAD goats synchronized with Estrumate®.

MATERIAL AND METHOD

Animals and Management. Twenty-one adult WAD goats consisting of 3 bucks and 18 does were involved in the study. The goats were kept in the small ruminant Unit of the Department of Veterinary Surgery and Reproduction, University of Ibadan. The three bucks were kept in a single pen while the does were randomly divided into two groups (i.e. Group A and B). Group A consisted of 10 does which served as control while Group B consisted of 8 does which served as test group. Each group was kept together in a pen and each doe wore a neck tag labeled between AG13 and AG30. The goats were fed with elephant grass in the mornings and commercial feedstuff containing 15.23% CP at the rate of 0.25kg/head in the afternoons. Freshwater was provided in their pens ad libitum. The goats were not allowed to graze the paddock throughout the study.

Source of Trypanosoma brucei. Trypanosoma brucei was obtained from passaged rats from the Department of Veterinary pathology, University of Ibadan. The strain was originally obtained from the Nigerian Institute for Trypanosomasis Research (NITR), Vom, Plateau State, Nigeria.

Parasite Estimation in albino rats and inoculation. Parasitaemia in albino rats was evaluated at 8.4 x 107 Trypanosomes per milliliter by haemocytometrical method using blood obtained from the tail vein. Further dilution of the blood was made with normal saline to obtain an inoculation dose of 4.8 x 105 trypanosomes/0.5ml.

Oestrous synchronization. All animals used in the study were synchronized using the method described by Leigh et al. (2010). Briefly, this involved the administration of a double injection, seven days apart of Estrumate® (Scherring-Plough Animal Health Corp; Germany) to each doe. Both Groups A (i.e. control) and B (i.e. test group) does received 125mcg of Estrumate® (i.e. chloprostenol sodium) via intramuscular route. The administration of the first dose of Estrumate® was done on day 12 after parasite inoculation.

Re-synchronization of oestrus. In both groups A and B, the does which failed to show estrus as detected by not standing to be mated were re-synchronized 18 days after the last intramuscular administration of Estrumate®. They were then closely observed for signs of estrus as before.

Receptivity to mating activity. The does in group A and B were brought into the pen of the bucks for observation of their response to mating activity. This procedure was carried out thrice daily (i.e. 0800 hours, 1200 hours and 1600 hours) starting from the 72nd hour following 2nd injection of Estrumate®. Observation was recorded on whether the does stood to be mated or butted the buck.

Collection of vaginal smear and cytology. Vaginal smear was collected from the does with the aid of improvised vaginal swabs which consisted of clean, soft and gentle, highly absorbent pure cotton buds from Thailand. The vulva and perineum were rinsed with clean water and gently wiped with a clean towel. Each doe was well restrained in standing position by an assistant and the swab was gently inserted into the anterior vaginal with the right hand while the left thumb and fore-finger were used to expose the vulva lips. Hands were gloved. At the anterior vagina, the swab was gently and briskly rolled against the vaginal mucosa and carefully withdrawn. The swab was immediately smeared on a warm gland slide and stained with Giemsa. This process was performed daily between 0800 hours and 0900 hours. The cells encountered in the vaginal smear were categorized as percentage of matured (i.e. superficial), immature (parabasal and intermediate) and leucocytes. Twenty five cells were counted from four fields of each slide and the percentage of each cell type was recorded.

Data analysis. Data collected in the mating receptivity experiment was recorded as “Accepted mating” or “Declined mating” while that on vaginal cytology was summarized as mean percentages of exfoliated cells and subjected at every stage of the cycle to Student –t- statistic (Elston & Johnson, 2008) with < = 5%.

RESULTS

Receptivity to mating activity. Table I below shows the variation in the mating responses of does in groups ‘A’ and ‘B’ to bucks in the experiment. Figure 5.1 also shows some portion of the ejaculate hanging from the vulva of one of the control does immediately after the male dismounted. Following the synchronization protocol involving the administration of 125mcg of Estrumate®, the does in both groups were introduced to the bucks for mating activity.
Seven (7) out of the 10 does in group ‘A’ accepted to be mated by the bucks between 72 and 96 hours following the 2nd intramuscular injection of Estrumate®, while 2 of the does accepted mating only between 192 and 216th hours after 2nd injection with Estrumate®. Throughout this period, one doe (AG 24) did not accept mating. AG 24 was later

### Table I. Variation in the mating responses of groups A and B does to different dosages of Estrumate® during the study.

|                  | Total no of does | Total no of does mounted during 72-96 hours after 2nd intramuscular injection of 125mcg Estrumate® | Total no of does mounted and mated during 192-216th hours following administration of 125mcg Estrumate® | Total no of does mounted and mated during 72-96th hours following single administration of 250mcg Estrumate® | Total no of does that responded to treatment with Estrumate® |
|------------------|------------------|--------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|----------------------------------------------------------|
| **Group ‘A’**    | 10               | 7                                                                                                 | 2                                                                                                | 1                                                                                                | 10                                                       |
| **Group ‘B’**    | 8                | 0                                                                                                 | 0                                                                                                | 0                                                                                                | 0                                                        |

### Table II. Mean % of cellular exfoliates encountered in groups ‘A’ and ‘B’ does during the study. The various mean percentage differences between groups ‘A’ and ‘B’ does during each stage of the study as shown in Table 5.II above were significant (P < 0.05).

|                  | Parabasal cells (Mean ± S.D) % | Intermediate cells (Mean ± S.D) % | Superficial cells (Mean ± S.D) % | Neutrophils (Mean ± S.D) % |
|------------------|---------------------------------|-----------------------------------|----------------------------------|----------------------------|
| Proestrus (group A) | 2.90 ± 0.03                     | 11.10 ± 0.03                      | 56.25 ± 0.75                     | 29.75 ± 0.04               |
| Expected proestrus (group B) | 22.07 ± 0.56                  | 27.27 ± 0.08                      | 0.00                             | 50.64 ± 1.68               |
| Estrus (group A)   | 3.05 ± 0.46                     | 11.10 ± 0.31                      | 63.7 ± 0.15                      | 22.14 ± 0.65               |
| Expected Estrus (group B) | 8.48 ± 0.05                   | 4237 ± 2.39                      | 3.39 ± 0.02                      | 45 76 ± 6.05               |
| Diestrus (group A)  | 2.42 ± 0.08                     | 1.21 ± 1.00                      | 7.37 ± 0.01                      | 86.70 ± 1.68               |
| Expected diestrus (group B) | 28.05 ± 1.09                   | 40.24 ± 1.06                     | 6.10 ± 0.01                      | 25.61 ± 2.85               |

![Fig. 1. Showing the perineal region of one of the control (group A) does with semen hanging from the vulva just after mating.](image1.png)

![Fig. 2. Cytology of proestrus in control does showing deep staining parabasal cell.](image2.png)

![Fig. 3. Cytology of estrus in control does showing superficial cells along with numerous sperm cells.](image3.png)
administered with a single injection of 250mcg Estrumate® on day 18 after completing the initial protocol of synchronization. Following re-introduction to the bucks between 72-96th hour after this treatment, AG 24 was observed to stand to be mounted and mated. None of the eight (8) does in group ‘B’ however was observed to stand to be mounted nor mated following introduction to the bucks starting from the 72nd hour through 216th hour after the second intramuscular injection of 125mcg Estrumate®. Following their treatment with a single intramuscular injection of 250mcg Estrumate®, they were also not observed to stand to be mounted or mated.

**Vaginal cytology.** Figures 2 to 7 below shows the pattern of exfoliated vaginal cells in both groups ‘A’ and ‘B’ does during the experiment. Figures 2, 3 and 4 are for group ‘A’ does while figure 5, 6 and 7 are for group ‘B’ does. The figures show differences in quantity and type of the different cells in the vaginal smear as the does pass through the different stages of the oestrous cycle in response to injection with Estrumate®. One of the does (i.e. AG 16) in the control group was however confirmed to be mated for a second time, six (6) days after an initial mating exercise as evidenced by the presence of sperm cells in its vaginal smear. As shown in Table II below, for group ‘A’ does, the mean percentages of each cell type during proestrus are 2.90 ± 0.03% (parabasal), 11.10 ± 0.03% (intermediate), 56.25 ± 0.75 (superficial) and 29.75 ± 0.04% (neutrophils). During estrus, 3.05 ± 0.46% (parabasal), 11.10 ± 0.31% (intermediate), 63.7 ± 1.05% (superficial) and 22.14 ± 0.65)% neutrophils. The estrus period was further shown (figure 3)% by presence of sperm cells in the vaginal smear in the study. In the immediate diestrus following at the end of mating activity, the cells chiefly observed included parabasal (2.42 ± 0.08)% intermediate (1.21 ± 1.00)% superficial (7.37 ± 0.01)% and neutrophils (86.7 ± 1.68)%.
For the eight (8) WAD does in group ‘B’, none of their vaginal smears was positive for sperm cells throughout the study. During the expected proestrus i.e. 48-72 hours post intramuscular administration of either 125mcg or 250mcg, Estrumate® the mean percentages of exfoliated cells were as follows: parabasal (22.07 ± 0.56)%; intermediate (27.27 ± 0.08)%; superficial (0)% and neutrophils (50.64 ± 1.68)%. During the expected estrus i.e. 72-96 hours, the mean percentages of the cells were parabasal (8.48 ± 0.05)%, intermediate (42.37 ± 2.39)%; superficial (3.39± 0.02)% and neutrophils (45.76 ± 6.05)%.

During the expected diestrus, the mean percentages of the vaginal cells were: Parabasal (28.05 ± 1.09)%; intermediate (40.24 ± 1.06)%; superficial (6.10 ± 0.01)% and neutrophils (25.61 ± 2.85)%.

The differences between the mean percentage values for each stage of the study between control and infected does were significant (P< 0.05).

DISCUSSION

The result of the present study on acceptance of mounting and mating by the does in both groups ‘A’ and ‘B’ showed that all does in group ‘A’ i.e. control, were brought into standing estrus by the intramuscular injection of Estrumate®. During 72-96 hours following the initial administration of 125mcg Estrumate®, 70% of the control does responded positively to mating. This observation is similar to our earlier report (Leigh et al., 2004). It is also interesting that the present result again attests to the reliability of the estrus synchronization protocol whereby two injections of a luteolytic agent (PGF2?) was administered 7 days apart to WAD does as described by Leigh et al. The administration of exogenous luteolytic agent during the mid-luteal phase results in premature luteolysis leading to a fall in peripheral progesterone concentration and an increase in gonadotropin and oestradiol -17 beta secretion in farm animals (Ball & Peters, 2004). These hormonal changes gradually culminate in ovulation in intact animals (Noakes et al., 2001). It is possible that the rate at which these processes occur differ in different individual animals or that they are affected by the availability of certain chemicals. This explanation may stand for the observation in the study where two of the control does where receptive to the bucks during 192-216th hours following the second injection of 125mcg Estrumate®.

The single doe which failed to come into estrus with two injections of 125mcg but responded positively to 250mcg Estrumate® suggests that indeed, ovarian threshold to luteolytic agents differ with different animals. While some respond to moderate/normal doses, others require higher doses to attain adequate blood level of the chemical substance needed for luteolysis (Greyling & van der Nest, 2000). In the case of group ‘B’ does i.e. infected group none of the does showed positive response to estrus as evaluated by standing to be mounted and or mated during the course of the study. This finding suggests that the infection prevented the does from exhibiting estrus in response to the administration of Estrumate® at both 125mcg and 250mcg dose per animal. Similar observations of anoestrus as the present one have been reported earlier (Elhassan et al., 1994; Sekoni). The present finding may suggests a possibility of affection of the normal physiological processes leading to behavioural and possibly fertile estrus in the female animal. Such affection may have occurred at the level of the hypothalamic regulation of anterior pituitary hormones through the production of gonadotropin releasing hormone (GnRH) or at the level of the anterior pituitary through the elaboration of follicle stimulating hormone (FSH) and luteinizing hormone (LH) or at the level of the ovary through the production of estrogens from growing follicles (Gordon, 2005). Some workers have reported acute coagulative necrosis of adenohypophysis in T. brucei infected sheep (Mutayoba et al., 1995) while others have observed a failure of the hypothalamic – pituitary –adrenal axis in response to insulin induced hypoglycaemia in Boran cattle experimentally infected with Trypanosoma congolense (Abebe et al., 1993).

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The observations in this study therefore ascertain the virulence of Trypanosoma brucei in the WAD goat with respect to reproductive pathology. It is concluded that experimental Trypanosoma brucei infection in the WAD does led to failure of estrus following synchronization with Estrumate®.

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