The aim of this study was to determine titers of antisperm antibodies (ASA) of Ig G and Ig A class in sera and cervical mucus of artificially inseminated Holstein cows in order to correlate these results with the duration of the open days period. Investigations were conducted on a total of 181 cows originating from three different dairy farms. Blood and cervical mucus samples for laboratory analyses were collected on the day of the last artificial insemination. Presence of ASA was determined by indirect immunofluorescence method (IIF) using bulls' sperm cells prepared for artificial insemination by suspending in TRIS - egg yolk or "Biociphos +" extenders prior to deep freezing.

Our results strongly confirm the hypothesis that immune mechanisms may be involved in reproductive disturbances due to high levels of ASA of Ig A class. In the sera and cervical mucus of cows, high levels of ASA were found in animals with longer open days period. In this study we were not able to demonstrate differences in ASA titers when sperm cells were suspended in different extenders and used for the IIF test.

Key words: sperm cells, cows, artificial insemination, subfertility, antisperm antibodies

INTRODUCTION

Despite numerous clinical studies and experimental efforts, over the past four decades, the precise role and significance of antisperm antibodies (ASA) in the pathogenesis of immunological sterility and subfertility of both males and females still remains unclear. There are plenty of data accumulated, so far, and now it is evident that ASA may impair fertility to certain extent. The reasons for our inability to explain immunological subfertility lies in the fact that the mechanisms involved in the gamete immune protection are very complex and the immune response to their antigens is never an "all or none" phenomenon. This mechanisms are in details reviewed elsewhere (Hogarth, 1982; Bronson et al., 1998, Landers et al., 1994). Spermatozoa are foreign cells to males as they appear
late into ontogenesis (in puberty), but their progenitors are separated from the immune cells by a blood-testis barrier during spermatogenesis (Furuya et al., 1978). Moreover, submucosa of the male reproductive tract is rich in T\textsubscript{s} cells and therefore, the immune response to spermatozoa is normally not elicited despite their isoantigenic properties (El Demiry and James, 1988). During passage through male and female genital tracts, spermatozoa are surrounded by seminal plasma that contains strong immunosuppressive molecules which block recognition and antigen processing (James and Hargreave 1984). Other molecules mask antigens important for the fertilization process on the surface of spermatozoa. In addition, cervical mucus, uterine and follicular fluid also contain immunosuppressive substances (Landers et al., 1994). One of the basic characteristics of sperm cells is a continuous change of their antigenic structure due to the loss of surface molecules during maturation and following insemination. Antisperm antibodies may be present in seminal plasma, cervical mucus (CM), uterine fluid, follicular fluid or blood sera (Stren et al., 1992). The role of ASA in infertility and subfertility in mammalian species is still not clearly understood but it is well known that auto-immunization of the male and iso-immunization of the female with sperm cells can lead to significant impairment of fertility. Experimentally generated bull antisperm antibodies significantly reduce fertilization in vitro (Kim et al. 1999). Much earlier, Wright (1980) described the negative influence of ASA, raised by intensive immunization, of adult animals on semen quality in the bull. This phenomenon has been recorded for numerous laboratory animals as reviewed by Hogarth (1982). Bratanov et al. (1980) showed that sera containing antisperm antibodies from infertile cows and women with unexplained infertility inhibit acrosomal proteolytic activity in vitro and therefore possibly may affect fertilization. Investigations in veterinary medicine regarding the influence of ASA in CM on fertility rates are very few. An elevation of sperm-agglutinating antibody titer in the CM and sera of artificially inseminated cows was clearly demonstrated by Jačević (1998).

In the technology of artificial insemination (AI), the antigenic structure of sperm cells is changed due to the addition of different extenders, freezing and thawing procedures and reduction of seminal plasma volume. In addition, the immunosuppressive activity of bull seminal plasma is significantly reduced during semen preparation for AI (Lazarević, 1991). Therefore, we have investigated the presence of ASA in CM and sera using sperm cells previously suspended in two different extenders and correlated these findings with the duration of the open days period in artificially inseminated cows.

One of the most important reproductive parameters in cows is “open days period” being defined as a period between calving and the next successful insemination. Normal duration of open days period is essential in dairy cows breeding and our goal is usually to get one calf per year from each cow. This is not a simple task and on dairy farms repeat breeding (RB) still represents one of the major problems. A repeat-breeder is a cow which shows a reduced probability of conception, while all other factors are optimal (Casida, 1961), and thus requires more AI attempts to achieve pregnancy. These cows are subfertile and for that reason, unless they have a very high milk yeald, their presence in the herd
elevates costs of milk production. If mated naturally, RB cows or heifers very often show improved reproductive results (Vukotić et al., 1982). It was postulated that enhanced immunological reactivity to sperm or semen extender antigens might be one of the reasons for RB (Park and Hunter, 1977). It has been documented (Lazarević et al., 2003; Jačević, 1998) by sperm agglutination and indirect immunofluorescence, that bull spermatozoa differ in antigenicity if different semen extenders are used for semen preparation for AI.

One of the major consequences of repeat breeding is a prolonged open days period and the aim of our study was to determine the levels of antisperm antibodies (ASA) of Ig G and Ig A class in the sera and cervical mucus of artificially inseminated Holstein cows and to correlate these results with the duration of the open days period.

**MATERIAL AND METHODS**

**Sera sampling:** Sera were collected from 181 Holstein cows at three regional dairy farms by jugular venipuncture on the day of artificial insemination. Sera were obtained following coagulation at room temperature and centrifugation at 3000 rpm for 20 min. All samples were kept frozen at -20 °C until use. The animals were divided into two, three or five groups according to the average titer values for ASA of Ig G or Ig A class in the sera or cervical mucus. All cows were inseminated with bull semen prepared for AI with TRIS egg-yolk or “Biociphos +” extender.

**Cervical mucus sampling:** Prior to AI, CM samples were collected by placing a sterile sponge swab in the near vicinity of the external cervix portion. Sponges were placed with a sterile plastic tube (1.5 cm width, 45 cm long) and the swab remained inside for the next 5 minutes. The swab was then removed and placed in an opened sterile syringe (20 ccm). The syringe was closed and the cervical mucus extracted by gentle pushing into a sterile plastic polystyrene tube. We were able to collect approx. 3 ccm of CM. All samples were kept frozen at -20 °C until use.

**Semen sampling:** Semen samples were collected from four black and white spotted bulls (Holstein breed) by means of an artificial vagina in the Regional Centre for Artificial Insemination. All semen samples possessed normal characteristics of motility, morphology and concentration. Ejaculates were pooled and then divided into two equal portions (split technique). The ejaculates underwent the standard procedure of preparation for AI. One half of each ejaculate was diluted with TRIS - egg yolk extender as described elsewhere (Lazarević et al., 1992) and one with “Biociphos +” extender (IMV, France). Ejaculates were diluted at an average ratio of 1:10 and kept frozen at -196 °C before use.

**Indirect immunofluorescence assay (IIF):** The IIF assay was performed according to Noel et al. (1974). After thawing the straws for AI (medium French straws 0.45 ml), sperm cells were separated and washed twice in PBS (pH 7.2) by centrifugation at 2000 rpm for 10 minutes. We always used straws originating from...
the same four bulls (3 from each bull). When the last supernatant was discarded the remaining cells were resuspended by Vortex and used for smear preparations. On the microscope slides with dried sperm cell smears, 10 µl of sera or CM sample (inactivated at 56 ºC for 20 minutes) was placed and incubated for 20 min at 37 ºC in a wet chamber. Sera and CM dilutions from 1:4 to 1:1024 (highest positive titer) were used for the test. Following incubation, the slides were washed three times (5 min) in PBS and dried at room temperature. In the second step, 10 ml of secondary FITC (fluorescein isothiocyanate) conjugated antibody (anti-bovine Ig A, ICN, USA, Cat No 641 751) was placed on the slide and incubated again under the same conditions. Anti Ig A antibodies were conjugated with FITC (ICN, USA, Cat No F 4274) according to The and Feltkamp (1970). After incubation, followed by the same washing procedure, the slides were kept in a dark and wet chamber until examined. As a positive control we used sera obtained by immunization of calves with the content of straws prepared with TRIS-egg yolk extender and “Biociphos +” extender as described in detail elsewhere (Lazarevic et al., 2000). Two 5 month-old calves were immunized for the first time with the straw content mixed with complete Freunds adjuvant and for the second time (after two weeks) with straw content mixed with incomplete adjuvant. Calf sera before immunization served as the negative control. Microscopic examination was performed on the NIKON EFD-3 microscope with the B-2A filter at 1600 X magnification. The appearance of fluorescence on the head, tail or neck of the sperm cell was considered as a positive result and the last dilution giving a positive reaction was taken into account. Titre values were expressed according to Sjurin et al. (1984) as - log2 n (1:2 = 1, 1:4 = 2 etc).

Statistical analyses was performed after calculating mean values and standard deviations. The significance of the differences between mean values was estimated by Mann-Whitney test.

RESULTS

Figure 1. shows the correlation between titres of Ig G ASA in the blood sera of cows and open days period. In this case we were not able to demonstrate statistically significant differences between groups of cows with different ASA levels.

The majority of animals had open days period between 150 - 200 days and no differences depending on the extender type were present. Similar results were obtained when titers of the Ig G ASA were determined in the cervical mucus (Figure 2.).

When titers of the Ig A ASA in cows sera were correlated to the duration of the open days period, statistically significant differences were obtained between cows with the highest titers of ASA (8.1 >) and all other groups of animals. In this test we used sperm cells previously suspended in TRIS-egg yolk extender (Figure 3.).
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Figure 1. Duration of the open days period according to the titre of the Ig G ASA in the blood sera of cows against spermatozoa suspended in TRIS - egg yolk and "Biociphos +" extender.

Figure 2. Duration of the open days period according to the titer of the Ig G ASA in the cervical mucus of cows against spermatozoa suspended in TRIS - egg yolk and "Biociphos +" extender.
Figure 3. Duration of the open days period according to the titer of the Ig A ASA in the blood sera of cows against spermatozoa suspended in TRIS-egg yolk extender.

Figure 4. Duration of the open days period according to the titer of the Ig A ASA in the cervical mucus of cows against spermatozoa suspended in TRIS-egg yolk extender.

Figure 4. represents results of the tests performed with sperm cells suspended in TRIS-egg yolk extender in samples of cervical mucus.
As can be seen (Figure 4.) we were able to demonstrate statistically significant differences between nearly all groups and open days period was prolonged in cows having higher titers of antisperm antibodies.

When sperm cells were suspended in "Biociphos +" extender we demonstrated the similar pattern comparing the duration of open days and titers of Ig A ASA (Figure 5.). In this case, cows with the highest Ig A ASA titer had slightly lower average values for the open days period and no statistically significant differences were obtained between them and other groups of cows. However, statistically significant differences were documented between cows with ASA titers from 6.1 - 8 and all other groups of animals.

Nearly same results were obtained in samples of cervical mucus when sperm cells also suspended in "Biociphos +" extender were used for IIF test (Figure 6).

We were, thus able, to conclude that titers of Ig G ASA in the sera and cervical mucus of cows did not correlate with the duration of the open days period. On the contrary, strong correlation was observed between titers of Ig A ASA and open days period. No differences were related to the type of the extender used.
DISCUSSION

Artificial insemination (AI) of cows is one of the useful models for studying ASA significance because in this case, natural conditions of insemination are significantly changed. This enables us to explore the role of certain mechanisms involved in the immunological protection of male gametes. Firstly, during preparation of semen for AI, seminal plasma is diluted several times thus reducing its immunosuppressive activities (Lazarević, 1991; Lazarević et al., 1992).

Secondly, new antigens are being added to sperm suspension in order to replace seminal plasma and to provide cryoprotection. We have also documented that mean titers of the Ig A ASA are higher in the repeat breeder cows (Lazarević et al., 2003). For these reasons, possibilities for an immune response elicitation are higher in artificially inseminated cows and problems in reproduction are most prominent in cattle breeding.

Interference of ASA with reproductive processes may occur by impairment of sperm migration through the cervix, uterus and tubes and by blocking adherence of spermatozoa to the surface of the zona pellucida of the oocyte as postulated by Schumacher (1998). The same author stated that serum antibody levels do not reflect properly the immunological situation in secretions of the genital tract especially in females, and that the secretory immunological system may be operational mainly in the cervical compartment of the genital tract which is in agreement with our findings. Our results are also in agreement with Marshburn and Kutteh (1994), Mazumdar and Levine (1998) and Stern et al. (1992) who proved that ASA in the blood and lymph belong predominantly to the

Figure 6: Duration of the open days period according to the titer of the Ig A ASA in the cervical mucus of cows against spermatozoa suspended in "Biociphos +" extender
immunoglobulin Ig G isotype, while those found in external secretions are predominantly of Ig A isotype. It is also proved that circulating ASA of Ig A and Ig G classes recognises different antigens on the human sperm plasma membrane (Auer et al., 1997). In our investigations higher titers of Ig A ASA were detected both in sera and cervical mucus of cows. Nonspecific ASA are also demonstrable in heifers and prepubertal calf’s sera but the titers are much lower (Lazarevic et al., 2003). In this study we were not able to demonstrate differences in ASA levels, both in sera and cervical mucus, when sperm cells suspended in two different extenders were used for IIF test. This data are in disagreement with Jačević (1998) but all cows included in her investigations were inseminated only with semen prepared for AI with TRIS - egg yolk extender which was not the case in our study.

Maas et al. (1998) demonstrated a close correlation between a positive sperm antibody test and a poor postcoital test in infertile couples. These authors concluded that determination of ASA in the CM must be regarded as an improvement in the diagnostic procedure in human infertility. We have also reviewed the possible negative influence of antisperm antibodies on reproduction processes (Jačević and Lazarević, 2000). In one survey (Shulman, 1977) it was found that in a large group of infertile women who were seronegative, 28% were positive in their CM. In our study we were also able to demonstrate that some sera samples had no detectable antisperm antibodies (data not presented here).

There is strong evidence that in humans sperm-mucus interactions can be affected by local ASA, especially of the Ig A class, both under in vitro and in vivo conditions (Eggert-Kruse et al., 1991). However, the significance of ASA in sera of infertile patients was not established, while those in seminal plasma or CM impaired the ability of sperm cells to penetrate CM (Eggert-Kruse et al., 1995). Check et al. (1994) demonstrated that the antifertility effect of ASA may be mainly due to immobilization of sperm in the CM and thus intrauterine insemination may effectively correct the problem. The role of antisperm Ig A antibodies in man has also been documented by Clarke et al. (1984) and Kremer and Jager (1988). Recently, in veterinary medicine valuable data were obtained by analysing sera of 537 cows by ELISA test for ASA (Zraly et al., 2003). The authors were able to detect higher concentrations of ASA in pluriparous cows and also in cows that were inseminated repeatedly. Their results are also in agreement with our findings. Vukotić et al. (1982), showed that reactivity of female blood serum towards sperm cells increases with repeated AIs and parallel to that, the probability of conception declines. This is also partly in agreement with our results. Presence of Ig A ASA, especially in cervical mucus is of great significance because the attachment of Ig A antibodies to the sperm surface lowers sperm penetration ability, while Ig G antibodies did not have the same effect. Wang et al. (1985). Moghissi et al. (1980) found antibody activity in the mucus of 26% in a group of infertile women and none in the control group. They detected such activity in serum in only 13% of infertile women. These findings are also in agreement with our results. Menge and Natz (1993) later demonstrated the presence of Ig A1 and Ig A2 subclasses in the CM of women.
Our data regarding correlation between open days period and levels of ASA in the cervical mucus and sera are a new contribution to the problem of "repeat breeder cow". We hope that this approach may contribute in overcoming it.

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REFERENCES
U ovom radu su izneti rezultati ispitivanja titra antitela Ig G i Ig A klase poreklom iz krvnog seruma i cervikalne sluzi veštački osemenjavnih holštajn krava protiv antigena spermatozoida bika. Ispitivanja su izvedena na ukupnom broju od 181 plotkinje sa tri regionalne farme muznih krava. Uzorci krvi i cervikalne sluzi su prikupljeni na dan poslednjeg veštačkog osemenjavanja. Titar antitela protiv antigena spermatozoida (ASA) je određivan metodom indirektnog imunofluoresca a za izvođenje testa su korišćeni spermatozoidi suspendovani u TRIS žumanjačnom ili "Biociphos +" razređivaču.

Naši rezultati potvrđuju hipotezu da su imunski mehanizmi uključeni u nastanak nekih reproduktivnih poremećaja jer su plotkinje sa visokim titrom ASA u serumu i cervikalnoj sluzi imala duži servis period. Osim toga, u ovim ispitivanjima nismo utvrdili postojanje razlika u titru ASA u zavisnosti od vrste korišćenog razređivača.