Antifertility Effects of Sperm Impairing Factors Isolated from Bacteria in Male Mice

Abstract

Sperm immobilizing (SIF) and sperm agglutinating factor (SAF) (already isolated in our laboratory) from Staphylococcus aureus and Escherichia coli, respectively, have shown 100% infertility in female mice. These microbial factors neither induce any abnormalities in the vaginal cell morphology nor do they affect structural integrity of vaginal epithelium. Therefore, the present study was carried out to check any adverse effect of these factors on the reproductive potential of male mice. Mice were administered with SIF or SAF (10μg/20μl) intraperitoneally for a period of 7 days on alternate days. No treatment related mortality was observed during the experiment. The mice were sacrificed 48h post inoculation on days 3, 5 and 9 after intraperitoneal administration and the results in terms of change in body weight, seminal parameters, TSI and histology were studied. The results showed that male mice receiving SIF and SAF showed significant decrease in seminal parameters viz. sperm count, motility and viability w.r.t control mice. Histological studies of male mice revealed hypospermatogenesis and inflammation in the reproductive organs of treated mice while non reproductive organs showed normal histology. In conclusion, SIF/SAF upon intraperitoneal administration can lead to a pronounced decrease in male reproductive potential.

Key words: SIF; SAF; Intraperitoneal; Seminal parameters; Histopathology

Introduction

Spermicides are known from centuries and the first written record of spermicide use is found in an Egyptian document namely the Kahun Papyrus, dating to 1850 BCE [1]. Spermicide is a substance that kills or immobilizes the sperm and is inserted deep in the vagina near the cervix prior to intercourse to prevent pregnancy. Spermicides come in several forms such as gel, foam, cream, film, suppository or tablet. Most widely used spermicide is Nonoxynol-9 while other includes benzalkonium chloride, chlorhexidine, menfegol, octoxynol-9, and sodium docusate. The spermicidal activities of these detergent-type contraceptives are associated with their structural affinity to the membrane lipids [2,3] therefore, the major drawback of using N-9 or other currently used surfactants is their detergent-type effect leading to an increased risk of vaginal or cervical infection, irritation or ulceration. In addition to this, alteration in the vaginal flora leads to an increased risk of opportunistic infections [4-6]. In a systematic search for developing a new spermicide and microbicide, an earlier study on a microbial origin antimicrobial peptide nisin has shown good contraceptive efficacy in vitro as well as in vivo [7]. The reproductive toxicity of nisin was also evaluated on the male rats where it was administered orally [8]. On the other hand, Qin et al. [9] demonstrated that when mice were given busulfan intraperitoneally, it resulted in deleterious effects on reproductive potential of male mice.

This intrigued us to evaluate the effect of microbial factors earlier isolated in our laboratory from Staphylococcus aureus and Escherichia coli, on the male mice which have shown excellent sperm impairment in vitro and caused 100% infertility in vivo in female mice [10,11]. Thus the present study was designed to evaluate the effect of these microbial-derived biologically active spermicidal proteins on the reproductive potential of male mice.

Materials and Methods

Microorganisms

The strains of both Staphylococcus aureus and Escherichia coli, isolated from the cervix of a women suffering from unexplained infertility and semen samples of infertile males undergoing semen analysis, respectively, were already available in the laboratory. The isolate of S. aureus showed sperm immobilization whereas E. coli showed sperm immobilization via agglutination in vitro.

Experimental animals

Sexually mature, 5-6 week old male (25±2g) BALB/c mice were used in the present study. The mice were maintained under standard laboratory conditions and were housed in propylene cages (430×270×150 mm³, six animals per cage) at 20-25°C, bedded with clean rice husk in well aerated animal room of the Department of Microbiology, Panjab University. All the animals were given standard pellet diet and water ad libitum. Animals were allowed to acclimatize to the new housing and experimental conditions for at least one week. All the experimental protocols were approved by the Institutional Animals Ethics Committee of the Panjab University Chandigarh vide letter no PU/IAEC/S/15/67.
dated 15.09.2015 and were performed in accordance with the guidelines of committee for the purpose of Control & Supervision of Experiments on Animals (CPCSEA), Government of India, on animal experimentation.

**Isolation and purification of sperm immobilization factor (SIF) from** *S. aureus*

SIF was extracted and purified from 72 h old cell culture of *S. aureus* by the method previously standardized in the laboratory [12]. Briefly, the cell culture of *S. aureus* grown in brain heart infusion (BHI) broth, under shake conditions (220 rpm) at 37°C for 72 h, was centrifuged at 10,000 rpm for 15 min at 4°C. SIF was purified from the supernatant by ammonium sulphate precipitation, gel permeation chromatography, and ion exchange chromatography.

**Isolation and purification of Sperm agglutinating factor (SAF) from** *E. coli*

The SAF was extracted and purified from 48 h old cell culture of *E. coli* by the method earlier standardized in the laboratory [13]. Briefly, the cell culture was washed with Phosphate buffered saline (PBS; pH 7.2, 50mM) and pellet was given salt treatment with 3M NaCl for 12 h which released the factor into the solution. After dialysis and concentration of salt treated sample, further purification of SAF was carried out using gel filtration chromatography and ion exchange chromatography.

**Impact of intraperitoneal administration on reproductive potential of male mice**

Mice were divided into 3 groups with 9 mice in each group. Group 1, administered intraperitoneally with 20 μl of PBS alone served as control. The remaining two test groups (Group 2 and 3) were intraperitoneally administered with 10μg/20μl of SIF (Sperm immobilization factor) from *S. aureus* and 10 μg/20μl of SAF (sperm agglutination factor) from *E. coli* without under any anesthesia. Each factor was administered on day 1, 3 and 7 and then the mice were sacrificed 48 h post-inoculation on days 3, 5 and 9.

**Body weight profile and tissue somatic indices (TSI %)**

Initial body weight of mice from each group was taken on the 1st day of experiment and final weight was on the last day of the experiment. On day 3, 5 and 9 mice from each group were sacrificed by cervical dislocation and the various reproductive organs (viz. testis, caudal epididymis and vas deferens) and non reproductive organs (spleen, kidney and bladder) were removed aseptically. The organs were grossly examined and weighed. The TSI (percent tissue/organ weight in relation to body weight) was evaluated [14].

**Assessment of seminal parameters**

**Sperm count**: Mice from each group were sacrificed by cervical dislocation and were dissected. The vas deferens was pulled out and placed in freshly prepared 500μl of PBS buffer (50mM, pH 7.2). Gentle teasing was done to enable the spermatozoa to swim out into the buffer in a glass plate. A fixed volume of 10 μl of the sample was placed on a glass slide and examined under the light microscope at 400X magnification. Around six to eight fields were scanned and the mean number of spermatozoa in all the fields was multiplied by 106. The slides were also assessed for the morphology of spermatozoa in each field to evaluate the respective abnormalities [15].

**Sperm motility**: Motility of the sperms extracted from the sacrificed mice was determined by the method of Emmens [16].

**Sperm Viability**: In order to estimate the percentage of viable sperms, an equal volume of mouse spermatozoa was mixed with 0.5% eosin and examined under the light microscope at 400X magnification.

**Histopathological studies**

Histological analyses of the reproductive organs (testis, caudal epididymis and vas deferens) of mice from each group were carried out. The various reproductive organs and nonreproductive organs were harvested, fixed in 10% formaldehyde for 24h and then embedded in paraffin according to standard histological methods. Serial paraffin sections of 4 mm were stained with hematoxylin eosin and observed at 400X magnification for any significant changes in reproductive organs.

**Results and Discussion**

The microbial factors capable of causing detrimental effects on spermatozoa, on one hand can cause infertility in female mice, but on the other hand show no adverse effects on reproductive performance of male mice. This indicates that the sperm impairing factors that have proved their mettle in one sex may or may not behave in a similar manner in their counterparts. A study citing the similar case has been reported by Reddy et al. [8] in which they have demonstrated that Nisin, an antimicrobial peptide with spermicidal properties, has shown excellent contraceptive efficacy when administered intravaginally in female, but displayed no adverse effects on the reproductive competence of male rats when administered orally. To the best of our knowledge, no other report citing such comparative work has been reported, though several reports regarding the adverse effect of some chemical compounds e.g. lead, busulfan, triptolide etc. on seminal parameters do exist. In this regard, previously in our laboratory we have also isolated strains of *S. aureus* and *E. coli* that have been found to produce microbial factors with sperm immobilizing and sperm agglutinating properties in-vitro, respectively [12,13]. They were also found to possess excellent contraceptive efficacy in female BALB/c mice when administered intravaginally [10,17]. On the lines of Reddy et al. [8], we also attempted to extrapolate the effect of these microbial factors in male mice. Hence, the present study was designed to evaluate the effect of both these microbial factors i.e. sperm immobilizing factor (SIF) from *S. aureus* and sperm agglutinating factor (SAF) from *E. coli* on the reproductive potential of male mice. For this, each factor was intraperitoneally administered on day 1, 3 and 7 and then the mice were sacrificed 48h post inoculation on days 3, 5 and 9. The various parameters which were evaluated included: body weight profile, seminal parameters, tissue somatic indices and histopathological studies.

**Isolation and purification of SAF and SIF**

Both the factors were isolated and purified by the protocol already standardized in our laboratory [12,13].
Weight profile

When the weight profile study of the different group of mice was carried out, the results showed that there was a consistent increase in the body weight in the control group during the course of the experiment. However, a mild decrease in body weight was observed in the case of mice in test groups. This mild decrease might be due to a decline in food consumption since weight change depends on a balance between energy expenditure and food consumption. This is in consistence with study by Qin et al. [9] wherein intraperitoneal administration of Busulfan in male mice led to a mild decrease in the body weight.

Tissue somatic indices (TSI %)

When TSI (%) of various reproductive (testes, cauda and vas deferens) and non-reproductive (spleen, kidney and bladder) organs was examined, it was found that there were no treatment-related changes in the absolute weights of all the organs, except spleen and vas deferens in case of SIF where a significant increase in weight was observed on all the 3 days. However, right cauda showed marked decrease in TSI on all the 3 days and testes showed a marked decrease in TSI on day 9 only. Further, in the case of SAF treated group, left vas deferens showed a considerable increase in TSI on day 3 while the right cauda epididymis showed a significant reduction on days 5 and 9 (Figure 1). The evaluation revealed that no clear pattern could be obtained, hence, reliance on more sensitive endpoints such as morphological and functional abnormalities of these organs could be helpful to better delineate the alterations.

Evaluation of seminal parameters

Some of the major determinants of fertilization potential of spermatozoa are based on their microscopic assessment of sperm concentration, motility, viability and morphology. In order to assess the treatment-related changes in seminal parameters, all the mice from each group were sacrificed on day 3, 5 and 9.

Sperm count: Since, the collective result of all the steps of sperm production is provided by sperm count analysis, hence it is considered as one of the most trusted tests for spermatogenesis. In this regard, when mice treated with SIF were assessed for changes in total sperm count, a significant reduction, with respect to control, was observed on all the days of experiment viz. 53% (on day 3), 80% (day 5) and 92% (on day 9). However, in the case of SAF treated group, day 3 witnessed an insignificant reduction in total sperm count whereas a complete inhibition of spermatogenesis was observed on day 5 and 9 (Figure 2a). The results showed that the factor-induced reduction/complete suppression in sperm count is indicative of the fact that reproductive function of male mice was compromised. Similar results have been reported by Lue et al. [18] where marked reduction in total sperm count was observed when triptolide was administered orally to male rats.

Motility: It is a well known fact that the sperm motility has a critical influence on successful fertilization, since the highly motile spermatozoa have a greater chance of fertilization. Hence, the % motility was evaluated in this study. The results showed that on day 3, in case of SIF and SAF treated mice, it was observed that there were no significant changes in percent motility of the sperms with respect to control (65%). Day 5 onwards, SIF rendered all the sperms non-motile (Figure 2b) whereas this parameter could not be assessed in case of SAF treated group because of complete loss of spermatogenesis. The results indicated that SIF and SAF not only deteriorate the quantity, but also the quality of spermatozoa, thereby, proving their mettle in reducing the reproductive vigour of male mice.

Viability: Although the true viability of spermatozoa is ultimately defined by their capacity to move and fertilise an egg, but here the viability of mice spermatozoa was investigated using live/dead sperm viability stains. Here, the viability refers really only to integrity of their membranes. When the spermatozoa were incubated with eosin dye, the dead sperm cells with damaged membranes stained pink whereas the live ones did not. The results revealed that the number of viable spermatozoa in both

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SIF and SAF treated groups was comparable to control (67%) on day 3. However, in the case of SIF, viability reduced to 43% and 13% on day 5 and 9 respectively. Since complete inhibition of spermatogenesis was observed on day 5 and 9 in SAF treated group, hence, viability could not be assessed for the same (Figure 2c), thus further indicating that both these microbial factors have the ability to induce functional deformities in spermatozoa. In support, Ayoka et al. [19] also reported a significant reduction in the sperm viability in test group given Lead (Pb) orally, as compared to control.

Morphology: When all the spermatozoa were examined for morphological changes upon treatment with SIF /SAF, it was observed that there were no morphological changes in spermatozoa on day 3. However, day 5 onwards, SIF treated mice showed separation of sperm heads from tails, termed as decapitation (Figure 3). Even in the absence of a thorough understanding of the mechanism of abnormality induction, however, it appears that the mere presence of abnormalities is indicative of the interference of SIF in reproductive competence of male mice. On the other hand, no morphological alterations were observed in SAF treated mice. In conclusion, SIF and SAF induced alteration of sperm count, motility, viability and morphology is indicative of the fact that reproductive function was severely compromised.

Histopathological examination

The control group revealed normal tissue histology in all the reproductive organs. Testis showed regular seminiferous tubule and germinal cell morphology (Figure 4a). Epididymis displayed the well-vascularized loose connective tissue present around the epididymal duct (Figure 4b). Further, the normal columnar epithelium was seen in vas deferens (Figure 4c). However, both SIF and SAF treated mice showed that seminiferous tubules in testes have vacant central part with no mature spermatozoa indicating hypospermatogenesis (Figure 4d, 4g). The caudal epididymis showed normal tissue histology (Figure 4e, 4h) whereas inflammation was observed in vas deferens (Figure 4f, 4i) in both the groups. Similar histological findings have been reported by Lu et al. [20] who observed severe vacuolization of the testicular interstitium when lead (Pb) was administered intraperitoneally in male rats.

However, in striking contrast to this, the tissue histology for non-reproductive organs viz. spleen, kidney and bladder; revealed no abnormalities. All the organs were found to be histologically normal, both in the case of control (Figure 4j,4k,4l) as well as treated groups (Figure 4m,4n,4o). Consistent findings were reported by Reddy et al. [8] wherein they declared that no treatment related alterations in the structural integrity of the reproductive and non-reproductive organs after Nisin treatment were observed compared to their respective controls.

Conclusion

From the present study, it can be concluded that both SIF and SAF can play an important role in reducing the male reproductive competence.

Conflict of Interest

The authors declare that there is no conflict of interest.
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