**INTRODUCTION**

*Jussiaea repens* (L), locally known as ‘Kesardam’ of onagraceae family, is a water creeping prime rose found in the wet lands of different parts of India as well as in other countries like Africa, China, Thiland, Malasya, Australia etc. This plant has great medicinal values as reported by others. It can be used as hepatoprotective, anti diabetic, anti helmintic, anti dysenteric, anti-inflammatory and anti bacterial agent. It has also some therapeutic uses i.e., in ulcer, fever, cough, diuretic, urinary tract infection etc. But there was no available scientific report of this plant on male reproductive system, only an ethno-biological report from Papua, New Guinea shows that leaves and stem of this plant can be used to prevent pregnancy. This plant, though it contains some antifertile agents like rutin, kaempferol, quercetin, triterpenes etc. as their active compounds, still unknowingly it is widely consumed by village people as vegetable and animal forage.

In our previous study, we have reported that aqueous extract of *Jussiaea repens* is a non-toxic antagonadal herb which affects the male reproductive system of rats when administered orally in a dose dependent manner for a period of two spermatogenic cycles. This plant extract reduced the epididymal sperm count, motility, viability and increased the percentage of abnormal sperm concentration significantly when compared with control and the effect was maximum at moderate dose (200mg/kg b.wt/day for consecutive 28 days).

The present study was aimed to evaluate the sperm characteristics including morphology and fertility activity of treated animals to ascertain the anti fertility nature of the plant extract after oral administration of the effective dose (200mg/kg b.wt/day for 28 days) to male albino rats. Drug withdrawal experiments were also performed to assess the nature of action of the extract i.e., reversible or irreversible. Since fertility depends on the viability, concentration and morphology of the sperm cells produced by the testes and this plant is widely consumed by the common people, the outcome of this study will be the basis for advising the ethnomedical practitioners and the general public on the usage of this herb.

**MATERIALS AND METHODS**

**Plant material**

The plant *Jussiaea repens* L. was collected from wet lands of Duttapukur, North 24 pargana district, WB, India, during the month of March - April. The plant was identified and authenticated by taxonomist of Central National Herbarium (Kolkata), Botanical Survey of India (BSI), Shibpur, Howrah, having voucher specimen number NP-01 dated 25.03.2011. The collected fresh plants were carefully washed repeatedly under running tap water and finally with distilled water then air dried at 35-40°C for 4-5 days and homogenized the whole plant except root to a fine powder using a sterilized mixer grinder and stored for extraction.
Preparation of extract

The plant extract (except root) was prepared as reported earlier. Briefly, the dried powdered sample (400gm) of J. repens was extracted in 4L boiled distilled water at 50°C for 30 minutes then allowed to cool and stay overnight at room temperature. The residue was further extracted twice similarly. Then it was filtered using a clean muslin cloth and ordinary filter paper finally by Whatman No.1 filter paper. The resulting filtrate was then concentrated using a rotary evaporator and further dried at 40°C. Finally yield 8% solid crude extract was stored in powdered form in an air tight container at (4°C) for further use in the experiment.

Animal

36 adult male albino rats (Rattus norvegicus L.) of Wistar strain weighing 130g ±10 were selected for this experiment. The animals were acclimatized to the laboratory environment for a period of one week before starting the experiment. The animals were maintained under standard laboratory conditions (12 hrs light: 12 hrs dark, 25±2°C and relative humidity 40-60%) with free access to standard normal diet (prescribed by ICMR, NIN, Hyderabad, India) and water ad libitum. All animal experiments were performed according to the ethical guidelines suggested by the Institutional Animal Ethics Committee (IAEC) guided by the Committee for the Purpose of Control and Supervision of Experiments on Animals (CPCSEA), Ministry of Environment and Forest, Government of India (Ref. no. PU 796/03/ac/CPCSEA).

Animal treatment

Animals were equally divided and randomly selected into 3 groups having 12 animals in each and were treated as -

Group I (Control): Given sterile distilled water (5 ml/ kg body weight/day) for 28 days.

Group II (Treated): Given aqueous extract (200 mg/kg body weight/day) for 28 days.

Group III (Recovery): Treated with aqueous extract (200 mg/kg body weight/day) for 28 days and allowed for next 28 days without any treatment to check the reversibility.

The daily dose was prepared by suspension the extract in 0.5 ml of sterile water and administered to each animal orally with the help of oral gavage needle. The initial body weight of animals was recorded. Animals were weighed twice weekly throughout the experiment and the dose was adjusted accordingly. Treatment schedule was selected for two seminiferous cycles (~ 28 days) consecutively. On the 29th day (24 hours after the last dose of treatment and 18 hours after fasting), 8 animals from control and treated groups, were anaesthetized by diethyl ether. Blood samples were collected and Serum samples were separated by centrifugation which was stored at - 20°C for different biochemical assay. The reproductive organs i.e. testes and epididymis of each animal were dissected out, freed from adherent tissues, blotted free of blood and wet weights were recorded on an electronic top pan balance to the nearest mg. Remaining 4 rats, each of control and treated group (group I, II) were subjected to fertility testing. In Group III, for recovery studies over a period of next 28 days after consecutive treatment, 8 animals were sacrificed. All spermatological and biochemical parameters along with the weight of testis and epididymis were repeated in order to ascertain the nature of action of extract, i.e. reversible or irreversible. The remaining 4 rats from Group III were left for fertility studies.

Sperm motility and total sperm count

One caudal epididymis of each animal in right side was rinsed and gently minced in 2 ml of phosphate buffered saline (PBS, pH 7.4). Epididymal sperm motility and sperm concentration was determined as per the method described in the WHO Manual. sperm suspension was placed on a Neubauer hemocytometer in WBC chamber and percentage motility was determined by counting both motile and immotile spermatozoa compared to total cells. The number of sperms was counted by Neubauer hemocytometer with a light microscope at X 400 magnification in a RBC counting 5 major squares and were expressed as million/ml of suspension.

Sperm viability

Sperm suspension (50µl) was mixed with eosin-nigrosin staining solution (50µl). The suspension was incubated for 30s at room temperature (20°C). Then incubated suspension was transferred with a pipette to a labeled glass slide and thin smear was prepared. Two such smears were prepared from each sample. The smears were air dried and examined directly under microscope. At least 200 sperms were studied at magnification of x400 under oil immersion with a bright field objective. Unstained sperms were considered as live and pink or red coloured sperm as dead.

Sperm morphology

Sperm morphology was studied from a total count of 200 spermatozoa in a smear prepared from above sperm suspension stained by eosin-nigrosin mixture (6.7 gm eosin and 100 gm nigrosin in 1l 0.9 gm % saline.) as observed under high power objective (magnification x 400). The defective shape and structure of either head and or tail were considered as abnormal and the data was presented as percentage incidence of total abnormalities. Sperm abnormalities were classified by the method of Bloom (1973).

Preparation of spermatozoa for SEM study

Rat spermatozoa for SEM study were prepared according to Mayuva Areekjiserre et al. Briefly, a drop of cauda epididymal spermatozoa suspension was fixed in 2.5% glutaraldehyde in 0.1 M phosphate-buffer (pH 7.4) for 2 h at 4°C and a thin film was applied on a cover slip then rinsed with same buffer. Preparation was dehydrated by graded alcohol and air dried, coated with gold, and finally observed under S-530 Hitachi SEM.
Fertility test

Fertility tests were carried out during the treatment and recovery periods (for 5 days prior to termination of treatment) along with control by allowing a male rat to mate with fertility proven females in a ratio of 1:2. Prior to mate, the females were isolated for one month to rule out pre-existing pregnancy. During mating the vaginal smear was checked daily in the next morning to observe the presence of sperm. When sperms were detected in the smear, which indicated the positive mating (sperm-positive) and the day was considered as zero day of pregnancy. The mated females were separated and observed the implantation sites on 16th day of pregnancy through laprotomy to assess the fertility rate with reference to the number of implantations. Percentage of fertility was calculated as number of pregnant female rats divided by the number of mated females multiplied by 100 according to WHO Protocol MB-50, 1983.23

Statistical analysis

The recorded values were expressed in mean ± SEM. The treated groups were compared to control using one way ANOVA with post hoc LSD test were performed using SPSS version 16 Software. The value of p<0.05 was considered to be statistically significant.

RESULTS

In the present investigation, the body weights were not affected after the treatment suggesting that the extract have no side effect of the animals throughout the experiment. But there is significant decrease in organ weights, viz., epididymis and testis at the dose of 200mg/kg b. wt/day for consecutive 28 days. The results were similar with the effects shown by J. repens extract treatment to rats in the earlier study.24 However, the organ weights were gradually recovered towards normal and the percentage of normal sperm increased towards control by 28 days after withdrawal of treatment (Table 1). Sperm analysis of treated group exhibited significant (p<0.01) decrease in total sperm count, motility, viability and increase in sperm abnormalities(58%) when compared to control (10%). However, the sperm count, motility and viability were gradually recovered to normal and the percentage of normal sperm increased towards control by 28 days after withdrawal of treatment (Table 2).

<table>
<thead>
<tr>
<th>Table 1: Body and relative weight of reproductive organs (gm) of male albino rats in different groups</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Treatment</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Group I: Control group</td>
</tr>
<tr>
<td>Group II: Treated with aqueous extract (200 mg/kg body weight)</td>
</tr>
<tr>
<td>Group III: Recovery group</td>
</tr>
</tbody>
</table>

Values are expressed as means ± S.E.M; N=8; *Significant (P < 0.05) Group II were compared to Group I (Control) & Group III (Recovery)

<table>
<thead>
<tr>
<th>Table 2: Cauda epididymal sperm characteristics of adult male albino rats of different groups</th>
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<tr>
<td><strong>Group</strong></td>
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<tr>
<td></td>
</tr>
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</tbody>
</table>

Values are expressed as means ± S.E.M; N=8; **Significant (P < 0.01), Group II were compared to Group I (Control) & Group III (Recovery).

The table 3 shows the morphological analysis of sperm cells in different groups of treatment and revealed that the extract treated group exhibits significant (p<0.01) increase in sperm abnormalities (58%) when compared to control (10%). Out of 58% abnormal sperms, the extract induced morphological alterations were found as predominantly of primary abnormalities (40%), where secondary and tertiary abnormalities were 10% and 8% respectively. Primary abnormalities include double head, hook less, pin head, banana head, head amorphous, double head, double tail, cytoplasmic droplets (light and heavy), bent neck, proximal and distal bent tail. While tail coiled around head, mid piece and below the mid piece level, headless tail and other multiple abnormalities were found as secondary abnormalities. The tertiary abnormalities were detached head and simple coiled tail. The primary abnormalities showed 5.25%, 39.4375%, 4.9375 % and secondary abnormalities were 3.5625%, 10.25%, 3.68 % where tertiary abnormalities were 2.0625%, 8.3125%, 2.3750% for Groups I –III respectively. The percentage of primary (39.4375 %), secondary (10.5%) and tertiary (8.3125%) abnormalities in Group - II (treated) were significantly (P<0.01) higher than Group I (control) and III (recovery). After 28 days of recovery, the extract induced altered sperm morphology was returned nearly to normal (Table-3, Figure 1 & 2).
### Table 3: Morphological analysis of sperm cells in different group

<table>
<thead>
<tr>
<th>Classification</th>
<th>Group - I (Control)</th>
<th>Group - II Treatment (200 mg/kg body wt./day)</th>
<th>Group - III (Recovery)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary Abnormalities</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Double Head</td>
<td>0.0000 ± 0.0000</td>
<td>0.6250* ± 0.0818</td>
<td>0.0000 ± 0.0000</td>
</tr>
<tr>
<td>Hook less</td>
<td>1.0625 ± 0.11329</td>
<td>7.4375** ± 0.29028</td>
<td>0.8125 ± 0.013153</td>
</tr>
<tr>
<td>Pin Head</td>
<td>0.3750 ± 0.08183</td>
<td>1.6875** ± 0.13153</td>
<td>0.2500 ± 0.09449</td>
</tr>
<tr>
<td>Banana Head</td>
<td>0.8750 ± 0.1250</td>
<td>4.1875** ± 0.20996</td>
<td>0.6875 ± 0.09149</td>
</tr>
<tr>
<td>Head amorphous</td>
<td>0.2500 ± 0.09449</td>
<td>2.5000* ± 0.28347</td>
<td>0.3125 ± 0.09149</td>
</tr>
<tr>
<td><strong>Secondary Abnormalities</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Double Head &amp; Double tail</td>
<td>0.0000 ± 0.0000</td>
<td>0.5000 ± 0.1889</td>
<td>0.1250 ± 0.08183</td>
</tr>
<tr>
<td>Cytoplasmic droplet (Light)</td>
<td>0.0000 ± 0.0000</td>
<td>0.6875* ± 0.09149</td>
<td>0.2500 ± 0.13363</td>
</tr>
<tr>
<td>Cytoplasmic droplet (Heavy)</td>
<td>0.0000 ± 0.0000</td>
<td>1.0000** ± 0.13153</td>
<td>0.1875 ± 0.13153</td>
</tr>
<tr>
<td>Bent neck</td>
<td>0.8750 ± 0.08183</td>
<td>10.1250** ± 0.83318</td>
<td>0.8750 ± 0.08183</td>
</tr>
<tr>
<td>Proximal bent tail</td>
<td>0.312 ± 0.09149</td>
<td>2.3125** ± 0.09149</td>
<td>0.3125 ± 0.09149</td>
</tr>
<tr>
<td>Distal bent tail</td>
<td>1.5000 ± 0.13363</td>
<td>8.3750** ± 0.68628</td>
<td>1.1250 ± 0.18298</td>
</tr>
<tr>
<td><strong>Total %</strong></td>
<td>5.2500 ± 0.35355</td>
<td>39.4375** ± 1.92363</td>
<td>4.9375 ± 0.41659</td>
</tr>
<tr>
<td><strong>Tertiary Abnormalities</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tail coiled around head</td>
<td>0.4375 ± 0.06250</td>
<td>1.3125** ± 0.09149</td>
<td>0.3125 ± 0.09149</td>
</tr>
<tr>
<td>Tail coiled around midpiece</td>
<td>0.7500 ± 0.09449</td>
<td>2.7500** ± 0.23146</td>
<td>0.6250 ± 0.08183</td>
</tr>
<tr>
<td>Tail coiled below</td>
<td>1.3750 ± 0.18298</td>
<td>2.3125** ± 0.1875</td>
<td>1.6250 ± 0.12500</td>
</tr>
<tr>
<td>Headless tail</td>
<td>1.0000 ± 0.13363</td>
<td>2.7500** ± 0.31339</td>
<td>1.0000 ± 0.13363</td>
</tr>
<tr>
<td>Multiple abnormalities</td>
<td>0.0000 ± 0.0000</td>
<td>1.1250** ± 0.35038</td>
<td>0.1250 ± 0.08183</td>
</tr>
<tr>
<td><strong>Total %</strong></td>
<td>3.5625 ± 0.31956</td>
<td>10.2500** ± 1.0220</td>
<td>3.6875 ± 0.20996</td>
</tr>
<tr>
<td><strong>Total abnormal %</strong></td>
<td>10.875 ± 0.789</td>
<td>58.000** ± 3.625</td>
<td>11.000 ± 0.597</td>
</tr>
</tbody>
</table>

Values are expressed as means ± S.E.M.; N=8. *Significant (P < 0.05), **Significant (P < 0.01), Group II was compared to Group I (Control) & Group III (recovery). Different superscript letters are significantly different; P<0.01 (a = Primary Abnormalities, b = Secondary Abnormalities, c = Tertiary Abnormalities).

**Figure 1:** Different types of sperm abnormalities - A) Normal sperm, (B) Bent neck, (C) Distal bent tail, (D) Hook less, (E) Banana head, (F) Amorphous head, (G) Pin head, (H) Proximal bent tail, (I) Cytoplasmic droplet (light), (J) Cytoplasmic droplet (heavy), (K) Tail coiled around midpiece, (L) Tail coiled around head, (M) Headless tail, (N) Detached head, (O) Multiple abnormalities, (P) Simple coiled tail, (Q) Double head and double tail. After Eosin-Nigrosin staining. Magnification: X400, Except (J) X1000.
In figure-3, the SEM (Scanning electron microscopic) observations of cauda epididymal spermatozoa in control rats showed the normal morphology (Figure 3A). The whole spermatozoon was intact with all the membranes and organelles. However, the animals fed with 200mg/kg b. wt./day of J. repens aqueous extract showed distortion in the plasma membrane and acrosomal membrane in most of the sperm heads. Serrations at the head region of the spermatozoa were observed. The shape and size of sperms were also changed considerably (Figure 3-B, D, E, F, M-S). There was acute dorsoventrally constriction in the mid-head region in most of the sperms. The sub-acrosomal material was bulged/swelled out (Figure 3-O, S). Spermatozoa showed a splitting of tail and distinct visibility of balloon-like cytoplasmic droplets in the mid region of tail. (Figure 3-I, J, K, L).

Figure 2: Percent abnormalities of different types of spermatozoa in different groups. **Significant (P < 0.01), Group II was compared to Group I (Control) & Group III (recovery).

Figure 3: Scanning electron micrographs (SEM) of cauda epididymal spermatozoa in control (Fig.A) and treated (J. repens L. extract of 200mg/kg b. wt/day for 28 days) rats (Figs. B - T) - A: Spermatozoa of control rat exhibiting normal parts of acrosome (a), post nuclear cap (c), plasma membrane(m), nucleus(n), sub-acrosomal material (p), basal plate(b) and tail region(t). Magnification: X 5000. B – T: Different morphological abnormalities observed in head and tail region of sperms in male albino rat. (→) yellow arrow = cytoplasmic droplet (light & heavy), (→) Red arrow = Absence of sub acrosomal material, swell and the constriction in middle region of sperm head, structural abnormalities in the acrosomal part and plasma membrane and also serration at the connective piece of spermatozoa.

Table 4: Fertility study on male rats of treated and recovery group.

<table>
<thead>
<tr>
<th></th>
<th>No. of mated male: female (1:2)</th>
<th>No. of Sperm-positive females</th>
<th>No. of Pregnant females</th>
<th>Mating index (%)</th>
<th>Implantation sites / rat</th>
<th>Fertility index (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I: Control group</td>
<td>4 / 8</td>
<td>8</td>
<td>8</td>
<td>100</td>
<td>10.70 ± 1.3</td>
<td>100</td>
</tr>
<tr>
<td>Group II: Treated with aqueous extract (200 mg/kg body weight)</td>
<td>4/8</td>
<td>4</td>
<td>0</td>
<td>50</td>
<td>0.00 ** ± 0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Group III: Recovery group</td>
<td>4/8</td>
<td>8</td>
<td>7</td>
<td>100</td>
<td>9.8 ± 1.1</td>
<td>87.5</td>
</tr>
</tbody>
</table>

**Significant (P < 0.01), Group II were compared to Group I (Control) & Group III (recovery).
Results of mating experiment for fertility study showed that, after oral administration of crude extract of the plant for a period of two spermatogenic cycle reduced mating index (50%) but fertility rate was ‘zero’ when compared to control (Table 4).

**DISCUSSION**

Fertility depends on the sperm quality like sperm cell concentration, motility, viability and also the morphology of spermatozoa.  

The epididymis plays an active role in sperm development and maturation.  

Our present studies showed that the *J* repens extract reduced the relative weights of the reproductive organs without affecting the body weight (Table-1) and it also altered the sperm parameters (Table-2) including sperm morphology (Table-3, Fig.1,3) significantly which support the non-toxic antagonadal activity of this extract as we reported earlier. 

Results from present studies showed that extract induced abnormal sperm percentage was 58% which was much more greater than the control group where only around 10% abnormality was found and it also altered other sperm parameters (Table-2). Similar effect with *L.breviflora* Roberts extract induced alterations of sperm parameters were also observed by Saba et al 2009. 

Chemicals, ions, and other herbal products induced alterations of spermatozoa morphology were also reported by others. 

The andrological parameters are usually considered to determine the fertility of the male individual. Zemjanis in 1977, reported that when critical percentage (>10%) of sperm cell abnormalities were present in the semen, the male subject was considered as infertile. Our studies showed that extract induced different types of sperm morphological abnormalities were 58% (Table-3) which was more than enough to develop infertility and it was confirmed by our fertility assessment study (Table-4).

According to classification by Noakes et al. 2004, the alteration of sperm cells morphology caused by *J*.repens in this study had been grouped into primary, secondary and tertiary abnormalities and found that the sperm cells abnormalities in treated group were mostly primary abnormalities (>39%) where significant abnormalities were found in head regions (Table-3, Fig.1,3). Bloom in 1977 proposed that primary defects were representing a failure of spermatogenesis in the seminiferous epithelium which was testicular in origin. Secondary abnormalities represented a failure of maturation and abnormal epididymal functions, where tertiary abnormalities develop in vitro.  

As *J*.repens developed all kinds of sperm abnormalities, it may be assumed that this extract affected the pathways for sperm production and maturation. Extract induced primary abnormalities might be due to aberrations in the process of spermatogenesis. Secondary and tertiary abnormalities found in treated group (Table 3, Fig.1,3) might be due to the defect in maturation stages. 

Presence of cytoplasmic droplets (Fig-3) supported the altered epididymal functions. Now it is known that all types of morphological abnormalities are due to the defect in spermatogenesis which is under the control of steroidogenesis processes. Abnormal sperm head morphology (Figure 3) could be due to the influence of the extract on the differentiation process of spermatogenesis and degenerative changes observed in the current study were deleterious for potential reproductive processes as reduced sperm count, viability and increase of morphological abnormalities of sperm likely to decrease fertility. So, any agent which affects these two pathways like spermatogenesis and steroidogenesis will be the cause of formation of abnormal spermatozoa which was further confirmed by our mating experiment (Table - 4) where significant reduction of mating index and zero fertility rate were observed. In our unpublished study, we have seen that oral administration of aqueous extract of *J*.repens (except root) herb inhibit both spermatogenesis and steroidogenesis pathways by inhibiting the factors responsible for these pathways, i.e., testicular ascorbic acid, cholesterol, fructose, Hydroxy steroid dehydrogenase (HSD), plasma and testicular testosterone level etc. (data communicated). It had also reported by others that the morphology of sperm was regarded to be controlled by genes  and most of the drugs, insecticides or environmental factors induced morphological alterations of spermatozoa were genotoxic or mutagenic, but non-genetic mechanisms were also known to induce morphological alterations where active metabolites of the extract directly act on germ cells.  

Our withdrawal of treatment on fertility experiment showed that *J*.repens induced infertility was reversible, as rapid restoration of sperm parameters (Table-1,2,3) and fertility rates occurred after withdrawal from treatment (Table - 4). So, as *J*. repens is a non toxic antagonadal herb and it is not supposed to cause any permanent damage to male reproductive tissues and shows reversal effects on sperm parameters and fertility index, this extract can be introduced as a potent non-toxic male herbal contraceptive in future.

**CONCLUSION**

From the above studies it may be concluded that the crude aqueous extract of *Jussavia repens* (except root) in a regulated dose and duration can be used as non-toxic, safe, herbal contraceptive for male in future.

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REFERENCES


2. Swapna MM, Prakashkumar R, Anoop KP, Manju CN, Rajith NP, A review on the medicinal and edible aspects of aquatic
and wetland plants of India, Journal of Medicinal Plants Research, 5(33), 2011, 7163-7176.


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