ANDROGENIC AND SPERMATOGENIC POTENTIAL OF METHANOLIC EXTRACTS OF TRIBULUS TERRESTRIS IN REPRODUCTIVELY DISRUPTED MALE ALBINO RATS

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Abstract

Pharmacologically, Tribulus terrestris (caltrop) holds the trait to cure sexual and reproductive disorders in Indian traditional medicinal system. To investigate the fact, effect of the Methanolic Extract of Tribulus terrestris (METt) on testosterone level and fecundity/fertility was evaluated in male rats after the incorporation of antifertility standard drug Sulphasalazine. Clomiphene citrate standard drug to enhance fertility was given at 10 mg/kg b.wt, for 60 days to compare the results. Sulphasalazine (SSZ) fed male rats showed a decrease in fertility in comparison to control group animals which were fed with the distilled water and later treated with clomiphene citrate and METt. METt treatment improved fertility (~75%), promote sperm motility and sperm density significantly. METt administration restored the serum LH, FSH and testosterone levels and reverse to standard range with a very a significant (p< 0.01) manner. Furthermore, histopathological evaluation confirmed improvement in spermatogenesis, increased seminal gland and interstitial cell count as well as restoration of testicular architecture observed in METt treated rats. METt showed a significant modulatory effect against SSZ induced reproductive disruption and proved to possess spermatogenic and androgenic potential thus justify its traditional use of the plant.

Keywords: Tribulus terrestris, methanolic extract, fertility enhancement, testicular, testosterone.

Introduction

WHO calculated approximately 60–80 million couple experience infertility worldwide at present and 10-40% are due to male factor. Human sperm quality declination increased globally in over past few decades noticeably (Sen et al., 2017). Multiple of factors influence male fertility and declination of semen quality that may be environmental as well as occupational factors along with lifestyle practices (Sharma et al., 2013). Normal Semen parameters include sperm concentration of 39 million sperm per ejaculate, 15 million spermatozoa/mL. (Nand and Singh, 2015) with progressive motility of 32%. Now attentions are shifted from synthetic drugs to natural plant products or herbal medicines and their combinations as potent fertility enhancer (Dada and Ajilore, 2009). The present investigation was carried out to evaluate the fertility enhancing effectiveness of Tribulus terrestris in experimentally induced infertility in male rats. Tribulus terrestris plant belong to family Zygophyllaceae commonly known as Gokhru a herbaceous perennial flowering plant inhabitant to temperate and tropical regions of South Europe, Asia, throughout Africa and Australia. Among all Tribulus species T. terrestris (Mahato et al., 1981), T. alatus (Temraz et al., 2006) and T. cistoides (Achenbach et al., 1994) have been phytochemically investigated for their isolated steroidal saponins. Tribulus terrestris is exclusively explored for isolated Kaempferol, kaempferol-3-glucoside, kaempferol-3-rutinoside and tribuloside (kaempferol-3-β-D-(6″-p-coumaroyl) glucoside) (Bhutani et al., 1969). T. terrestris has been used as tonic, aphrodisiac, astringent, analgesic, anti-hypertensive, diuretic and urinary anti-septic (Khanbakh and Jahanani, 2003).

The present investigation was designed to evaluate the fertility enhancing effect of the different levels of methanolic fruit extract of the T. terrestris in male albino rats.

Materials and Methods

Plant material and preparation of plant extract

T. terrestris fresh fruits were collected from Campus of Rajasthan University during period of March to May 2010 and identified and authenticated by the Department of Botany, University of Rajasthan, Jaipur, India. The shed dried fruits (500g) were crushed, powdered and extracted with the 70% methanol for 72 hours at b.p 60-80 in a soxhlet. A viscous brown material was obtained after the removal of methanol under reduced pressure, which turned dark in brown solid after washing with petroleum ether. Five hundred gram dried fruits yielded 25gm of dark brown solid.

Standard drugs

Sulphasalazine (SSZ) and Clomiphene citrate of BAL Pharm Ltd, India were used as standard drugs. Biochemical Parameters were estimated by using kits (Accurex Biomedical Pvt. Ltd, Mumbai, India).

Animals

Adult, Male albino rats of Rattus norvegicus, Wistar strain, 16-18 weeks old weighing between 150-160gm (Sengupta et al., 2013) colony bred in polivpropylene cages under standardized condition of humidity, temperature and 12 hr light/12hrdark period. Rats pallets feed (Ashirwaad Ltd.) and tap water ad libitum. Body weight of each animal in all groups was measured weekly to see the possible body weight changes throughout the experiments.

Ethical aspects

The Indian National Sciences Academy, New Delhi (INSA,2000), (26) guidelines were followed for research ethics, maintenance and the use of experimental animals. The approval of the study (Reg. No. 1678/GO/RE/S/12/CPCEA) was done by ethical committee, Center for Advanced Studies, Department of Zoology, University of Rajasthan, Jaipur (India).

Experimental Methodology

Fertile healthy male rats were divided into 4 groups of 10 rats each. The daily dose of the plant extract was freshly dissolved in 0.5 ml of distilled water and administered to each treated animal every morning for 60 days. Group 1 had Control rats received 0.5ml/day of the distilled water. Group
2 Rats treated with the sulphasalazine standard drug to produce antifertility 100 mg/kg/day. Group 3 Rats of (group 2) treated with the clomiphene citrate standard drug for fertility enhancement at 50 mg/kg/day. Group 4 Rats of (group 2) treated with the Tribulus terrestris seed methanolic extract (MeTt) at 50 mg/kg/day. All groups were administered orally after treatment with SSZ for the duration of 60 days.

Fertility Test

On day 55-60, for mating experimentation the male rats were cohabited with proestrus females at a ratio of 1:3 (Rahman et al., 2014). The presence of vaginal plugs and sperms in the vaginal smear in the next morning were considered positive mating. The mated females were sacrificed on day 16 of pregnancy to find the implantation sites.

Autopsy

Twenty-four hours after the last dosing, the males were weighed and autopsied by mild anaesthesia as per CPCSEA guideline, 2003 and all reproductive organs such as testes, cauda, seminal vesicles, ventrak prostrate and vas deferens dissected out, weighed and preserved in a deep freezers at −20°C for further biochemical and histopathological parameters analysis. Blood samples were collected by the cardic puncture.

Fertility, Sperm density and motility

The fertility percentage was calculated from formula (Number of pregnant/Number of mated) × 100. The sperm density was estimated in cauda epidedymides and testis. Spermatozoa were collected by from epidedymal tissues and then homogenized in 100ml normal saline (0.9% NaCl) and incubated at 37°C for 5-10 min to make the sperms fully detached. Sperm counting (no.x10^6/ml) done by Neubaur haemocytometer. The sperm motility was determined by counting both motile and nonmotile spermatozoa per unit area (Prasad et al., 1972).

Biochemical studies

Testicular tissue was assayed for protein, sialic acid, glycogen and cholesterol. Fructose analysis was done in the seminal vesicle. R.B.C, W.B.C counts, hemoglobin, haematocrit and the sugar was analyzed in Blood (Lowry et al., 1951; Montgomery et al., 1957; Zlatkis et al., 1953).

Serum hormonal studies

Serum protein, cholesterol, triglycerides, phospholipids and HDL-Cholesterol were determined.

Commercially available kits for enzyme-linked immunosorbent assay (ELISA) were used as standard protocol for estimation of sex hormone i.e testosterone, luteinizing hormone (LH), and follicle-stimulating hormone (FSH).

Histopathological studies

Tissues were fixed in Bouin’s fluid, passed through series of ethanol and then through xylene and then embedded in paraffin. Paraffin embedded tissues were sectioned and stained with haematoxylin and eosin to assist in discrimination of the stages of the spermatogenesis. The sections were scanned under ×400 magnifications (Abercrombie et al., 1946).

Statistical Analysis

Data are expressed as mean ± SEM and are analyzed by using one-way analysis of variance (ANOVA) followed by Duncan’s multiple comparison test (Statistical Package for Social Sciences, SPSS, v21.0), at p < 0.01 values were considered to be significant.

Results

Effect of MeTt Toxicological studies

No abnormal signs or death of animal was noticed in any of the treated group. Most of the animals treated with 50 mg/kg/day MeTt exhibited enhanced growth and appetite with no apparent conditions of toxicity such as mortality, weakness, Lethargy and negative response for physiological behavior. Therefore, the METt can be safely used for therapeutic use.

Effect of MeTt body and reproductive organ weight

This weight gain of body and testis was however significantly increased treated in (METt) treated groups as shown in Table I. Significant (P≤0.01) weight gain was observed in the testicular weights in 50mg/kg bw group . SSZ administration did not cause any significant body weight gain when compared to control rats. The relative testicular weight was significantly (P≤0.01) higher in (METt) treated group when compared with chlomiphene citrate treated animals.

Table I : Body and Reproductive organ weight

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Body Weight</th>
<th>Testes</th>
<th>Epididymides</th>
<th>Seminal Vesicles</th>
<th>Ventral Prostate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Initial</td>
<td>Final</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group-I : Control</td>
<td>184.65±3.42</td>
<td>205.67±2.13</td>
<td>1530.26±16.84</td>
<td>554.67±3.80</td>
<td>461.17±4.35</td>
</tr>
<tr>
<td>Group-II : Sulphasalazine 100mg/kg/day</td>
<td>183.56±3.12</td>
<td>189.33±2.65</td>
<td>1396.67±18.56</td>
<td>469.17±6.45</td>
<td>389.17±3.12</td>
</tr>
<tr>
<td>Group-III : Chlomiphene Citrate 10mg/kg/day</td>
<td>179.33±2.18</td>
<td>193.26±1.62</td>
<td>1491.67±10.84</td>
<td>539.17±7.80</td>
<td>416.17±4.34</td>
</tr>
<tr>
<td>Group-IV: Tribulus terrestris fruit extract 50mg/kg/day</td>
<td>189.02±2.01</td>
<td>199.5±1.55</td>
<td>1502.50±10.84</td>
<td>498.17±2.61</td>
<td>439.67±3.37</td>
</tr>
</tbody>
</table>
Effect of MeTt on Sperm Dynamics, Fertility

Fertility rate decreased drastically to (~35%) in SSZ treated group along with the other decreased fertility dynamics sperm density & motility. The progressive sperm motility in 50mg (MeTt) treated group significantly (P≤0.01) increased up to 24% when compared to Sulphasalazine treated group in table II. The cauda sperm count was highly significant (P≤0.01) by 19% in treated group when compared with SSZ aided group as shown in Table II.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Sperm Motility (Cauda epididymides) (%)</th>
<th>Sperm Density (Million/ml)</th>
<th>Fertility (%)</th>
<th>Testosterone (ng/dl)</th>
<th>L.H</th>
<th>F.S.H</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Testes</td>
<td>Cauda epididymides</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Group-I : Control</td>
<td>73.67 ±1.12</td>
<td>5.22 ±0.07</td>
<td>5.03 ±0.05</td>
<td>100%</td>
<td>4.55 ±0.10</td>
<td>4.32 ±0.07</td>
</tr>
<tr>
<td>Group-II : Sulphasalazine 100mg/kg/day</td>
<td>54.67** ±2.20</td>
<td>3.18** ±0.16</td>
<td>2.45** ±0.15</td>
<td>35.25%</td>
<td>2.88** ±0.05</td>
<td>3.21** ±0.07</td>
</tr>
<tr>
<td>Group-III : Chlomephene citrate 10mg/kg/day</td>
<td>70.50* ±0.99</td>
<td>4.95** ±0.14</td>
<td>4.18** ±0.08</td>
<td>90.65%</td>
<td>4.25** ±0.06</td>
<td>4.05** ±0.04</td>
</tr>
<tr>
<td>Group-IV : T. terrestris fruit extract 50mg/kg/day</td>
<td>68.67 ±1.45</td>
<td>4.87* ±0.08</td>
<td>3.97* ±0.10</td>
<td>75.25%</td>
<td>3.95* ±0.09</td>
<td>3.79* ±0.69</td>
</tr>
</tbody>
</table>

Effect of MeTt on Serum Hormonal assay

Hormonal assay shows significant (P≤0.01) and highly significant increase in serum hormonal concentration testosterone, LH and FSH level of the treated group compared to controlled rats as shown in Table II that was dramatically reduced when treated with SSZ previously.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Glycogen</th>
<th>Cholesterol</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Testis</td>
<td>Testis</td>
<td>Testis</td>
</tr>
<tr>
<td>Group-I : Control</td>
<td>3.06 ±0.12</td>
<td>8.30 ±0.10</td>
<td>229.61 ±6.10</td>
</tr>
<tr>
<td>Group-II : Sulphasalazine 100mg/kg/day</td>
<td>2.16** ±0.10</td>
<td>11.98** ±0.30</td>
<td>177.50 ±3.82</td>
</tr>
<tr>
<td>Group-III : Chlomephene citrate 10mg/kg/day</td>
<td>2.98* ±0.15</td>
<td>9.76** ±0.18</td>
<td>219.83** ±3.37</td>
</tr>
<tr>
<td>Group-IV : T. terrestris fruit extract 50mg/kg/day</td>
<td>2.50* ±0.12</td>
<td>10.90* ±0.14</td>
<td>196.67* ±3.33</td>
</tr>
</tbody>
</table>

Data are represented as mean ± SEM (n = 10). *(p < 0.01) statistically significant difference when compared with Normal control. (a) (p < 0.01) (*) statistically significant and highly significant (p < 0.001) difference when compared with SSZ control.

Effect of MeTt on tissue Biochemical parameters

Significant (p<0.01) reduction in Testicular sialic acid, protein, glycogen and fructose parameters were found while cholesterol concentration increased significantly in SSZ treated rats as compared to normal control rats. MeTt administration significantly altered concentrations back in to nearly normal ranges of glycogen, protein, cholesterol, fructose and sialic acid in reproductive tissues after comparing with SSZ treated rats shown in Table III.

Effect of MeTt on Histopathological parameters

The histopathological findings on the photomicrographs of testis showed degenerated shrunk seminiferous tubules with scanty spermatocytes and lumen with few leydig cells in sections treated with SSZ as shown in figure 2 compared with control sections figure 1. However, chlomephene citrate (figure 3) and MeTt treated (figure 4) group show dense spermatocytes within the tubular lumens along with the recovered leydig cells.

Discussion

Methanolic extract of T. terrestris is claimed to increase the body’s natural testosterone levels i.e. positive effect on sperm qualities and thereby investigation showed the fertility enhancing activity of the plant fruits. All data confirmed the increased sperm motility counts, testosterone and positive changes in protein, cholesterol, ascorbic acid and the significant weight gain in all reproductive organs could be due to increased androgen biosynthesis i.e. rise in serum testosterone levels (Shetty et al., 1997; Prins et al. 1991). The rats SSZ treated not only increased structural deformity but also altered sperm motility and sperm density thus leads to male reproductive disruption (Sharma & Kalla, 1994; Rangi et al, 2003). MeTt treatment leads to rise in LH that consequently increase the number and proper functioning of Leydig cells. Leydig cell in turn support to increase testosterone level. The rise in level of testosterone enhances reproductive organs weight and functions (O’Donnel et al., 1994). After treatment, enrichment of spermatogenic activities i.e. increase in sperm activities may be due to higher level of testosterone, which also supported the stimulation of Leydig cells (Zhang et al., 2001). MeTt
increased sialic acid secretion by the epididymal epithelium that is required for the maintenance of accurate functioning of sperm by facilitating them downward movement without friction (Singh and Chakravarthy, 2001). The improved seminal fructose also exhibits the enhancement of seminal parameters and eventually assisting better reproductive capacity. (Gonzales et al., 1989). Sertoli cells proper activity is essential for accurate spermatogenesis. (Grootegoed & Baarends, 2004). In our Indian traditional medicinal system natural plant remedies are widely used at the place of synthetic drugs for the treatment of male with reproductive disorders (Sharma et al., 2017). After screening in male albino rats, it could be concluded that it had effective spermatogenic and androgenic properties and T. terrestris fruits could be useful in fertility enhancing activity in males. These observations may be due to the peculiar compound present in the plant which is still need to observed more.

Conclusion

The whole experiment confirmed that oral administration of Tribulus terrestris fruit extract act significantly in case of fertility against drug (SSZ) induced male reproductive disorders and accredit for androgenic and spermatogenic nature just same as standard marketed drug clomiphene. Moreover the above done treatment also evidenced to improve all fertility parameters with no toxicity thus validating to be beneficial in fertility related disorders in close species such as human also.

Testicular photomicrographs of sections showing spermatogenesis stages (viewed under light microscope at ×400 magnification).

- (Figure 1) Normal control rats show a normal histoarchitecture of seminiferous tubule with all progressive stages of spermatogenesis. Sertoli cell and Leydig cell are also present
- (Figure 2) Sulphasalazine treated rats with reduced size of seminiferous tubule and eruption in germ cells along with the degeneration in Leydig cells are also seen
- (Figure 3) Clomiphene citrate treated rat with recovered with increase in size of seminiferous tubule and in the number of spermatogenic cells
- (Figure 4) MeTt treated rat with mature spermatozoa in lumen of seminiferous tubule. A significant recovery in Sertoli and Leydig cells

Reference


