Postweaning Consumption of Soy isoflavones Induced Alterations on Some Reproductive Parameters of Prepubertal and Postpubertal Male Wistar Rats

G Ssimbwa1, ED Eze**, O S Sheu1, OA Okpanachi1, AM Afodun2, P Nganda3 and E T Ayikobua1
1Department of Physiology, Faculty of Biomedical Sciences, Kampala International University, Western Campus, Ishaka, Bushenyi, Uganda
2Department of Anatomy, Faculty of Biomedical Sciences, Kampala International University, Western Campus, Ishaka, Bushenyi, Uganda

Abstract

Background: Soy contains phytoestrogens which are potent endocrine disruptors. Soy mainly contains phytoestrogens called isoflavones predominantly daizein and genistein.

Main body: The present study determined the effects of isoflavones consumed post-weaning on prepubertal and postpubertal Leydig and Sertoli cell numbers, spermatozoa parameters and body weight to paired testicular weight ratio of Wistar rats. In this study, three diets were formulated containing different amounts of isoflavones. The diets were formulated by adding to the base diet different quantities of novasoy which is an isoflavone concentrate.

The formulated diets contained 74.5, 235.6 and 1046.6 mg total isoflavones/kg pelleted diet, representing the isoflavones content less, equivalent or greater than that found in soy infant formula respectively. The results obtained showed that; administration of all doses of isoflavones significantly (p<0.05) increased Leydig and Sertoli cell numbers in both prepubertal and post pubertal rats as compared to the control groups. Administration of all doses of isoflavones significantly (p<0.05) decreased sperm count as compared to the control group. Administration of low doses (74.5 mg/kg) of isoflavones significantly increased sperm motility as compared to the control group. Administration of moderate (235.6 mg/kg) and high (1046.6 mg/kg) doses of isoflavones significantly (p<0.05) increased sperm deformation as compared to the control group. Administration of moderate (235.6 mg/kg) and high (1046.6 mg/kg) doses of isoflavones significantly (p<0.05) increased body weight to paired testicular weight ratio as compared with the control groups.

Conclusion Overall, the alterations brought about by isoflavones at all doses are indications of adverse effects on the male rat testicular function and this may adversely affect the functional capacities of the testes.

Keywords: Soy isoflavones; Post weaning; Sertoli cell; Leydig cell; Sperm; Prepubertal; Postpubertal; Testicular function

Introduction

Soy isoflavones have estrogenic activities and are structurally and functionally related to 17β-estradiol [1]. Genistein and daidzein plus their derivatives tend to mimic estrogen through binding to estrogen receptors [1]. Different studies have been carried out in humans indicating that exposure to endocrine disruptors such as phytoestrogens where isoflavones belong has led to decline in the quality of semen [2]. The studies have further shown that a decline in sperm count and semen volume among men have been associated with phytoestrogens [3].

Phytoestrogens have been associated with different reproductive conditions in male animals, exemplified by metaplasia of male accessory gland in cattle due to consumption of high levels of coumestrol [4], a decrease in testosterone levels and disorders of erection in rats due to consumption of daizein [5], the effects being associated with steroid regulation disruption of the epididymis [6]. Consumption of genistene and vinclozolin caused a decline in sperm motility and count as well as litter size in rats [7].

Other studies carried out in mice and humans have shown that genistene and daidzein could lead to acrosome loss and thus impair fertility in the males [8]. The alpha estrogen and androgen receptors in the testes of adult rats had their expression reduced following the neonatal exposure of the rats to genistene but sperm count and motility were not affected [9], while penile erection was reported to be affected by administration of daizein to the rats [10]. Utero-trophic effects of soy-based diets were compared to cow milk formula by Ashby JH [11], where all experimental feeds were presented in drinking bottles to the rats from PND 21/22 to 24/25. In addition, animals were also allowed ad libitum feed and fluid intake. The results showed that the rats had more preference for the soy-based diets in comparison to the normal rodent feeds and it was then concluded by the authors that the high preference for the soy-based diets increased the rats' isoflavone intake about three times the recommended amount for human infants. Thus, that limited the application of the results to what is seen in human infants. Soy infant formula was fed to marmosets during the period of lactation, an experiment carried out by Sharpe [12] and Tan [13].

After the experiment, soy formula fed rats had significantly lower plasma testosterone and increase Leydig cell abundance per testis in the absence of significant testicular weight. Since the animals were allowed to nurse from their mothers while being fed on soy infant formula, the authors suggested that the studies may underestimate the effects of soy formula on testicular development. Thus, the NTP experts concluded...
that the research did not give sufficient evidence to what is seen in infants fed with soy based infant formula [14] and this suggested further research. The NTP consideration was as a result of technical limitations in the carrying out of the researches which made them not to depict the conditions (period, concentration, and route of exposure) as seen in human infants [14].

This study was therefore aimed at determining the effect of oral administration during the post-weaning period on the pre-pubertal and post-pubertal testicular cells as well as sperm parameters of Wistar rats.

The route and mode of administration as well as age of consumption would simulate the real situation that occurs in humans, and the parameters determined later in life after the cessation of soy consumption, for both prepubertal and post pubertal stages.

Materials and Methods

Experimental design

Forty male rats aged 21 days were used in this experiment. The rats were fed on the diets containing different amounts of soy isoflavones, formulated by adding novasoy, an isoflavones concentrate to the rat base diet. The experiment was carried out at kampala international university animal house. These rats were provided with water and respective feeds ad libitum.

Four experimental diets were formulated and these included one which was isoflavone free and three of which contained varying doses of isoflavones that were 74.5, 235.6 and 1046.6 mg/kg of rat base diet representing low, moderate and high isoflavone doses respectively. These concentrations were in line with those used by McVey [15].

The rats were randomly divided into eight groups each containing five rats (Table 1). These were fed on the respective experimental diets from post-natal day 22 up to post-natal day 29. The rats were fed as follows:

With the dead rat on dorsal recumbence, the incision was made along the linear alba down the abdomen into the pelvic cavity using a pair of scissors. The surrounding tissues were removed to expose the testes on each side. The testes were then retrieved, spermatic cords severed and the testes removed. The testicular weights were then taken using a digital weighing scale. These were then stored in tissue containers with Davidson fixative and transported for testicular histology. Sperm analysis was done with the aid of a microscope, a method employed by Fathi [16]. Processing of testicular tissues was done by a method previously described by Nagao [17]. The Sertoli and Leydig cell numbers per testis were obtained by procedure clearly described by McCoard [18] and Wreford [19].

Data Analysis

The means were written with standard deviations and one way ANOVA was used to get the statistical analysis across the groups at 0.5% level of significance. The students’ t-test was then used to determine the statistical significance within groups.

Table 1: Experimental Groups

<table>
<thead>
<tr>
<th>Pre-pubertal groups</th>
<th>Group 1</th>
<th>Group 2</th>
<th>Group 3</th>
<th>Group 4</th>
<th>Group 5</th>
<th>Group 6</th>
<th>Group 7</th>
<th>Group 8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quantity of isoflavones given</td>
<td>0 mg/kg</td>
<td>74.5 mg/kg</td>
<td>235.6 mg/kg</td>
<td>1046.6 mg/kg</td>
<td>0 mg/kg</td>
<td>74.5 mg/kg</td>
<td>235.6 mg/kg</td>
<td>1046.6 mg/kg</td>
</tr>
<tr>
<td>Post-natal day of sacrifice</td>
<td>30</td>
<td>30</td>
<td>30</td>
<td>30</td>
<td>45</td>
<td>45</td>
<td>45</td>
<td>45</td>
</tr>
</tbody>
</table>

Results

Leydig cell numbers of prepubertal rats fed on different amounts of isoflavones

The Leydig cell numbers (Mean ± SD) were 0.122 ± 0.011, 0.222 ± 0.019, 0.300 ± 0.122 and 0.660 ± 0.114 ×10^6 cells for groups 1-4 respectively. The results indicated that isoflavones caused significant (p<0.05) increase in the Leydig cell numbers as shown in Figure 1.

Sertoli cell numbers of prepubertal rats fed on different amounts of isoflavones

The Sertoli cell numbers (Mean ± SD) were 15.00 ± 0.71, 15.60 ± 0.55, 20.60 ± 0.89, 25.40 ± 1.14 ×10^6 cells for groups 1-4 respectively. The results indicated that the moderate and high amounts of isoflavones led to a significant (p<0.05) increase in sertoli cell numbers as shown in Figure 2.

Leydig cell numbers for post pubertal rats fed on different amounts of isoflavones

The Leydig cell numbers (Mean ± SD) for the post pubertal rats were 0.560 ± 0.114, 0.940 ± 0.089, 1.180 ± 0.084 and 1.42 ± 0.084 ×10^6 cells for groups 5-8 respectively. The results indicated that consumption of isoflavones led to the post-pubertal increase (p<0.05) in Leydig cell numbers, as shown in Figure 3.

Sertoli cell numbers post pubertal rats fed on different of isoflavones

Sertoli cell number (Mean ± SD) for the prepubertal rats were 32.000 ± 0.707, 35.400 ± 0.89, 42.000 ± 0.707 and 45.000 ± 0.707 ×10^6 cells for groups 5-8 respectively. The results in the current study indicated the consumption of soy isoflavones led to a significant increase in the post-pubertal Sertoli cells as shown in Figure 4.

Figure 1: Leydig cell numbers of prepubertal rats fed on different amounts of isoflavones. Bars with the same superscript (p, q, r, or s) have no significant differences between them (p>0.05). KEY: GP1=Prepubertal group fed on isoflavone free diet (control), GP2=Prepubertal group fed on diet containing 74.5mg isoflavone/kg base diet, GP3=Prepubertal group fed on diet containing 235.6 mg isoflavone/kg base diet, GP4=Prepubertal group fed on diet containing 1046.6 mg isoflavone/kg base diet.

Figure 2: Sertoli cell numbers of prepubertal rats fed on different amounts of isoflavones. Bars with the same superscript (p, q, r, or s) have no significant differences between them (p>0.05). KEY: GP1=Prepubertal group fed on isoflavone free diet (control), GP2=Prepubertal group fed on diet containing 74.5mg isoflavone/kg base diet, GP3=Prepubertal group fed on diet containing 235.6 mg isoflavone/kg base diet, GP4=Prepubertal group fed on diet containing 1046.6 mg isoflavone/kg base diet.

Figure 3: Leydig cell numbers of postpubertal rats fed on different amounts of isoflavones. Bars with the same superscript (p, q, r, or s) have no significant differences between them (p>0.05). KEY: GP1=Postpubertal group fed on isoflavone free diet (control), GP2=Postpubertal group fed on diet containing 74.5mg isoflavone/kg base diet, GP3=Postpubertal group fed on diet containing 235.6 mg isoflavone/kg base diet, GP4=Postpubertal group fed on diet containing 1046.6 mg isoflavone/kg base diet.

Figure 4: Sertoli cell numbers of postpubertal rats fed on different amounts of isoflavones. Bars with the same superscript (p, q, r, or s) have no significant differences between them (p>0.05). KEY: GP1=Postpubertal group fed on isoflavone free diet (control), GP2=Postpubertal group fed on diet containing 74.5mg isoflavone/kg base diet, GP3=Postpubertal group fed on diet containing 235.6 mg isoflavone/kg base diet, GP4=Postpubertal group fed on diet containing 1046.6 mg isoflavone/kg base diet.
Sperm count of post pubertal rats fed on different amounts of isoflavones

Sperm counts (Mean ± SD) for the post-pubertal rats were 87.400 ± 2.302, 74.000 ± 1.414, 71.200 ± 1.643 and 52.600 ± 3.130 million/mL for groups 5-8 respectively. The results of the study indicated that post-weaning consumption of isoflavones led to a significant decrease (p<0.05) in sperm count, as shown in Figure 5.

Sperm motility of post pubertal rats fed on different amounts of isoflavones

Sperm motility (Mean ± SD) for post-pubertal rats were 94.300 ± 2.427, 94.880 ± 2.149, 90.360 ± 2.158 and 81.700 ± 5.347% for groups 5-8 respectively. The results obtained in this study showed that post-weaning consumption of low and high amounts led to a significant (p<0.05) decrease in sperm motility as shown in Figure 6.
Sperm deformation of postpubertal rats fed on different amounts of isoflavones

The sperm deformation percentages (Mean ± SD) for the postpubertal rats were 1.820 ± 1.018, 3.14 ± 0.835, 4.460 ± 0.586 and 10.200 ± 2.270% for groups 5-8 respectively. The results in the current study indicated that post-weaning consumption of moderate and high amounts of isoflavones led to a significant (p<0.05) increase in sperm deformation, as shown in Figure 7.

Body weight to paired testicular weight ratio of prepubertal rats fed on different amounts of isoflavones

The ratios of body weight to paired testicular weight (Mean ± SD) were 212.340 ± 141.100, 197.150 ± 57.191, 220.250 ± 63.834, 298.320 ± 216.672 for groups 1-4 respectively. The results obtained in the current study indicated that post-weaning consumption of moderate and high amounts of soy isoflavones led to a non-significant (p>0.05) changes in the body weight to paired testicular weight ratios in all groups, as shown in Figure 8.

Body weight to paired testicular weight ratio of postpubertal rats fed on different amounts of isoflavones

The body weight to paired testicular weight ratios (Mean ± SD) for the post pubertal rats were 105.870 ± 9.374, 92.386 ± 10.941, 143.680 ± 23.193 and 127.360 ± 15.710 for groups 5-8 respectively. The results obtained in the current study indicated that post-weaning consumption of moderate and high amounts of soy isoflavones led to a significant (p<0.05) increase in the body weight to paired testicular ration of postpubertal rats, as shown in Figure 9.
Discussion

Leydig cells form 20% of mass of adult testis and they secrete androgens [20]. These cells are clustered near blood vessels in the testicular interstitium [21]. They are characterized by possession of many mitochondria, large smooth endoplasmic reticulum, much lipids and prominent droplets [22]. These cells are subjected to changes both in number and function as the animal matures [22]. Results of the current study have indicated a significant increase (p<0.05) in Leydig cell numbers following soy isoflavones administration. The findings are in line with those reported by Kumi-Diaka [23].

Sertoli cells are described as nurse cells of the testis and these are the supporting cells for the developing spermatozoa [24]. They provide attachment to the semiferous tubules where the spermatozoa develop from as well as provision of nutrients [25]. The Sertoli cells have fundamental importance to the development and maintenance of spermatogenesis, as well as numerical relationship to sperm production [25].

The present study showed that consumption of low doses of isoflavones (74.5 mg/kg) produced a non-significant increase in Sertoli cell numbers while the moderate (235.6 mg/kg) and high (1046.6 mg/kg) recorded a significant increase in Sertoli cell numbers in prepubertal rats.

The results have also indicated an increase in sertoli cell numbers in post-pubertal rats following isoflavones administration. The increase in both the seroli and leydig cell numbers may be explained by the proliferative effect of isoflavones as stated by Socorro [26].

In males, sperm count is one of the parameters considered during assessment of male fertility [27]. Many factors including ejaculate failure, genetic tract obstruction, genetic disturbances, exposure to too much heat as well as some drugs can lead to reduction in sperm count [28]. The complete absence of spermatozoa, which is also described as azoospermia may be brought about by factors such as hinderance of transportation of spermatozoa due to obstruction of the passages, inadequate androgens, drugs, as well as genetic causes [29].

Results of this study have revealed a significant (p<0.05) decrease in sperm count of the post-pubertal rats. The findings are supported by the study carried out [30] and [31] which showed a reduction in sperm count following administration of isoflavones.

Normal linear progressive sperm motility and normal morphology are major factors on which male fertility depends [32]. Sperm motility is the best predictor of fertility potential in males [32].

Sperm motility is considered low when more than 50% of the sperm in the semen sample cannot move progressively within 60 mins [33]. Factors such as mistakes during semen processing, contaminated equipment, and excessive heat can be predisposing to reduction in sperm motility [34]. Several confounding factors related to diet, life style, stress and socioeconomic status also affect semen quality [34]. In the current study, a significant (p<0.05) decrease in sperm motility was shown following consumption of moderate (235.6 mg/kg) and high (1046.6 mg/kg) amounts of soy isoflavones.

Sperm morphology is among factors that affect male fertility. Among factors that may bring about abnormalities in sperm morphology include toxins such as tar in tobacco, genetic abnormalities, as well as idiopathic causes [34]. Sperm morphology is another important indicator of the reproductive ability in males since very high morphological abnormalities is an indicator of progressive reproductive failure [35]. Researchers have got different views about the magnitude of sperm morphology as an indicator of male fertility. The study done by Dada [36] described sperm morphology as a very good indicator of male fertility while the one carried out by Schmahl [37] showed it to be a poor indicator.

In the poststudy, moderate (235.6 mg/kg) and high (1046.6 mg/kg) are significantly (p<0.05) increased sperm deformity. The fact that isoflavones have been associated with decreased synthesis of steroidal hormones [38] and high levels of phytoestrogens being associated with cell death and inhibiting spermatogenesis, these factors may have contributed to the low sperm count, low sperm motility and high sperm deformation following administration of isoflavones.

The increase in testicular weights improves male fertility [39]. The reduced size and weight of testes is associated with inability in semen production [40]. However, once testicular weight becomes too great, improvement in fertility tends to be small [41]. Therefore, it is important to carefully control the weight gains to achieve optimum body and testicular weights [41].

The results of the current study indicated that a significant (p<0.05) decrease in the body weight to paired testicular weight ratio in pre-pubertal rats was caused by low doses of soy isoflavones (74.5 mg/kg) while in post-pubertal rats, moderate (235.6 mg/kg) and high (1046.6 mg/kg) amounts caused significant (p<0.05) increase in body weight to paired testicular weight ratios.

Conclusion

Our results have indicated that isoflavones variously altered the testicular function indices investigated and this may adversely affect the functional ability of the rat testes.

Acknowledgements

The authors thank the financial support provided by Kampala International University management towards the completion of this research work. Great thanks also go Mr. Bukunya Robert, a laboratory technician at Mulago, Kampala, Uganda for all support rendered. The authors also thank all staff of physiology department, kampala international university, Uganda for all their input in this research.

References

biphenoCA causes few alterations on measures of postweaning activity and learning. Neurotoxicol Teratol 34: 598-606.


