INTRODUCTION

Mammalian testicular biology is complex next to the nervous system. There are two biosynthetic events significantly contribute to male reproductive processes are spermatogenesis and spermatogenesis, thus under control of endocrine and exocrine factors peptides or steroids. Testicular steroid metabolism is complex biochemical pathway can be influenced by biologically active dietary signalling factors present at low abundance such as phytosteroids, phytotaxoids, polyphenols, and terpenoids are being considered for their role in human health and disease recently [1]. Studies with phytosteroid showed testicular Liver X Receptor (LxR) transactivation resulted to modulation of several testicular functions such as steroidogenesis, cholesterol homeostasis, and proliferative apoptosis balance affected through regulation of gene expression [2]. Two Lxr isoforms have been reported, Lxr-β was found to be expressed in all the tissues and Lxr-α expression restricted in liver, intestine, kidney, and spleen. LxRs named orphan nuclear receptor until the oxysterol was recognized as a ligand. Oxysterol are an oxygenated derivative of cholesterol, synthesized in mammalian tissues (endogenous oxysterol) at lower levels example, 24-hydroxycholesterol, 24, 25-hydroxycholesterol and thus regulating cholesterol, glucose homeostasis in animal tissues [3]. Upon the ligand binds of the binding pocket and induces receptor structural conformation changes leads to dimerization with retinoid X receptor (RxR) than translocated into nuclease were its binds to gene promoter sites for regulation of Lxs targeted gene expression in the cell [4]. Fatim-Zorah et al. cell culture studies reported that LXR-α isoform regulates sertoli cell cholesterol metabolism, whereas LXR-β regulates Leydig cell cholesterol metabolism and testosterone biosynthesis, in contrast, LxRs knockout mice based study the independent role of LxRs in testicular tissue was reported that the LXR-α is highly expressed in the Leydig cells and LXR-β in sertoli cells and both isoforms are found to be expressed in the germ cells [2]. However, which isoform of LxRs and its specific types of ligands (oxysterol) responses for testicular testosterone production was unclear.

Earlier studies with plant oxysterol 28-homobrassilinoide (28-HB) an aldo steroid induces testosterone production, 3β-HSD, 17β-HSD, Lxr-α and β gene expression in normal and diabetic rat testis, proposed that phytooxysterol modulating rat testicular steroid metabolism through LxRs activation. This study Premalatha et al. failed to specified which isomer of LxRs upregulated by 28-HB, that given the observed effect on diabetic rat testicular tissues [5]. However, 28-HB and 28-HC are active plant hormones comes under the brassilinoide family aldo-keto isomers, both isomers retained anti-glycemic property in diabetic male rat [6]. Nevertheless, elevated plasma triglyceride level was reported in 28-HB (aldo isofrom) treated diabetic rat in contrast with 28-HC (keto isofrom) reduced plasma triglyceride level in diabetic rat [7, 8]. Those observed differential influences on lipid metabolism by orally fed with 28-HB and 28-HC (100µg/150 gm bwt) in diabetic male rat that given attraction towards to study 28-HC role on testicular steroid metabolism.

MATERIALS AND METHODS

Materials

All chemicals used were of analytical grade and purchased from Sigma-Aldrich, USA. Testosterone assay ELISA kit purchased from Genomics, Bangalore, India. All chemicals used were of analytical grade and purchased from Sigma-Aldrich, USA. Testosterone assay ELISA kit purchased from Genomics, Bangalore, India.

Experimental design

Male albino wistar rats 8-10 w old and weighing about 150-180 gm were purchased from certified laboratory animal supplier Sri Raghavendra Enterprises, Bengaluru, India. They were housed in plastic cages and given atmospheric temperature (25±5 °C) with a photoperiod of 12 h light/dark cycle. They were allowed freely access water and standard diet ad libitum throughout of experiment. Animal use and care were strictly followed as per the CPCSEA regulations and Institutional Animal Ethics Committee (IAEC),Indian Council of Medical Research.

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Preparation of testicular tissue homogenate

10% (w/v) tissue homogenate was prepared in phosphate buffer (0.1 M, pH 7.4) solution. The testicular tissues were homogenized in a motorized Teflon tissue homogenizer (Remi RQ-127A, Remi Motors, Mumbai, India). The homogenates were centrifuged at 10,000 g in a Sorvall RC 5C refrigerated centrifuge and the supernatant used for the determination of enzyme activity, testosterone, cholesterol, lipid peroxidation and anti-oxidants status employing standard methods.

Estimation of plasma and testicular tissue testosterone

Plasma and testicular tissue testosterone concentration were determined by using an ELISA kit (Omega diagnostics, Scotland, UK). The assay procedure was strictly followed according to manufacturer guidelines. The intensity of the colour developed was measured using ELISA reader and the level of testosterone expressed as ng/ml.

3-beta hydroxyl steroid dehydrogenase (3β-HSD), 17-Beta hydroxyl steroid dehydrogenase (17β-HSD) activity

The activities of 3β-HSD was measured by the method of Bergmeyer (1983). 10% testicular tissue homogenate supernatants were used to determination of the activity of the enzyme. The 2.0 ml of reaction volume contained 100 μmol of sodium pyrophosphate buffer (pH 9.0), 0.5 μmol NADP, 0.08 μmol dehydroepiandrosterone (substrate), and 100 μl of tissue homogenate (enzyme source). Change in absorbance at 340 nm was a monitor at 20 sec. the interval for a 3 min. in the shinsadu UV-VIS double beam spectrophotometer. The activity of the enzymes was expressed as nmol of NADPH converted to NADP/mg protein/min.

The activities of 17β-HSD were measured by the method of Bergmeyer (1983). 10% testicular tissue homogenate supernatants were used to determination of the activity of the enzyme. The 2.0 ml of reaction volume contained 100 μmol of sodium pyrophosphate buffer (pH 9.0), 0.5 μmol NADPH, 0.08 μmol androstenedione (substrate) and 100 μl of tissue homogenate (enzyme source). Change in absorbance at 340 nm was a monitor at 20 sec. the interval for a 3 min. in a spectrophotometer. The activity of the enzymes was expressed as nmol of NADPH converted to NADP/mg protein/min [10].

Determination of lipid peroxidation and reduced glutathione level

Lipid peroxidation was measured by the method of Ohkawa et al. [1979] and reduced GSH was assayed by the method of Beutler et al. [1963].

Testicular histology

Following anaesthesia, tissues were immediately surgically removed from the rat and thoroughly washed with 0.1 M phosphate buffer pH 7.4 and transferred to 10% buffered formalin. After an overnight tissue fixation, further tissues process was carried out in the following serial steps are dehydration with alcohol, clearing with xylene, paraffin infiltration, paraffin embedding the tissues and 4 μm size section were cutting with the help of a digitalized microtome (Rotary microtome). The sections (4 μm size) were stained with haematoxylin and eosin (H/E) stain. The morphological changes of tissues were analysed under camera attached binocular microscope at 10x and 40x magnifications.

Scanning electron microscopy

Paraffin embedding tissues section (4 μm size) were used, post section processing steps are deparaffinization with xylene, dehydration with ascending grades of ethanol 20, 40, 60, 80 and 100% each step given 10 min, followed by 12 h drying in room temperature. The section was then mounted on stubs, sputter-coated with carbon and visualized under the scanning electron microscope in different magnifications.

Reverse transcriptase polymerase chain reaction

RNA was isolated from testicular tissues using the TRizol reagent protocol as for manufacturer guidelines. The cDNA was synthesized from isolated RNA employing reverse transcriptase polymerase chain reaction (RT-PCR) technique by using AMV reverse transcriptase and a total RNA (100 ng) as a template. The cDNA was as a used template to amplify the target gene of interest using gene-specific primers were LXR-α, Forward primer (5'-3') CTTGTTGCAACGGGATTTGT, Reverse primer (5'-3') CAGCCT- TGATATGAGCGCAGAG, LXR-β, Forward primer (5'-3') CTCAGGTCATCATGTTGTCATCT, Reverse primer (5'-3') AGTGAGGACATCATCTGCAAGGT. PCR amplified products were then separated by electrophoresis on a 2% (w/v) agarose gel with ethidium bromide stain. The polynucleotide band intensities were measured by densitometry [11].

Statistical analysis

Experimental results were analysed by one way ANOVA followed by post hoc test employed with the statistical software SSPS version 16 (USA). Results were expressed mean±SD.

RESULTS

Measurement of plasma testosterone level in normal and diabetic rat indicated the elevation of this steroid by 20% following administration of 28-HC to the normal control rat and 67% increase in the treated diabetic animal using the same amount of 28-HC (table 1). Diabetic control rat, however, registered plasma testosterone level as being 89% below that of the normal control. Testicular testosterone level was also determined for the control and treated groups of animals. Diabetic rat showed 93% decrease in testicular testosterone level whereas 28-HC treated diabetic group yielded 570% in this steroid level in the tissue. Treated normal control exhibited 93% increase in testosterone level in the tissue. The cholesterol content of testicular tissues showed enhancement of greater than 50% in the 28-HC treated normal control animal tissues, whereas 28-HC treatment diabetic animal yielded testicular tissues cholesterol level increased 98% compared to diabetic control. Diabetic control rat was found to reduce cholesterol level in testicular tissue by 19% (table 1).

Table 1: Testosterone and cholesterol level in 28-HC treated rat

<table>
<thead>
<tr>
<th>Group</th>
<th>Plasma testosterone ng/ml</th>
<th>Testis testosterone ng/ml</th>
<th>Testis cholesterol mg/gm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>4.99±0.06</td>
<td>1.38±0.03</td>
<td>5.74±0.2</td>
</tr>
<tr>
<td>C+100µg 28-HC</td>
<td>6.0±1*</td>
<td>2.66±0.06*</td>
<td>8.74±0.3*</td>
</tr>
<tr>
<td>Diabetic</td>
<td>1.8±0.01</td>
<td>0.10±0.02</td>
<td>4.65±0.2</td>
</tr>
<tr>
<td>D+100µg 28-HC</td>
<td>3.0±0.08*</td>
<td>0.67±0.07*</td>
<td>9.22±0.1*</td>
</tr>
</tbody>
</table>

Values are expressed±SD. Group n=6. *Group Indicates statistical significance against normal control (p<0.05). †Indicates statistical significance against diabetic control (p<0.05). 3β hydroxyl steroid dehydrogenase (3β-HSD) activity noted a 31% decrease in diabetic control compared to normal control and 95% increase in the treated diabetic rat. The treated control rat group exhibited 22% increases in this enzyme activity was also noted.
Table 2: 3βHSD and 17βHSD enzyme activity in normal, diabetic and 28-HC treated rat

<table>
<thead>
<tr>
<th>Group</th>
<th>Testis 3βHSD activity</th>
<th>Testis 17βHSD activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>5.26±0.48</td>
<td>3.63±0.45</td>
</tr>
<tr>
<td>C+100µg 28-HC</td>
<td>6.44±0.72*</td>
<td>6.94±0.93*</td>
</tr>
<tr>
<td>Diabetic</td>
<td>3.63±0.29</td>
<td>2.92±0.29</td>
</tr>
<tr>
<td>D+100µg 28-HC</td>
<td>7.01±0.95*</td>
<td>9.16±1.32*</td>
</tr>
</tbody>
</table>

Values nmol of NAD converted to NADH/mg protein/min. Values are expressed±SD. Group n=6. *Group Indicates statistical significance against normal control (p<0.05). †Indicates statistical significance against diabetic control (p<0.05).

17β hydroxyl steroid dehydrogenase (17β-HSD) activity observed 20% reduction of this enzyme activity in diabetic control rat and registering 214% increased enzyme activity in 28-HC treated a diabetic rat. The treated control showed 91% increased due to 28-HC.

Table 3: Lipid peroxidation and reduced GSH level in 28-HC treated rat testicular tissue

<table>
<thead>
<tr>
<th>Group</th>
<th>MDA (nmol/min/mg protein)</th>
<th>Reduced GSH mg/mg protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.156±0.02</td>
<td>26.6±1.92</td>
</tr>
<tr>
<td>C+100µg 28-HC</td>
<td>0.123±0.05*</td>
<td>31.52±1.2*</td>
</tr>
<tr>
<td>Diabetic</td>
<td>0.417±0.08</td>
<td>21.37±1.98</td>
</tr>
<tr>
<td>D+100µg 28-HC</td>
<td>0.172±0.04*</td>
<td>54.18±1.9*</td>
</tr>
</tbody>
</table>

Values are expressed±SD. Group n=6. *Group Indicates statistical significance against normal control (p<0.05). †Indicates statistical significance against diabetic control (p<0.05).

The LxR gene amplicons of rat testicular tissue yielded 90bp band size for LxR-α and 80bp band size for LxR-β and intensity difference of 11% increase in 28-HC treated control bands of α and 25.50% increase in 28-HC treated control bands of β isoforms was registered. In diabetic treated 12%and 14.60% that of LxR-α and LxR-β (fig. 3).

![mRNA Expression](image1)

**Fig. 3:** Testicular LxR-α and β mRNA expression following 15 d oral administration of 28-HC. Values are the means±SD. Group n=6.

*p<0.01 considered to be significant

Spermatogenesis was seen in most of the seminiferous tubules in the normal and normal treated group. Disruption of the architecture of the tubules seen frequently and foci of necrosis seen in few tubules near the periphery and rest of the tubules shows spermatogenesis in the diabetic control group. In a diabetic treated group, most of the seminiferous tubules shows spermatogenesis, few tubules shows disruption of lining membrane and focal areas of necrosis (fig. 1).
Scanning electron microscopic (SEM) examination of the testicular sections (4 µm) of the control group showed seminiferous tubules with rounded, regular outlines and all types of germ cells noted close to each other, tubules were completely occupied with the mature spermatids and facing towards the lumen (fig. 2. a). In the control treated group, there is no significant alteration compared to control and observed different stages of spermatogenic cells of spermatogenesis in most of the tubules and abundantly occupied by mature elongated spermatids (fig. 2. b). Diabetic control group was noted that the seminiferous tubules compressed abnormally, irregular intercellular space was observed in between the spermatogenic cells in the tubules (fig. 2. c) and decreased mature spermatids number compared to normal control group. In diabetic treated group shows a significant alteration compared to diabetic control, observed different stages of spermatogenic cells in most of the tubules and abundantly occupied mature elongated spermatids (fig. 2. d).

Fig. 1: Histological sections of testicular tissue (a) Control, (b) Control+28-HC, (c) Diabetic, (d) Diabetic+28-HC following 15 d oral administration of 100 µg 28-HC Hematoxylin and Eosin Staining at X 400 magnification

Fig. 2: SEM for testicular tissue (a) Control, (b) Control+28-HC, (c) Diabetic, (d) Diabetic+28-HC following 15 d oral administration
DISCUSSION

Testicular sex steroid concentration has been implicated as a marker in evaluating the expression of steroid metabolizing enzymes in a testicular tissue. Since hyperglycemia was known to suppress testis steroid metabolism in the diabetic rat, it provided a basis for evaluating 28-HC influences on rat testicular steroid metabolism. In the present study, orally fed with 28-HC induces changes on testicular 3β-HSD and 17β-HSD enzyme activity in STZ induced diabetic rat, parallelly rat plasma testosterone level is also elevated by 28-HC treated normal and diabetic rat suggesting that the elevation in testicular steroidogenic enzyme activity is probably a consequence of the observed elevation in rat plasma testosterone level [12, 13]. The observed sever decreases in testosterone level and steroidogenic enzyme activity in the diabetic animal considered as a consequent destruction of gonadic function due to hyperglycemia [14, 15]. 28-HC greatly augmented rat testicular and plasma testosterone level both in treated normal and diabetic rats even though diabetic rat testicular testosterone level was greatly attenuated conforming that this brassinosteroid keto isofrom functioned verse similar to the hydroxy form 28-HB in inducing testicular testosterone levels as reported by Premalatha et al. [16].

The plant hormone brassinosteroid isofrom 28-HC is now being considered as an isofrom in mammalian gonads influencing LxR and LXR-β function and showing relatively greater expression of LxR-β gene expression, than that of Lxr-α. Earlier reported studies Premalatha et al. using the brassinosteroid isofrom 28-HB had implicated the involvement of Lxr-α and LXR-β transcriptional regulators in rat testicular steroidogenesis. The observed increase of the biomarkers cholesterol, 3β-HSD, 17β-HSD and testosterone is therefore considered as a cellular response of 28-HC, through LxRs receptor downstream signalling cascade in the rat testicular cells. The remarkable ability to increase testosterone level by 28-HC in the present study reflects the significant biological potency associated with this compound. However, Premalatha et al. failed to specified which isofrom of LxRs gene and protein upregulated by 28-HB, thus given the observed effects on diabetic rat testicular tissues [5, 17, 18]. On the other hand Fatim-Zorah et al. reported that LxR-α isoform regulates sertoli cell cholesterol metabolism, whereas LxR-β regulates leyding cell cholesterol metabolism and testosterone biosynthesis [2]. In the present study a relatively greater LxR-β isoform mRNA expression, than that of Lxr-α in testicular tissue, suggested that LxR-β gene transcription modulating plasma and testicular testosterone levels in rat [19]. Seemingly plant oxysterol 28-HC retained a differential effect on LXR-β gene expression in a mammalian testicular cell. That’s suggestive of attempts for drug development for improving diabetic subject testicular steroidogenesis by using oxysterols, LXR-β transcriptional regulator a potent target.

Cell membrane lipid peroxidation (LPO) is a well-known process of cellular injury and marker for oxidative stress. Cell membrane peroxidation generates MDA and measurement of MDA level have been used as a marker of peroxidation status in the cells. Hyperglycemia resulted to increases reactive oxygen species (ROS) generation and induced LPO and protein glycation inactivation of the antioxidant enzymes and diminished GSH in the testicular tissues [20, 21]. Thus effects can be prevented by induction of antioxidants enzymes by oral administration of antioxidant preparation. In the present study, we noted that the increased GSH content and reduction of MDA level following 28-HC treated diabetic rats increased oxidative stress and decreased reduced GSH level were also observed in diabetic rat tissues. That’s the indicative of the 28-BC improved antioxidant status and concomitantly reduced testicular tissue LPO.

CONCLUSION

Several studies carried out in the past on effect of brassinolide family phytohormones on diabetic rats showed beneficial effects in terms of antihyperglycemic and antilipidemic. In the present study demonstrated that 28-HC increases 3βHSD and 17βHSD enzyme activity, the testosterone level in plasma and testicular cells, thus indicative of the steroidogenic potential and capable of transactivating LxR-α and β molecular operative that elucidating the observed responses in rat testicular tissue. Sublimely intake of this phyto oxysterol may improve testicular functions in animal and human.

CONFLICT OF INTERESTS

Declared none

ACKNOWLEDGEMENT

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AUTHORS CONTRIBUTIONS

Research original idea and manuscript preparation was carried out by corresponding author Prof. K. Srikumar. Experimental design and work, data analysis and manuscript draft preparation were carried out by primary author Dr. V. Athithan. Data analysis, scientific discussion and manuscript proofreading was carried out by second author Prof. R. Ramesh.

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REFERENCES


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