CYP2E1 TESTIS EXPRESSION AND ALCOHOL-MEDIATED CHANGES OF RAT SPERMATOGENESIS INDICES AND TYPE I COLLAGEN

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This study is a complex investigation of alcohol-mediated changes in CYP2E1 mRNA and protein expression in the testes, as well as spermatogenesis indices and type I collagen amino acid contents, in male rats. Wistar albino male rats were divided into two groups: I – control (intact animals), II – experimental (chronic alcoholism, exposure to a 15% ethanol aqueous solution during 150 days). The destructive changes in the spermatogenic epithelium were accompanied by a decrease in sperm number and motility time. CYP2E1 mRNA and protein expression were elevated in the testes 3 and 1.4 times, respectively. Also, significantly lower contents of lysine, glutamic acid, serine, proline, alanine, valine, and phenylalanine residues accompanied by an increase of hydroxyproline, glycine, and threonine residue contents were detected in the skin type I collagen of the experimental group. Chronic ethanol consumption caused testicular failure along with an overexpression of CYP2E1 mRNA and protein in the testes as well as quantitative changes in type I collagen amino acid contents. The profound alcohol-mediated changes in collagen type I amino acid contents may have affected the spermatogenic epithelium state. The modulation of testicular cytochrome P450 2E1 mRNA and protein expression could change the functioning of this isozyme in target organs and take part in the mechanism of ethanol gonadotoxicity.

KEY WORDS: chronic alcoholism, ethanol, gonadotoxicity, motility

Alcohol abuse has a negative effect on all three factors that influence the male reproductive function: hypothalamus-hypophys-gonads system, endocrine glands, and hormones (1). It is well-known that chronic alcohol dependence causes testicular abnormalities and sexual dysfunctions both in humans and animals (2, 3); however, the exact mechanisms of ethanol-induced damage to the male reproductive organs remain unclear. A great number of epidemiological studies have demonstrated that alcohol abuse is associated with low testosterone production and testicular atrophy (4). Ethanol-induced oxidative stress results from the combined impairment of antioxidant defences as well as from the production of reactive oxygen species by the mitochondrial electron transport chain, the alcohol-inducible cytochrome P450 (CYP) 2E1, and activated phagocytes. Furthermore, during the metabolism of ethanol, CYP2E1 also generates hydroxyl ethyl free radicals (HER). The mechanisms by which oxidative stress contributes to alcohol gonadotoxicity are still not fully clear (5).

It has also been shown that the negative effects of alcohol on the testes could set in through the initiation...
of oxidative damage as a result of the increased generation of free radicals and the exhaustion of the antioxidant pool (6). Taking into account that testicular membranes are rich in fatty acids, which are sensitive to oxidative damage, it could be suggested that lipid peroxidation is also a factor in the development of gonad dysfunction (7). Alcohol-induced changes in the testes are often connected with severe cell damage, which causes their necrosis or apoptosis (1, 8).

Some investigations have demonstrated that acetaldehyde possesses greater toxicity than ethanol during testosterone production, therefore directly inhibiting protein kinase C – a key enzyme in testosterone biosynthesis (9). It can also cause changes in the pro- and antioxidants balances of testis cells (10). Ethanol conversion to acetaldehyde and free radicals may take place directly at the testes. CYP2E1, P450 reductase, and other enzymes with lipoxygenase/peroxidase-like behaviour could also be involved (11).

Ethanol induces changes in the protein metabolism of probably every organ or tissue system; e.g., increased hepatic collagen resulting from cirrhosis, myosin reduction resulting from cardiomyopathy, or the loss of skeletal collagen resulting from osteoporosis (12). Oxidative mechanisms can contribute to changes in collagen biosynthesis by triggering the release of pro-fibrotic cytokines and collagen gene expression in target cells (5).

Chronic alcoholism (6 g kg\(^{-1}\) per day \(\times\) 60 days) has been associated with fatty liver and collagen accumulation (13). Collagens VI and XIV, procollagen-III-N-propeptide, and hyaluronic acid appear to be sensitive markers of fibrotic transformation in alcoholics. The correlation between procollagen-III-N-propeptide and TGF-beta1 emphasizes their role in hepatic fibrogenesis (14). Collagen obtained from ethanol-fed rats showed alterations in solubility properties, increased fluorescence, peroxidation, and aldehyde content (13). Certain quantitative changes in the collagen contents and qualitative changes of its structure have been demonstrated (15).

Considering that quantitative changes in collagen structure correlate with male reproductive system parameters such as percentage of normal sperm and sensitivity to testicular degeneration (16), an analysis of the effects that alcoholism has on collagen and the reproductive system is of great importance.

The aim of this study was to investigate the alcohol-mediated changes of testes CYP2E1 mRNA and protein expression, spermatogenesis indices, and type I collagen amino acid contents in male rats.

**MATERIALS AND METHODS**

Wistar albino male rats, body weight (b.w.) of 150 g to 170 g, were used in the study. They were kept under a controlled temperature (from 22 \(^{\circ}\)C to 24 \(^{\circ}\)C), relative humidity of 40 % to 70 %, lighting (12 h light-dark cycle), and on a standard pellet feed diet (“Phoenix” Ltd., Ukraine). The study was performed in accordance with the recommendations of the European Convention for the Protection of Vertebrate Animals used for Experimental and other Scientific Purposes and approved by the Institutional Animal Care and Use Committee.

For the experimental (chronic alcoholism) model, reproducing male rats were selected according to the method for measuring voluntary alcohol self-administration in rats, which provides a continuous choice between an alcohol solution and water (two-bottle preference test) (17). The six selected rats were used for chronic alcoholism modelling by replacing water with a 15 % ethanol solution during 150 days (18). Six intact male rats (of the same age and weight) were used as controls. From the beginning of the experiment, they were kept in the same conditions as experimental animals but were given only water *ad libitum*.

After 150 days, both the experimental and control rats were sacrificed under a mild ether anaesthesia by decapitation. The skin, testes and epididymis were used for investigation.

Epididymal sperm quality was evaluated according to generally accepted methods (19). The right testicle was used for the evaluation of morphologic and morphometric parameters and spermatogenesis. It was fixed in a 10 % solution of neutral formalin, dehydrated in ethanol solutions, and embedded in paraffin. Histologic sections (6 mm) were stained by haematoxylin and eosin. The histological examination was performed under a light microscope (100 x). The determination of the spermatogenic index was carried out according to a four-point system (19). Simultaneously, the qualitative changes to the spermatogenic epithelium were evaluated: cells desquamation (shedding of epithelial elements), epithelium exfoliation (detachment) from the tubule basal membrane, and the presence of cell-free regions (“windows”).
Skin and bone type I collagen was extracted and purified according to Bondarenko et al. (20). All of the procedures were carried out in cold regime (+4 °C). The fractionation of pure type I collagen was carried out with NaCl according to Trelstad et al. and Rubin et al. (21, 22). Collagen preparations purity was controlled electrophoretically (23). Collagen fractions were hydrolysed for 24 h with 6 mol L⁻¹ HCl at 105 °C (24). Their amino acid compositions were analysed by ion exchange chromatography with the AAA-881 amino acid analyser (Czech Republic).

The expression of CYP2E1 mRNA in the testes was determined by a reversed transcriptase polymerase chain reaction (RT-PCR). Testes samples (50 mg) were collected, quickly frozen in liquid nitrogen, and stored at -80 °C before RNA extraction. The isolation of total mRNA was carried out with a TRI-Reagent (Sigma, USA). The integrity and concentration of RNA was analysed in a 2 % agarose gel. First-strand complementary DNA was synthesized using a First-Strand cDNA Synthesis Kit (Fermentas, Germany). The reaction mixture contents for PCR, amplification protocol, and specific primers for the CYP2E1 gene were chosen according to S.M. Lankford et al. (25). The primer sequences were: sense, 5’-CTTCGGGcccAGTGTCAC-3’ and anti-sense, 5’-CCCATATCAGAGTTGTGC-3’. RT-PCR with primers of β-actin (sense, 5’ – GCTCGTCGTCGACAACGGCTC – 3’ and anti-sense 5’ – CAAACATGAT CTGGGTCATCTTCT – 3’) was carried out for internal control. All of the primers were synthesized by “Metabion” (Germany). The MyCycler thermocycler (BioRad, USA) was used for amplification. PCR products (CYP2E1-744 bp and β-actin-353 bp) were separated in a 2 % agarose gel, stained with ethidium bromide, and visualized under a UV transilluminator (BIORAD, USA). Data analysis was carried out with Quantity One Software (USA) and presented in relative units as the ratio of CYP2E1 mRNA contents and β-actin mRNA contents.

CYP2E1 protein concentration was determined by Western Blot analysis, using specific polyclonal antibodies for mouse CYP2E1 protein, produced in the Institute of Molecular Biology & Genetics of the National Academy of Sciences of Ukraine. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as an internal loading control with polyclonal antibodies (Sigma, USA). The protein concentration in the cell lysate was quantified by the Bradford method (26). Equal amounts of protein per sample (50 μg) were used. The electrophoretic fractionation of proteins was carried out in 12 % PAAG (with 0.1 % SDS) according to the Laemmli method (27). The proteins were transferred to the nitrocellulose membrane. After blocking in a TBST buffer, which contained 5 % defatted dry milk, the membrane was incubated with CYP2E1 antibodies (1/400, v/w), and then with anti-rabbit IgG secondary antibodies (Sigma, 1/500, v/w). The CYP2E1 protein was visualized by chemiluminescence. Blots were photographed and densitometric analysis performed using the ImageJ software. CYP2E1 protein data was presented in relative units as the ratio of CYP2E1 protein contents and control GAPD protein contents in the same gel band.

The obtained data were calculated by one-way analysis of variance (ANOVA) and expressed as the mean±standard error of the mean (M±S.E.M.). Data were compared using Tukey’s test. Differences were considered to be statistically significant at *p*<0.05 (28).

RESULTS

Cauda epididymal sperm quantitative and qualitative parameters significantly deteriorated after 150 days of 15 % ethanol treatment (Table 1). The sperm count and motility time decreased in comparison with the control group by 36 % and 14 %, respectively. Simultaneously, a tendency toward a lower sperm osmotic resistance was also registered.

Almost regardless of the cellular target within the reproductive system, the most common morphological negative consequence was a disturbance in spermatogenesis. In our experiment, the impairment of sperm quality in alcohol-dependent rats was accompanied by a development of destructive changes in their spermatogenic epithelium. According to the data from Table 2, the spermatogenic index in the experimental group decreased in comparison with controls (along with mitotic activity and spermatogonia number). At the same time, the number of cells during the 12th stage of meiosis, characterized by the meiotic division of primary spermatocytes, demonstrated only a tendency toward decreasing.

Apart from the abovementioned quantitative changes, qualitative changes in the spermatogenic epithelium were also present in the seminiferous tubules. Increased (4.1 times) epithelial cell desquamation levels were observed in the experimental
group (Table 2). Extreme degenerative changes in the
testes, such as epithelium exfoliation from the tubule
basal membrane and the presence of cell-free regions
(“windows”), were also present in a large quantity of
experimental animals (Figure 1, Table 2).

To evaluate the effect of ethanol exposure on the
transcriptional activation of CYP2E1 in the testes, an
RT-PCR was performed. As the ethanol-mediated
induction of CYP2E1 is known to occur through
protein stabilization (29), we used Western Blot
analyses to assess the protein levels. We found that
the amount of the CYP2E1 protein in the testes of
alcohol-treated rats increased 1.4 times (Figure 2).
The increase in CYP2E1 protein was accompanied by
a corresponding increase of CYP2E1 mRNA content,
as determined by the RT-PCR (Figure 3).

According to the data from Figure 2, chronic
alcoholisation caused a three-fold increase in CYP2E1
mRNA expression in rat testes in comparison with
controls.

We also analysed the effect of chronic alcoholism
on collagen amino acid content. The changes in rat
skin type I collagen amino acid content induced by
continuous alcohol consumption are shown in Table
3. Statistically significant changes were registered for
10 amino acids.

Skin type I collagen of alcoholic rats contains
significantly lower content of lysine (-12.4 %),
glutamic acid (-9.04 %), serine (-11.9 %), proline
(-3.7 %), alanine (-9.8 %), valine (-22.3 %),
phenylalanine (-16.7 %) residues, accompanied by
higher contents of hydroxyproline (+25.1 %), glycine
(+2.7 %), and threonine (+10.1 %) residues. It is clear
that chronic alcohol consumption caused certain
quantitative changes in collagen molecules, as
previously demonstrated for other pathologies
associated with CYP2E1 induction (16, 20).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Rat sperm parameters after 150 days of 15 % ethanol consumption (M±S.E.M., n=6)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Indices</strong></td>
<td><strong>Control</strong></td>
</tr>
<tr>
<td>Sperm number / x10⁶ mL⁻¹</td>
<td>82.17±7.07</td>
</tr>
<tr>
<td>Sperm motility time / min</td>
<td>423.16±17.70</td>
</tr>
<tr>
<td>Sperm osmotic resistance / % KCl</td>
<td>2.83±0.22</td>
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</table>

* p<0.05 in comparison with control
DISCUSSION

In our experiment, chronic ethanol self-administration in male rats caused the development of testicular failure manifested by the deterioration of the quantitative and qualitative parameters of cauda epididymal sperm and destructive changes in the spermatogenic epithelium. These results are in accord with data from other authors, who have demonstrated that chronic alcohol intoxication is accompanied by testicular injury (2, 3). Ethanol oxidation is a necessary stage for its toxic realization. Quintans et al. (11) proved that the metabolic transformation of ethanol to acetaldehyde and free radicals could take place directly in the testes.

The molecular regulation of CYP2E1 induction is realized through transcriptional, post-transcriptional, and post-translational mechanisms and depends on the dose, duration, manner of administration, and specific structure of the inducing agent (30). CYP2E1 induction by ethanol has been suggested to include two stages: a post-transcriptional mechanism for low doses and a transcriptional mechanism for high doses (31). Our results may have proven that chronically administered ethanol induces CYP2E1 expression not only in the liver [as was previously described in a number of reviews (35, 37-39)], but also in the testes, possibly through each of the abovementioned mechanisms. The increase in CYP2E1 mRNA content from our experiment suggests a transcriptional

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Spermatogenic epithelium indices in control and experimental rats after 150 days of 15 % ethanol consumption (M±S.E.M., n=6)</th>
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<tr>
<td>Indices</td>
<td>Animal groups</td>
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<td></td>
<td>Control</td>
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<tr>
<td></td>
<td>Experimental</td>
</tr>
<tr>
<td>Spermatogenic index (stages of spermatogenesis total / number of examined tubules)</td>
<td>3.623±0.012</td>
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<td>3.523±0.027*</td>
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<td>Spermatogonia / number per tubular cross section</td>
<td>67.605±0.890</td>
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<td>57.845±1.864*</td>
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<tr>
<td>12th stage of meiosis / %</td>
<td>2.667±0.715</td>
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<td>2.000±0.258</td>
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<tr>
<td>Desquamated epithelium / %</td>
<td>1.167±0.447</td>
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<tr>
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<td>4.833±1.537</td>
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<tr>
<td>Exfoliation of epithelium / %</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>0.833±0.307*</td>
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<tr>
<td>“Windows” / %</td>
<td>0.500±0.342</td>
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<td>3.333±1.085</td>
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*p<0.05 in comparison with control

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<tr>
<th>Table 3</th>
<th>Rat skin type I collagen amino acid contents (residues/1000 residues) in control and experimental animals after 150 days of 15 % ethanol consumption (M±S.E.M., n=6).</th>
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<tr>
<td>Amino acid</td>
<td>Animal groups</td>
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<tr>
<td></td>
<td>Experimental</td>
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<tr>
<td>Hydroxylysine</td>
<td>4.30±0.28</td>
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<td>4.60±0.90</td>
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<tr>
<td>Lysine</td>
<td>29.80±0.70</td>
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<td></td>
<td>26.10±0.62*</td>
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<tr>
<td>Histidine</td>
<td>4.69±0.40</td>
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<td>4.70±0.20</td>
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<tr>
<td>Arginine</td>
<td>49.80±3.60</td>
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<td>48.80±1.24</td>
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<tr>
<td>Hydroxyproline</td>
<td>90.90±1.30</td>
</tr>
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<td></td>
<td>113.0±2.60*</td>
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<tr>
<td>Aspartic acid</td>
<td>46.70±4.10</td>
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<td>48.90±3.96</td>
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<td>Threonine</td>
<td>17.80±0.20</td>
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<tr>
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<td>19.60±0.40*</td>
</tr>
<tr>
<td>Serine</td>
<td>38.80±0.20</td>
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<td>34.20±0.20*</td>
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<tr>
<td>Glutamic acid</td>
<td>75.20±0.50</td>
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<td></td>
<td>68.40±2.40*</td>
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<td>Proline</td>
<td>127.10±0.70</td>
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<td>122.40±0.80*</td>
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<td>Glycine</td>
<td>323.70±2.70</td>
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<td>332.56±1.22*</td>
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<tr>
<td>Alanine</td>
<td>104.40±3.10</td>
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<td>94.20±1.36*</td>
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<tr>
<td>Valine</td>
<td>27.10±0.60</td>
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<td>21.05±0.95*</td>
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<tr>
<td>Methionine</td>
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<tr>
<td>Leucine</td>
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<td>29.10±5.25</td>
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<tr>
<td>Tyrosine</td>
<td>3.70±0.20</td>
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<td>3.80±0.32</td>
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<tr>
<td>Phenylalanine</td>
<td>12.00±0.50</td>
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<td></td>
<td>10.00±0.30*</td>
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*p<0.05 in comparison with control
regulation of its expression, or mRNA stabilization. However, taking into account the considerable increase of CYP2E1 protein content, its stabilization by ethanol and the prolongation of its half-life time could also be presumed (30, 33, 34).

The CYP2E1 expression increase in the testes of alcoholic rats could have had a pronounced importance as a mechanism that contributes to the development of testosterone-dependent testicular malfunction. At the present, data regarding the mechanism of alcohol-mediated testosterone biosynthesis inhibition in the testes are both controversial and contradictory (1). Considering that the chronic consumption of ethanol led to a significant induction of the testicular CYP2E1 isoform, the stimulation of free radicals could be a subsequent negative stage (30, 34). Other authors have confirmed this conclusion by demonstrating changes in pro- and antioxidant balances in rat testes after repeated administrations of ethanol (35). Lipid peroxidation is a key factor in damaging cell membranes, which means that cytotoxic action could be realized not only by oxygen reactive forms but also by the terminal products of the peroxide catabolism. Testis CYP2E1 is localized in the Leydig cells, where testosterone biosynthesis takes place (36, 37). Consequently, these structures and the damage of their microenvironment by free radicals resulting from ethanol biotransformation could be a reason for the decrease in steroidogenesis enzyme activity. The cell membrane damage caused by enhanced lipid peroxidation is one of the causes for the decrease in the membrane-related enzymatic activity involved in steroid biosynthesis. In our opinion, the activation of CYP2E1-dependent ethanol-metabolizing systems in steroidogenic cells could determine at least part of the negative effects on the testes caused by alcohol. Furthermore, the CYP2E1-dependent generation of reactive oxygen species could play a role in the inhibition of sperm motility and loss of fertility (38).

In our experiment, changes mediated by chronic alcohol consumption could have also partially resulted from the degenerative processes in the reproductive organs caused by disturbances in the protein, lipid, and nucleic acid metabolisms of the rats. As a result, acetaldehyde was able to produce adducts upon reaction with amino groups, nucleic acid bases, or phosphatidylethanolamine and others (39-41).

An important aspect of spermatogenesis is the detachment of germ cells from the basement membrane and their subsequent migration towards the tubule lumen. Procollagen I, a precursor of type I collagen, is a trimer consisting of two α1 chains and one α2 chain whose sequences are encoded by two different genes; COL1A1 and COL1A2, respectively (42). The distribution of procollagen I within the seminiferous tubules of immature and adult mice correlates with the process of germ cell attachment and detachment from the basement membrane. The unique distribution pattern of procollagen I in adult mouse testes implies a possible role for COL1A1, COL1A2, and procollagen I in regulating the adhesion of spermatogonia and preleptotene spermatocytes to the basement membrane and the detachment and migration of spermatocytes and spermatids towards the lumen during spermatogenesis (43).

The alcohol-mediated changes in collagen amino acid composition reported here may induce disturbances in physicochemical qualities. Lysine residues, along with hydroxylisine and histidine (44), participate in collagen cross-linking. Changes in the ratio of hydroxylysine : lysine : histidine residues could seriously influence the number and type of cross-links in the collagen fibrils. This might induce changes in the mechanical strength and elasticity/rigidity of the extracellular matrix. Our results regarding changes in lysine, proline, and hydroxyproline residues are in accordance with data from other authors (45-47).

Changes in the number of glutamic acid, threonine, serine, and threonine residues could cause changes in the surface charge of collagen molecules (49, 53), whereas changes in the quantity of valine, alanine, and phenylalanine residues could influence the level of collagen helix rigidity (49). In addition, changes in glycine residues could affect the number of Arg-Gly-Asp domains responsible for the processes of cell adhesion to collagen structures (50-52). Glycine residues in collagen molecules are also part of the special loci responsible for interactions with chaperones as well as for procollagen to collagen processing (53).

Our study indicates the presence of qualitative changes in the skin type I collagen of rats exposed to chronic alcohol consumption. These changes could lead to alterations in the helix structure, surface charge, rigidity, number, types of cross-links and specific loci responsible for cell adhesion, interaction with chaperons, and the processing of procollagen to collagen.

We could hypothesize that these changes are caused by the impact of ethanol and the products of its metabolism on collagen synthesis (54). The pathological changes in the amino acid metabolism...
could also influence the collagen metabolism (55). The ability of most amino acids to regulate protein biosynthesis by the stimulation of 70kD-ribosomal proteinS6-kinase has been established in vitro (56).

On the other hand, taking into account the existence of collagen gene polymorphisms (57, 58), the changes could have also been the result of disturbances in the transcription rates of different genes from the same collagen type I superfamily, as it was previously demonstrated for osteogenesis imperfecta (59). Such changes to collagen molecules could thus affect the properties and correct functioning of the spermatogenic epithelium and other reproductive organ tissues.

It could also be hypothesized that the changes in collagen structure mediated by chronic ethanol consumption were partially caused by oxygen reactive forms produced by cytochrome P450 2E1 (30, 34). As it was previously mentioned, these reactive oxygen species were reported to mediate the paracrine stimulation of type I collagen synthesis in different stages of this process (60) and caused genotoxic effects on rodent germ cells (61).

CONCLUSION

The results obtained in this investigation demonstrate that chronic ethanol administration in male rats caused testicular failure accompanied by an overexpression of CYP2E1 mRNA and protein in the testes, as well as quantitative changes in type I collagen amino acid contents. It could be suggested that profound alcohol-mediated changes in collagen type I amino acid contents unfavourably affected the spermatogenic epithelium state. The modulation of testicular cytochrome P450 2E1 mRNA and protein expression could change the functioning of this isozyme in target organs and thus take part in the mechanism of ethanol gonadotoxicity. Our data add to the understanding of the pathogenetic mechanisms of male infertility associated with the overconsumption of alcohol.

REFERENCES


Sažetak

EKSPRESIJA CYP2E1 U TESTISIMA ŠTAKORA I ALKOHOLOM PROUZROKOVANE PROMJENE INDEKSA SPERMATOGENIZE I KOLAGENA TIPA I

Ovo istraživanje proučava alkoholom uzrokovane promjene u ekspresiji CYP2E1 mRNA i bjelančevina iz testisa, indeksu spermatogeneze i aminokiselinskom sastavu kolagena tipa I u muških štakora. Albino štakori tipa Wistar podijeljeni su u dvije skupine: I – kontrolna, II – eksperimentalna (kronični alkoholizam, izloženi 150 dana 15-postotnoj vođenoj otopini etanola). Destruktivne promjene u spermatogenetskom epitelu popraćene su smanjenjem broja i pokretljivosti spermija. Ekspresija mRNA gena CYP2E1 i bjelančevina bila je povišena u testisima 3, odnosno 1,4 puta. Također, u kolagenu tipa I ustanovljene su značajno manje količine lizina, glutaminske kiseline, serina, prolina, alanina, valina i fenilalanina, te veće količine ostataka hidroksiprolina, glicina i treonina. Kronična konzumacija etanola uzrokovala je otkazivanje testisa uz izraženu ekspresiju mRNA CYP2E1 i bjelančevina u testisima, te kvantitativne promjene u aminokiselinama kolagena tipa I. Izražene alkoholom prouzrokovane promjene mogle su utjecati na spermatogenetski epitel. Modulacija ekspresije mRNA testikularnog citokroma P450-2E1 i bjelančevina mogla bi promijeniti djelovanje ovoga izozima u ciljnim organima te sudjelovati u mehanizmu gonadotoksičnosti etanola.

KLJUČNE RIJEČI: etanol, gonadotoksičnost, kronični alkoholizam, pokretljivost

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