Research Article

Toxic Effects of Trazodone on Male Reproductive System via Disrupting Hypothalamic-Pituitary-Testicular Axis and Inducing Testicular Oxidative Stress

Sinem İlgin, Gözde Aydoğan-Kılıç, Merve Baysal, Volkan Kılıç, Mina Ardiç, Şeyda Uçarcan, and Özlem Atlı

1Faculty of Pharmacy, Department of Pharmaceutical Toxicology, Anadolu University, Eskisehir, Turkey
2Faculty of Science, Department of Biology, Anadolu University, Eskisehir, Turkey

Correspondence should be addressed to Sinem İlgin; silgin@anadolu.edu.tr

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Depression and anxiety are recognized as public health problems. Epidemiological studies have shown that depression and anxiety often occur during reproductive ages between 20 and 60 years of age in males. Trazodone is one of the most frequently prescribed drugs in the treatment of depression and anxiety. Drugs used in repeated doses also play a role in the etiology of infertility. In our study, it was aimed to identify the possible toxic effects of trazodone on male rats and elucidate the underlying mechanisms. Vehicle or trazodone (5, 10, and 20 mg/kg/day) was administered to rats for 28 consecutive days (n = 8 per group). At the end of that period, sperm concentration, motility, morphology, and DNA damage were determined and testicular morphology was assessed histopathologically in rats. Additionally, we investigated hormonal status by determining serum testosterone, FSH, and LH levels and oxidative stress by determining glutathione and malondialdehyde levels in testicular tissue to elucidate mechanisms of possible reproductive toxicity. According to our results, sperm concentration, sperm motility, and normal sperm morphology were decreased; sperm DNA damage was increased in trazodone-administered groups. Degenerative findings on the testicular structure were observed after trazodone administration in rats. Additionally, serum FSH, LH, and testosterone levels were elevated in the trazodone-administered groups. Increased MDA levels were the signs of enhanced oxidative stress after trazodone administration in testis tissues. Thus, we concluded that trazodone induced reproductive toxicity in male rats; this reproductive toxicity was accompanied by oxidative stress and hormonal changes, which are considered as important causes of reproductive disorders.

1. Introduction

Reproductive health affects the quality of life of the individual as well as both maternal and fetal health during pregnancy, newborn, infant, and child health after pregnancy. In this respect, reproductive health has become an important research area in recent years [1, 2]. Infertility, which is defined as the failure to achieve a clinical pregnancy after 12 months or more of regular unprotected sexual intercourse, is an important disease of the reproductive system. Although it is considered as a widespread health problem worldwide, its incidence and prevalence is very difficult to determine [2–4]. At least 30 million people are stated to be infertile worldwide. It is even more difficult to determine the incidence of male infertility due to cultural constraints [3]. The etiology of infertility arises from female-derived factors, male-derived factors, and both of female- and male-derived factors, which are 40%–50%, 30%, and 20% of all cases, respectively [5, 6]. It could be said that direct or indirect male-derived infertility constitutes approximately 30–50% of the infertility cases [3, 7].

In the etiology of the male infertility, several factors such as varicocele, accessory gland infection, immunological factors, malignancies, genetic abnormalities, endocrine disturbances, and congenital abnormalities have been reported [8–10]. However, idiopathic causes account for 30–40% of...
male infertility [10, 11]. Obesity, radiation, climate, environmental factors, occupation, and various chemical agents such as pesticides, environmental pollutants, industrial products, and drugs may play a role in the etiology of male infertility [12–15]. The effects of many drugs on male reproductive system have been evaluated in numerous studies. Particularly, drug exposure in repeated doses causes infertility by affecting sperm parameters such as sperm count, motility, and morphology, hypothalamic-hypophyseal axis, Sertoli/Leydig/germ cells in testicular structure, and auxiliary sex organs [16–19].

Depression is a common disease affecting about 300 million people worldwide [20]. Selective serotonin reuptake inhibitors (SSRIs) such as sertraline, fluoxetine, citalopram, escitalopram, paroxetine, fluvoxamine, and trazodone (TRZ) are often prescribed for the treatment of depression due to their high efficacy, safety, and tolerability. SSRIs are known to cause sexual adverse effects (erectile dysfunction, decreased libido, and anorgasmia) [21]. It is also known that serotonin plays a role in reproductive hormonal regulation and spermatogenesis [22, 23]. Some studies have indicated relationship between infertility and hypersecretion [22]. Additionally, clinically studies, between SSRI and sperm parameters, have also been reported that sperm count, sperm motility, and normal sperm morphology were decreased in patients who were undergoing SSRI treatment [24–31].

TRZ is an antidepressant of the serotonin antagonist and reuptake inhibitor class. TRZ also has anxiolytic and sleep-inducing effects [32, 33]. It is the most prescribed drug for the treatment of insomnia [34]. Otherwise, it is noteworthy that there has been no published study on reproductive toxicity of TRZ which is frequently used during reproductive ages in males. Therefore, in this study, it was aimed to investigate male reproductive toxicity by determining the sperm concentration, motility and morphology, DNA damage, and histopathologic examination of testis tissues after TRZ exposure at repeated pharmacological doses in rats. In addition, a possible mechanism of action of the adverse reproductive effects was evaluated via the determination of the oxidative status of testis tissue and serum testosterone and follicle-stimulating hormone (FSH) and luteinizing hormone (LH) levels.

2. Materials and Methods

2.1. Materials. Desyrel® (50 mg tablet, Angelini Pharma) was used for experimental studies. Testosterone, FSH, LH, MDA, and GSH levels were determined using ELISA kits from Cusabio Biotech Co. Ltd. (Wuhan, China).

2.2. Animal Model and Experimental Design. Male Sprague-Dawley rats (12 weeks old, 200–250 g) were obtained from Anadolu University Research Center for Animal Experiments. The rats were housed in a room with controlled temperature (24°C) with a 12 h light/12 h dark cycle (lights on at 08:00 h) with free access to standard rat food and water. Temperature, sound, and light conditions of the laboratory were maintained during the course of the experiments. Animals were acclimatized to the laboratory environment for at least 48 h before experimentation. The experimental protocol was approved by the Local Ethical Committee on Animal Experimentation of Anadolu University, Eskisehir, Turkey (file registration number 2015-08). The rats were assigned randomly into the following treatment groups:

(i) Control group: animals received distilled water orally for 28 days ($n = 8$).

(ii) 5 mg/kg TRZ-treated group: animals received 5 mg/kg dose TRZ orally for 28 days ($n = 8$).

(iii) 10 mg/kg TRZ-treated group: animals received 10 mg/kg dose TRZ orally for 28 days ($n = 8$).

(iv) 20 mg/kg TRZ-treated group: animals received 20 mg/kg dose TRZ orally for 28 days ($n = 8$).

Pharmacological doses, which were determined in previous studies, were chosen as the study doses of TRZ [35–40]. Furthermore, the clinical dose of TRZ for the treatment of depression is 250–600 mg per day and the proper dose for the treatment of insomnia is 25–100 mg before sleep [41]. The doses we have chosen were in accordance with the guidelines extrapolating human doses to animal doses [42]. All drugs were administered at a volume of 1 ml/100 g by dissolving in distilled water. Concentrations were adjusted to deliver the intended dose levels of the base compound. The treatment period was in accordance with the guideline OECD 407: repeated dose oral toxicity study in rodents [43]. Additionally, 28-day period is considered as suitable for determining xenobiotic-induced reproductive toxic effects in male rats [44].

At the end of 28 days, the animals were anesthetized by intraperitoneal injection of 1.5 mg/kg urethane [45]. Blood samples for hormonal analysis (FSH, LH, and testosterone) were collected from the right ventricle of the animals via syringe. The animals were euthanatized via withdrawal of large amounts of blood from the heart.

Testis and epididymis tissues were removed. The left testis and epididymis were cleaned of blood and contaminants in PBS and fixed for histological examination. The cauda of the right epididymis was used to evaluate sperm parameters [46–48].

2.3. Collection and Evaluation of Sperm Samples. Spermatozoa obtained immediately after euthanizing the rats from the right epididymis, which was placed in a Petri dish containing DMEM/Hams F-12 at 37°C. The cauda epididymis was transferred to a new Petri dish with 1 ml of the same medium, and the blood vessels and fat tissue were removed. A section of the cauda epididymis (0.5 cm) was cut out and placed in another Petri dish containing 1 ml of the same medium, and spermatozoa were allowed to swim out for 1 min to obtain a cloud of spermatozoa [46–50].
2.4. Assessment of Sperm Concentration and Motility. Five microliters of concentrated spermatozoa cloud was collected and placed on a Leja slide (Leja Products BV, Nieuw Vennepe, Netherlands). The Leja slide was placed onto a temperature-controlled stage of the Nikon E200 microscope (37°C). A 4x negative phase-contrast objective in conjunction with a phase-contrast condenser was used to determine sperm motility and concentration via the motility/concentration module of the Sperm Class Analyzer® version 5.4.0.1 software (Microptic SL, Barcelona, Spain) at 50 frames/s. Data were collected by capturing images with a digital camera (Basler, A78075gc, Germany). For motility analysis, eight fields were captured with the SCA system until 200 motile spermatozoa were analyzed, as recommended by WHO (1999) [46–48,51].

2.5. Assessment of Sperm Morphology. Fresh sperm smears were prepared for morphometric analysis by placing 5 μl of the fresh semen on the clear end of a frosted slide by dragging the drop across the slide. The smears were air-dried before staining. Three semen smears were prepared and stained with Spermblue® (Microptic Automatic Diagnostic System, Barcelona, Spain) according to Van der Horst and Maree (2009) [52]. Stained slides were used to perform morphology evaluation using the morphometry module of the Sperm Class Analyzer version 5.4.0.1 software (Microptic SL, Barcelona, Spain). The machine was equipped with a Nikon Eclipse model 50i (Nikon Corporation, Tokyo, Japan) microscope with a 60x bright-field objective and a video camera (Basler, A78075gc, Germany). A total of 200 sperms/animal were analyzed. The morphometric parameters of head and tail were determined, and abnormal sperms were detected based on previous criteria [46–48, 53–57]. Sperms with banana-shaped head, amorphous head, bent neck, or two-headed and headless sperms were classified as sperms with head abnormalities, whereas sperms with a bent or broken tail were classified as sperms with tail abnormalities (Figure 1).

2.6. Detection of Sperm DNA Damage by Using Comet Assay. Frosted microscope slides were covered with 1% normal melting point agarose in Ca²⁺- and Mg²⁺-free PBS. A sperm sample (10 μl) containing 1 x 10⁵ sperm/ml was suspended in 75 μl of 1% (w/v) low melting point agarose. This suspension (85 μl) was placed on the surface of a microscope slide (precoated with 1% normal melting point agarose) to form

Figure 1: Classification of sperm morphology in rats: (a) normal sperm, (b) banana-shaped head, (c) amorphous head, (d) bent neck, (e) bent neck, (f) headless, (g) tailless, (h) broken tail, and (i) banana-shaped head and bent tail (multiple abnormalities).
a microgel and allowed to set at 4°C for 5 min. Slides were
dipped in cell lysis buffer (2.5 M NaCl, 100 mM EDTA,
10 mM Tris-HCl, pH 10.0, containing 1% Triton X-100
added just before use and 40 mM dithiothreitol) for 24 h at
20 to 22°C. Following the initial lysis, proteinase K was added
to the lysis solution (0.5 mg/ml) and additional lysis was per-
formed at 37°C for 24 h. After cell lysis, all slides were washed
three times with deionized water at 10 min intervals to
remove the salt and detergent from the microgels. Slides were
placed in a horizontal electrophoresis unit and were allowed
equilibrate for 20 min with running buffer (500 mM NaCl,
100 mM Tris-HCl, 1 mM EDTA, pH 9) before electrophore-
sis (0.60 V/cm, 250 mA) for 30 min. After electrophoresis,
slides were then neutralized with 0.4 M Tris (pH 7.5), stained
with SYBR Green I (1 : 10,000) for 1 h and covered with cover
slides were then neutralized with 0.4 M Tris (pH 7.5), stained
slides were analyzed using Leica DM1000 fluorescence
microscope (Leica Microsystems, Wetzlar, Germany) and
Comet Assay IV Windows XP Pro software (Perceptive
Instruments, Suffolk, United Kingdom). At least 100 cells
were analyzed per sample [46–48, 58].

2.7. Histological Analysis of Testis Tissue. The right testis
tissues were sliced into small pieces (2 mm3) and then fixed
in paraformaldehyde (4%) in phosphate buffer pH 7.2 for 2 h
at 20 to 22°C. They were dehydrated in a graded series of alco-
hol. In order to improve infiltration, the samples were treated
with a mixture of LR White (Electro Microscopy Sciences, FT
Washington, PA) and 70% ethanol (2:1) (v:v) for 1 h at 20 to
22°C. The samples were then embedded in LR White and sec-
tioned at 700 nm (0.7 microns) thickness by using a Leica EM
U7 ultramicrotome. Semithin sections were stained with 1%
OsO4 for 1 h and covered with cover slips. Slides were analyzed using a Leica DM1000 fluorescence
microscope (Leica Microsystems, Wetzlar, Germany) and
Comet Assay IV Windows XP Pro software (Perceptive
Instruments, Suffolk, United Kingdom). At least 100 cells
were analyzed per sample [46–48, 58].

2.8. Determination of Serum FSH, LH, and Testosterone
Levels. After 30 min of drawing the blood to allow clotting,
blood samples from rats were centrifuged at 1000 g for
15 min at 4°C, and serum was separated. The hormonal
analyses were performed using the commercially available
kits according to the manufacturer's instructions.

2.9. Determination of GSH and MDA Levels in Testis Tissue.
The right testis was divided into equal parts and stored
at −20°C after freezing in liquid nitrogen. The GSH and
MDA levels in the testis were determined by using commer-
cially available kits according to the instructions of the
manufacturer.

2.10. Statistical Analysis. All data were expressed as mean ±
standard error. Statistical analyses of the groups were per-
fomed using the SigmaPlot v.10 package program (Systat
Software, USA). All values were verified to be normally
distributed. For the sperm comet assay, the Dunnnett T3 test
was performed as a post hoc test. In the other experiments,
one-way analysis of variance following the Tukey test as a post
hoc test was performed. p < 0.05 was considered statistically
significant.

3. Results

3.1. Effects of TRZ Treatment on Testis and Epididymis
Weights in Rats. When relative testis and epididymis weights
were compared among groups, relative testis and epididymis
weights obtained from the TRZ-administered groups were
indistinguishable from the control group. No significant dif-
fferences were observed among the TRZ-administered groups
in terms of relative testis and epididymis weights (Table 1).

3.2. Effects of TRZ Treatment on Sperm Concentration,
Motility, and Morphology in Rats. When the groups were
compared in terms of sperm concentration, significant and
dose-related decreases in sperm concentration were observed
in all TRZ-administered groups compared to the control
group. No significant differences were observed among the
TRZ-administered groups (Table 2).

Sperm motility percentages of TRZ-administered groups
were decreased significantly and dose-dependently when
compared to control group. No significant differences were
observed among the TRZ-administered groups. (Table 2).

A significant increase in the percentage of the sperm
abnormalities was observed in the TRZ-administered groups
at all concentrations compared to control. Among the TRZ-
administered groups, the percentages of the sperm abnor-
malities did not show any significant differences (Table 2).

These abnormalities were found to be more in the tail, as
represented by bent and broken tail abnormalities (Figure 1),
in the 5, 10, and 20 mg/kg TRZ-administered groups at
64.50%, 70.92%, and 78.30%, respectively. The percentages
of sperm head abnormalities, including banana-shaped head,
amorphous head, bent neck, and two-headed and headless sperms (Figure 1), were determined to be 33.29%, 27.83%, and 20.34% in the 5, 10, and 20 mg/kg TRZ-administered groups, respectively. The percentages of multiple abnormalities were 2.21%, 1.25%, and 1.36% in the 5, 10, and 20 mg/kg TRZ-administered groups, respectively.

3.3. Effects of TRZ Treatment on Sperm DNA in Rats. Results of the comet assay are expressed as extent tail moment, a product of the tail length and the tail DNA% (extent tail moment = tail length x tail DNA% /100) (Lee et al., 2004). Values were as the following: 1.65 ± 0.17, 1.91 ± 0.18, 5.55 ± 0.41, and 10.66 ± 0.92 (mean ± SE) for control, 5 mg/kg, 10 mg/kg, and 20 mg/kg TRZ-administered groups, respectively. Control group and 5 mg/kg TRZ-administered group did not show any significant damage. Exposure to 10 and 20 mg/kg TRZ increased the tail moment over 3-fold and 6-fold, respectively, when compared to control (Figure 2).

3.4. Effects of TRZ Treatment on the Testicular Histology in Rats. Histological alterations as a result of TRZ administration at different doses are shown in Figures 3 and 4. Control group animals manifested regular feature of the seminiferous tubules, germinal epithelial cells, and interstitial cells (Figures 3 and 4).

5 mg/kg TRZ administration resulted in mild central degeneration of the tubules. Cells at the center of the tubules were separated from each other and some of them accumulated in the lumen. Thickening of basement membrane and vacuolation in Sertoli cells were also observed (Figures 3 and 4).

In 10 mg/kg TRZ-administered group, many of the germ cells abnormally accumulated in the lumen. Large vacuoles were observed in the germinal epithelium. Deformed Sertoli cells were partially detached from the basement membrane. Leydig cells showed intense vacuolation and deformation (Figures 3 and 4).

20 mg/kg TRZ administration resulted in total degeneration of the seminiferous tubules. Cells of adluminal compartment disintegrated and desquamated into the lumen. Architecture of germinal epithelium was disorganized. Necrosis was observed in Sertoli cells and neighboring germ cells. Degenerating cells were showing nuclear pyknosis. Lysis of the cytoplasm of Leydig cells was also observed (Figures 3 and 4). Johnsen’s scores and semiquantitative comparison of pathology at the cellular level are shown in Table 3.

3.5. Effects of TRZ Treatment on the Serum Hormone Levels in Rats. When the groups were compared in terms of serum FSH levels, statistically significant dose-related increases were observed in the TRZ-administered groups compared to the control group. Compared to the control group, a statistically significant increase was found in the serum LH levels in the 10 and 20 mg/kg TRZ-administered groups. When the groups were compared in terms of serum testosterone levels, statistically significant increases were found in the TRZ-administered groups. No significant differences were observed among the TRZ-administered groups in terms of serum FSH, LH, and testosterone levels (Table 4).

3.6. Effects of TRZ Treatment on GSH and MDA Levels in Testis Tissue of Rats. GSH levels of testis tissues did not show any significant difference in the TRZ-administered groups in comparison to the control group. No significant differences were observed among the TRZ-administered groups in terms of GSH levels in testis tissues. When the groups were compared in terms of the MDA levels of testis tissue, dose-related increases were observed in the TRZ-administered groups compared to the control group (Table 5).

4. Discussion

According to our study results, which we performed independently of other risk factors related to reproductive toxicity, TRZ administration decreased sperm concentration, motility, and normal sperm morphology, increased sperm DNA damage, and induced degeneration of testicular structure. Detected reproductive toxicity findings were accompanied by increases of serum FSH, LH, and testosterone levels and oxidative stress in the testicular tissue.

The reproductive function of men is evaluated via semen analysis, by assessing sperm concentration, motility, and morphology. These parameters provide information about sperm quality [61, 62]. Spermatozoa must be produced in sufficient numbers and exhibit normal motility and shape for normal sperm function [61]. It was noteworthy that sperm concentration, motility, and normal sperm morphology decreased in TRZ-administered groups dose-dependently. As known, pregnancy rates by intercourse and intrauterine insemination decline as sperm density decreases. The efficient passage of spermatozoa through cervical mucus is dependent on rapid progressive motility. Persistent poor motility is a good predictor of failure in fertilization [63]. Sperm morphology measurement still has a very important role in the clinical evaluation of sperm fertilization capacity [62]. In general, pregnancy is possible with low morphology scores and that both motility and morphology have also demonstrated prognostic value [63]. Otherwise, it is emphasized that tail
anomalies are positively correlated with infertility and are an important parameter affecting motility [55, 64, 65]. When tail and head anomalies were evaluated in groups, tail anomalies were found to be higher than head anomalies in TRZ groups. These increased tail anomalies in TRZ groups may also reflect decreased sperm motility.

Recently, there has been a focus on the analysis of sperm DNA damage, as an indicator of sperm quality and fertility. The most common types of identified sperm DNA damage are single and double DNA strand breaks, the chemical modification of a base, inter- or intrastrand cross-linkage, and DNA-protein cross-links [62, 66]. Increased sperm DNA damage can cause a lower fertilization potential, a lower blastocyst formation rate, a lower implantation rate, and adverse effects on embryo development [67–69]. The limited repair capacity of sperm DNA makes it more susceptible to damage [70, 71]. The Comet method, which is often used as a reliable method for determining sperm DNA damage, is especially precise when determining DNA errors associated with double-strand breaks [72–75]. The tail moment calculated in this method is a parameter that is frequently used in terms of the comparability of DNA damage among groups. When the tail moment was compared between the groups, it was notable that DNA damage increased in TRZ-treated groups dose-dependently. Additionally, there are positive correlations between abnormal sperm morphology and sperm DNA damage, but not DNA fragmentation [3, 72]. At this point, previous findings supported our results, which showed that induced sperm abnormalities were accompanied by sperm DNA damage in TRZ-administered groups.

Histopathological analysis is frequently used as an important biomarker in toxicity research [76]. Reduced sperm quality and quantity can also be accompanied by histopathological changes in the testes [77]. The first pathological findings observed were slight vacuolization in seminiferous tubules and Leydig cell deformation in the low-dose TRZ group. While basal lamina irregularity, vacuolar enlargement, and Leydig cell deformation were increased in 10 mg/kg TRZ-administered group, testicular degenerative findings

![DNA damage in rat sperms in control and experimental group animals exposed to physiological saline (0.9%) or different doses of trazodone. (a) Sperm comet assay photo of control rats. (b) Sperm comet assay photo of 5 mg/kg trazodone-administered rats. (c) Sperm comet assay photo of 10 mg/kg trazodone-administered rats. (d) Sperm comet assay photo of 20 mg/kg trazodone-administered rats. (e) Tail moment graph: *Different from C (p < 0.05); †Different from TRZ5 (p < 0.05); ‡Different from TRZ10.](image)
progressing to lysis and necrosis in the cells were notable in high-dose TRZ group.

The spermatogenesis process is regulated by endocrine activity of the hypothalamus-pituitary-testicular axis. FSH and LH are released from the anterior pituitary to maintain spermatogenesis. While LH mediates release of testosterone from Leydig cells, FSH mediates the release of androgen-binding protein from Sertoli cells which is required for sperm maturation [78, 79]. In our study, serum FSH, LH, and testosterone levels were increased in TRZ-administered groups dose-dependently. At this point, it could be stated that secondary to increased levels of LH, testosterone plasma levels increased in TRZ-administered groups. On the other hand, previous studies investigating the effects of serum hormone levels on sperm parameters demonstrated that FSH and LH levels showed negative correlation with sperm concentration, motility, and morphology [80–82], testosterone level did not affect sperm parameters or only positively correlated with motility [81–84]. In our study, significant reductions in sperm quality with TRZ administration were also correlated with increases in serum FSH and LH levels. Additionally, the degenerative findings identified in the testis tissues may also be associated with the increased serum LH levels as well as the oxidative stress induced by TRZ. Some studies have shown that increased LH levels cause degeneration of the germinal cells, which negatively affect spermatogenesis [85].

Oxidative stress, which occurs as a result of oxidant/antioxidant imbalance in favor of oxidants, can cause deteriorations in testicular structure and spermatogenesis process and, correspondingly, infertility [86–88]. Testis tissue is vulnerable and also highly dependent on oxygen to drive spermatogenesis process and highly susceptible to the toxic effects of reactive oxygen metabolites; in this context, the testis is very similar to the brain [86]. Furthermore, cell membranes of spermatozoa are rich in polyunsaturated fatty acids which makes them more susceptible to oxidative damage [89]. Animal models demonstrated a causal relationship between the induction of oxidative stress in the testes and the impairment of male reproductive function [86]. Oxidative stress is defined as either an excessive production of reactive oxygen species/reactive nitrogen species and/or a deficiency of enzymatic and nonenzymatic antioxidants in the biological system. We evaluated the oxidative status by measuring the level of MDA which is the end product of lipid peroxidation and the level of GSH which is an important nonenzymatic antioxidant in testis tissue [90]. In our study, MDA levels increased significantly and dose-dependently, but no significant difference was found in terms of GSH levels in TRZ-administered rats. According to these findings, it could be asserted that TRZ induced oxidative stress in the testicular tissue. A wide variety of different xenobiotics have also been shown to induce oxidative stress in the testes [86]. Toxicity studies indicated TRZ-induced hepatotoxicity via oxidative stress in isolated hepatocyte. Also, these studies emphasized that the bioactive intermediate metabolites of TRZ might cause hepatotoxicity [32, 33, 91]. These metabolites are the reactive quinoneimine and epoxide species formed by the
cytochrome P4503A4 (32). Excessive extrinsic reactive metabolites cause plasma membrane damage and negatively affect sperm parameters such as sperm concentration, motility, and morphology [62, 71, 89, 92]. Otherwise, excessive intrinsic reactive metabolites cause DNA damage and induce sperm DNA strand breaks [62, 66, 71]. Additionally, ongoing oxidative stress in testicular tissue is known to cause histopathological changes [87, 93, 94]. Oxidative stress can induce testicular atrophy and degeneration of seminiferous tubules by disrupting membrane integrity [95]. In our study, decreased sperm quality, increased sperm DNA damage, and testicular degenerative...
findings could be the result of TRZ-induced testicular oxidative stress.

In conclusion, our study outlines the reproductive toxicity of TRZ, commonly used to treat depression and insomnia, in male rats with respect to certain reproductive parameters. TRZ-induced reproductive toxic effects may be a consequence of increased serotonin associated with this agent and/or direct toxic effects of the agent/metabolites in the reproductive system. We emphasize that clinical researches are very important in patients under TRZ treatment. Determination of sperm parameters in patients before, during, and after TRZ treatment will contribute to the identification of its reproductive toxicity in males.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

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