Dose-Dependent Effect of Polystyrene Microplastics on the Testicular Tissues of the Male Sprague Dawley Rats

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Abstract
Due to the continuous increase in polystyrene microplastics (PS MPs) incorporation in the environment, growing number of adverse effects on living organisms and ecosystem have become a global concern. Therefore, current study was planned to elucidate the impacts of 5 different concentrations control, 2, 20, 200, and 2000 µgL⁻¹ of PS MPs on testicular tissues of rats. PS MPs significantly reduced the activities of antioxidant enzymes (catalase, superoxide dismutase and peroxidase) as well as total protein contents, while elevated the level of lipid peroxidation and reactive oxygen species. Moreover, expressions of steroidogenic enzymes (3β-hydroxysteroid dehydrogenase, 17β-hydroxysteroid dehydrogenase and steroidogenic acute regulatory protein) as well as the levels of follicle-stimulating hormone (FSH), luteinizing hormone (LH) in plasma, intra-testicular testosterone and plasma testosterone were reduced and a significant (P < 0.05) reduction was noticed in the sperm count, motility and viability. Furthermore, PS MPs significantly up-regulated the expressions of Bax and caspase-3, while down-regulated the Bcl-2 expression. The histomorphological assessment revealed significant damages in the testicles as well as decrease in the number of germ cells (spermatogenic, spermatocytes and spermatids). Collectively, PS MPs generated oxidative stress (OS) and caused potential damage to the testicles of rats in a dose-dependent manner.

Keywords
polystyrene microplastics (PS MPS), oxidative stress, testicular damage, spermatogenesis, male reproductive system

Introduction
Environmental pollution and industrial exposure to various contaminants play a significant role in the induction of male infertility.¹ Global plastic production and its utilization in making various products were increased from 1.7 to 335 million tons during 1950-2016.² Plastic manufacturing is relatively convenient and inexpensive.³ Microplastics (MPs) are the emerging environmental pollutants consisting of plastic particles (<5 mm), which are small fragments of macroplastics.⁴ MP pollution also results in other universal threats, such as ocean acidification, climate change and ozone depletion.⁵

Microplastics are formed of various materials, such as polyvinyl chloride (PVC), polyethylene (PE) and polystyrene (PS). PS is one of the most abundantly used aromatic polymer amongst these plastics.⁶ A reason behind its excessive presence in the ecosystem is its low cost and exceptional physical properties. It is used in the manufacturing of disposable cups, trays, bowls, plates, pegs, toys, office supplies, paper clips, cleansing agents, cosmetics, pharmaceuticals.⁶ Moreover, it is reported to induce a number of damages in living organisms. At the biochemical and cellular level, PS exposure leads to stimulation of inflammatory reactions and oxidative stress (OS) in crab liver.⁷

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Received 21 January 2021; received revised 30 April 2021; accepted 3 May 2021

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In mice, exposure of PS induces the OS, as well as reduces weight of liver and body. PS derived MP is a serious environmental hazard, which is known as a white pollutant of the environment. Few studies have been conducted on the impacts of MPs on male reproductive system of the mice but the effect of MPs and their possible damage inducing mechanisms in the testicular tissues of mammalian species demand a further investigation. It is an established fact that steroidogenic enzymes play an important role in maintaining the normal physiological functions of testis. Thus, by keeping this fact under consideration, the current study was designed to elucidate the dose-dependent effects of PS MPs on rat’s testicular toxicity especially by considering the steroidogenic enzymes.

Materials and Methods

Chemical Reagents

PS MP particles of 10μm diameter were bought from Sigma Aldrich, USA. Other chemicals were of high analytical grade and bought from Merck and Sigma Aldrich, USA.

Animals

Sexually mature male Sprague Dawley rats (n = 60) were collected from breeding and rearing section in the Animal House of the University of Agriculture, Faisalabad. The animals were housed in steel cages at standard temperature (24 ± 2°C) and light conditions (10 h light and 14 h dark). Food pellets (comprising free soy, alfalfa, 40%-50% carbohydrates, 20%-25% protein and 4%-7% adipose tissue) were fed to the rats. Tap water ad libitum was provided in bottles made up of polysulfone. Animals were treated in compliance with the European Union of Animal Care and Experimentation (CEE Council 86/609) approved protocol.

Experimental Design

Sexually mature male Sprague Dawley rats (n = 60) were distributed into 5 groups (n = 12/group). Various concentrations Control (0.9% saline), 2, 20, 200, and 2000 μg/L of PS MPs were administered to group 1 to 5 respectively. The animals of the control group were treated with normal saline as a vehicle. Various concentrations of PS MPs were dissolved in culture media and given to rats by oral gavage for 60 days. The dose selection of PS MPs was based on OECD protocol 423 for various complexes with anonyymous effect (OECD 2001). On the 61st day, rats were anesthetized with diethyl ether and killed by decapitating. To separate plasma, trunk blood was diluted using 2 mL of phosphate buffer with 7 pH. Diluted homogenate (2 mL) was added with phosphate buffer (1 mL) of pH 7 comprising 30 mM of H2O2 in the test tube and then, diluted water was added to blanks. After instantaneously mixing, absorbance was noted at 240 nm. CAT activity was expressed as an absorbance alteration of 0.01 U/min.

Biochemical Assay

The homogenization of testicular tissues was carried out in PBS. Finally, centrifugation was performed at 3,000 rpm for 10 minutes. The supernatant obtained was used for the biochemical assay, protein evaluation, lipid profile assessment, and hormonal analysis. Biochemical assessment of tissues was performed via supernatant obtained from testicular tissues.

Analysis of catalase (CAT). The activity of catalase was determined by following the technique of Aebi. 50μL of tissues homogenate was diluted using 2 mL of phosphate buffer with pH 7. Diluted homogenate (2 mL) was added with phosphate buffer (1 mL) of pH 7 comprising 30 mM of H2O2 in the test tube and then, diluted water was added to blanks. After instantaneously mixing, absorbance was noted at 240 nm. CAT activity was expressed as an absorbance alteration of 0.01 U/min.

Analysis of superoxide dismutase (SOD). Superoxide dismutase activity was measured by following technique of Kakkar et al. Reaction solution contained of 1.2 mL of sodium pyrophosphate buffer (0.052 mM; pH 7.0) and 0.1 mL of phenazine methosulphate (186 μM). 0.3 mL of supernatant after centrifugation (1500 × g for 10 min followed by 10000 × g for 15 min) of homogenate was added to the reaction mixture. Then, 0.2 mL of NADH (780 μM) was added to start the enzyme reaction, which was later on ended by adding 1 mL of glacial acetic acid. Finally, chromogen was assessed by noticing the change in color intensity (at 560 nm). The values of SOD activity were presented as unit/mg protein.

Analysis of peroxidase (POD). In homogenate, the activity of peroxidase was evaluated by following the technique of Chance and Maehly, but with few amendments. First of all, homogenate mixing was carried out with 0.1 mL of guaiacol, 2.5 mL of phosphate buffer (pH 5) and 0.3 mL of hydrogen peroxide. At 470 nm, the variations were detected after 1 min. 1 unit of POD activity was noted at 0.01 absorbance change as U/min.

Analysis of reactive oxygen species (ROS). ROS were assessed from homogenate as per the process explained by Hayashi et al. Homogenate (5μL) and 0.1 M sodium acetate buffer (140μL) with pH 4.8 were mixed and dispensed in 96 well-plate. After incubating at 37°C for 5 minutes, 100μL of mixed solution of ferrous sulfate and N, N-diethyl-para-phenylenediamine was dispensed to each plate, and then incubated at 37°C for 1 minute. At 505 nm, the absorbance was noticed with the help of a microplate reader for 180 s with a 15 s interval. Finally, the standard curve was plotted. ROS was recorded as Unit/mg tissues and 1 unit of ROS was equivalent to 1.0 mg/L of H2O2 in the sample.

Analysis of lipid peroxidation (LPO) by thiobarbituric acid reactive substances (TBARS). The analysis of malondialdehyde in the homogenate was performed by reacting it with thiobarbituric acid, as per the procedure described by Iqbal et al. TBARS are produced as a result of lipid peroxidation (LP), therefore,
degree of LP was indicated by TBARS level. 1.0 mL total volume of reaction mixture consisted of 0.2 mL homogenate sample, 0.02 mL ferric chloride (100 mM), 0.58 mL phosphate buffer (0.1 M pH 7.4) and 0.2 mL ascorbic acid (100 mM). Incubation was carried out at 37°C for about 1 hour in a shaking water bath. Addition of 1.0 mL 10% trichloroacetic acid terminated the reaction. After adding 1.0 mL 0.67% thiobarbituric acid, tubes were boiled in water bath for about 20 minutes. Then, mixture was transferred to crushed ice-bath prior centrifuging at 2500 g for about 10 minutes. The TBARS level was determined by measurement of optical density of supernatant at 535 nm with spectrophotometer against reagent blank. Its final values were shown as nM TBARS/min/mg tissue at 37°C using molar extinction coefficient of 1.56 x 10⁵ M⁻¹ cm⁻¹.

**Analysis of Total Protein Content**

Estimation of total protein constituents was carried out by the protein kit (Cat No. BR5202-S, AMEDA Labordiagnostik GmbH, Krenngasse, Graz, Austria). Results were computed by plotting absorbance of standard vs sample absorbance on the graph. Final data was shown in mg/g of tissues.

**Real-time Polymerase Chain Reaction (qRT-PCR)**

qRT-PCR was used to assess expression levels of 3β-hydroxysteroid dehydrogenase (3β-HSD), 17β-hydroxysteroid dehydrogenase (17β-HSD), steroidogenic acute regulatory protein (StAR), Bax, Bcl-2 and caspase-3. TRIzol reagent (Invitrogen, Carlsbad, CA) was used for isolation of total RNA (Takara Bio, Japan). RNA concentrations were determined by NanoDrop 2000c spectrophotometer (Thermo Fisher, USA). Total RNA was with 260/280 ratio between 1.8-2.2 was used for reverse transcriptase PCR. Reverse transcription kit (Promega, USA) was employed to transform total RNA into complementary DNA. qRT-PCR was carried out using SYBR@ Premix Ex TaqTMII kit. β-actin was used as internal control and relative expressions were assessed by 2⁻ΔΔCT. Primer sequences of β-actin and target genes are shown in Table 1 as reported previously in Ijaz et al.17

**Analysis of Hormones**

The levels of LH (serial number-H206), FSH (serial number-H101), and plasma testosterone (serial number-H090) were assessed by ELISA kits (Los Angeles, CA USA) according to manufacturer’s procedures. 50 μL of assay diluent and 10 mL of plasma were added to 96-well ELISA plate and incubated for approximately 2 hours at room temperature. Then, plates were rinsed with the deionized water and before adding 100 mL of peroxidase-conjugated immunoglobulin G (IgG) anti-FSH solution, anti-LH, or anti-testosterone in each well, incubation was carried out for maximum 2 hours. Plates were again rinsed with the deionized water, substrate solution was added in wells and incubated for about 25 minutes at room temperature. 50 μL of stop solution was added into each well to terminate the reaction. Finally, the absorbances of FSH, LH and plasma testosterone were recorded at 450 nm. All samples were run in triplicates and conducted at same time under same conditions to avoid inter-assay variation.

**Spermatogenic Analysis**

Epididymal spermatozoa were collected by separating the caudal part of the epididymis on both sides. Spermatozoa were separated from epididymal tubules by splitting the caudal part of the epididymis in 5 mL of Hams F10 solution. The solution was incubated for 5 min at 37°C. After pipetting, 1 drop of sperm suspension was placed on a microscope slide and cover slipped. We observed at least 5 microscopic fields at 400X magnification and calculated the % of in situ motile, progressive and immotile spermatozoa as a fraction of the total counted spermatozoa according to the WHO recommendations. Sperm count. Epididymal sperm count was evaluated with a hemocytometer by following the method described by Ciftci et al18 but with few amendments. The right epididymal portion was crushed by the help of anatomical scissors in 5 mL saline. After incubation at room temperature, sperms present in the supernatant fluid were counted under a light microscope.

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**Table 1. Primers Sequences for RT-qPCR.**

| Gene       | Primers 5' -> 3' Accession number | Product size | Temperature |
|------------|-----------------------------------|--------------|-------------|
| 3β-HSD     | Forward: GCATCCTGAAAAATGGTG GCCG Reverse: GCCACATTGCCCTACATACAC | NM_001007719 | 135         | 57          |
| 17β-HSD    | Forward: CAGGGTTCAAGGCTTTGTG GTG Reverse: CAGGGTTTCAGCTCAATCGT | NM_054007 | 161     | 59          |
| StAR       | Forward: AAAAGGCTTGGGCACTACTC Reverse: CATAGGAGTTGCTGTTGAGGCG | NM_031558 | 113        | 58          |
| Bax        | Forward: GGGCTTTTGTCAAGGTTT GTT Reverse: AGTCTCAGTTTGTGAGGCG | NM_017059.2 | 119       | 58          |
| Bcl-2      | Forward: ACAAACATCGCTCTGTGAT Reverse: TCAGAGACAGCCAGGAGAA | NM_016993.1 | 103       | 57          |
| Caspase-3  | Forward: ATCCATGGAAGCAGTCGAT Reverse: CTTTTTGTGATGATGCTCCTT | NM_012922.2 | 233       | 57          |
| β-actin    | Forward: TACAGCCTTCCACACACG Reverse: GGAAACCGCTCATACGCCGATA | NM_031144 | 135       | 58          |

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Table 2. Effect of Different Concentrations (Control, 2, 20, 200, and 2000 µgL⁻¹) of PS MPs on Body Weight, Testes, Epididymis, Seminal Vesicles, and Prostate Gland Weight in Control and Treated Groups.*

| Groups       | Body weight gain (g) | Left testes weight (g) | Right testes weight (g) | Epididymis (g) | Seminal vesicles (g) | Prostate gland (g) |
|--------------|----------------------|------------------------|-------------------------|----------------|----------------------|-------------------|
| Control      | 55.4 ± 1.08          | 1.35 ± 0.03            | 1.40 ± 0.03             | 0.67 ± 0.02    | 0.76 ± 0.02          | 0.63 ± 0.01       |
| 2 µgL⁻¹      | 51.8 ± 0.89          | 1.33 ± 0.02            | 1.38 ± 0.02             | 0.64 ± 0.01    | 0.73 ± 0.01          | 0.63 ± 0.02       |
| 20 µgL⁻¹     | 50.4 ± 0.98          | 1.32 ± 0.03            | 1.38 ± 0.01             | 0.62 ± 0.01    | 0.74 ± 0.04          | 0.63 ± 0.01       |
| 200 µgL⁻¹    | 50.3 ± 1.46          | 1.31 ± 0.04            | 1.34 ± 0.02             | 0.64 ± 0.03    | 0.67 ± 0.07          | 0.60 ± 0.02       |
| 2000 µgL⁻¹   | 50.5 ± 1.80          | 1.30 ± 0.04            | 1.35 ± 0.03             | 0.64 ± 0.03    | 0.70 ± 0.07          | 0.62 ± 0.04       |

* Values are shown as Mean ± SEM (n = 12/group). Means within same row (for each parameter) carrying different superscripts are significantly different at P < 0.05.

Sperm motility. For evaluation of the percentage of sperm motility, a slide was kept under light microscope equipped with a prewarned (37°C) stage. A drop of semen was placed on slide and motility was estimated from 3 random fields per sample. Finally, a mean of 3 estimated values was noted as sperm motility.¹⁹

Sperm viability. For determination of sperm viability, eosin/nigrosin stain was dropped on a semen sample on a prewarned slide and smear was formed. After it dried, the slide was examined under a light microscope. Unstained/white sperms were considered as alive, whereas red sperms were considered dead. 300 spermatozoa were examined, and the percentage of dead sperm was estimated.²⁰

Tissue Histology

Tissues were preserved in a solution of formalin (10%) for 48 h for fixation. In the next step, tissues were shifted to increasing alcohol grades for dehydration, followed by double washing with xylene. For the formation of blocks, tissues were shifted into paraffin wax for microtomy. 7 µm thick sections of tissues were obtained. Each of the 10th slices of tissues were cut down from the ribbon and transferred to warm water for stretching. Slices were shifted carefully to alburnum slides, which were air-dried for half an hour and placed overnight in paraffin oven at 38 ± 2°C for complete deparaffinization. These deparaffinized slices were then passed through different grades of alcohol for the dehydration and later on, stained with hematoxylin in eosin.

Lastly, these slides were examined under Leica Microscope (DMLB Leica microscope, Leica Microsystems Ltd, UK), and a digital camera (Canon, Japan) was installed with it. After capturing images at 20× and 40×, the histomorphometric assessment was carried out by ImageJ Software. From 20× images, about 30 photographs per rat were chosen and regions of seminiferous tubules, epididymal tubule and interstitial spaces were assessed through the free selection tool of software. Following formula was used to measure area: % As = As × 100/T. Where As known as the seminiferous tubules covered area and T is the total area of the field.

Mean and average area percentages were observed to compare control and treated groups. Count of different germ cell numbers was taken from 50 seminiferous tubules per rat at 100× and the average number of spermatocytes, spermatogonia and spermatid in each seminiferous tubule was computed.

Statistical Analysis

Final data was shown as Mean ± SEM. After applying 1-way analysis of variance (ANOVA), Tukey test was employed using Minitab software. The significance level was P < 0.05.

Results

Effect of PS MPs on Testicular, Epididymis, Seminal Vesicles, and Prostate Gland Weight

There was no significant change in animal weight gain among all groups. PS MPs treatment did not alter the weights of the both testis (Right and Left), epididymis, seminal vesicles, and prostate gland (Table 2).

Effect of PS MPs on Antioxidant Capacity and Total Protein Content

After PS MPs administration, CAT, SOD and POD activity was significantly (P < 0.05) reduced only at higher doses (200 and 2000 µgL⁻¹) in PS MPs-intoxicated rats compared with control rats (Table 3).

On the other hand, ROS level was significantly (P < 0.05) increased only in higher doses of PS MPs treated groups compared to control. LPO also showed the similar trend, significant (P < 0.05) elevation in LPO was noticed in groups treated with higher doses (200 and 2000 µgL⁻¹) in PS MPs treated rats compared with control rats. Similarly, protein contents were significantly (P < 0.05) decreased at higher doses (200 and 2000 µgL⁻¹) in PS MPs groups in comparison to control (Table 3).

Effect of PS MPs on Expression of Steroidogenic Enzymes

As shown in Figure 1, PS MPs led to a significant down-regulation in expression of 3β-HSD and 17β-HSD at 2 higher-doses (200 and 2000 µgL⁻¹) in PS MPs-intoxicated groups compared to control. While, a significant reduction was seen in expression of StAR at 3 higher-doses (20, 200 and 2000 µgL⁻¹) in PS MPs-intoxicated rats compared with control rats.
Effect of PS MPs on Hormonal Concentrations

Oral exposure to different doses of PS MPs significantly ($P < 0.05$) suppressed the concentration of plasma LH in all PS MPs-induced groups when compared to control (Table 4). A significant ($P < 0.05$) suppression was observed in plasma FSH concentration, but only at the highest dose ($2000 \mu gL^{-1}$) in PS MPs groups compared to control. In the same manner, the concentration of plasma testosterone was significantly ($P < 0.05$) suppressed by PS MPs only at highest dose ($2000 \mu gL^{-1}$) compared with control. A significant ($P < 0.05$) reduction was noticed in intra-testicular testosterone concentration at higher dose ($200$ and $2000 \mu gL^{-1}$) PS MPs-intoxicated rats compared to control rats (Table 4).

Effect of PS MPs on Sperm Indices

The exposure of PS MPs led to a significant ($P < 0.05$) reduction in quantity of sperms (Table 4). The PS MPs significantly ($P < 0.05$) reduced the sperm count and viability at higher doses ($200$ and $2000 \mu gL^{-1}$) in PS MPs groups compared to control. Whereas, sperm motility was significantly ($P < 0.05$) reduced at highest ($2000 \mu gL^{-1}$) PS MPs treated group compared to control (Table 4).
Table 4. Effect of Different Concentrations (Control, 2, 20, 200, and 2000 μgL−1) of PS MPs on Hormones Luteinizing Hormone (LH), Follicle-Stimulating Hormone (FSH), Testosterone and Sperm Parameters in Rat Testicles.*

| Parameters                              | Control  | 2 μgL−1 | 20 μgL−1 | 200 μgL−1 | 2000 μgL−1 |
|-----------------------------------------|----------|---------|----------|-----------|------------|
| LH (ng/mL)                              | 2.91 ± 0.02a | 2.70 ± 0.01b | 2.64 ± 0.02b | 2.41 ± 0.01c | 2.18 ± 0.04d |
| FSH (ng/mL)                             | 3.75 ± 0.04a | 3.72 ± 0.02a | 3.70 ± 0.01a | 3.68 ± 0.02a | 3.37 ± 0.04b |
| Plasma testosterone (ng/mL)             | 4.74 ± 0.15a | 4.77 ± 0.13a | 4.67 ± 0.06a | 4.54 ± 0.15a | 4.17 ± 0.12b |
| Intra-testicular testosterone (ng/g tissue) | 45.76 ± 1.94a | 47.14 ± 1.04a | 44.19 ± 1.24ab | 37.52 ± 1.12bc | 36.10 ± 1.57c |
| Sperm count (x 106/gm of cauda)         | 163.15 ± 2.55a | 163.28 ± 1.88a | 163.11 ± 1.59a | 139.92 ± 3.03b | 104.19 ± 1.79c |
| Sperm viability %                        | 78.78 ± 1.50a | 78.75 ± 0.82a | 78.81 ± 0.87a | 63.51 ± 0.41b | 58.45 ± 0.88c |
| Sperm motility %                         | 69.93 ± 3.81a | 69.87 ± 2.84a | 69.96 ± 1.11a | 68.75 ± 0.76a | 55.86 ± 0.81b |

* Values are shown as Mean ± SEM (n = 12/group). Means within same row (for each parameter) carrying different superscripts are significantly different at P < 0.05.

**Effect of PS MPs on Apoptotic Markers**

As shown in Figure 2, PS MPs significantly increased the gene expression of Bax at all doses (2, 20, 200 and 2000 μgL−1) of PS MPs, while expression of caspase-3 was increased at 3 high-doses (20, 200 and 2000 μgL−1) in PS MPs groups compared to control. On the contrary, the expression of Bcl-2 showed a significant reduction at all concentrations (2, 20, 200 and 2000 μgL−1) in PS MPs-induced group compared to control (Figure 2).

**Effect of PS MPs on Histopathology**

In the morphometric analysis of testis, 2 higher-dose groups of PS MPs (200 and 2000 μgL−1) exhibited a significant (P < 0.05)
decrease in area and diameter of seminiferous tubules and epithelial height when compared with control rats (Table 5). While, the area of interstitium was significantly ($P < 0.05$) decreased by PS MPs in 3 groups (20, 200 and 2000 $\mu$gL$^{-1}$) compared to control (Table 5).

A significant ($P < 0.05$) decrease was observed in number of spermatogonia and spermatocytes in seminiferous tubules by PS MPs in 2 higher-dose groups (200 and 2000 $\mu$gL$^{-1}$) compared to control (Table 5). A similar trend was shown by spermatids. Spermatid number in seminiferous tubules displayed a significant ($P < 0.05$) decrease by PS MPs administration in 3 higher-dose groups (20, 200 and 2000 $\mu$gL$^{-1}$) in comparison to control (Table 5). Histopathological changes have been shown in Figure 3.

**Discussion**

In the past 2 decades, MPs have received increased attention due to their wide distribution, persistence and toxic impacts on living beings and ecosystems.\(^{21}\) The toxic effects of PS MPs are investigated in various animal models due to a possible threat to marine biota as well as human health.\(^{22}\) A recent study reported that PS MPs exposure led to a reproductive disturbance in oysters and specifically affected their larval stages.\(^{23}\) Lönstedt and Eklov reported that PS derived MPs are ingested by a variety of living beings, which disturb multiple physiological events in the body.\(^{24}\) Therefore, the current research was planned to evaluate PS MPs-induced reproductive toxicity in male rats by evaluating the biochemical status, hormonal concentrations, spermatogenic indices, morphometry and histopathological changes were determined.

PS MPs decreased the activities of antioxidant enzymes (CAT, SOD and POD) and total proteins, while elevated the level of ROS and LPO in PS MPs-intoxicated rats compared to control rats. Antioxidant enzymes are the first line of a defense that protects the biological molecules (DNA, lipids, and proteins) from impairment by reducing ROS production.\(^{25,26}\) CAT, POD and SOD form the antioxidant defense system in semen.\(^{27}\) CAT is considered as the central enzyme of the antioxidant system as it plays a significant role in $H_2O_2$ catabolism.\(^{28}\) SOD catalyzes superoxide radicals to $H_2O_2$ and $O_2$.\(^{29}\) Normally, antioxidant enzymes nullify the toxicities produced due to ROS. However, when ROS are generated excessively, they overtake the antioxidant defense system of the body, which leads to the generation of OS.\(^{30}\) The previous study by Xie et al\(^{10}\) has reported oxidative stress by using 5$\mu$m size microplastic in rat testes. Therefore, the current study endorsed the findings of previous study that PS MPs caused toxicity by altering the activities of antioxidant enzymes and generating higher levels of ROS and LPO in testicular tissues of rats, but in the current study these effects were pronounced in the rats treated with only higher doses of 10$\mu$m size PS MPs.

In the current investigation, the concentrations of luteinizing hormone (LH), follicle-stimulating hormone (FSH), plasma testosterone as well as intra-testicular testosterone were significantly ($P < 0.05$) decreased at high doses in PS MPs-intoxicated animals as compared to control. Previously Xie et al\(^{10}\) and Jin et al\(^{11}\) also reported the decrease in testosterone level followed by PS MPs exposure, but the current investigation expanded the horizon and studied the complete feedback loop of hypothalamo-pituitary-gonadal axis to reveal the effect of PS MPs exposure on FSH, LH and subsequently on testosterone level. According to Wisniewski et al spermatogenesis is dependent on ratio of FSH, LH and testosterone.\(^{31}\) Synthesis of testosterone is a vital indicator of male reproductive health since the testosterone is involved in healthy sperm development and regulation of spermatogenesis.\(^{32}\) The pituitary gonadotropic cells were stimulated by gonadotropin-releasing hormones (GnRH) in the hypothalamus, which releases LH and FSH to regulate spermatogenesis in animals.\(^{33}\) LH acts on Leydig cells to produce testosterone, while FSH helps in proliferation of Sertoli cells.\(^{34}\) Karami et al reported the downregulation of GnRH in the brain of African catfishes due to exposure of MPs.\(^{35}\) Sun et al have also evinced the MPs-induced reproductive disruption due to disturbance in normal functioning of GnRH.\(^{36}\) As the hypothalamic–pituitary–gonadal (HPG) axis mainly controls the functioning of reproductive system, therefore, PS MPs-induced reduction in level of plasma and intra-testicular testosterone, as well as LH and FSH probably happened due to the disturbance of HPG axis.

To evaluate the mechanism underlying the low concentration of testosterone after PS MPs intoxication, the steroidogenic enzymes expression was determined. PS MPs reduced the expression of steroidogenic enzymes, $\beta$-HSD, $\beta$-HSD and StAR, in a dose-dependent manner. $\beta$-HSD and $\beta$-HSD mediate steroidogenic activities and play major androgenic role.

**Table 5. Effect of Different Concentrations (Control, 2, 20, 200, and 2000 $\mu$gL$^{-1}$) of PS MPs on Histopathological Parameters of Testes in Rats.***

| Parameters | Control | 2 $\mu$gL$^{-1}$ | 20 $\mu$gL$^{-1}$ | 200 $\mu$gL$^{-1}$ | 2000 $\mu$gL$^{-1}$ |
|-----------|---------|-----------------|------------------|-------------------|-------------------|
| Area of seminiferous tubules ($\mu$m) | 70.18 ± 1.15$^a$ | 71.12 ± 1.53$^a$ | 69.80 ± 0.92$^{ab}$ | 65.11 ± 0.54$^{bc}$ | 60.49 ± 0.87$^c$ |
| Area of interstitium ($\mu$m) | 19.21 ± 0.61$^a$ | 19.02 ± 0.48$^a$ | 16.52 ± 0.35$^b$ | 16.32 ± 0.19$^b$ | 14.59 ± 0.39$^b$ |
| Seminiferous tubules diameter ($\mu$m) | 190.01 ± 5.52$^a$ | 182.14 ± 2.45$^a$ | 177.30 ± 3.91$^{ab}$ | 162.53 ± 3.33$^{bc}$ | 155.22 ± 4.09$^c$ |
| Epithelial height ($\mu$m) | 64.39 ± 1.58$^a$ | 63.90 ± 1.11$^{ab}$ | 63.74 ± 1.18$^{ab}$ | 58.37 ± 1.54$^{ab}$ | 57.70 ± 1.33$^b$ |
| Spermatogonia (n) | 55.21 ± 1.07$^a$ | 53.41 ± 0.44$^{ab}$ | 52.85 ± 0.52$^{ab}$ | 51.40 ± 0.52$^{ab}$ | 49.79 ± 1.32$^b$ |
| Spermatocytes (n) | 69.92 ± 1.44$^a$ | 68.55 ± 1.03$^{ab}$ | 67.35 ± 0.45$^{ab}$ | 62.29 ± 0.61$^{bc}$ | 60.92 ± 1.29$^b$ |
| Spermatids (n) | 229.99 ± 3.06$^{a}$ | 224.67 ± 2.31$^{ab}$ | 219.52 ± 1.17$^{bc}$ | 216.75 ± 2.06$^{bc}$ | 213.26 ± 2.34$^c$ |

* Values are shown as Mean ± SEM (n = 12/group). Means within same row (for each parameter) carrying different superscripts are significantly different at $P < 0.05$. 
in testes. StAR is a rate-limiting steroidogenic enzyme which regulates the shifting of cholesterol inside mitochondria, to stimulate production of testosterone. As described by Raucci et al, testicular steroidogenesis is the vital event for production of testosterone (insert citation number) which is certainly mediated by steroidogenic enzymes and proteins. In current experiment, PS MPs exposure resulted in reduction of expressions of testicular steroidogenic enzymes, 3β-HSD, 17β-HSD and StAR, which eventually decreased the concentration of testosterone. These adverse changes are attributed to the anti-androgenic nature of PS MPs.

Various sperm parameters, such as sperm count, motility and viability are indicators of normal spermatogenesis and male reproductive health. As previously stated, the exposure of PS MPs substantially reduced the concentration of intratesticular and plasma testosterone. This reduction in

Figure 3. Seminiferous tubules after the exposure of various concentrations of PS MPs: (A) Control group shows compactly arranged seminiferous tubules having thick epithelial height and lumen filled with mature sperms, (B) 2 μg/L-1 group displays normal state of tubules with thick epithelial height and lumen filled with spermatids, (C) 20 μg/L-1 group also shows thick epithelium and lumen with less spermatids. Interstitial area is slightly decreased, (D) 200 μg/L-1 group exhibits loosely arranged tubules with thin epithelial height and a smaller number of all germ cells; and (E) 2000 μg/L-1 group shows most disordered condition, i.e. loosely packed tubules with thin epithelium and less germ cells. H&E (x40). TA: Tunica Albuginea; EH: Epithelial Height; TL: Tubular Lumen; IS: Interstitial Spaces; SG: Spermatogonia; ST: Spermatids; and SC: Spermatocyte.
concentration of testosterone is considered one of the major factors behind the decrease in sperm count. Male gametes are vulnerable to ROS owing to their continuous divisions. According to Oborna et al OS has been proved to affect the membrane fluidity and spermatogenic motility. As previously described, testicles are specifically susceptible to OS due to the presence of excessive polyunsaturated fatty acids in lipids. Polyunsaturated fatty acids are pivotal to maintain functions and membrane integrity of sperms. However, excess generation of ROS leads to disruption in viscosity and permeability of the spermatozoa membrane, thereby resulting in peroxidation of polyunsaturated fatty acids. OS-generated toxicity decreases the production of ATPs by imparting a direct impact on mitochondria of sperm cells. The reduction in ATPs in spermatozoa impairs the flagellar function, subsequently resulting in sperm immobility and apoptotic death. According to Ko et al, elevated levels of ROS stimulate a cascade of damages in the body and specifically cause toxic effects on spermatological parameters and male fertility. Therefore, these spermatogenic damages are possibly due to the elevated ROS levels and reduced concentration of testosterone. Previously Xie et al and Jin et al also reported similar results, but in the current dose-dependent study no effects were seen with lower doses of PS MPs, only higher doses deteriorated the sperm quality.

PS MPs increased the expression of Bax and caspase-3, while reduced the expression of Bcl-2. Apoptotic processes are particularly mediated by proteins of Bcl-2 and caspase family. Bax and Bcl-2 belong to Bcl-2 family. Bcl-2 is considered as an anti-apoptotic protein which mainly prevents apoptotic cell death. While, Bax is a pro-apoptotic protein which performs the antagonist action and promotes cell apoptosis. These proteins affect mitochondrial membrane permeability, thereby releasing cytochrome C in cytoplasm, which subsequently activates caspase 3. Caspase-3 belongs to cysteine proteases family which is essential for cleaving cellular proteins, thus leading to structural alterations in cells which culminates in apoptotic cell death. According to Wang and Luo, caspase-3 over-expression is the key event behind the regulation of mitochondrial apoptotic-pathway. Therefore, PS MPs exposure resulted in apoptosis due to elevation in the expression of apoptotic proteins, Bax and caspase-3, while reduction in the expression of anti-apoptotic protein, Bcl-2.

In the current research, the histopathological assessment revealed a decrease in diameter as well as the epithelial height of seminiferous tubules in the PS MPs-induced rats. In addition, seminiferous tubules having decreased interstitial spaces were noticed in testes. PS MPs significantly \((P < 0.05)\) reduced the number of various germ cells in a dose-dependent manner. Higher doses of PS MPs led to the even excessive generation of ROS and histological damages in testes. Previous investigations have demonstrated that the reduced epithelium height might be due to decreased levels of testosterone, which also reduces the number of germ cells. It was deduced that PS MPs exposure interrupted spermatogenesis, probably by disrupting the differentiation of spermatozoa to its mature stages (spermatocytes and spermatids). These damages may be attributed to disrupted integrity of blood-testis barrier followed by PS MPs exposure. Collectively, apoptotic cell death due to elevated levels of ROS as well as alterations in hormonal levels, ultimately affected the number of germ cells and induced morphometric damages within testicular tissues. Therefore, findings of the current study confirmed the endocrine disruptive property and toxic nature of PS MPs especially at higher doses.

**Conclusion**

Our findings indicated that high doses of PS MPs induced OS not only by elevating the degree of ROS and LPO but also by decreasing the antioxidant enzymes (CAT, SOD, POD). It was observed that the lipid profile was disturbed, and total protein contents were seen to be reduced. PS MPs reduced the expression of steroidogenic enzymes and hormonal concentrations as well as decreased sperm count, motility and viability. In addition to it, the apoptotic profile was changed, and histopathological changes were noticed in testicular tissues. It was observed that higher concentrations of PS MPs caused significant damages as compared to low concentrations. Taken together, PS MPs administration exhibited an anti-androgenic effect, which led to an overall testicular dysfunction in rats.

**Declaration of Conflicting Interests**

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

**Funding**

The author(s) received no financial support for the research, authorship, and/or publication of this article.

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