EXPERIMENTAL STUDY

Morphometric and stereological assessment of the effects of titanium dioxide nanoparticles on the mouse testicular tissue

Khorsandi L1,2, Orazizadeh M2, Mansouri E1,2, Hemadi M3, Moradi-Gharibvand N2

Cell & Molecular Research Center, Faculty of Medicine, Ahvaz Jundishapur University of Medical Sciences, Ahvaz, Iran. layasadat@yahoo.com

ABSTRACT
OBJECTIVE: To evaluate the effects of titanium dioxide nanoparticles (TNPs) on morphometric and stereological parameters of mouse testis.

BACKGROUND: TNPs are increasingly used in sunscreens, biosensors, food additives, pigments, rubber manufacture, and electronic materials. However, the potential toxicity of these nanoparticles is not well understood.

METHODS: Experimental Groups (TNP-1, TNP-2 and TNP-3) received one of the following treatments daily for 35 days: 75, 150 and 300 mg/kg TNPs respectively. Right testis from each animal was fixed in bouin’s solution for measurement of total volume of testsis, total volume of seminiferous tubules, total volume of interstitial tissue and total number of Leydig cells. The left testes were homogenized for measurement of testosterone concentration.

RESULTS: There was a significant decrease in the weight of testis in TNP-3 groups. The stereological and morphometrical parameters were significantly changed in TNP-2 and TNP-3 groups. TNP-2 and TNP-3 groups also showed a significant decrease in testosterone concentrations (p < 0.05).

CONCLUSION: This study had demonstrated that TNPs could change stereological and morphometrical parameters of the seminiferous tubules and reduce the number of Leydig cells and testosterone concentration (Tab. 3, Fig. 3, Ref. 35). Text in PDF www.elis.sk.

KEY WORDS: zinc oxide nanoparticles, seminiferous tubules, Leydig cells, stereology.

Introduction

Spermatogenesis is a complex process of germ cell proliferation and differentiation, which leads to the production and release of spermatozoa from the testis. This elaborate process depends on hormonal and dynamic interactions between somatic cells and germ cells of the testes. The intricate regulation and cellular interactions that occur in the testis provide multiple distinct targets, by which toxicants can disrupt spermatogenesis (1, 2). Many recent studies have demonstrated that most nanoparticles (NPs) have an adverse or toxic effect on male germ cells (3, 4). NPs are materials with at least one dimension ≤ 100 nm, and this large surface-to-volume ratio results in unique characteristics compared to their corresponding bulk materials (5). The administration of NPs to mice resulted in their accumulation in the various tissues including the brain and the testis, indicating that they could easily pass through the blood-brain and blood-testis barriers (6, 7).

Metal NPs and their oxides have a considerable number of present and future applications in the medical and industrial fields (8). Among the various metal nanomaterials, titanium dioxide nanoparticles (NTiO2) are commonly used in industry, surface coating materials, consumer products including food and biocides. Exposure to NTiO2 is not only common within the workplace, but also occurs through water, food, and cosmetics and so on (9). In vitro studies also showed that NTiO2 induces cellular toxicity of reproductive system (10-14). It has been revealed that NTiO2 could be absorbed by Leydig cells in male offspring, and induce histological changes in seminiferous tubules (15).

There are many qualitative studies of testis after NPs treatment, but the stereological aspects of NPs effects on seminiferous tubule and Leydig cells have received little attention. Thus, the present study investigated the TNPs effects on total volumes of testis, seminiferous tubules and interstitial tissue as well as total Leydig cell numbers.

Materials and methods

Animals

In this study, 32 healthy adult male NMRI (Naval Medical Research Institute) mice (6–8 weeks old, 25–30 g) were used. We performed this study according to the guidelines of the insti-
tution’s Animal Ethics Committee (approve number: IR.AJUMS.REC.1394.541). The animals were kept under standard laboratory conditions (12 h dark and 12 h light cycle, relative humidity of 50 ± 5% and 22 ± 3 °C) for at least 1 week before the experiment and those conditions were preserved until the end of the experiment. Animal cages were kept clean, and commercial food (pellet) and water were provided ad libitum.

**Experimental design**

The mice were randomly divided into the four groups, all of which contained eight animals. The doses of TNP (Sigma) were selected according to previous studies that demonstrated a significant toxicity in rodents (16, 17).

Experimental groups (TNP-1, TNP-2 and TNP-3) received 75, 100 and 300 mg/kg TNP for 35 consecutive days, respectively. The duration time of treatment was selected according to the timing of mouse spermatogenesis (18). The control group received saline orally for 35 consecutive days. One day after the last administration, after blood sampling, the mice were sacrificed by cervical dislocation under ether anaesthesia. Body and testis weight were recorded at the time of the sacrifice. Right testis from each animal was fixed in Bouin’s solution for measurement of total volumes of testis, seminiferous tubules and interstitial tissue as well as total number of Leydig cells by stereological methods. The left testicles were homogenized for measurement of testosterone concentration.

After dissection, the testis was weighed, and the primary volume (V primary) was measured using the immersion method (19). Briefly, a container with distilled water was placed on the scale and weighed, and then the testicles suspended by a thin thread were immersed in the container so that it was fully covered by water and did not touch the bottom of the container. The new weight in grams, minus the weight of the container and water, divided by the specific gravity of distilled water (1.0) was the volume of the testis in cubic centimetres. The final volume of the testis should be estimated in a stereological study to prevent reference trap.

**TNP preparation**

One gr of NTO2 nanopowder (Sigma) was added to 100 ml BSA (bovine serum albumin) solution. BSA solution at a concentration of 40 mg/ml was prepared by dissolving the powder in Milli-Q water at pH 4.7 (isoelectric point of BSA). The mixture was then ultrasonicated for 1 h. A stock solution of 1 mg/ml TNP was prepared in the BSA solution. The mixture was dispersed for 10 min by using an ultrasonicator and kept at 4 °C. The stock solution was stable for at least 7 days at 4 °C. Just before each administration, the stored solution was diluted and sonicated for 15 minutes. Atomic force microscopy (AFM) was used to characterize the prepared TNP.

**Testosterone assay**

The blood samples were collected in heparinized centrifuge tubes and centrifuged to obtain serum. Serum testosterone concentration was measured by radio immunoassay (RIA) method. Testosterone was also extracted from testes as previously described (20). Briefly, testes were homogenized by sonication and centrifuged at 5900 × g for 5 min. The supernatant was combined with an equal volume of ethyl acetate, and the organic phase was dried under a stream of N2 gas at room temperature and reconstituted in 1× PBS. The concentration of testosterone by this procedure was estimated by RIA.

**Morphometry**

Diameters of the seminiferous tubules and the lumen diameter were measured by using Motic Images Plus 2.0 image analysis software. The height of the seminiferous epithelium was calculated by subtracting the lumen diameter from the tubule diameter. For each animal, 150 tubules were analysed. Finally, a “blind” method has been used for slides reading (21).

**Stereological assessments**

1. **Histology and sampling of sections**

Each testis was embedded randomly in paraffin and sectioned exhaustively into 5μm-thick sections. Figure 1 illustrates the sampling of sections. Three sections were collected onto each glass slides. With a random start between the first 20 sections, every 20th section was sampled (the primary sections). In addition, two sections ahead of every primary section were sampled as the reference section. Because every section was 5 μm thick, it follows...

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**Fig. 1. Sampling method of histological sections is shown.**
that there was 100 μm between the primary sections and 10 μm between a primary section and the corresponding reference section. All primary and reference sections were stained with hematoxylin and eosin (H & E).

2. Total volumes of testis, seminiferous tubules and interstitial tissue

Using step-lengths of 750 μm in the x-direction (Δx) and 550 μm in the y-direction (Δy), all primary sections from each testis were systematically examined. A point-counting grid with 108 points, 1 of them encircled, was applied (Fig. 2A). Moving through all primary sections from the testis, we counted how many times 1 of the 108 points hit a seminiferous tubules or interstitial tissue. Simultaneously, we counted how many times the encircled point hit testis tissue (seminiferous tubules and interstitial tissue). The values for the total volume of testis, seminiferous tubules and interstitial tissue were then calculated based on the Cavalieri principle (22, 23).

1) V (Tes) = a/p (Tes) × N (p - p) × T × ΣP (Tes) = 0.095 mm³

Where V (Tes) is the total volume of testis, a/p (Tes) is the area per point (in this case Δx× Δy because only one point in the grid was used to count points that hit testis), N (p – p) is the number of sections between the primary sections (20 sections in this case), T is the section thickness (5 μm), and ΣP (Tes) is the total number of points that hit testis.

2) V (Sem) = a/p (Sem) × N (p - p) × T × ΣP (Sem) = 0.095 × 103 mm³

Where V (Sem) is the total volume of seminiferous tubules, a/p (Sem) is the area per point and ΣP (Sem) is the total number of points that hit the seminiferous tubules.

3) V (Int) = a/p (Int) × N (p - p) × T × ΣP (Int) = 0.095× 103 mm³

Where V (Int) is the total volume of interstitial tissue, a/p (Int) is the area per point and ΣP (Int) is the total number of points that hit the interstitial tissue.

Tissue shrinkage influences all stereological size estimators whether it is distance, surface area, or volume. There is no exact unbiased way to obtain information about tissue deformation during tissue fixation and processing. The area of a piece of testis tissue before and after fixation/processing may be estimated, and the tissue shrinkage can be calculated as (24):

\[ \frac{1 - (\frac{A - \text{after}}{A - \text{before}})^{1/5}}{ } \]

3. Total number of Leydig cells

In another session, the sampling within the primary sections was performed, but an unbiased counting frame (Fig. 2B) was now attached to the monitor. The rules of the counting frame define objects completely outside the frame or objects that touch the exclusion lines (the full lines in the Figure 2B) as being outside the frame, whereas objects that are completely within the frame or touch only the inclusion lines (the dashed lines in the Figure 2B) are defined as being within the frame. We applied the dissector principle (Sterio, 1984) to count the Leydig cells.

Whenever the Leydig cell was sampled by the counting frame, the corresponding position in the reference section was located with the other microscope, and we determined whether the Leydig cell was also visible in the reference section. The Leydig cell was counted if it appeared in the primary section, but not in the reference section. Because the sampling of sections as well as the within section sampling were performed with known sampling fractions, the total number of Leydig cells can be calculated according to the fractionator’s principle (22, 23) from:

\[ N (\text{Ley}) = \frac{N(p-p) \times \Delta x \times \Delta y}{N(p-r)} \times \Sigma Q (\text{Ley}) = 34.53 \times \Sigma Q (\text{Ley}) \]

Where N (Ley) is the total number of islets in the pancreas, N (p – p) is the number of sections between the primary sections, N (p – r) is the number of sections between a primary section and the corresponding reference section (two in this case), Δx and Δy are the step lengths, A(frame) is the area of the counting frame corrected for magnification (412.674 μm²), and ΣQ (Ley) is the total number of Leydig cells counted in one testis (23).

Results

Organ weight

No significant changes were observed in body weight of the experimental groups. Weight of testicles in the TNP-1 and TNP-2 groups was slightly lower than in the control group (p > 0.05). A significant reduction in testicular weight was observed in the TNP-3 (p < 0.05) group (Tab. 1). Relative testes weight was obtained by dividing testes weight by body weight. Relative testes weight

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**Tab. 1. Body and testis weight of control and experimental groups.**

| Group    | Body weight (g) | Testis weight (mg) | Testis/body weight (mg/g) |
|----------|-----------------|--------------------|--------------------------|
| Control  | 29.4±3.3        | 131.7±6.3          | 4.47±0.83                |
| TNP-1    | 29.2±5.2        | 130.1±10.4         | 4.45±0.95                |
| TNP-2    | 29.3±4.6        | 104.7±7.8          | 3.57±0.34                |
| TNP-3    | 29.5±3.9        | 90.5±9.9           | 3.06±0.25                |

Values expressed as mean±SD for 8 mice. * p < 0.05

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in TNP-3 group was significantly lower than in the control (p < 0.05). These results are shown in Table 1.

Testosterone assay
To determine whether TNP affect Leydig cells function, the serum and testis testosterone concentrations were assessed. As illustrated in Figure 3, concentration of the serum and testis testosterone in TNP-1 group was slightly lower than in the control group (p < 0.05). In TNP-2 and TNP-3 animals, testosterone levels were significantly decreased compared to the control group.

Morphometry
TNP-1 group showed no significant alteration in testicular parameters compared to the control group. Diameters of the seminiferous tubules and height of the seminiferous epithelium were significantly decreased in both TNP-2 and TNP-3 groups. The results of morphometric studies are shown in Table 2.

Stereology
The present study confirmed a 21% of tissue shrinkage in paraffin embedding. This shrinkage was considered when the final results were reported. In TNP-1 group, total volumes of testis, seminiferous tubules and interstitial tissue were similar to the control group. Total Leydig cell numbers were slightly lower than the control group.

In TNP-2 group, total testis volume was slightly decreased (p > 0.05). Total volume of seminiferous tubules was significantly decreased, while total volume of interstitial tissue was significantly increased. Total Leydig cell numbers were also significantly decreased when compared to the control group.

In TNP-3 group, total testis volume and total seminiferous epithelium volume were significantly lower than in the control group (p < 0.05). Total interstitial tissue volume showed a significant increase in comparison to control group (p < 0.05). Total Leydig cell numbers were significantly decreased (p < 0.01). These results are reported in Table 3.

Discussion
Based on stereological methods (such as fractionators sampling and disector counting) we found a reduction in the total volume of seminiferous tubules in TNP treated mice. This reduction was significant in TNP-3 group. TNP treatment also induced a reduction in seminiferous tubule diameter and seminiferous epithelium height. Talebi et al showed that zinc oxide nanoparticles could decrease seminiferous tubule diameter and seminiferous epithelium height (21). It is well known that increase in seminiferous tubule diameter is indicative of fluid retention resulting from an impaired emptying through the efferent ducts, whereas decrease in seminiferous diameter may indicate germ cell loss (25).

Previous studies revealed that NPs had the capacity to penetrate the blood–testis barrier and some of them had toxic action on male germ cells (26, 27). Braydich-Stolle et al showed that mammalian spermatogonial stem cells were sensitive to Ag-NPs (4). Thus, the reduction in the total volume of seminiferous tubules may have been a consequence of the germ cells loss induced by TNPs.

The total volume of interstitial tissue was increased in experimental groups. This may be a consequence of the reduction in the volume of seminiferous tubule or seminiferous tubules diameter. Another possibility is that TNPs may induce inflammation or

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**Tab. 2. Morphometrical parameters in control and experimental groups.**

| Group     | STD (μm) | SEH (μm) |
|-----------|----------|----------|
| Control   | 210.4±10.0 | 69.2±4.6 |
| TNP-1     | 209.3±11.2 | 68.9±5.9 |
| TNP-2     | 167.6±14.6** | 59.4±6.1** |
| TNP-3     | 110.1±13.7*** | 55.3±4.1*** |

Values expressed as mean±SD for 8 mice. * p < 0.05, ** p < 0.01

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**Tab. 3. Sterological parameters in control and experimental groups.**

| Group     | Leydig cell number (10⁶) | Volume |
|-----------|--------------------------|--------|
|           |                         | Testis | Seminiferous tubules | Interstitial tissue |
| Control   | 2.43±0.31                | 126.3±12.2 | 102.6±11.2 | 13.4±3.08 |
| TNP-1     | 2.36±0.43                | 125.8±9.8 | 100.4±8.7 | 13.7±0.65 |
| TNP-2     | 1.65±0.84*               | 110.7±11.8* | 81.7±7.4* | 27.3±4.1* |
| TNP-3     | 0.76±0.26**              | 84.1±9.9** | 54.7±7.2** | 30.6±3.6** |

Values expressed as mean±SD for 8 mice. * p < 0.05, ** p < 0.01
edema in interstitial tissue. However, any unusual leukocyte infiltration in this area was not observed. In spite of an increase in the total volume of interstitial tissue, the number of Leydig cells was considerably decreased in present study. Leydig cells synthesise and secrete testosterone to blood stream and testicular tissue. The presence of testosterone is essential for normal function and survival of the germ cells in seminiferous tubules (28). When the testicular environment cannot support spermatogenesis, specific mechanisms leading to germ cell death are activated (29). TNP's also caused a significant decrease in serum and testis tissue testosterone concentrations. Komatsu et al reported that titanium oxide and carbon black nanoparticles were taken up by mouse Leydig TM3 cells, and affected their viability, proliferation and gene expression (30). Yoshida et al reported that exposure to diesel exhaust nanoparticles induce Leydig cell degeneration, increase the number of damaged seminiferous tubules, and reduce daily sperm production (31).

Additionally, the reduction in testis weight also indicates that TNP could induce testicular cell loss including germ cells and somatic cells. As mentioned above, Leydig cell numbers were decreased in TNP-treated mice. The reduction in the number of Leydig cells can lead to decrease in testosterone levels and consequently germ cell apoptosis.

Apoptotic effects of TNP have also been reported in other researches. Kang et al (2009) have shown that TNP induced apoptosis in phytohemagglutinin-stimulated human lymphocytes (32). Orazizadeh et al have demonstrated that TNP induced apoptosis in liver and testicular tissues (33, 34). Park et al. showed that TNP induced apoptosis and oxidative stress in cultured BEAS-2B cells (35).

Diminution in the testis weights and relative testis weight in TNP-treated mice indicates toxic action of TNP on mouse testis. As the body growth was not significantly altered in TNP-treated mice, the effect of TNP on the testis may be due to its specific toxic effect on the target organ and not the result of its general toxicity.

In summary, this study had proved that TNP caused dose-dependent reduction in testis weight, total volume of testis, total volume of seminiferous tubules, seminiferous tubule diameter, seminiferous epithelium height and total number of Leydig cells. The reduction in the number of Leydig cells lead to decrease in testosterone concentrations and this can induce germ cell death. Increase in germ cell loss can lead to the decrease in seminiferous tubule volume and diameter, testis volume and weight. However, to state the mechanism by which TNP exerts its effects needs more investigations.

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