Transgenerational Genetic Effect of Trichloroethane (TCE) on Phenotypic Variation of Acrosomal Proteolytic Enzyme and Male Infertility Risk

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Abstract: Exposure to trichloroethane (TCE), a ambiguous environmental toxicant, has been negatively associated with male reproductive performance. The objective was to investigate, in-vivo, the mutagenic, carcinogenic or teratogenic effect of TCE maternal exposure on sperm quality and testicular cytoarchitecture of F1 generation of mice. A motile sperm separation technique was used to estimate sperm motility and a gelatin slide technique was used to measure the number of the halo around the acrosome of individual sperm as an acrosomal proteolytic enzyme (APA). Animals were followed up for signs of toxicity and mortality. Alterations in testicular tissues have been histopathology investigated. No adverse signs, symptoms and mortality were observed in the animals treated with TCE. Moreover, significant changes were seen in body and testis weight. Results of semen analysis revealed that TCE lead to low sperm count, abnormal sperm morphology, and frequently of sperm motility. These results were correlated with decrease in APA when pre-leptotene or spermatogonial cells were tested, indicating a transgenerational toxic effects. Histopathological examination revealed that TCE insult marked alterations in the microstructures of testicular tissues appeared as severe morphological abnormal spermatozoa and vacuoles. Taken together, these results suggest that early exposure to TCE causes testicular toxicity and poor semen quality. The sperm phenotypes utilized in this study may increase the value of sperm for detection mutagenic developmentally active agents, and agent with anti-fertility effects in mammals. This in-vivo animal model represents a unique platform for assessing human reproductive toxicity potential and genetic risk of various environmental mutagens, carcinogens and teratogens in a rapid, efficient, and unbiased format.

Keywords: Trichloroethane, Transgenerational Genetic Effect, Sperm Quality, Testicular Toxicity, Mice

1. Introduction

Environmental influences and insults by reproductive toxicant exposure can lead to impaired spermatogenesis or infertility. Understanding how toxicants disrupt spermatogenesis is critical for determining how environmental factors contribute to impaired fertility (8). Numerous environmental toxicants (e.g. cadmium, mercury, bisphenol A (BPA) and dioxin) have been reported to adversely affect spermatogenesis in rodents and humans, which can lead to low sperm count, abnormal sperm morphology and poor semen quality (9-11).

Trichloroethane (TCE), is widely used as an industrial solvent and a degreasing agent (12-14). It is reported that TCE is well absorbed by all exposure routes (15). Several studies with developmental exposure to lower doses than the “safe” dose suggest that TCE exposure can cause various detrimental defects, such as low fetal weight, birth defects, developmental disorders. Recently, it has been documented that chemical exposure environmentally or occupationally on a daily basis is associated with increase a risk of infertility, low fetal weights, and birth defects (15).

In general, the exposure during early embryogenesis or postnatal stage is important in the transgeneration of toxic effects (3). It has been indicated that following the direct exposure of F0 (F0 generation) pregnant rat about 90% of four generation males (F1-F4) had abnormal phenotype in tests and male germ cells, these effects increased by age to kidney disease, tumor development, prostate disease, immune abnormalities, and severe infertility (3, 5, 7). These
abnormalities were passed through male germ line, and not from females.

Exposure of mouse germ cells to radiation and chemicals results in mutation, malformation, cancer and other adverse effects (e.g., functional disorders) in the offspring, though these findings have not been proven in human studies (1). Traditionally, it is known that individual phenotypes result primarily from inherited genetic variants together with environmental exposures. However, several studies showed that a remarkable variety of factors including parental behaviors, maternal physiology, xenobiotics, nutritional supplements, environmental agents and others lead to epigenetic changes that can be transmitted to subsequent generations without continued exposure (2). The transfer of acquired heritable material or disease from parents to multiple subsequent generations through germ line is known as “transgeneration” (3). Many studies have reported that exposure to environmental toxicants may promote the transition of abnormal phenotype between generations (4-5).

The transgenerational phenomena of toxic effects involves the transmission of abnormal phenotype independently of direct exposure (6). Two possible mechanisms induce abnormal phenotype through transgeneration of toxic effects, the first is alteration in genetic material structure such as mutation and DNA sequence change, the second one includes the changing in epigenetic information such as histone acetylation and methylation, or DNA methylation (3, 5, 7).

However, to our knowledge, until now there are little/ or no studies covering the transgenerational genetic effects of environmental toxicants on male reproductive function. Thus, the aim of this study was to investigate possible transgenerational toxic effects of TCE on microstructures of testicular tissues as well as sperm parameters of F1 mice.

2. Materials and Methods

2.1. Animals, Housing and Treatment

Four-week old female albino mice (19 - 23 g) were inbred and housed in plastic cages on a bedding of wood chips at the Zoology Department (animal house) facility maintained at ~22 °C, 50–60% relative humidity, and a 12 hours light/dark cycle. The animals were provided with standard lab chow and drinking water ad libitum and were acclimated for 1 week prior to the treatment. The animals were divided into four groups of six animals each, were treated with TCE [100 and 400 µg/kg in corn oil, i.p., every 3 days (13, 16)]. The doses were calculated and delivered in 80 - 100 µl corn oil based on their body weight. The vehicle controls were received an equal volume of corn oil only. However, the sham controls were not received any treatment. TCE at 100 and 400 µg/kg was selected according to the EPA (Environmental Protection Agency) as a safe dose (17). The treatment window was selected because this is the critical development window in the mouse (15).

During the course of the exposure period, animals were observed twice per day for any adverse clinical signs or abnormal behavior that may result from toxicity. The animals were weighed daily during the dosing period to adjust the administered dose for body weight changes.

At eight weeks of age, treated F0 females were mated with fertility confirmed control males. Mating was confirmed by the presence of vaginal plug. The day the vaginal plug was observed, females were removed from males and individually caged. F0 pregnant dams were observed daily and body weight was measured daily to further confirm pregnancy. The dams were allowed to deliver naturally and the delivery day was designated as PND-0.

After delivery, the size of each litter was standardized on PND-4 by eliminating extra pups through random selection within sex from litters with more than 10 pups to yield 10 pups, with five females and five males per litter. Natural litters with 10 or fewer pups were not standardized. F1 litter sizes, and weights, sex ratios, stillbirths, and percent of dead pups were recorded.

At three weeks of age, two F1 males from each litter were selected for examination of semen quality and tests. Moreover, at least one F1 male from each litter was selected to examine fertility. F1 males were housed with fertility confirmed control females. During the mating phase, all the females were weighed twice per week. Once the plug was observed, females were weighed and individually caged. Then, these females were monitored twice daily, and weighed twice per week until parturition. F2 litter sizes and weight as well as sex ratios and percent of dead pups were recorded on PND-1. After the three-week mating period, unmated males were singly housed and were considered as infertile males.

2.2. Measurement of Body and Testis Weight

Body and testis from the four groups were weighed before and at the end of the exposure period.

2.3. Sperm Function Evaluation

Spermatozoa were collected by mincing the epididymides in 37ºC normal saline from F1. Epididymal sperm counts and evaluation of the motility of epididymal sperms were performed as previous described (18-19) and the WHO manual for semen analysis (20). The examination of sperm morphology was carried out by dropping a drop of sperm suspension on a slide. After air dried, stained with 1% eosin Y, washed and air dried again, the smears on the slides were observed under a light microscope (Leica, Germany).

2.4. Acrosomal Proteolytic Activity (APA)

A gelatin substrate film assay was used to assess the acrosomal proteolytic activity (APA) of individual spermatozoa, according to the method described previously (18, 21). This assay provides a visual demonstration of the release of enzyme from the acrosome of individual sperm. This technique is based on the principle that when sperm are incubated after smearing on the gelatin, the proteolytic enzyme in the acrosome digests the gelatin, making a halo
around the acrosome as an indication of APA. The release of proteolytic enzyme from the intact acrosome of sperm is due to the evaporation procedure that causes ruptures of the plasma membrane, with consecutive disintegration of the acrosomal matrix. In brief, sperm suspension was washed thrice in 0.9% saline solution with 0.5% bovine serum albumin. On drop of sperm suspension was spread on the surface of glass slide coated with a thin layer of gelatin. The slides with sperm were incubated for one hour at 37°C in a wet chamber to allow proteolysis of the gelatin by acrosin and the surface of glass slide coated with a thin layer of gelatin. The slides with sperm were incubated for one hour at 37°C in a wet chamber to allow proteolysis of the gelatin by acrosin and then stained with toluidine blue. APA was indicated by a halo of unstained degraded protein around individual spermatozoa heads, sperm without activity lack an unstained halo. The percentage of sperm showing a halo was assessed by counting 200 sperm.

2.5. Histological Studies

At three weeks of age, to avoid litter effects, two F1 males per litter from each treatment group was euthanized by decapitation and the testis were collected for analysis. Following dissection, 10%-formalin-fixed testis were processed in a series of graded ethanol solutions and embedded in paraffin wax. Paraffin sections were cut at 6-8 µm thickness, deparaffinized, hydrated, stained with hematoxylin and eosin (H&E) (22) and examined under a light microscope (Leica, Germany). All testicular histology was assessed blind by a histopathologist.

2.6. Statistics

Data were expressed as means ± SEM from six female mice of each treatment group using SPSS software, version 20. A computerized Kolmogorov-Smirnov test was used to determine whether the data fitted a normal distribution. One-way ANOVA test followed by Tukey's post hoc comparisons was used to make multiple comparison between treatment groups. Student's t-tests were used to make comparisons between two groups. Mann-Whitney U-test was used for nonparametric samples. Statistical significance was assigned at *P* ≤ 0.05.

3. Results

3.1. Effect of TCE on Body and Testis Weight

The statistical data in Table 1 showed that body and testis weight significantly (*P* < 0.05) changed in the TCE-treated groups compared to controls. The results showed that body weight in F1 males in 400 µg/kg TCE-treated group, but not 100 µg/kg TCE-treated group (*P* = 0.45) was significantly increased (*P* = 0.0013) comparing to controls (Table 1). However, testis weight in F1 males in 400 µg/kg TCE-treated group, but not 100 µg/kg TCE-treated group (*P* = 0.24) was significantly decreased (*P* = 0.025) comparing to controls (Table 1). No significant changes were observed in body and testis weight in sham controls and vehicle controls (data not shown).

<table>
<thead>
<tr>
<th></th>
<th>Body weight (g)</th>
<th>Testis weight (g)</th>
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<tbody>
<tr>
<td>Control</td>
<td>23.33 ± 1.03</td>
<td>0.25 ± 0.007</td>
</tr>
<tr>
<td>100 µg/kg</td>
<td>25.22 ± 0.43</td>
<td>0.23 ± 0.008</td>
</tr>
<tr>
<td>400 µg/kg</td>
<td>29.4 ± 1.03##</td>
<td>0.18 ± 0.021##</td>
</tr>
</tbody>
</table>

Table 1. The effects of TCE exposure on body and testis weight.

Data are mean ± SEM (n = 6 per group). Significance at *P* ≤ 0.05. #Significantly different from the controls.

3.2. TCE Reduces Sperm Quality of F1 Mice

Here we explored the effects of TCE insult on the sperm quality, count, motility, and morphology. The results showed that TCE at a dose of 100 µg/kg had no significant (*P* = 0.388; Figure 1) effect on the count of cauda epididymal sperm comparing to controls. However, TCE exposure at a dose of 400 µg/kg has significantly (*P* = 0.033) decreased the sperm count of cauda epididymis comparing to controls (Figure 1).

![Figure 1](image1.png)

Figure 1. Effect of TCE insult on the count of cauda epididymal sperm of F1 mice. Quantification of sperm count. Data are expressed as the mean ± SEM (n = 6). #Significantly different from the controls (*P* ≤ 0.05).

Next, we investigated the effect of TCE insult on the sperm motility of caudal epididymis *in-vitro*. Sperm motility seemed significantly (*P* < 0.05) changed in TCE-treated groups compared to controls (Figure 2). Specifically, 100 µg/kg TCE treatment group showed lower (*P* = 0.049) percent of motile sperm compared to controls (Figure 2). The results also showed that the percent of immotile sperm induced by 400 µg/kg TCE (~ 55%) was greater than observed with 100 µg/kg TCE (~ 41%). Moreover, there was no significant difference in the percent of motile sperm between 100 and 400 µg/kg TCE treatment groups (Figure 2).

![Figure 2](image2.png)

Figure 2. Effect of TCE insult on the motility of cauda epididymal sperm of F1 mice. Quantification of sperm motility. Data are expressed as the mean ± SEM (n = 6). #Significantly different from the controls (*P* ≤ 0.05).

Regarding the morphology of cauda epididymal sperm of F1 mice, TCE insult had a harmful effect on the sperm...
morphology in TCE-treated groups compared to controls (Figure 3). The sperm morphology were shown in Figure 3A from which no head sperm, sticky sperm and sperms with coiled tail were observed in the TCE-treated groups. Sperm head-tail detachment and sperm tail fragmentation were most commonly observed abnormalities by microscopy in cauda epididymal spermatozoa of TCE-exposed F1 mice (Figure 3 A). Sperm fragmentation was not commonly observed after TCE treatment (Figure 3A). Common defects in the TCE groups/set included sperm head malformations (Figure 3A) and a pattern of longitudinal bundling of seemingly morphologically normal spermatozoa (Figure 3A).

The statistical analysis showed that the percent of abnormal cauda epididymal sperm in the TCE-treated groups was significantly increased compared to controls (Figure 3B). Specifically, in 100 µg/kg TCE-treated group showed lower ($P = 0.000025$) abnormal sperm compared to 400 µg/kg TCE-treated group (Figure 3B).

3.3. Effects of TCE on Sperm Acrosome Reaction

Here we characterized, in-vitro, the impact of TCE insult on the caudal epididymis sperm. Sperm from the caudal epididymis produced halos on gelatin substrate slides (Figure 4A). The motility of cauda epididymal sperm seemed unchanged in sham controls compared to vehicle controls (data not shown). Gelatin substrate film assay was conducted to assess APA of individual sperm. The heads of ~ 75% of sperm from controls had protein-digested halos generated by acrosin activity on the gelatin substrate films. By contrast, only the heads of ~ 37% and 30% of sperm from 100 and 400 µg/kg TCE-treated groups, respectively, had protein-digested halos generated by acrosin activity on the gelatin substrate films (Figure 4B), meaning that proteolytic activity is inhibited by TCE insult.

Figure 4. Sperm motility after TCE insult. (A) Photomicrograph of motility of the cauda epididymal sperm of F1 mice treated parents showing haloes of unstained degraded protein around individual spermatozoan heads (arrow) and sperm with acrosomal proteolytic activity (no haloes; arrowhead). The magnification is X100. Scale bar = 20 µm. (B) Quantification of halo formation in response to TCE exposure. Data are expressed as the mean ± SEM (n = 6). # Significantly different from the controls ($P ≤ 0.05$).

3.4. TCE Insult Alters Cytoarchitecture of the Testicular Tissue of F1 Mice

To explore the effect of TCE insult on the microstructures of testicular tissues of F1 mice, we performed histopathological evaluations. Microscopic observation of the testicular tissues by H&E staining revealed that normal morphology and cellular arrangement of seminiferous tubules appeared in the control (Figure 5A-B). Moreover, closely packed seminiferous tubules, separated from each other by narrow interstitial spaces containing interstitial Leydig cells around blood capillaries. These seminiferous tubules were lined by spermatogenic and Sertoli cells. The spermatogenic cells were formed of spermatogonia, primary spermatocytes and spermatids. The spermatogonia appeared as small cells Spermatogonia with dark, ovoid nuclei, these cells are located basally in the epithelium next to the basement membrane. Complete spermatogenesis and regular structure, many spermatozoa present in disorganized tubules.

In contrast, there was morphological abnormal spermatooza and vacuoles in the TCE-treated groups. Testicular tissues from the 100 µg/kg TCE-treated group showed disorganization in some seminiferous tubules, cellular irregularity and large vacuolization between spermatogenic cells and intracellular, also a thickness in basement membrane of spermatogenic epithelium were observed in some tubules (Figure 5C-D). Further analysis showed that the testicular tissues of 400 µg/kg TCE-treated group showed degenerative changes in all seminiferous tubules with a few germ cells in the lumen and marked intercellular and basal vacuolation, detachment of spermatogonia from the basement membrane and separation between germinative cells in the seminiferous tubules (Figure 5E-F). In some seminiferous tubules there are slightly reduction in density of germinal cells. There was also disruption to the arrangement of Sertoli cells and germinal cells and increasing space between them. Degeneration in
Leydig cells were also observed. No spermatogenesis in the lumen of seminiferous tubules were observed.

**Figure 5.** The effect of TCE insult on microstructures of testicular tissues of F1 mice observed by H&E staining. (A-B) A photomicrograph of a testicular tissue of control showing (A) closely packed seminiferous tubules (ST), lined by normal spermatogenic cells (↔), Leydig cells (LY) within interstitial spaces (IS). (B) The STs were lined by spermatogenic cells and Sertoli cells (SC), spermatozoa (SP) present in the lumen. (C-D) A photomicrograph of a testicular tissue of 100 µg/kg TCE-treated group showing (C) detachment of spermatogonia from the basement membrane (→), Leydig cells (LYD) within interstitial tissue of control showing (A) closely packed seminiferous tubules (ST), lined by normal spermatogenic cells (↔), Leydig cells (LYD) within interstitial spaces (IS). (D) Detachment of spermatogonia from the basement membrane (→) and intercellular vacuolations (VAC). (E-F) A photomicrograph of a testicular tissue of 400 µg/kg TCE-treated group showing; (E) Affected ST with detachment of spermatogonia from the basement membrane (→) and intercellular VAC. (F) complete loss of spermatogenic cells (VAC). (A, C, E, X10). (B, D, F, X40). Scale bar = 50 µm.

4. Discussion

Infertility caused by exposure to many environmental toxicants is a global problem, particularly in industrialized countries (23). The environmental toxicants, such as vinclozolin, bisphenol A and phthalates have the ability to induce changes in the genetics and the epigenetics of male germ line, and promote the transgenerational inheritance that affect male fertility (3, 7, 24).

The male reproductive system has emerged as one of the major toxicity targets of environmental toxicants. Although acute exposure of toxicants contributes to apoptosis and necrosis of testicular cells, chronic and sub-lethal exposure is prevailing in the general public (23, 25). Due to the unusual long half-lives of some of these toxicants in mammalian body (e.g. cadmium has a mean half-life of 15 years (26)), chronic and low level exposure to humans could cause long-term unwanted health effects. **In-vivo** studies are crucial to assess toxicants with adverse effects on human health.

The purpose of the present study was to evaluate, **in-vivo**, the long-term effect of TCE exposure on sperm APA, sperm motility, sperm morphology and sperm count. Although, TCE has been classified as non carcinogenic (group-3), because there is inadequate evidence for carcinogenicity in both human and animals (27-29), many reports indicated that TCE bound with DNA, RNA and proteins in many organs such as liver, kidney, lung and stomach in F0 mice and rats following a single intraperitoneal injection, but did not induce any abnormal changes (30-31). Based on these finding, this study suggested that TCE might induce transgenerational inheritance in the genetic and epigenetic of male germ line. The result of this study indicated that the spermatogenesis of F0 was slightly affected by TCE, it significantly decreases the total count of sperms in treated groups, these finding similar to previous result of many reports (32). However, in F1 the figure of spermatogenesis parameters was completely different, the result showed important histopathologic finding in testis, decreases in sperm count, abnormal acrosome reaction and teratogenic sperms.

The proper functioning of the flagellum and acrosome of sperm is vital for the process of fertilization and thus determines fertility potential of the sperm (33). One of the more elusive questions in genotoxic research is whether a change in sperm phenotype following treatment with a suspected mutagen is due to mutation or developmental disturbance. However, the effects of TCE exposure on male reproduction in later reproductive life are unclear. The results of this study showed that the induced loss sperm motility and APA in single spermatozoa derived from TCE-treated mice is caused by mutational or developmental effects. Gene mutation/s induced in spermatogonial cells and spermatogonial stem cells could readily be transmitted through cell cycle to be induced and expressed in spermatozoa. That environmental toxicant TCE is capable of inducing gene and chromosomal mutations in spermatogonial cells are unclear. The findings of this study suggest that TCE treatment of these stages therefore probably interferes with the expression of existing gene products (18).

Exposure to different classes of environmental toxicants can disrupt male reproductive function by affecting the endocrine system, by changing gene expression that is pertinent to spermatogenesis and steroidogenesis and by exerting epigenetic effects, which can result in abnormalities in the reproductive system of male offspring up to four generations following in **utero** exposure (23).

When referring to the testicular toxicity induced by environmental toxicant TCE, (34-35) indicate that the administration of TCE has no impact on the weights of body and testis in mice but can induce spermatogenic damage and affect sperm counts and sperm motility. In contrast, in this study we found that early exposure to TCE had significant effect on the body and testis weight. Therefore, histopathologic analysis of testis and epididymis showed that an extensive emptying of the tubular germinal elements into the epididymis. Since 90% of the volume of the testis is seminiferous tubules (36), the extensive loss of germinal cells from these tubules may account for the 50% decrease in testis weight. Along with their
indications, the sperm toxicity tests also reveal that sperm count, motility and normal sperms were sharp reduction in the TCE-treated mice. The decrease in the body weight may be caused by the decline of food intake and the reduction in the testicular weight is attributed to the necrotic changes in the testis (37-38). Accordingly, we consider that the observed abnormal features in the testis tissues by H&E analysis are responsible for the decreased testicular weight.

It is well known that histological assays are reliable tools to detect morphological changes due to toxicants; hence, the histopathology of various treated tissues was examined. The abnormal features in the testis and the aberrant sperm parameters are reported to result from the reactive oxygen environmental toxicants to male reproductive health.

This study offers a framework to possibly develop approaches to therapeutically treat and/or manage the damaging effects of toxicants and have no efficiently alleviating effects on the sperm toxicity. Concerning the situation in the paper, the factors can be complicated. One of them may be that the ROS induced by TCE exposure may be not the sole reason leading to the damage of testis. Additional studies comparing low level chronic exposure versus high level acute exposure to environmental toxicant TCE are also required to elucidate fully the underlying molecular mechanism by which this toxicant disrupts male reproductive function.

In conclusion, the present study finds that the TCE exposure can cause serious testicular toxicity. Since a low sperm count is correlated with decreased sperm quality and acrosomal integrity, which is essential for penetration of the zona pellucida by sperm. The presence of these sperm phenotypes may help to detect chemicals with anti-fertility effects. This study offers a framework to possibly develop approaches to therapeutically treat and/or manage the damaging effects of environmental toxicants to male reproductive health.

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References


