# **Protective Effect of N-Acetyl Cysteine on Chlorpyrifos-Induced Testicular Toxicity in Mice**

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Abstract <sub>-</sub>

**Background:** Chlorpyrifos (CPF), an organophosphate pesticide, is widely used in farms in order to preserve crops and fruits. Previous studies have shown that CPF exposure might cause chronic toxicity in male genital system. The present study investigated the protective effect of N-Acetyl Cysteine (NAC), a potent antioxidant against testicular toxicity of CPF in male mice.

*Archivels* **Example 1 I** maile mice.<br> **Methods:** In this experimental study, 42 adult male mice were divided into seven group<br>
display (5 mg/kg, b.w), NAC+CPF 0/5 mg/kg<br>
dimethyl sulfoxide (DMSO, 0.75% solution mg/kg, b **Materials and Methods:** In this experimental study, 42 adult male mice were divided into seven groups, CPF low (0.5 mg/kg.b.w) and high (5 mg/kg.b.w) doses groups, NAC group (35 mg/kg.b.w), NAC+CPF 0/5 mg/kg.b.w, NAC+CPF 5 mg/kg.b.w, dimethyl sulfoxide (DMSO, 0.75% solution mg/kg.b.w) and control group. All treatment were done intraperitoneally. Treatment was conducted for four consecutive weeks (five days each week). However NAC was injected to NAC+CPF groups five days before initiation of the treatment procedure. One week after the last injection, mice were sacrificed using anesthetic gas to evaluate alterations in testicular histology and sperm parameters.

**Results:** Seminiferous tubules area and diameter were significantly diminished in the group treated with 5 mg/kg CPF (P<0.05). CPF also statistically reduced sperm parameters (count and motility) and damaged sperm morphology) at both doses (P<0.05). However, NAC significantly improved spermatogonia, spermatocytes, spermatid cell counts as well as sperm parameters in mice treated with both CPF concentrations (P<0.05).

**Conclusion:** According to our results, NAC may significantly ameliorate CPF-induced damages to spermatogonia, spermatocytes, spermatids cell counts and sperm parameters.

*Keywords:* Chlorpyrifos, N-acetylcysteine, Protective, Sperm

**Citation: Kheradmandi R, Jorsaraei SGA, Feizi F, Moghadamnia AA, Neamati N. Protective effect of N-acetyl cysteine on chlorpyrifos-induced testicular toxicity in mice. Int J Fertil Steril. 2019; 13(1): 51-56. doi: 10.22074/ijfs.2019.5494.**

# Introduction

As reported by the World Health Organization (WHO), unsuccessful pregnancy has been globally increased. Re searchers found that 48.5 million couples worldwide were unable to have a child after five years of unprotected regu lar sexual intercourse (1).

Almost all people working on agricultural fields are exposed to various toxins that may cause reproductive toxicity. Pesticides are widely used for eliminating pests to protect corps and fruits. Organophosphate pesticides are regarded as dangerous types of pesticides for the en vironment as they can affect humans and animals health (2). Chlorpyrifos (O,O-Diethyl O-3,5,6-trichloropyridin-2-yl phosphorothioate), is an organophosphate pesticide which can cause adverse effects on the reproductive system of both males and females (3). For instance, seminiferous tubules were significantly degenerated in chlorpyrifos (CPF)-treated mice (4). In addition, sexual hormones disturbance and also defects in sperm production have been reported following CPF exposure (5). CPF was also shown to increase DNA impairment (6, 7) and induce harmful effects in different organs such as the thyroid (8) and lung (9). CPF permanently binds acetylcholinesterase and inhibits deactivation of acetylcholine in the synapses. So, acetylcholine signaling may last longer. This process is irreversible unless new acetylcholinesterase enzymes are synthesized. It has been reported that CPF also induc es oxidative stress (8, 10).

Acetylcysteine, also known as N-acetyl cysteine (NAC) is widely used in management of acetaminophen over dose, cystic fibrosis and also chronic obstructive pulmo -

**Received: 19/March/2018, Accepted: 8/July/20-18**

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**Royan Institute International Journal of Fertility and Sterility Vol 13, No 1, April-June 2019, Pages: 51-56**

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nary disease. NAC may be useful in toxins treatments, since it can escalate glutathione levels and prevent further injuries caused by lipid peroxidation (11). Furthermore, a significant improvement of sperm motility and morphol ogy were observed by NAC treatment in varicocele and also other models induced by synthetic drugs such as par acetamol (12, 13). Moreover, NAC may protect male gen ital system against strong toxins such as arsenic trioxide in (14). It has been observed that NAC has more marked effects compared to vitamin C in improvement of sperm parameters (15). Therefore, this study was conducted to investigate the protective effect of NAC on histopathol ogy of testis and sperm parameters in CPF-treated mice.

## Materials and Methods

## **Chemicals**

Chlorpyrifos (99%) and NAC (99%) technical grade were purchased from Sigma-Aldrich (St. Louis, MO, USA) [lot No. LC13116V and 616-91-1, respectively]. Also dimethyl sulfoxide (DMSO) was provided from Sig ma too (St. Louis, MO, USA) [Lot No. 67-68-5].

## **Experimental design**

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spectring to diffusion method (1<br>
13116V and 616-91-1, respectively], cubator at 37°C for allmost Forty-two healthy adult BALB/c mice (6-8 weeks old) were obtained from the Animal Research Unit, Babol Medical University, Babol. Animal care and handling was done based on Animal Research Unit and follow ing approval of Ethics Committee (MUBABOL.HRI. REC.1395.73). The animals were habituated to labora tory conditions for 1 week before initiation of the experi ment. Mice were maintained on 12 hours light-dark cycle at 21-24°C with 50-60% humidity. Mice had free access to normal diet and water, ad libitum. The animals were di vided into seven groups: group I (control group) received normal saline, group II (sham group) received DMSO (0.75% solution), group III received NAC 35 mg/kg.b.w, group IV (high CPF) received CPF 5 mg/kg. b.w, group V (low CPF) received CPF 0.5 mg/kg.b.w, and group VI and VII received CPF at low (0.5 mg/kg.b.w) and high (5 mg/ kg.b.w) doses, respectively along with NAC on a daily basis. In groups VI and VII, NAC was given intraperito neally from five days before the experimental timeline, in order to acclimate mice with this antioxidant. All groups were treated intraperitoneally except the control group. Treatment was conducted for 4 weeks and injections in all groups were administrated on five consecutive days each week. One week after the last injection, mice were sacrificed using anesthetics to evaluate sperm parameters and testis histopathological alterations.

### **Chemical solution preparation**

Here, 15 µL DMSO was added to 1985 µL distilled wa ter to prepare 2 ml DMSO solution to be administered to the sham group. Also, 1 mL DMSO was added to CPF powder vial (1 mg) in order to prepare CPF stoke solution ( $1 \text{mg}/1 \text{m}$ L). Afterward, 15 µL CPF was added to 135 µL distilled water and after pipetting, the whole solution was added to 1850 µL distilled water to prepare 2 ml High CPF (5 mg/kg.b.w) solution. Eventually, 200  $\mu$ L of high CPF solution was added to 1800 µL distilled water to pre pare low CPF (0.5 mg/kg.b.w) solution. NAC was dis solved in water at 35 mg/kg.b.w. It should be noted that fresh CPF solutions were daily prepared.

#### **Sperm motility, count and morphology assessment**

Seven days after the last day of treatment, mice were anaesthetized via an inhalation induction chamber and sacrificed. Right testis of each animal was excised and put in 10% formalin solution for histopathological evalu ations. Afterward, the caudal of left epididymis of each animal was excised and put in petri dish containing 3 mL Ham's F10 (St. Louis, MO, USA) [Lot No. 87120401]. According to diffusion method (16), for assessment of sperm parameters, epididymis was tattered to smaller pieces using sterile needle syringe and kept in a  $CO<sub>2</sub>$  incubator at 37°C for almost 30 minutes. Then, sperm pa rameters including sperm count, motility and morphology were evaluated under light microscopy.

From semen samples prepared by diffusion method, almost 50 µL semen from each mouse was smeared by a pipette on a slide. Afterwards, maximum 100 sperms were observed on right upper quarter of each slide to examine sperm count, motility and morphology. Sperm normality percentage for each mouse was easily calcu lated using a counter by knowing about mice sperm ab normalities (16).

It was very important that well-mixed semen sample was spread at appropriate thickness on each slide to evaluate sperm parameters. During sperm assessment, room temperature was maintained at 21-24°C because increased temperature may enhance semen degeneration speed.

### **Histopathological examinations**

Testis specimens were kept in 10% neutral buffered formalin. For testis histopathological evaluations, 5 µm sections were prepared from each testis, stained with haematoxylin and eosin (H&E) and observed under a light microscope. Images were captured by Olympus optical microscope equipped with a Canon HD camera at magnifications  $\times$ 4,  $\times$ 10 and also  $\times$ 40 at four random points. Afterwards, data were evaluated on a proper per sonal computer using Motic software instruction (17). Numbers of spermatogonia, spermatocytes and sperma tid cells and also seminiferous tubules area and diameter were observed by using Motic histomorphometric utility options.

### **Statistical analysis**

Data were presented as mean  $\pm$  standard error (SE). Statistical analysis was performed in SPSS (version 22, SPSS Inc., Chicago, IL) using one-way analysis of vari ance (ANOVA) followed by Tukey as the post hoc test.



The data are presented as mean ± SE (n=6). Sperm count is expressed as number×10<sup>s</sup> per caudal epididymis. <sup>a</sup>; Indicates a significant difference as compared to control group (P<0.05), **<sup>b</sup>; Indicates a significant difference as compared to CPF group (P<0.05), NAC; N-Acetyl Cysteine, CPF; Chlorpyrifos, and DMSO; Dimethyl sulfoxide,** 

#### Results

#### **Morbidity and mortality**

Male mice that received CPF (0.5 and 5 mg/kg.b.w/day) for 35 days showed signs of toxicity such as salivation, diarrhea and tremor. No death was recorded throughout the study period.

#### **Sperm characteristics**

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<i>Archive of CPF* group,  $P > 0.05$ ). Admi According to our data, no significant differences were found in sperm characteristics between DMSO and con trol group (Table 1, P>0.05). Administration of CPF 0.5 mg/kg.b.w/day (low CPF) showed significant decreases in sperm motility, count and also morphology (P<0.0001). In addition, sperm characteristics considerably decreased following administration of CPF 5 mg/kg.b.w/day as com pared to control group (P<0.0001). Treatment with NAC alone made no significant changes to motility, counts and morphology. However, NAC treatment in combination with low CPF caused significant increases in motility and counts and markedly improved sperm morphology as compared to control CPF-induced groups (P<0.0001). NAC also caused significant increases in motility and improvements in mor phology when co-administered with high CPF (P<0.0001), while sperm count showed no significant increases.

#### **Histomorphometry**

Average count of spermatogonia, spermatocytes and spermatid in DMSO group slightly decreased but it was not significant; however, a significant increase was observed in NAC group (P<0.001, Fig.1). It was demonstrated that mean number of spermatogonia cells significantly decreased in low CPF group  $(P<0.04)$ . Meanwhile in High CPF group, spermatogonia, sper matocytes and spermatids were considerably decreased (P<0.0001). Treatment of CPF groups with NAC re sulted in significant increases in the average number of spermatocytes and spermatids (P<0.001 and P<0.007, respectively). However, mean of spermatogonia cells counts in high CPF+NAC group had no significant in crease  $(P<0.05)$ .

Based on data given in Figures 2 and 3, there was no significant increase in mean seminiferous tubules area and diameter in DMSO group compared to con trol (P> $0.\overline{05}$ ); but, NAC showed a significant increase in both variables (P<0.001). While high CPF treat ment significantly diminished seminiferous tubules, low CPF treatment (0.05 mg/kg.b.w) caused no considerably damage in seminiferous tubules shape. NAC could not ameliorate the effects caused by high CPF  $(P>0.05)$ .



Fig.1: Bars presents mean ± SE of spermatogonia, spermatocytes and spermatid cells counts in different groups. <sup>\*</sup>; Indicates a significant difference in spermatogonia counts as compared to control group (P<0.05), \*\*; Indicates a significant difference in spermatocytes counts as compared to control group<br>(P<0.05), \*\*\*; Indicates a significant difference in spermatids counts (P<0.05), \*\*\*; Indicates a significant difference in spermatids counts as compared to control group (P<0.05), NAC; N-Acetyl Cysteine, CPF; Chlorpyrifos, and DMSO; Dimethyl sulfoxide.







Fig.3: Bars presents mean ± SE of seminiferous tubules diagonal length in different groups. \*; Indicates a significant difference as compared to control group (P<0.05), NAC; N-Acetyl Cysteine, CPF; Chlorpyrifos, and DMSO; Dimethyl sulfoxide.



**Fig.4:** Histopathological difference is shown between experimental groups. It is demonstrated a massive destruction in CPF groups. However NAC considerably improved histopathology of testis. **A.** Control, **B.** DMSO, **C.** NAC, **D.** Low CPF, **E.** High CPF, **F.** Low CPF+NAC, and **G.** High CPF+NAC. NAC; N-Acetyl Cysteine, CPF; Chlorpyrifos, and DMSO; Dimethyl sulfoxide.

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Based on data given in Figure 4, there was no significant increase in mean seminiferous tubules area and diameter in DMSO group compared to control (P>0.05); but, NAC showed a significant increase in both variables (P<0.001). While high CPF treatment significantly diminished semi niferous tubules, low CPF treatment (0.05 mg/kg.b.w) caused no considerably damage in seminiferous tubules shape. NAC could not ameliorate the effects caused by high CPF  $(P>0.05)$ .

# **Discussion**

In CPF-exposed mice, a considerable reduction in sperm parameters was found. Meanwhile, NAC could not significantly improve sperm motility, morphology and count. NAC in combination with CPF 5 mg/kg.b.w. considerably prevented further damages to sperm motil ity and morphology; However, NAC could not improve sperm counts caused by CPF-induced toxicity. In a simi lar study on CPF reproductive toxicity, a significant de crease in sperm motility and counts was observed by CPF gavage at 20 mg/kg.b.w (18).

In addition, CPF considerably decreased the level of antioxidant enzymes and glutathione in plasma. *Nigella sativa* oil can act like NAC as a potent protective agent which statistically improved sperm parameters, antioxi dant enzymes activity and testosterone level (18). Ac cording to our results, adverse effects of CPF is likely irreversible and has negative effects on genitalia systems.

For decades, CPF devastating effects on spermatogen esis process was unclear. Other scientists investigated various pesticides at different doses for their negative effects on spermatogenesis process (19). Organophosphates or any other chemicals, such as different toxins, that have adverse effects on tissues and cells may be inadvertently absorbed through the skin or digestion system. In order to confront these harmful effects, consumption of antioxi dant substances like syrup of Malva sylvestris (20) along with hydroalcoholic extract of *Fumaria parviflora* (21) or products such as propolfenol (22) is highly recommended to protect against damages induced by toxins, particularly against those cause in the genital system. However, some antioxidant materials such as catechin and quercetin did not have significant influences in this regard (9). Never theless, natural nutrients such as ginger and cinnamon could be effective on male genital dysfunction due to their anti-oxidant efficacy (23).

Vitamin C and E have been widely used in previous stud ies and were introduced as beneficial protective materials to compensate damages induced by organophosphates such as malathion, a broad spectrum organophosphate pesticide that could decrease sperm parameters and induce histo pathological alteration (24). Although vitamin C and E are potent protective materials against various toxins, a lower dose of intraperitoneal NAC possibly has a more marked impact on sperm parameters based on our findings but use of food or fruits overfilled with these vitamins is suggested for people who are daily exposed to pesticides (25). An -

other study showed that vitamin C only resulted in a sig nificant improvement of sperm motility (26).

The present study indicates that NAC at 35 mg/kg.b.w somehow significantly increased seminiferous tubules area, diagonal diameter, and spermatogonia, spermato cytes and spermatids counts. Meanwhile CPF 0.5 mg/ kg.b.w could not considerably reduce seminiferous tu bules area and diagonal diameter. Furthermore, CPF 5 mg/kg.b.w significantly diminished seminiferous tubules.

Based on these findings, NAC in combination with CPF ameliorates the pesticide's adverse effects on testis. It seems that intraperitoneal injection of NAC, even at a low dose has a more marked effect on sperm parameters than gavage administration (15). It has been proven that NAC can affect lipid peroxidation (LPO) (27). Therefore, NAC might decrease ROS elevation caused by CPF. However, in the present study, NAC exact effects on sexual hor mones or anti-oxidant enzymes such as superoxide dis mutase, catalase, or glutathione in treated groups, were not evaluated. But considering significant reductions in testis germinal cells, oxidative stress level was probably elevated by CPF and NAC ameliorated the adverse effect of CPF on the testis.

computation with CP+ 5 mg/g<sub>1</sub> b.w.<br> *Area* computed further damages to sperm motil-<br>
an affect lipid peroxidation (LPO)<br> *ARC* exact study, NAC could not improve<br>
any deligit decrease ROS glevation cause<br>
aby CPF-induced According to our results, seminiferous tubules area and diagonal diameters were not affected by CPF. It sug gests that resting times at the end of each week and also seven days after the last injection of CPF might provide a chance for the immune system to recover and regener ate genital and possibly other tissues. Therefore we did not expect NAC to protect these two unaffected vari ables. Meanwhile, we assume that CPF at the dose of 0.5 mg/kg.b.w could not significantly diminish seminiferous tubules area following four-week administration. Maybe by longer treatment periods, CPF could induce more de structive effects at the dose of 0.5 mg/kg.b.w It is clear that NAC is able to confront negative effects of CPF toxicity in male genital system but what if we could use NAC at doses higher than 35 mg/kg.b.w? In this case, we probably observe NAC protective effects against CPF typical tissue toxicity. Further *in vivo* studies us ing intraperitoneal injections, are highly recommended to affirm our data.

# Conclusion

Both low and high doses of CPF can decrease sperm parameters. Also, this pesticide at 5 mg/kg.b.w dose sig nificantly diminishes the length and diagonal diameter of seminiferous tubules. NAC significantly improved CPF adverse effects on sperm parameters and spermato genesis cells except spermatogonia. However, this anti oxidant could not statistically ameliorate the histopatho logical alterations of seminiferous area induced by CPF.

# Acknowledgements

We gratefully appreciate the efforts of Dr. Ali Asghar Ahmadi and Dr. Maryam GholamiTabar Tabari in this *<www.SID.ir>*

study. We also appreciatively acknowledge the contri butions of the Cellular and Molecular Biology Research Center at Babol University of Medical Science for the grant and full support. This study was supported by Babol University of Medical Science, Islamic Republic of Iran (grant number: 3904). There is no conflict of interest in this study.

## Authors' Contributions

S.G.A.J.; Contributed to conception and built an ideal design. R.K.; Contributed to all experimental work, wrote the manuscript, and also performed statistical analysis. A.A.M, N.N.; Contributed to pharmacological and chem ical approaches, respectively. F.F.; Contributed to histopa thology part of this study and revised the manuscript. All authors read and approved the final manuscript.

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