169

Nurulain SM et al. ANTIOXIDANTS AND ORGANOPHOSPHORUS POISONING Arh Hig Rada Toksikol 2013;64:169-177

Review

DOI: 10.2478/10004-1254-64-2013-2294

# ANTIOXIDANTS IN ORGANOPHOSPHORUS COMPOUNDS POISONING

Syed M NURULAIN<sup>1</sup>, Peter SZEGI<sup>2</sup>, Kornèlia TEKES<sup>2</sup>, and Syed NH NAQVI<sup>3</sup>

Department of Pharmacology and Therapeutic, Faculty of Medicine and Health Sciences, UAE University, AlAin UAE<sup>1</sup>, Department of Pharmacodynamics, Semmelweis University, Budapest, Hungary<sup>2</sup>, Department of Pharmacology, Baqai Institute of Pharmaceutical Sciences, Baqai Medical University, Karachi, Pakistan<sup>3</sup>

Received in January 2012 CrossChecked in August 2012 Accepted in August 2012

Oxidative stress has recently been implicated as a factor in the mortality and morbidity induced by organophosphorus (OP) compound poisoning. An overwhelming number of research papers are based on studying at the cellular and organ level. Such studies have concluded that antioxidants can be used as an adjunct compound in the treatment of both chronic as well as acute OP poisoning. Still, the role of antioxidants in reducing the mortality and morbidity induced by OP compounds has scarcely been verified, as well as their role as adjunct treatment compounds for both structurally and functionally different OP compounds. The present review of the literature was undertaken to establish the role of antioxidants in survival studies following acute exposure to OP compounds. The review found no substantial evidence that antioxidants demonstrate any positive effect following extremely toxic poisoning. However, for a more comprehensive and rational conclusion, further research needs to be conducted.

**KEY WORDS:** acute poisoning, oxidative stress, survival study

# Intoxication by organophosphorus compounds

The history of organophosphorus compounds (OP) and their poisonous effect stretches throughout more than a century. These compounds manifest their toxicity by irreversibly inhibiting the enzyme acetylcholinesterase at the nerve synapse. Despite decades-long research, mortality caused by acute OP poisoning continues to be high (1), while no new standard therapies are being introduced. There are over 150 different types of synthesized OPs, though their generalized structure is much the same. Each OP has a unique profile of toxicity and behaviour. For instance, death due to dichlorvos poisoning occurs very rapidly, while dimethoate toxicity takes several hours to develop (2), even though both belong to the same OP class. From the standpoint of chemistry, OP compounds comprise organophosphates, organophosphonates and organophosphinates, each of which is further divided into sub-groups. Other classifications of OPs are based on the lethality of a compound. According to the classification of the World Health Organisation (WHO) (3), Class Ia belongs to extremely toxic OPs, Ib are highly toxic, Class II comprises moderately toxic, whereas Class III consists of mildly toxic OP compounds. Some examples are shown in Table 1. In addition, there are also the deadly organophosphorus chemical weapons (OPWs), called nerve agents.

Acute organophosphate insecticide poisoning manifests itself through three different phases of toxicity; namely, acute cholinergic crisis, which occurs from within a few minutes to twenty-four hours;

Organophosphorus compounds (WHO's hazardous level)	Structures
Paraoxon- ethyl (Extremely hazardous; Class Ia)	$H_3C$ O O O O O O O O O O
Dichlorvos (Highly hazardous; Class Ib)	
Chlorpyrifos (Moderately hazardous; Class II)	CI CI CI CI CI CI CI CI CI CI CI CI CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CI CI CI CI CI CI CI CI CI CI CI CI CI
Malathion (Slightly hazardous; Class III)	$H_3C$ O $H_3C$ O $H_3C$ O $H_3C$ O O O O O O O O

Table 1 Structure of organophosphorus compounds belonging to different class of OP insecticides

intermediate syndrome (IMS), which sets in 48 h to 96 h after exposure; and delayed neuropathy (4). Acute cholinergic crises and IMS have been considered a major contributing factor in organophosphate-related morbidity and mortality because of their frequent occurrence and probable causative role in respiratory failure. Appropriate therapy leads to a complete recovery within 5 to 18 days (4).

# Standard and non-standard therapy

Standard therapeutic treatment of OP poisoning includes atropine, oximes, and benzodiazepines accompanied by supporting measures (5). The use of oximes has frequently been deemed controversial (6). Supporting measures include proper ventilation and the decontamination of skin and body parts by an alkali solution, by specific decontamination kits, etc. Petroianu (5) named the treatment of OP poisoning AFLOP; an abbreviation for atropine, fluid, oxygen, and pralidoxime (oxime). Atropine relieves muscarinic signs and symptoms, while oximes (pralidoxime/ obidoxime/HI-6, etc.) shorten the duration of respiratory muscle paralysis through acetylcholinesterase reactivation. Benzodiazepines are used to control OPinduced seizures. In warfare, alongside regular therapy, pre-treatment with pyridostigmine is recommended (7). Some of the non-regular antidotes include clonidine, fresh frozen plasma, magnesium sulphate (8), N-acetylcysteine (9), activated charcoal, milk and certain other home remedies (10, 11), but their effectiveness has not yet been sufficiently established (12). Other experimental approaches include the use of NMDA receptor antagonists such as gacyclidine (13), haemoperfusion (10) and the nanocarrier of magnetic magnesium (14). Non-regular antidotes for some reason usually do not receive attention from the scientific community, so the related scientific reports are negligible. Approaches such as the alkalization of blood plasma, use of weak inhibitors against strong inhibitors or use of bioscavengers are very popular but have not gained validity.

# Oxidative stress and use of antioxidants in OP poisoning

The imbalance between the production of free radicals and antioxidant defences in the body is called oxidative stress and has significant health implications. Oxidative stress is a major mechanism in the pathophysiology of several toxins and diseases. In addition, oxidative stress is also a process related to xenobiotic exposure and different levels of environmental contamination. It has recently been

Reference	<b>OP</b> compounds	Organ studied	
	Dichlorvos	0	
Srivastava and Shivanandappa 2011 (37)	(Highly toxic)	Rats brain	
	Ethion	Erythrocytes	
Bhatti et al., 2011 (38)	(Moderately toxic)		
Amara et al., 2011 (39)	Dimethoate	T inter	
	(Moderately toxic)	Liver	
Derived is and Flow 2011 (40)	DDVP and monocrotophos	Blood, brain,	
Dwivedia and Flora 2011 (40)	(Highly toxic)	liver	
Dirican and Kalender 2011 (41)	Dichlorvos	Testes	
Diricali and Kalender 2011 (41)	(Highly toxic)	Testes	
	Chlorpyrifos, malathion, methyl		
Ojha et al., 2011 (42)	parathion	Liver, brain, kidney, spleen	
	(Moderately/mildly/extremely toxic)		
Ehrich et al., 2011 (43)	Paraoxon and DDVP	Human neuroblastoma	
	Extremely/highly toxic	SH-SY5Y cells	
Lua et al., 2010 (44)	Omethoate	Liver	
בעת כו מו., 2010 (דד)	(Highly toxic)	Liver	
	Review: oxidative stress	Oxidative stress by OP leads to	
Lukaszewicz-Hussain 2010 (24)	and its role in toxicity of	organ damage.	
	organophosphate insecticides		
Cemek et al., 2010 (45)	Fenthion	Different rats tissues	
Centek et al., 2010 (45)	(Moderately toxic)	Different rats ussues	
Shah and Iqbal 2010 (46)	Diazinon	Kidney	
	(Moderately toxic)	Klulley	
Kose et al., 2010 (33)	Dichlorvos Blood and Cardiac muscle		
Kose et al., 2010 (55)	(Highly toxic)		
$U_{7}$ Uzun et al. 2010 (47)	Chlorpyryfos	Lung	
Uzun et al., 2010 (47)	(Moderately toxic)	Lung	
Kalender et al., 2010 (48)	Malathion	Liver	
Mansour and Mossa 2010 (49)	Chlorpyrifos	Suckling pups	
	(Moderately toxic)	Sucking pups	
Togun et al., 2010 (50)	Dichlorvos	Cardiac tissues	
	(Highly toxic)		

**Table 2** The oxidative stress in chronic/sub-chronic OP poisoning at organ level studies.

**Note** Only two reports are related to extremely toxic OPs, whereas the majority of the study covers moderately or highly toxic OP compounds

Reference	Type of study / article Tested compounds Organ studied	Conclusion
Dandapani et al., 2003 (17)	Clinical study	Severe and prolonged AChE inhibition is associated with oxidative stress and may contribute to the development and severity of intermediate syndrome.
Kovacic, 2003 (22)	Review	Toxic manifestations of OP are apparently due in part to oxidative stress.
Vidyasager et al., 2004 (18)	Clinical study	The increased level of MDA in OP poisoned patients who failed to survive was probably reflective of oxidative stress, but the patients who did survive after specific treatment did not show change in antioxidant status.
Sharma et al., 2005 (51)	Dimethoate Rat liver and brain	The organophosphate increases the generation of certain free radicals in the liver and brain by alterations to antioxidant status.
Ranjbar et al., 2005 (20)	Clinical study	Oxygen free radicals and their related interactions like lipid peroxidation are present in acute OP poisoning.
Cankayali et al., 2005 (28)	DDVP Rat serum	In addition to classic treatments, drugs with antioxidants might be promising in the treatment of OP poisoning.
Fortunato et al.,2006 (52)	Malathion Rat brain	Malathion induced oxidative stress and modulated SOD and CAT in selective brain regions.
Venkatesh et al., 2006 (35)	Clinical study	Occurrence of oxidative stress in severe acute OP poisoning was evident; however the development of type II paralysis is not associated with the level of oxidative stress.
Gunay et al., 2007 (32)	Survival study on rats DDVP	There was no evidence for increased oxidative stress due to DDVP.
Possamai et al., 2007 (19)	Malathion Rat organs, muscle and serum	Oxidative stress, particularly lipoperoxidation, is involved in OP toxicity.
Yurumez et al., 2007 (30)	Survival study on mice Fenthion	NAC used as an antioxidant improved the survival rate in mice.
Lukaszewicz-Hussain, 2008 (24)	Review OP insecticides	Supplementation with antioxidants may be beneficial in OP poisoning however the rat models do not completely reflect clinical trials.
Jiang et al., 2010 (53)	Methyl parathion Rat plasma and liver	It is also important to administer antioxidants in acute OP toxicity, in addition to standard therapy.
Zhang et al., 2010 (54)	Clinical study	Effective antioxidant therapy may be a therapeutic option following acute organophosphorus poisoning.
Kose et al., 2010 (33)	DDVP Serum Rat cardiac cells	Acute DDVP administration did not cause marked oxidative stress.
Rastogi et al., 2009 (55)	Epidemiological study	Pesticide sprayers exposed to insecticides including OP display more oxidative stress.
Hundekari et al., 2011 (56)	Clinical study	Antioxidant supplementation may be useful to reduce toxic effects in acute OP poisoning in addition to regular therapy.

 Table 3 Some of the references of acute OP poisoning and status of antioxidants.

postulated that OPs produce oxidative stress through the formation of reactive oxygen species (ROS) (15). ROS such as hydrogen peroxide, superoxide anions and hydroxyl radicals are produced in a number of

Name of OP compounds

Paraoxon-ethyl

Chlorfenvifos

Metasystox

Anilophos

Chlorpyrifos

Dimethoate

Diazinon

Ethion

Fenthion

Fenthion

Malathion

Sumithion (Fenitrothion)

Methidathion

Monocrotofos

DDVP

Parathion-methyl

OP cl

Ia; extrem

Ia; extrem

Ib; highl

Ib; highl

Ib; highl

Ib; highly toxic

Ib; highly toxic

II; Moderately toxic

III; Slightly toxic

studies (25).		Antioxidant
lass	Type of study	used in the study
ely toxic	Survival study	NAC, Glutathione (reduced)
ely toxic	Cellular/organ-level study	Not available for acute study
ly toxic	Cellular/organ-level study	Not available for acute study
ly toxic	Cellular/organ-level study	NAC
ly toxic	Cellular/organ-level study	Vitamin E

Table 4 The antioxidants investigated (last co survival and cellular & organ level s

Cellular/organ-level study

Survival study

Cellular/organ-level study

Cellular/organ-level study

DFP Not listed Cellular/organ-level study Vitamin E Menazon Not listed Cellular/organ-level study Not available for acute study Note It is evident from the table that in vivo survival studies were conducted only with paraoxon-ethyl and fenthion. All the other studies were carried out to evaluate the oxidative responses at cellular and organ level only, which may or may not have implications in pathophysiological conditions but cannot be correlated with mortality/morbidity. There is no correlation study

to predict the oxidative response by OP at cellular level and mortality/morbidity.

cellular reactions by enzymes such as lipoxygenases, peroxidases and dehydrogenases (16). ROS are part of the normal oxidative metabolism, but when produced in excess, they cause tissue injury. The role of oxygen-free radicals has been established in many chronic disorders, but their significance in acute conditions has not been given much attention.

During recent years, oxidative stress has been described as a co-lethal factor in OP-induced poisoning (17-21). Many reviews have stressed the role of oxidative stress in OP poisoning (15, 22-26). Bayrami et al. (27) found oxidative stress and acetylcholinesterase inhibition, along with many other parameters, in farmers chronically exposed to OP, but the name and class of OP exposure was not mentioned and the observed effect was asserted to be due to chronic exposure. Antioxidants have been suggested as adjunct to OP antidotes (28, Tables 2 to 4). One oxime has been reported to possess antioxidant property (29), but among the impressive volume of published articles, only two survival studies documented the benefits of antioxidants. Most of the studies were done

on moderately or highly toxic compounds. One paper published in 2007 by Yurumez et al. (30) determined the beneficial effect of N-acetylcysteine (NAC) in counteracting the organophosphate fenthion (a moderately toxic OP according to the classification of the WHO) in mice, and demonstrated that NAC has prophylactic as well as therapeutic activity in OP poisoning and clearly improves survival rates in mice. Pena-Llopis et al. (31) showed that NAC increased fish survival following exposure to lethal doses of dichlorvos. These papers described the effectiveness of the antioxidant NAC in acute poisoning. It is also possible that NAC improved the survival rate for another reason; it may have prevented lung-related pathological conditions such as shortness of breath or obstructive pulmonary conditions. Possami et al. (19) showed that the most sensitive targets of oxidative damage after acute treatment with malathion (a mild toxic OP) were the kidneys, lungs and diaphragm, as well as the liver, quadriceps and serum after subchronic treatment. Moreover, mortality by OP intoxication is mainly caused by respiratory obstruction,

Vitamin C and E

Not available for acute study

Not available for acute study

Vitamin C, E, melatonin Vitamin C, E, theophylline,

pentoxifylline

Vitamin C and E

Not available for acute study

Vitamin C, E, NAC, melatonin

NAC

Not available for acute study

Vitamin E, Ginger, ZnCl.

while most of the mortality in acute OP poisoning results from acute respiratory failure due to central respiratory depression, respiratory muscle weakness, and/or direct pulmonary effects (bronchospasm and bronchorrhea) (2). NAC may also have other roles which can contribute to improving the survival of animals. The protective effect of NAC may only extend to experimental OP fenthion and not to other structurally different OPs. In an acute study on rats, Gunay et al. (32) reported no evidence of oxidative stress due to dichlorvos. Kose et al., (33) concluded that acute dichlorvos administration does not cause marked oxidative stress and probably does not play a major role in dichlorvos-induced poisoning. Nurulain et al. (34) found no survival effect for glutathione reduced (GSH) and NAC with an acute dose of paraoxon (extremely hazardous OP) induced intoxication in Wistar rats. It was evident that NAC and GSH had a negative effect, instead of being protective (unpublished data). In a clinical trial, Venkatesh et al. (35) corroborated that the in-hospital morbidity and mortality of OP poisonings are mostly associated with type II paralysis (intermediate syndrome) and the development of type II paralysis is not associated with the level of oxidative stress. However, there was an early occurrence of oxidative stress in severe acute OP poisoning. This shows that an antioxidant has no role at phase II of acute toxicity. Indirect evidence of oxidative stress in IMS has been reported in a review by Abdollahi and Karami-Mohajeri (36). Other antioxidants used for OP-induced toxicity include vitamin C and E, date palm, etc. (Table 4), but where mixed results were noted, the outcomes describe only the cellular-level damage and biochemical estimation of oxidative stress parameters.

# SUMMARY

- 1. It is evident from the literature that oxidative stress occurs in acute organophosphorus poisoning, but there is no convincing evidence that antioxidants may prevent mortality in acute OP poisoning.
- 2. Whether all antioxidants are beneficial or merely NAC in certain acute OP poisonings is unclear.
- 3. Can antioxidants be effective for all classes of OPs? This is an issue that has been completely ignored.
- 4. Can antioxidants be effective for all phases of acute toxicity? This issue has also been completely ignored.

- 5. The use of antioxidants might be effective (speculation) in long-term pathophysiological conditions induced by OP compounds through chronic or sub-chronic exposures, or even in the delayed phase of acute OP poisoning.
- 6. Since antioxidants might not be useful for survival, their inclusion into standard therapy cannot be warranted.

# Acknowedgement

Sincere thanks to Prof. Huba Kalasz from the Semmelweis University in Budapest, Hungary for help and encouragement in preparing the manuscript.

# REFERENCES

- Eddleston M, Eyer P, Worek F, Mohamed F, Senarathna L, von Meyer L, Juszczak E, Hittarage A, Azhar S, Dissanayake W, Sheriff MH, Szinicz L, Dawson AH, Buckley NA. Differences between organophosphorus insecticides in human self-poisoning: a prospective cohort study. Lancet 2005;366:1452-9.
- Rosenbaum C, Bird SB. Non-muscarinic therapeutic targets for acute organophosphorus poisoning. J Med Toxicol 2010;6:408-12.
- The WHO Recommended Classification of Pesticides by Hazard and Guidelines to Classification 2009 [displayed 6 September 2012]. Available at http://www.inchem.org/ documents/pds/pdsother/class\_2009.pdf
- 4. Yang CC, Deng JF. Intermediate syndrome following organophosphate insecticide poisoning. J Chin Med Assoc 2007;70:467-72.
- 5. Petroianu GA. Poisoning with organophosphorus compounds (OPC): Mythology vs Reality. The Middle East J Emer Med 2007;6:3-8.
- 6. Rahimi R, Nikfar S, Abdollahi M. Increased morbidity and mortality in acute human organophosphate-poisoned patients treated by oximes: a metaanalysis of clinical trials. Hum Exp Toxicol 2006;25:157-62.
- Wetherell J, Price M, Mumford H, Armstrong S, Scott L. Development of next generation medical countermeasures to nerve agent poisoning. Toxicology 2007;233:120-7.
- Pajoumand A, Shadnia S, Rezaie A, Abdi M, Abdollahi M. Benefits of magnesium sulfate in the management of acute human poisoning by organophosphorus insecticides. Hum Exp Toxicol 2004;23:565-9.
- 9. Shadnia S, Ashrafivand S, Mostafalou S, Abdollahi M. Nacetylcysteine a novel treatment for acute human organophosphate poisoning. Int J Pharmacol 2011;7:732-5.
- Peter JV, Moran JL, Graham PL. Advances in the management of organophosphate poisoning. Expert Opin Pharmacother 2007;8:1451-64.
- Balali-Mood M, Saber H. Recent advances in the treatment of organophosphorus poisonings. Iran J Med Sci 2012;37:74-91.
- 12. Blain PG. Organophosphorus poisoning (acute). Clin Evid (Online) 2011;2102.

- 13. Lallement G, Baubichon D, Clarençon D, Galonnier M, Peoc'h M, Carpentier P. Review of the value of gacyclidine (GK-11) as adjuvant medication to conventional treatments of organophosphate poisoning: primate experiments mimicking various scenarios of military or terrorist attack by soman. Neurotoxicology 1999;20:675-84.
- 14. Mohammadi H, Karimi G, Seyed Mahdi Rezayat, Ahmad Reza Dehpour, Shafiee H, Nikfar S, Baeeri M, Sabzevari O, Abdollahi M. Benefit of nanocarrier of magnetic magnesium in rat malathion-induced toxicity and cardiac failure using non-invasive monitoring of electrocardiogram and blood pressure. Toxicol Ind Health 2011;27:417-29.
- Abdollahi M, Ranjbar A, Shadnia S, Nikfar S, Rezaie A. Pesticides and oxidative stress: a review. Med Sci Monit 2004;10:RA141-7.
- Halliwell B, Gutteridge JMC. Free Radicals in Biology and Medicine. 3<sup>rd</sup> ed. Oxford: Oxford University Press; 1999.
- Dandapani M, Zachariah A, Kavitha MR, Jeyaseelan L, Oommen A. Oxidative damage in intermediate syndrome of acute organophosphorus poisoning. Indian J Med Res 2003;117:253-9.
- Vidyasagar J, Karunakar N, Reddy MS, Rajnarayana K, Surender T, Krishna DR. Oxidative stress and antioxidant status in acute organophosphorus insecticide poisoning. Indian J Pharmacol 2004;36:76-9.
- Possamai FP, Fortunato JJ, Agostinho J, Quevedo D, Wilhelm FF, Dal-Pizzol F. Oxidative stress after acute and sub-chronic malathion intoxication in Wistar rats. Environ Toxicol Pharmacol 2007;23:198-204.
- Ranjbar A, Solhi H, Mashayekhi FJ, Susanabdi A, Rezaie A, Abdollahi M. Oxidative stress in acute human poisoning with organophosphorus insecticides; a case control study. Environ Toxicol Pharmacol 2005;20:88-91.
- Shadnia S, Dasgar M, Taghikhani S, Mohammadirad A, Khorasani R, Abdollahi M. Protective effects of alphatocopherol and N-acetylcysteine on diazinon-induced oxidative stress and acetylcholinesterase inhibition in rats. Toxicol Mech Methods 2007;17:109-15.
- 22. Kovacic P. Mechanism of organophosphates (nerve gases and pesticides) and antidotes: electron transfer and oxidative stress. Curr Med Chem 2003;10:2705-9.
- Peña-Llopis S. Antioxidants as potentially safe antidotes for organophosphorus poisoning. Curr Enzyme Inhib 2005;1:147-56.
- Lukaszewicz-Hussain A. Role of oxidative stress in organophosphate insecticide toxicity – Short review. Pest Biochem Physiol 2010;98:145-50.
- 25. Soltaninejad K, Abdollahi M. Current opinion on the science of organophosphate pesticides and toxic stress: a systemic review. Med Sci Monit 2009;15:RA75-90.
- Karami-Mohajeri S, Abdollahi M. Toxic influence of organophosphate, carbamate, and organochlorine pesticides on cellular metabolism of lipids, proteins, and carbohydrates: a systematic review. Hum Exp Toxicol 2011;30:1119-40.
- Bayrami M, Hashemi T, Malekirad AA, Ashayeri H, Faraji F, Abdollahi M. Electroencephalogram, cognitive state, psychological disorders, clinical symptom, and oxidative stress in horticulture farmers exposed to organophosphate pesticides. Toxicol Indl Health 2012;28:90-6.
- 28. Cankayali I, Demirag K, Eris O, Ersoz B, Moral AR. The effects of N-acetylcysteine on oxidative stress in

organophosphate poisoning model. Adv Ther 2005;22:107-16.

- Puntel GO, de Carvalho NR, Gubert P, Palma AS, Dalla Corte CL, Avila DS, Pereira ME, Carratu VS, Bresolin L, da Rocha JB, Soares FA. Butane-2, 3-dionethiosemicarbazone: an oxime with antioxidant properties. Chem Biol Interact 2009;177:153-60.
- Yurumez Y, Cemek M, Yavuz Y, Birdane YO, Buyukokuroglu ME. Beneficial effect of N-acetylcysteine against organophosphate toxicity in mice. Biol Pharm Bull 2007;30:490-4.
- Peña-Llopis S, Ferrando MD, Peña JB. Fish tolerance to organophosphate-induced oxidative stress is dependent on the glutathione metabolism and enhanced by N-acetylcysteine. Aquat Toxicol 2003;65:337-60.
- Gunay N, Kose A, Tarakcioglu M, Gunay NE, Demiryurek AT. Evaluation of cardiac oxidative stress parameters and mortality in a rat model of organophosphate poisoning. J Emerg Med 2007;33:338.
- Kose A, Gunay N, Kose B, Ocak AR, Erel O, Demiryurek AT. Effects of atropine and pralidoxime pretreatment on serum and cardiac oxidative stress parameters in acute dichlorvos toxicity in rats. Pest Biochem Physiol 2010;97:249-55.
- Nurulain SM, Kalasz H, Adem A, Petroianu G. Antioxidant has no role in survival of rats after intoxication with acute dose of paraoxon. In: 5<sup>th</sup> International Online Medical Conference (IOMC 2012); 3-11 March 2012; Abstract 147.
- 35. Venkatesh S, Kavitha ML, Zachariah A, Oommen A. Progression of type I to type II paralysis in acute organophosphorus poisoning: is oxidative stress significant? Arch Toxicol 2006;80:354-61.
- 36. Abdollahi M, Karami-Mohajeri S. A comprehensive review on experimental and clinical findings in intermediate syndrome caused by organophosphorus poisoning. Toxicol Appl Pharmacol 2012;258:309-14.
- Srivastava A, Shivanandappa T. Differential cholinesterase inhibition in the rat brain regions by dichlorvos and protective effect of Decalepis hamiltonii roots. Neurotoxicology 2011;32:931-4.
- Bhatti GK, Bhatti JS, Kiran R, Sandhir R. Biochemical and morphological perturbations in rat erythrocytes exposed to ethion: protective effect of vitamin E. Cell Mol Biol (Noisyle-grand) 2011;57:70-9.
- Amara IB, Soudani N, Troudi A, Bouaziz H, Boudawara T, Zeghal N. Antioxidant effect of vitamin E and selenium on hepatotoxicity induced by dimethoate in female adult rats. Ecotoxicol Environ Saf 2011;74:811-9.
- 40. Dwivedia N, Flora SJS. Concomitant exposure to arsenic and organophosphates on tissue oxidative stress in rats. Food Chem Toxicol 2011;49:1152-9.
- 41. Dirican EK, Kalender Y. Dichlorvos-induced testicular toxicity in male rats and the protective role of vitamins C and E. Exp Toxicol Pathol 2012;64:821-30.
- 42. Ojha A, Yaduvanshi SK, Pant SC, Lomash V, Srivastava N. Evaluation of DNA damage and cytotoxicity induced by three commonly used organophosphate pesticides individually and in mixture, in rat tissues. Environ Toxicol 2011; doi: 10.1002/tox.20748 [Epub ahead of print].
- 43. Ehrich M, Van Tassell R, Li Y, Zhou Z, Kepley CL. Fullerene antioxidants decrease organophosphate-induced

acetylcholinesterase inhibition in vitro. Toxicol In Vitro 2011;25:301-17.

- 44. Lua L, Xiuli Wangb X, Liwei Langa L, Fu F. Protective effect of reduced glutathione on the liver injury induced by acute omethoate poisoning. Environ Toxicol Pharmacol 2010;30:279-83.
- 45. Cemek M, Emin BM, Yürümez Y, Yavuz Y, Aslan A, Büyükben A, Aymelek F. Tissue trace and major element levels in organophosphate insecticide fenthion (Lebaycid) toxicity in rats: prophylactic and therapeutic effect of exogenous melatonin. Ecotoxicol Environ Saf 2010;73:206-12.
- Shah MD, Iqbal M. Diazinon-induced oxidative stress and renal dysfunction in rats. Food Chem Toxicol 2010;48:3345-53.
- Uzun FG, Demir F, Kalender S, Bas H, Kalender Y. Protective effect of catechin and quercetin on chlorpyrifosinduced lung toxicity in male rats. Food Chem Toxicol 2010;48:1714-20.
- Kalender S, Uzun FG, Durak D, Demir F, Kalender Y. Malathion-induced hepatotoxicity in rats: the effects of vitamins C and E. Food Chem Toxicol 2010;48:633-8.
- 49. Mansour SA, Mossa AH. Adverse effects of lactational exposure to chlorpyrifos in suckling rats. Hum Exp Toxicol 2010;29:77-92.
- Togun N, Kose A, Gunay N, Tarakcioglu M, Demiryurek AT. Formulation of effects of atropine, pralidoxime and magnesium sulfate on cardiac tissue levels of nitric oxide,

malondialdehyde and glutathione in organophosphate poisoning using artificial neural network. Comput Biol Med 2010;40:29-36.

- Sharma Y, Bashir S, Irshad M, Gupta SD, Dogra TD. Effects of acute dimethoate administration on antioxidant status of liver and brain of experimental rats. Toxicology 2005;206:49-57.
- 52. Fortunato JJ, Feier G, Vitali AM, Petronilho FC, Dal-Pizzol F, Quevedo J. Malathion-induced oxidative stress in rat brain regions. Neurochem Res 2006;31:671-8.
- 53. Jiang N, Lu L, Wang T, Zhang L, Xin W, Fu F. Reduced glutathione attenuates liver injury induced by methyl parathion in rats. Toxicol Mech Methods 2010;20:69-74.
- 54. Zhang JW, Lv GC, Zhao Y. The significance of the measurement of serum xanthenes oxidase and oxidation markers in patients with acute organophosphorus pesticide poisoning. J Int Med Res 2010;38:458-65.
- 55. Rastogi SK, Satyanarayan PVV, Ravishankar D, Tripathi S. A study on oxidative stress and antioxidant status of agricultural workers exposed to organophosphorus insecticides during spraying. Indian J Occup Environ Med 2009;13:131-4.
- 56. Hundekari IA, Suryakar AN, Rathi DB. Oxidative stress and antioxidant status in acute organophosphorus pesticides poisoning cases of North Karnataka (India). J Environ Health Res 2011;11:39-44.

#### Sažetak

#### ANTIOKSIDANSI I TROVANJE ORGANOFOSFORNIM SPOJEVIMA

Oksidacijski stres u novije je vrijeme označen kao faktor pri mortalitetu i morbiditetu uzrokovanom trovanjem organofosfornim spojevima. Sve veći broj studija zasnovan je na proučavanju na razini stanice i organa i takve su studije većinom zaključile da se antioksidansi mogu rabiti kao dodatne tvari pri liječenju kroničnog, ali i akutnog trovanja organofosfornim spojevima. No uloga antioksidansâ u smanjenju mortaliteta i morbiditeta izazvanog trovanjem organofosfornim spojevima još nije u dovoljnoj mjeri potvrđena. Štoviše, funkcija antioksidansâ kao dodatnih tvari pri liječenju i dalje je uvelike nerazjašnjena za strukturalno i funkcionalno različite vrste organofosfornih spojeva. Ovaj pregledni rad napisan je s namjerom određivanja uloge antioksidansâ u studijama preživljavanja zbog akutne izloženosti organofosfornim spojevima. Pregledom se nije utvrdio nijedan čvršći dokaz da antioksidansi imaju pozitivan učinak nakon ekstremno toksičnog trovanja. Međutim za sveobuhvatniji i racionalniji zaključak nužno je dalje proučavanje.

KLJUČNE RIJEČI: akutno trovanje, oksidacijski stres, studija preživljavanja

CORRESPONDING AUTHOR:

Dr Syed M. Nurulain P.O. Box 17666, AlAin, UAE E-mail: nurulain@uaeu.ac.ae