Presence of organochlorine pesticides and characterization of biomarkers in wild mice living in crop fields

MOISES ANDRADE HERRERA¹, GRISELDA ESCALONA SEGURA¹, MAURICIO GONZÁLEZ JÁUREGUI², RAFAEL REYNA HURTADO¹, JORGE A. VARGAS CONTRERAS³, JAIME RENDÓN VON OSTEN^{4*}

¹ El Colegio de la Frontera Sur. Unidad Campeche. Avenida Rancho Polígono 2-A, Ciudad Industrial, CP. 24500, Lerma. San Francisco de Campeche. Campeche, México. Email: <u>moandrade@ecosur.edu.mx</u> (MAH), <u>gescalon@ecosur.mx</u> (GES), <u>rafaelcalakmul@gmail.com</u> (RRH).
 ² Instituto de Ecología. Carretera antigua a Coatepec 351, El Haya, CP. 91070, Xalapa. Veracruz, México. Email: <u>mauglezj@gmail.com</u> (MGJ).
 ³ Facultad de Ciencias Químico Biológicas. Universidad Autónoma de Campeche. Avenida Agustín Melgar S/N entre calle 20 y Juan de la

Barrera, Buenavista, CP. 24039, San Francisco de Campeche. Campeche, México. Email: <u>jalbino64@hotmail.com</u> (JAVC). ⁴ Instituto EPOMEX. Universidad Autónoma de Campeche. Campus VI. Av. Héroe de Nacozari 480. CP. 24070, San Francisco de

Campeche. Campeche, México. Email: jarendon@uacam.mx (JRvO).

* Corresponding Author.

The use of pesticides in crops bordering conservation areas poses risks for wildlife incidentally exposed; its effects in the Yucatan Peninsula, Mexico, are still unknown. Wild mice that inhabit farming land play a key ecological role, and can also be used as bioindicators of wildlife exposure to pollutants. The objectives of this work were to determine the presence of organochlorine pesticides (OC) in liver and evaluate the seasonal response of enzymatic biomarkers (BM) such as acetylcholinesterase (AChE), glutathione-S-transferase (GST), and catalase (CAT). Wild mice (*Mus musculus*) were captured between June 2015 and April 2016 in a watermelon crop of a rural community in Quintana Roo, Mexico. Individual mice were sacrificed *in situ*, followed by tissue dissection (liver, brain and skeletal muscle). Pesticides were determined by gas chromatography; BM activity was estimated by spectrophotometry. We captured 35 individuals, with a capture success of 2.33%. The prevailing OCs detected were drines in both climatic seasons. The rainy season influenced the activity of biomarkers to a greater extent, since AChE showed a lower activity (16 % and 40 % in brain and liver, respectively). GST was activated during the same season (77 % higher), while CAT did not show significant differences between seasons. There was no significant correlation between OC concentrations and biomarker activity, except for drines and AChE in the brain. OC concentrations recorded in the present work are below (2- and 20-fold lower) those reported in other works on rodents under controlled conditions. BM activity suggests that rainfall seem to exacerbate the effects of pesticides on mice; however, it seemingly does not pose a risk for their survival. The use of wild mice as bioindicators is a valuable and practical tool to detect disturbances derived from the use of pesticides in agricultural areas. Further research is recommended using a broader BM battery to identify those pollutants with the most severe effect on the physiology of wild

El uso de plaguicidas en cultivos aledaños a áreas de conservación, significa un riesgo para la fauna silvestre expuesta accidentalmente cuyos efectos son desconocidos, particularmente en la Península de Yucatán, México. Los ratones silvestres que habitan las regiones agrícolas, desempeñan un papel ecológico importante, además pueden servir como bioindicador de exposición a contaminantes. Los objetivos del presente trabajo fueron determinar la presencia de plaguicidas organoclorados (OC) en hígado y evaluar la exposición a partir de la respuesta temporal de los biomarcadores enzimáticos (BM), tales como acetilcolinesterasa (AChE), glutatión-S-transferasa (GST) y catalasa (CAT). Ratones silvestres (Mus musculus) fueron capturados entre junio de 2015 y abril de 2016 en un cultivo de sandía de una comunidad agrícola, en Quintana Roo, México. Los individuos fueron sacrificados in situ y se le diseccionaron los tejidos (hígado, cerebro y músculo esquelético). La presencia de plaguicidas se determinó por cromatografía de gases. La actividad de los BM se determinó por espectofotometría. Se capturaron 35 individuos, con un éxito de captura de 2.33%. Los OC más detectados fueron los drines en ambas temporadas climáticas. La temporada lluviosa influyó más en la actividad de los biomarcadores, ya que la AChE mostró menor actividad (16 y 40 % en cerebro y músculo respectivamente), la GST aumentó su actividad para la misma temporada (77 % mayor), y finalmente la CAT no mostró diferencias significativas entre temporadas. No hubo correlación significativa entre las concentraciones de OC y la actividad de BM, a excepción de los drines y la AChE en cerebro. Las concentraciones de OC encontradas en el presente trabajo, están por debajo (entre 2 y 20 veces) de las reportadas en otros trabajos con roedores en condiciones controladas. La actividad de los BM refleja que las lluvias parecen acentuar los efectos de los plaguicidas sobre los ratones, sin embargo, puede no representar un escenario de riesgo para su sobrevivencia. La utilización de ratones silvestres como bioindicadores, es una herramienta útil y práctica para vislumbrar disturbios ocasionados por el uso de plaquicidas en zonas agrícolas. Se recomiendan trabajos complementarios y ampliar la batería de BM, para poder determinar con mayor precisión cuales son los contaminantes que afectan en mayor medida la fisiología de los animales silvestres expuestos accidentalmente.

Keywords: Acetylcholineterase; catalase; glutathione-S-transferase; biomarkers; Mus musculus; organochlorine pesticides; wildlife

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Introduction

Technology-dependent agriculture (hereinafter called tech agriculture) involving the use of high amounts of pesticides poses a risk to wildlife that comes into contact with these toxic compounds, which may cause still unknown health effects. The continued application of pesticides and other agrochemicals to crop fields causes acidification, loss of soil biodiversity, and soil toxicity (<u>Givaudan *et al.* 2014</u>). Pesticides not only affect the target pest species, but also impact wildlife living in and around agricultural fields (<u>Johnston</u>,

2000). The damage caused by pesticides on exposed organisms depend mainly on their toxicokinetics (uptake route and distribution) and toxicodynamics (effects on physiology and metabolism) and is influenced by external factors such as pesticide nature, chemical composition, persistence, dose, and time of exposure, as well as by the health status and susceptibility of the exposed organism (<u>Ramírez and</u> <u>Lacasana 2001</u>). The main characteristics of organochlorine pesticides include high persistence, mobility across great distances from their source of application, bioaccumulation, and long-term adverse effects (<u>Mackay et al. 2001</u>).

In this sense, the vertebrate group with the most severe adverse effects from incidental poisoning in the wild documented to date (Köhler and Triebskorn 2013), since they are deemed highly sensitive to pesticide exposure. In contrast, a group that has received little attention as to harm caused by pesticides is rodents, mainly because these are also considered as pests; however, this neglects the important ecological roles they play, mainly regarding seed dispersal and control of other pests such as insects and other invertebrates, besides being preyed by higher predators (Ceballos 2005).

In general, small mammals have been used as proxies to validate human exposure models by various environmental protection organizations. For example, the USEPA (United States Environmental Protection Agency) has commonly used rodents as indicators of human and environmental protection (Sheffield et al. 2001). Particularly, murine models have been used in the characterization of various effects of pesticides under controlled conditions. These comprise the evaluation of tolerance thresholds by determining the lethal dose 50 (LD_{so}) — the dose that kills 50% of the test population (Dell'Omo et al. 2003; Hirano et al. 2017) and adverse effects by endocrine disruption (Bhaskar and Mohanty 2014; Ghodrat et al. 2014), oxidative stress (Zhao et al. 2016; Morales-Prieto and Abril 2017; Latchoumycandane and Mathur 2002), genotoxic effects (Peris-Sampedro et al. 2016), or neurotoxic effects (Chen et al. 2016; Dell'Omo and Shore 1996; Westlake et al. 1983). All these studies have involved the application of various doses of one or more (mixtures) pesticides.

However, studies that consider mice as bioindicators of environmental conditions for wildlife are scarce. Population dynamics of rodents has been used as an indicator of disturbance in polluted areas. These studies have focused mainly on accidental deaths and population declines caused by acute exposure to organophosphate pesticides (OF) and carbamates (CB) (Barrett and Darnell 1967; Giles 1970; Sheffield *et al.* 2001). For example Block *et al.* (1999) assessed the long-term population impact of the pesticide OF Counter^{*} (terbufos), based on population parameters in two wild mice species (*Peromyscus maniculatus* and *P. leucopus*) in an agricultural field in Iowa, USA, and concluded that OF has no apparent influence on reproduction and other population parameters that would affect their long-term demography. Other studies conducted in free-living mice evaluated the accumulation of persistent organic pollutants (POPs) in tissues (<u>Perez-Gonzalez et al. 2017</u>), or the effect of various pollutants at the histological level (<u>Gomez-Ugalde 2003</u>), or with a biochemical approach by analyzing enzymatic biomarkers (BM; <u>Andrade-Herrera 2011; Chi Coyoc et al. 2016</u>).

In Mexico, studies reporting the influence of crop pesticides on wild rodents are scarce. Specifically in the Yucatan peninsula, these studies focus on the presence and accumulation of pesticides (Rendon-von Osten et al. 2005; Charruau et al. 2013), and relatively few studies report harmful effects to the fauna surrounding farming areas where pesticides are used (Noreña-Barroso et al. 2004; Cobos et al. 2006; Buenfil-Rojas et al. 2016). Accordingly, in the state of Quintana Roo, particularly in the Maya area, the main tech crops are sugar cane, Maradol papaya and watermelon (SAGARPA 2015). These involve the use of various technological packages based on the use of agrochemicals to achieve high crop yields. For example, Endosulfan 35 Le Cag[®] (endosulfan), Furadan® (carbofuran), Herbipol® (2.4-D Amine) and Cerillo® (paraguat), Lannate[®] (methomyl) or Velfosato[®] (glyphosate) were recorded during field trips as the main pesticides used. It was noted that these pesticides are applied in excessive amounts, and farmers apply them disregarding the safety requirements (personal protection equipment).

Therefore, the lack of studies in the region analyzing the effects of tech agriculture on wildlife health, and in particular as regards small mammals, made it necessary to generate knowledge to provide evidence about the potential risks for these animals. Thus, the aim of the present study was to analyze the influence of the pesticides used in a tech watermelon crop on wild mice (*Mus musculus*). This was done by first determining the presence of pesticides in liver tissue of individuals captured, followed by characterizing the toxic effects through enzymatic biomarkers (AChE in brain and muscle; GST and CAT in liver), and finally by exploring the relationship between the pesticides detected and the activity of biomarkers.

Materials and Methods

Field Work. The sampling site is located at coordinates 19° 35' 24.70" N, -89° 10' 51.20" W, in the Municipality of José María Morelos, Quintana Roo, in a tech watermelon crop within the Mayan community named *X noh Cruz* (Figure 1). The area is surrounded by rainfed maize fields and forest patches, and is adjacent to the *Bala'Ka'ax* Wildlife Protection Area (APFFBK).

Specimens were captured between June 2015 and April 2016 using 50 Sherman[®] traps placed every five meters on both sides along a 125 m transect across the crop field. Traps were baited with a mixture of oats and vanilla and left overnight. The individuals captured were identified using field guides, and standard morphometric measurements were recorded. Then, these individuals were killed *in situ* by cervical dislocation according to NOM-033-SAG/ZOO- 2014, which stipulates the humane killing of domestic and wild animals; brain, liver and skeletal muscle samples were taken



Figure 1. Sampling area, located in coordinates 19° 35′ 24.70″ N, -89° 10′ 51.20″ W, within the Mayan community of Xnoh Cruz, municipality of José María Morelos, Quintana Roo, which is adjacent to the *Bala'an 'k'aax* APFF.

for subsequent analysis. Tissues were stored in liquid nitrogen, transported to the laboratory, and stored at -80 °C.

Laboratory work. Laboratory work was divided into two phases: the first determined the presence of organochlorine pesticides (OC) in mouse liver tissue; the second characterized the effect of exposure to the various xenobiotics by assessing three biomarker enzymes: acetylcholinesterase (AChE) that reflects neurotoxicity, glutathione-S-transferase (GST) that indicates detoxification processes (phase II biotransformation), and catalase (CAT), an oxidative stress biomarker.

Determination of Pesticides in Liver Tissue. High-purity solvents were used (98 % HPLC); glass material was washed with Extran[®] and rinsed with distilled water, then dried for 4 hours at 200 °C and rinsed with acetone and hexane.

The analyses of pesticides in liver tissue of mice were performed according to the method described by Zhang et al. (2007). A fraction of liver tissue was collected, ovendried for 24 h at 45 °C, and weighed for subsequent analysis. Any chemicals in samples were extracted with a mixture of acetone:hexane (1:1) in a sonicator for 20 min. After extraction, the amount of fat was determined to estimate the concentrations of pesticides on fat basis ($\mu q/q$ of fat). Samples were purified through a glass column packed with 7 g of silica gel and 1 g of sodium sulfate. Afterwards, samples were eluted with 20 ml of hexane, 20 ml of a 1:1 mixture of dichloromethane and hexane, and 20 ml of dichloromethane; then, samples were immediately evaporated to dryness and resuspended in 50 µl of hexane for testing through gas chromatography. Pollutants were quantified using a Varian 3800 gas chromatograph with a Ni⁶³ electron capture detector (GC-ECD) and equipped with a HT8 capillary column (60 m x 0.25 mm; 25 µm film thickness; SGE Analytical Science, USA). Injector and detector temperatures were 150 °C and 350 °C, respectively. Pollutant concentrations were calculated through the area under the curve with the software Star Chromatography Workstation

version 6, and through the calibration of standards. Pesticides were identified and quantified using a mixture of 20 OC pesticide standards, CRM No. SUPELCO[®] 47426 EPA CLP Organochlorine Pesticide Mix, and the Sigma-Aldrich standards Mirex No. 45887, 2.4'-DDE No. 36663, 2.4'-DDD No. 35485 and 2.4'-DDT No. 45839, which were pooled into families for testing (Table 1).

Activity of enzymatic biomarkers (BM). The three BM were analyzed in specific tissues according to their target organ, with AChE tested in brain and skeletal muscle, while GST and CAT were analyzed in liver. For the three BM, all samples were homogenized for 20 sec. in a homogenizer (PRO Sci-

 Table 1. Organochlorine (OC) pesticides analyzed in liver tissue of Mus musculus; LD

 = limit of detection, LR = Limit of Report

Isomer and/or metabolites	LD	Group	LR		
analyzed	μg/g	Gloup	μg/g		
αHCH	0.003				
βΗCΗ	0.004	нсна	0.007		
χHCH	0.003	TICHS	0.007		
δΗCΗ	0.002				
Aldrin	0.004				
Dieldrin	0.004				
Endrin	0.009	DRINES	0.0018		
Endrin aldehyde	0.010				
Endrin ketone	0.005				
Endosulfan I (alpha)	0.005				
Endosulfan II (beta)	0.006	Endosulfans	0.007		
Endosulfan sulfate	0.005				
o,p' DDE	0.005				
p,p' DDE	0.004				
o,p' DDD	0.031	DLL	0.001		
p,p' DDD	0.025	Diad	0.001		
o,p' DDT	0.013				
p,p' DDT	0.010				
Trans Chlordane	0.004	Chloritan	0.000		
Cis Chlordane	0.004	Chlordanes	0.009		
Methoxychlor	0.018	Methoxychlor	0.01		
Mirex	0.006	Mirex	0.006		
Heptachlor	0.004	Hontachlar	0.012		
Heptachlor epoxide	0.004	Heptachior	0.013		

entific 250[®]), using the specific buffer for each BM according to the method used. AChE activity was determined as per the method of Ellman et al. (1961), adapted to microplate (Guilhermino et al. 1996); samples were prepared with a 0.1 M potassium phosphate buffer, pH 7.2. Enzyme kinetics was recorded at 414 nm and expressed in units (U) of hydrolyzed acetylcholine per minute per mg of protein using an extinction coefficient of 13.6 x 10³ M⁻¹ cm⁻¹. GST activity was determined according to the method described by Habig et al. (1974). A 0.1 M solution of potassium phosphate buffer, pH 6.5, was used for sample preparation, and enzyme kinetics was read at 340 nm, expressing activity as U of S-(2.4-dinitrophenyl) glutathione per minute per milligram of protein, using an absorption coefficient of 9.6 x 10³ M⁻¹ cm⁻¹. CAT activity was measured through the decrease in H₂O₂ at 240 nm using the method by <u>Aebi (1984)</u> for example, methanol, ethanol, formic acid, phenols, with the consumption of 1 mol of peroxide (peroxide activity, using a 50 mM solution of potassium phosphate buffer of pH 7, with activity expressed as U of hydrogen peroxide produced per minute per milligram of protein. Protein concentration was determined by the method of <u>Bradford (1976)</u> at 595 nm, expressed as mg/ml. In all cases, activity was determined using microplates (Corning[®]) in a spectrophotometer (Thermo Scientific Multiskan Spectrum[®]).

Statistical analyses. All analyzes were performed using the software R version 3.2.4 (<u>R Core Team, 2016</u>). Compliance of BM activity with the assumption of a normal distribution was tested using a Shapiro-Wilk test, and differences were evaluated by comparing BM activity between climatic seasons (raint-dry), using a Student's *t*-test when the data were normally distributed, or with a Wilcoxon test otherwise (Table 2); the significance level used was $\alpha \le 0.05$ (Zar, 2010). Subsequently, the measures of central tendency and dispersion of pesticides were determined with the package NADA 1.6-1 (Lopaka 2013), taking into account left-censored data according to the chromatograph detection limit.

Table 2. Average activity of BMs per climatic season, with values expressed as U/ min/mg protein. Also shown are the statistical *t* values (parametric), W (non-parametric); SD = standard deviation; max = maximum value; min = minimum value; DF = degrees of freedom; *p* = statistical probability; *< 0.05, **< 0.01, ***<0.01.

Season	n	AChE in brain											
		Mean	SD	Max	Min	t	W	DF	р	α = 0.05			
Rainy	17	16.81	4.55	26.86	7.43	2 0752		21.00	0.046	*			
Dry	17	20.07	4.63	29	1255	-2.0752	-	31.99	0.046				
		AChE in	muscle										
Rainy	17	5.50	4.32	15.42	0.87	2 1050		20.40	0.004	**			
Dry	18	9.07	1.68	12.43	6.01	-3.1639	-	20.49	0.004				
		GST											
Rainy	17	98.36	74.80	208.60	15.75		252		0.001	***			
Dry	18	30.53	9.48	52.60	10.63	-	233	-	0.001				
		CAT											
Rainy	17	6.15	5.76	17.99	1.19		127		0.404				
Dry	18	6.24	3.94	15.72	1.32	-	127	-	0.404	-			

The "vegan" package (Oksanen et al. 2017) was used to perform correlations between the concentrations of pesticides detected (classified by chemical family) in liver and the BM analyzed. These correlations were explored using rank-transformed data (Spearman correlations), as concentration data were left-censored by the limits of detection and quantification (Helsel 2012).

Results and Discussion

This work reports the concentrations of pesticides in liver tissue (on fat, dry weight, and wet weight bases), as well as their potential effects characterized from the expression of three enzymatic BMs in three different wild-mice tissues. Adult individuals of *Mus musculus* were chosen as target species, because this was the dominant mouse species captured in sampling, as well as for its abundance in the study area, being a natural inhabitant in maize crop fields that surround watermelon crops (Alcérreca *et al.* 2009). The remaining animals were not deposited in any biological collection, given the poor condition of skin and skeleton (mainly the skull) after dissections. A total of 35 individuals were captured during field work (17 in the wet season and 18 in the dry season), over 30 nights of trapping, with a 2.33 % capture success.

Presence of OC Pesticides in Liver Tissue. Of the 24 OC pesticides analyzed, those most frequently detected in the wet season belong to the "drines" family (aldrin, endrin, endrin aldehyde, endrin ketone, and dieldrin), which were detected in 10 individuals, followed by hexachlorocyclohexanes (HCHs), detected in 7 specimens; in contrast, the least detected pesticide was methoxychlor, found in just two individuals. In the dry season, drines were also the pesticides most frequently detected in the 18 individuals analyzed, along with derivatives of dichloro-diphenyl-trichloroethane (DDT) and heptachlorine, with each detected in eight individuals; similar to the wet season, methoxychlor was the least detected pesticide, identified in only three specimens (Table 4).

On the other hand, the chemicals found in higher concentrations in both seasons were DDT isomers and methoxychlor, with peak concentrations of 702 and 524 μ g/g fat, respectively, both in individuals caught in the dry season. The chemical recorded in the lowest concentration was chlordane in both seasons, with a mean concentration of 6.63 ± 11.3 μ g/g fat and 15.02 ± 29.3 μ g/g fat in the wet and dry seasons, respectively.

The concentrations found in this study are below the exposure-related toxicity thresholds (mainly *via* ingestion) reported in various investigations on wild rodents under controlled laboratory conditions (Jefferies *et al.* 1973; Morris 1968; Sheffield *et al.* 2001; Wolfe and Esher 1980), which are 2- to 20-fold higher versus those reported here. These works were conducted primarily to set lethal doses and tolerances to various OC including drines, HCHs and DDTs; these works report both physiological and behavioral impairment, but only some report OC content in tissues.

A fact worth considering is that bioaccumulation in wild mammals is mainly driven by ingestion, although pollutants may also be absorbed through the skin or by inhalation (Linder and Joermann 2001). In this regard, the work on wild rodents in free-living conditions focus mainly on evaluating variations in population size or recording presence/absence in areas where OC pesticides have been used (Sheffield et al. 2001). The few studies that quantify the presence of OC in tissues have been carried out primarily to identify DDT, dieldrin and heptachlor residues. In the case of DDT, the values found in mice in this study (between 0.001 and 1.834 μ g/g dry weight) are similar to those reported by Lincer and Sherburne (1974) wet weight, ranging between 0.01 and 0.13 µg/g dry weight. In addition, Laubscher et al. (1971), in crop fields in Tucson, Arizona, found DDT and its metabolites at concentrations between 0.006 to 0.92 µg/g wet weight in liver of Peromyscus eremicus and P. man**Table 4.** Summary of pollutant concentrations (µg/g) in the liver of *Mus musculus*. OC= Organochlorine pesticide; (D= number of data detected; ND= number of data not detected; SD = standard deviation.

	Weight Fat															
	Rainy									Dry						
OC	D	ND	ND %	Min	Max	Mean	SD	D	ND	ND %	Min	Max	Mean	SD		
HCHs	7	10	58.820	0.007	82.213	10.340	28.370	5	13	72.222	0.007	303.116	23.370	75.940		
DRINES	10	7	41.176	0.002	135.532	21.690	43.936	8	10	55.555	0.002	279.350	22.691	66.579		
ENDOSULFAN	4	13	76.470	0.007	89.165	23.127	27.844	5	13	72.222	0.007	195.783	18.027	48.257		
DDTs	4	13	76.470	0.010	303.812	39.522	111.434	8	10	55.555	0.010	702.315	45.947	170.208		
CHLORDANES	9	8	47.058	0.009	32.109	6.633	11.318	6	12	66.666	0.009	120.856	15.023	29.304		
HEPTACHLOR	5	21	70.588	0.013	54.537	10.606	17.970	8	10	55.555	0.013	227.825	18.473	54.433		
METHOXYCHLOR	2	15	88.235	0.010	232.528	232.530	0.010	3	15	83.333	0.010	524.625	60.177	137.962		

Wet Weight

	Rainy							Dry						
OC	D	ND	ND %	Min	Max	Mean	SD	D	ND	ND %	Min	Max	Mean	SD
HCHs	7	10	58.820	0.005	0.023	0.009	0.007	5	13	72.222	0.004	49.930	2.770	12.780
DRINES	7	10	58.820	0.002	0.042	0.009	0.013	8	10	55.555	0.002	0.079	0.012	0.024
ENDOSULFAN	4	13	76.470	0.007	0.031	0.028	0.001	5	13	72.222	0.007	0.080	0.015	0.022
DDTs	4	13	76.470	0.001	0.094	0.019	0.031	8	10	55.555	0.001	0.200	0.021	0.057
CHLORDANES	9	8	47.058	0.004	0.357	0.028	0.087	6	12	66.666	0.003	0.064	0.010	0.016
HEPTACHLOR	5	21	70.588	0.003	0.017	0.007	0.009	8	10	55.555	0.003	0.065	0.009	0.014
METHOXYCHLOR	2	15	88.235	0.010	0.072	0.072	0.000	3	15	83.333	0.008	0.148	0.016	0.040

Dry Weight

				Rain	у						Dry			
OC	D	ND	ND %	Min	Max	Mean	SD	D	ND	ND %	Min	Max	Mean	SD
HCHs	7	10	58.820	0.007	0.035	0.011	0.009	5	13	72.222	0.007	0.125	0.019	0.029
DRINES	7	10	58.820	0.002	0.057	0.012	0.018	8	10	55.555	0.002	0.943	0.066	0.229
ENDOSULFAN	4	13	76.470	0.007	0.037	0.037	0.001	5	13	72.222	0.007	1.040	0.070	0.263
DDT	4	13	76.470	0.001	0.127	0.025	0.043	8	10	55.555	0.001	1.834	0.122	0.449
CHLORDANES	9	8	47.058	0.006	0.765	0.054	0.189	6	12	66.666	0.009	0.821	0.060	0.202
HEPTACHLOR	5	21	70.588	0.004	0.023	0.008	0.010	8	10	55.555	0.004	0.094	0.013	0.021
METHOXYCHLOR	2	15	88.235	0.010	0.097	0.097	0.000	3	15	83.333	0.010	0.216	0.028	0.056

iculatus; these levels are up to five-fold higher than those recorded in this study (0.0001 to 0.200 μ g/g wet weight).

However, the concentrations found here are far lower than those reported by <u>Chi Coyoc et al. (2016)</u> for DDTs, HCHs and drines (0.876, 0.546, and 0.451 µg/g dry weight basis respectively) in liver of wild rodents (*Oryzomys couesi, Peromyscus leucopus* and *Reithrodontomys gracilis*). These concentrations were up to 35 (DDT), 49 (HCHs) and 37 (Drines) times higher than those recorded in this study. These results suggest that the concentrations observed in this case may not pose a risk for the rodents exposed; however, the effects of the various pollutants are influenced by a number of factors — both environmental and physiological intrinsic to each species and even to each individual (<u>Peakall and McBee 2001</u>).

Finally, in general terms, there is a higher concentration of all OCs in the dry season than in the rainy season (Table 4). This could be due to the fact that rain leads to the partial leaching of toxic compounds to greater soil depths, thus reducing the exposure of organisms that live at the surface (González Valdez *et al.* 2013). An additional factor is the clayey silt soil in the sampling area (Andrade Herrera *et al.* 2018), which favors the leaching of pollutants given its low organic matter content, which adsobs OCs and makes them more persistent in upper soil layers (Cárdenas and Márquez 2015). Furthermore, factors associated with temporality may also account for the higher concentration of OCs in the dry season; for example, this coincides with the winter season, during which organisms move less due to the low temperatures (most reproductive periods occur in the rainy season), so that fat content of organisms increases (Caldas *et al.* 1999).

Activity of Enzymatic Biomarkers. As regards BMs, significant differences were found between climatic seasons in AChE and GST, while CAT did not record these differences (Table 2). AChE was determined in both brain and muscle, registering a higher mean activity (18.32 U/min/mg protein) in the brain relative to the muscle (7.35 U/min/mg protein), representing a difference slightly exceeding two orders of magnitude between the two tissues. However, both tissues show the same pattern, *i.e.* the activity recorded was higher in the dry season vs. the rainy season (Figure 2). In the brain, AChE showed an inhibition slightly above 16 % ($t_{(32)} = -2.075$, p < 0.05) in the wet season; in the muscle, this inhibition was about 40 % ($t_{(20)} = -3.18$, p < 0.01), also in the rainy season (Table 2).



Figure 2. Activity of BMs in tissues of *Mus musculus* captured in agricultural fields. Data were compared between climatic seasons; AChE was evaluated in (A) brain and (B) muscle, while GST (C) and CAT (D) were analyzed in liver. Results are shown as median values, quartiles (box); activity is expressed as U/minute/mg protein.

AChE is considered to be the best BM for assessing exposure to organophosphate and carbamate pesticides — the main anticholinergic pesticides. However, their characteristics (poorly persistent in the environment, highly soluble in water) make it difficult to quantify them in both the environment and animal tissues (Sheffield *et al.* 2001). Rattner and Hoffman (1984) report the inhibition of AChE after the administration of sublethal oral doses of the organophosphate (OF) insecticide acephate to wild mice (*M. musculus, P. leucopus* and *M. pennsylvanicus*); inhibition rates were similar to those reported in this study (16 to 40 %) at doses between 25 mg/kg and 100 mg/kg of food for 5 days. In addition, the inhibition rates recorded in our results are seemingly harmless for the mouse populations, as in other investigations an inhibition rate of more than 40 % was found, which was not lethal in exposed mice (<u>Dell'Omo and</u> <u>Shore 1996; Meyers and Wolff 1994</u>).

Studies based on field work that characterize the inhibition of AChE in wild rodents are scarce. In this sense, our results on AChE inhibition (16 to 40 %) are consistent with some studies carried out in the wild in other rodent species such as *Microtus pennsylvanicus* (Jett 1986), *M. pinetorum* (Durda *et al.* 1989), *P. maniculatus* and *P. leucopus* (Block *et al.* 1999), *M. musculus* (Edward *et al.* 1983; Custer *et al.* 1985), and particularly in Mexico in *O. couesi* and *R. gracilis* (Chi Coyoc *et al.* 2016). In these works, it is concluded that the degree of inhibition found did not cause lethal acute poisoning, and population stability was not affected.

In the present work, AChE activity decreases during the wet season, when the main crops are maize, squash, and beans, involving the use of herbicides such as glyphosate or paraquat in soil preparation, in addition to insecticides for the protection of crops, including carbofuran, endosulfan or imidacloprid. The data recorded point to a higher exposure to anticholinergic chemicals in the rainy season, which, although not analyzed here, were reported by farmers as used for pest control.

Various xenobiotics cause oxidative stress from the production of reactive oxygen species (Limon-Pacheco and Gonsebatt 2009). To avoid oxidative stress, cells activate detoxification processes through antioxidants in order to reduce toxic effects, involving the two enzymes used as BMs, GST (phase II of xenobiotic metabolism) and CAT (antioxidant), *i. e.*, these increase their activity when exposed to toxic compounds (van der Oost *et al.* 2003). GST activity in the mice captured exhibited significant differences between climatic seasons (W = 253, *p* < 0.001), showing a higher activity in the wet vs. dry season (98.36 and 30.53 U/ min/mg protein, respectively), representing a difference of nearly 77 % in BM expression between both seasons (Table 2, Figure 2).

When GST is activated, it acts on toxic chemicals by increasing their water solubility, thereby facilitating their excretion from the body (van der Oost *et al.* 2003). The influence of pesticides on GST in mice in agricultural areas is poorly documented. A study characterized its activation in agricultural workers after exposure to carbamates and γ HCH (Lukaszewicz-Hussain 2010); the latter is one of the pesticides found in tissues of some mice in this work. In this regard, GST activity increased significantly in the rainy season, suggesting that rainfall leads to the release of some xenobiotics trapped in soil, leading to an increased reaction of detoxification enzymes such as GST. A similar pattern was found in wild mice (*Peromyscus* spp.) in forested areas of Mexico City where GST activity was significantly higher during the rainy season (Andrade-Herrera 2011).

Contrary to all the BMs discussed above, CAT yielded no significant differences between the dry and rainy seasons (6.15 and 6.24 U/min/mg protein, respectively). However, the average values recorded in this work are higher than those reported in other (experimental) studies, where mean activity ranges from 0.2 to 2.5 U/min/mg protein (Ayed-Boussema *et al.* 2012; Panemangalore and Baby 2000; Yilmaz *et al.* 2004). In this regard, catalase is a biomarker that can react in different ways, *i. e.*, inhibition or activation, depending on various factors, mainly the dose of the xenobiotic, time of exposure, or a combination of various compounds (van der Oost *et al.* 2003).

In laboratory experiments, dermal exposure to various pesticides leads to increased CAT activity. For example, Panemangalore and Baby (2000) recorded a 13 % increase in activity relative to its controls after 4 weeks of exposure to a mixture of OFs. Similar effects were found in other studies where rats and mice were exposed to a variety of pesticides (OF, OC, pyrethroids and neonicotinoids, among others) via ingestion, with average increases in activity of 25 % (Jin et al. 2011, 2013). When pesticides are administered intraperitoneally, both effects occur. For example, Ayed-Boussema et al. (2012), recorded increased activity levels of up to three orders of magnitude in mice after the administration of multiple doses of dimethoate for 30 days; in contrast, <u>Yilmaz et al. (2004)</u> found a significant CAT inhibition after administering a systemic hormone herbicide for three days in albino mice (Mus musculus).

Correlation between Pesticides and BM Activity. Spearman's correlations were performed between the OC pesticides detected (per family) and the activity of BMs recorded in mice to explore the influence of these pollutants in the activity of these BM in exposed organisms. The analyses showed a single significant correlation (rho = -0.489, p < -0.489) 0.01), between drine pesticides and AChE activity in brain tissue (Table 3). In this sense, although exposure to OC pesticides causes multiple toxic effects on organisms, AChE inhibition is caused primarily by exposure to OF and CB pesticides (Rodríguez-Castellanos and Sánchez-Hernández 2007), which were not analyzed in the present work. However, farmers mentioned the use of mixtures of various insecticides and acaricides, including organophosphates, pyrethroids, carbamates, and neonicotinoids, particularly in watermelon crops; in addition, empty containers of these pesticides were found in agricultural fields, suggesting that these pollutants (or their mixtures) may account for the results observed.

In Mexico, few works report the effect of xenobiotic agents in wild rodents. For example, <u>Andrade-Herrera</u> (2011) correlated the concentrations of air pollutants in Mexico City with the activity of enzymatic biomarkers, finding a greater impact in the dry season, likely due to thermal inversions, as airborne pollutants may cause adverse effects. In contrast, this study suggests that rainfall seemingly affects to a greater extent the physiology of mice, as AChE and GST were particularly affected in the wet season. Rainfall carries and deposits toxic chemicals in soil, hence increasing their availability for rodent living in the local environment.

Table 3. Spearman's correlations between concentrations of organochlorine pesticides and characterization of the activity of the BMs analyzed in mice. Figures in bold represent significance at $p \le 0.05$.

	AChE in brain		AChE in	muscle	GS	т	CAT		
	Rho	р	Rho	р	Rho	р	Rho	р	
HCHs	0.036	0.842	-0.014	0.937	0.043	0.805	-0.260	0.131	
DRINES	-0.489	0.003	-0.084	0.630	0.118	0.501	0.172	0.323	
ENDOSULFAN	-0.240	0.172	-0.259	0.133	-0.036	0.839	-0.079	0.653	
Dldd	-0.093	0.601	0.022	0.899	0.008	0.965	0.061	0.730	
Chlordanes	-0.392	0.022	-0.260	0.131	0.062	0.724	0.010	0.954	
Heptachlor	-0.186	0.293	-0.119	0.497	-0.058	0.741	0.033	0.849	
Methoxychlor	-0.192	0.277	-0.068	0.697	0.099	0.573	-0.072	0.680	

Conclusions

This work is the first in Mexico to show the presence of pesticides and their biochemical effects (neurotoxicity and defense response mechanisms and antioxidants) in wild mice. The study shows that mice bioaccumulate pesticides in their tissues, which poses a risk to wildlife living in agricultural fields. The use of BMs is a valuable tool for determining sublethal hazard in wildlife due to exposure to pollutants. In this work, the activity of BMs shows a higher influence of pesticides in the rainy season, likely due to the presence of various pesticides in the environment that are deposited in soil with rain. Also, the use of wild mice as bioindicators is suitable for identifying effects associated with the use of pesticides in farming areas. For future studies, it is recommended to use blood as a non-destructive matrix, besides to the fact that blood contains chemicals that were recently ingested. Other types of pesticides should also be analyzed, in addition to broadening the battery of biomarkers used, to determine other potential adverse effects and, in turn, determine in further detail which pollutants are the major disruptors of the physiology of animals exposed, as the indiscriminate use of synthetic pesticides is likely to impact the surrounding wildlife.

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