Acute Toxicity of Insecticide Thiamethoxam to Crayfish (Astacus leptodactylus): Alterations in Oxidative Stress Markers, ATPases and Cholinesterase

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Abstract
Thiamethoxam (Thmx) is a globally used neonicotinoid pesticide contaminated in freshwater ecosystems with residues detected in fishery products. Astacus leptodactylus is a popular freshwater crustacean that is cultivated and exported in many countries. In this study, we investigated the acute toxic effects of Thmx on A. leptodactylus using various biomarkers (acetylcholinesterase, carboxylesterase, glutathione S-transferase, glutathione, superoxide dismutase, glutathione peroxidase, glutathione reductase, and adenosinetriphatases). The 96-h LC50 value of Thmx was calculated as 8.95 mg active ingredient L–1. As the dose of Thmx increased, oxidative stress was induced by the inhibition/activation of antioxidant enzymes, while the activities of acetylcholinesterase, carboxylesterase and adenosinetriphatases were inhibited. As a result, it can be said that Thmx has highly toxic effects on crayfish, therefore they are under threat in the areas where this pesticide is used.

Keywords: Acetylcholinesterase; Antioxidant enzymes; Crustacean; Insecticide; Metabolic enzymes; Toxicity

1. Introduction
Among the insecticides widely used in agriculture, it is necessary to focus on neonicotinoids, which are chemically similar to nicotine.1 Neonicotinoid insecticides have been the fastest growing insecticide class due to their safe use of biochemical properties, broad spectrum activities, and systemic distribution mechanism in plants.2,3 Thiamethoxam (Thmx) 3-(2-chloro-1,3-thiazol-5-yilmethyl)-5-methyl-1,3,5-oxadizinan-4-yldene (nitro) amine is one of the second generation neonicotinoid insecticides and is used against a wide target population of insects.4 Thmx is a potential pollutant that is mixed with surface and ground water due to its low absorption from the soil, high leakage capacity and high water solubility.5 Thmx, like other neonicotinoid insecticides, bind agonistically with high affinity to nicotinic acetylcholine receptors, which are target sites in insects.6 There is much information in the literature specific to the exposure profiles of neonicotinoids in aquatic ecosystems, but there is little information about second-generation neonicotinoids such as Thmx in published studies on the effects of neonicotinoids on non-target aquatic organisms. Knowing the effect of neonicotinoids on aquatic invertebrates provides important data for aquatic risk assessment.7 Although low-risk for some non-target organisms, Thmx is a potential pollutant for surface and groundwater due to its low absorption, low infiltration, high water solubility and resistance to biological treatment, therefore it poses a danger to aquatic organisms.8,9 Thmx has been found to be generally around 0.001–225 ppb in surface waters.10 The persistence in the soil (229 days) and high-water solubility (4100 mg L–1) of Thmx mean there is high potential to be transported into surface waters.11 The results of a comprehensive review of laboratory and semi-field microcosm studies show that aquatic invertebrates are highly susceptible to neonicoti-
noids. However, the most studied of neonicotinoids in aquatic ecosystems is imidacloprid, the effects of a newer neonicotinoid, Thmx, on aquatic organisms have been less studied.

Turkey’s natural freshwater crayfish species, *A. leptodactylus*, one of the most popular species in Europe is due to the presence of a wide range of areas outside of Anatolia and economic importance. Crayfish are part of the ecological balance in their natural freshwater areas. Due to the important role they play in the processing of all kinds of organic materials, they are active on energy balances in the ecosystem, therefore they are seen as key species for still and fluvial habitats. Indicator species in aquatic ecosystems are considered to be a suitable way of demonstrating environmental quality. Not all organisms are suitable for use as an indicator. Crayfish are benthic, solitary, constantly in contact with objects, omnivorous, long-lived, slow-moving, narrow habitat, large enough to easily sample from different body tissues, and can accumulate pollutants increases its value as an indicator species.

Many xenobiotics, including pesticides, can trigger the production of reactive oxygen species by various biochemical mechanisms, such as disruption of electron transport across the cell membrane, facilitation of the Fenton reaction, inactivation of antioxidant enzymes, and depletion of free radical scavengers. Antioxidant defense systems have been developed in organisms to scavenge these reactive oxygen species, and by evaluating the activation / inhibition level of these antioxidant systems, the oxidative damage caused by xenobiotics to the organism is estimated. The aim of this study was to investigate the acute toxic effects of Thmx on *A. leptodactylus*. For this, we tested the effect of different doses of Thmx on the enzymes responsible for ion homeostasis in the cell (Na+/K+ -ATPase, Mg2+ -ATPase, Ca2+ -ATPase), neurotoxicity biomarker acetylcholinesterase (AChE), antioxidant defense system parameters [superoxide dismutase (SOD), glutathione (GSH), glutathione peroxidase (GPx), glutathione reductase (GR)], an oxidative damage marker [malondialdehyde (MDA)], phase II biotransformation enzymes [glutathione S-transferase (GST), carboxylesterase (CaE)] of aquatic invertebrate crayfish *A. leptodactylus*.

2. Materials and Methods

2.1. Test Animals and Experimental Design

Crayfish used in this study were obtained from the Crayfish Breeding Unit at the Firat University Fisheries Faculty, Elazığ, Turkey. During the study, glass aquariums with a capacity of 30 liters with tubular shelters were used. Studies were done at room temperature (23 ± 1 °C) and in natural daylight (12 h dark /12 h light). Adequate ventilation was provided with the air pump. Rested tap water was placed in the aquariums. Before applying the pesticide, the crayfish were adapted to the laboratory environment for 15 days. Matured crayfish were used regardless of their gender. In order to achieve standardization, crayfish weighing around 20 ± 5 g were preferred. Crayfish were not given food during the applications. The pesticide sold under the trade name Actara 25 WG was obtained from Syngenta. The Active Ingredient (AI) of Thmx is 240 g L⁻¹. Water prepared according to ASTM standards was used in the study. Stock solution of 5000 mg L⁻¹ was prepared freshly by dissolving Thmx in tap water. Test waters containing Thmx solution were left in the containers with static renewal every 24 hours and the pH values of these waters were recorded daily. A total of five groups were formed, four of which were the pesticide-treated groups and one was the non-pesticide-applied group (control). Four crayfish were placed in each aquarium and the study was done in three replicates, so fifteen aquariums and sixty animals were used in total.

2.2. Determination of LC50 Values and Application Concentrations of Thiamethoxam

Dose ranges of 0.50–400 mg L⁻¹ of the commercial stock solution were used to determine the 96-h LC50 value of Thmx. Among live animals, those who were immobilized over time and showed signs of death were considered dead. The number of dead animals was recorded at 24, 48, 72 and 96 hours and accordingly 96h-LC50 was determined as 8.95 mg AI L⁻¹ using SPSS 24 probit. This determined LC20 dose and its three sub-doses of Thmx (LC50/2, LC50/4, LC50/8) were administered to the crayfish. No Thmx application was applied to the control group. The experiment was repeated three times for each group of four animals (N = 12). After applying solutions containing Thmx at its own concentration to each group for 96 hours, the animals were sacrificed and the hepatopancreas, muscle and gill tissues were removed and stored at −80°C until analyzed. An ice bath was used for anesthesia of the animals, and the abdominal areas of the animals between the thorax and tail were dissected.

2.3. Biochemical Assays

Analysis of biochemical markers was performed in tissues of surviving animals after a 96-h acute toxicity test. The numbers of animals by groups are as follows: Control: 12, LC50/8: 12, LC50/4: 12, LC50/2: 9, LC50: 8. Homogenisation of the tissues was carried out in homogenization buffer (0.1 M, pH 7.4 in potassium phosphate buffer; 0.15 M KCl, 1 mM EDTA, 1 mM DTT) and on ice using a polytron homogenizer (Heidolph RZ 2021 Germany). The homogenates were centrifuged at 16,000 × g for 20 min at 4°C (Hettich 460 R). Total protein and all enzyme readings were done in triplicate on a microplate reader (Thermo Varioscan Flash 2000). The total protein level was meas-
ured according to Bradford method (1976). The protein levels of the samples were determined using the standard curve constructed from measurements of the following bovine serum albumin standard solutions. In hepatopancreas tissue, GST, GR, AChE, CarE, GPx, SOD, GSH and MDA analyses were performed. ATPases analyses were done in gill and muscle tissues. All enzyme activities were expressed as specific activity (nmol min⁻¹ mg protein⁻¹).

2. 3. 1. Cellular Redox Status

The GST activity was determined by a spectrophotometric method according to protocol described by Habig et al. (1974) using CDNB as substrate. The change in absorbance was measured at 344 nm for 2 min. The GR activity was detected according to Cribb et al. (1989) by microplate assay with modifications. The reaction was initiated by the addition of GSSG into the reaction solution. Due to formation of GSH from GSSG, the decrease in the amount of DTNB was monitored at 405 nm for 3 min. The CarE activity was determined according to a modified procedure of Santhoshkumar and Shivanandappa (1999) for a microplate reader. The reaction was initiated by the addition of PNPA as substrate to the reaction solution. The liberated p-nitrophenol was monitored at 405 nm for 2 min. In the determination of GPx activity, the method developed by Bell et al. (1985). Based on using hydrogen peroxide (H₂O₂) as substrate and sodium azide (NaN₃) as catalase inhibitor, was used. The specific activity value of the enzyme was calculated based on the change in absorbance at 340 nm based on the oxidation of NADPH in a microplate reader. Superoxide dismutase (SOD) activity was determined by the method (Sun et al., 1988) based on subtracting the Mg²⁺ ATPase (containing Ouabain) activity from the total ATPase (without Ouabain) activity. The Mg²⁺ ATPase activity arises from the inhibition of Ouabain's activity by binding to Na⁺/K⁺ ATPase. Ca²⁺ ATPase activity was calculated by subtracting the enzyme activity measured in the absence of enzyme activity in the presence of CaCl₂.

2. 3. 2. Neurotoxicity (AChE)

The AChE activity was determined following the Ellman and Andres (1961) method using ACTI as a substrate, modified for the microplate reader by Ozmen et al. (1998). Enzyme activity was monitored at 412 nm for 1 min.

2. 3. 3. Ion Transport

The methods of Atlı and Canlı (2011) were used to determine ATPase activities (Na⁺/K⁺ -ATPase, Mg²⁺-ATPase, Ca²⁺-ATPase ) in gill and muscle. Analyzes were performed in a microplate reader in triplicate. 5 µL of sample and 60 µL of incubation medium consisting of 1 mM ouabain, 40 mM Tris-HCl, 4 mM MgCl₂, 20 mM KCl and 100 mM NaCl were pipetted into each microplate well and incubated at 37 °C for 5 minutes. 10 µL of 3 mM ATP was added to the top of the mixture in these wells and incubated at 37 °C for 30 minutes, so the reaction was initiated. After incubation, 35 µL of cold distilled water (+4 °C) was added to these wells to stop the reaction. The value of the inorganic phosphate (Pi) released from ATP at the end of the reaction was calculated by measuring the absorbance at 390 nm of the yellow compound formed by the main reagent consisting of polyoxyethylene 10 lauryl ether and ammonium molybdate (Atkinson et al. 1973). 190 µL of main reagent was added to microplate wells containing 60 µL of incubation medium, 5 µL of supernatant and 35 µL of cold distilled water, and after incubating at room temperature for 10 minutes, absorbance values were measured at 390 nm. The results were evaluated based on the standard curve obtained using different concentrations of KH₂PO₄ solution. Enzyme activities were expressed as specific activity (µmol P₁ min⁻¹ mg protein⁻¹). Na⁺/K⁺ ATPase activity was calculated by subtracting the Mg²⁺ ATPase (containing Ouabain) activity from the total ATPase (without Ouabain) activity. The Mg²⁺ ATPase activity arises from the inhibition of Ouabain's activity by binding to Na⁺/K⁺ ATPase. Ca²⁺ ATPase activity was calculated by subtracting the enzyme activity measured in the absence of enzyme activity in the presence of CaCl₂.

2. 4. LC-MS/MS Analysis of Thiamethoxam in the Test Water

The actual Thmx concentrations in the test waters were determined using a liquid chromatography tandem mass spectrometry (LC-MS/MS) in Adiyaman University Central Research Laboratory. The retention time of Thmx was approximately 3.84 min. The calibration curve constructed from the standards for the calculation of Thmx concentrations was in the range of 1–100 µg L⁻¹. The limits of detection, quantification, and coefficient of determination (r²) were determined as 0.07 µg L⁻¹, 0.32 µg L⁻¹, and 0.999, respectively. Thmx was detected through the transitions 292.1 → 211.0 mass-to-charge ratio (m/z) (collision energy (CE); –24 V) and 292.1 → 181.0 m/z, CE; –24 V. The Thmx standard was purchased from Dr. Ehrenstorfer GmbH with 99.8% purity. Each water sample was analyzed in triplicate.

2. 5. Data Analyses

In the statistical analysis of the data, computer software package SPSS 22 was used. Data normality was evaluated using Shapiro-Wilk test (p < 0.05). Kruskal Wallis
test was used to determine the comparison of data between groups. Mann Whitney U test was used to determine whether there was a significant difference within the groups. The statistical significance level was based on \( p < 0.05 \).

The integrated biomarker response (IBR) was used to incorporate all the biochemical marker reactions assessed into a single overall stress index to determine the risk potential of thiamethoxam. The IBR indexes were calculated according to the method defined by Arzate-Cárdenas and Martínez-Jerónimo (2012). The IBR index was calculated based on the mean and standard deviation for each biomarker. The average value for each response was standardized separately using the formula \( Y = (X_m) / SD \); where \( Y \) is the standardized value, \( X \) is the average value, and \( m \) is the average of the biochemical markers. Depending on the biochemical responses, \( Z \) values were calculated as \( Z = Y \) (inhibition) or \( Z = -Y \) (activation). Score (\( S \)) was evaluated with the formula \( S = |\min| + Z \); where \( |\min| \) is the absolute value of the minimum of all biochemical markers. The scores were utilized were \( [(S_1 \times S_2) / 2 + (S_2 \times S_3) / 2 + \ldots (S_{n-1} \times S_n) / 2] \) to give a normalized IBR, and estimated values were divided by the number of biochemical markers calculated.

3. Results and Discussion

3.1. The Actual Thiamethoxam Concentrations in the Test Waters

Data on the actual concentrations of Thmx in solutions applied to crayfish as determined by LCMSMS are shown in Table 1. A difference of approximately 15%, 12%, 10% and 11% was found between the nominal and actual concentrations, respectively. These differences may be because Thmx is not sufficiently soluble in water due to surfactants, solvents, and preservatives found in this commercial form (Korkmaz et al. 2018).

| Nominal Dose | N | Mean ± SE |
|--------------|---|-----------|
| 1.12         | 3 | 0.95 ± 0.03 |
| 2.24         | 3 | 1.97 ± 0.04 |
| 4.48         | 3 | 4.01 ± 0.06 |
| 8.95         | 3 | 7.98 ± 0.04 |

3.2. Acute Toxicity Assay

In our search, and to the best of our knowledge, no peerreviewed studies examining Thmx toxicity to \( A. leptodactylus \) have been published. In our study, the 96-hour acute lethal concentration value (96 h-LC\(_{50}\)) of Thmx for \( A. leptodactylus \) was determined as 8.95 mg AI L\(^{-1}\). The 96 h LC\(_{50}\) value for crayfish, \( Procambarus clarkii \) was determined as 0.967 mg AI L\(^{-1}\) by Barbee and Stout (2009) and 10 mg AI L\(^{-1}\) by Maloney et al. (2018) in two separate studies. In a study, 48-h LC\(_{50}\) value of Thmx for water louse \( Asellus aquaticus \) was found as 2.3 mg L\(^{-1}\). For crustacean \( Gammarus kischineffensis \), the 96-h LC\(_{50}\) value of Thmx determined as 8.985 mg L\(^{-1}\) and 3.751 mg L\(^{-1}\). The reason that these acute LC\(_{50}\) values of Thmx determined for crustaceans differ from each other may be due to the differences in the experimental conditions and the parameters such as application period, physiological status, life stage, age and body weight of the animals used in the experiment.

3.3. Mortality Rates of Crayfish Determined During 96-h of Study

The mortality rates of crayfish exposed to Thmx at different concentrations for 24, 48, 72 and 96-h are shown in Table 2. No death was observed at any Thmx concentration at 24th hour. At 48th hour, only 1 death was observed for each of the LC\(_{50}/2\) and LC\(_{50}\) doses. At 72th hour, 1 animal died at the LC\(_{50}/2\) dose and 2 animals died at the LC\(_{50}\) dose. At 96th hour, 1 animal died at both the LC\(_{50}/2\) and LC\(_{50}\) doses. Mortality rates were 25% and 33% at the LC\(_{50}/2\) and LC\(_{50}\) doses, respectively and the difference between these groups from control was statistically significant (\( p < 0.05 \)).

Table 2. The mortality of crayfish exposed to Thmx at different concentrations for 24, 48, 72 and 96-h.

| Concentration (mg AI L\(^{-1}\)) | N  | 24 h | 48 h | 72 h | Mortality 96 h | Total death | Mortality rate (%) |
|---------------------------------|----|------|------|------|---------------|-------------|-------------------|
| Control                         | 12 | 0    | 0    | 0    | 0             | 0           | 0                 |
| LC\(_{50}/4\)                   | 12 | 0    | 0    | 0    | 0             | 0           | 0                 |
| LC\(_{50}/2\)                   | 12 | 0    | 1    | 1    | 1             | 3           | 25*               |
| LC\(_{50}\)                     | 12 | 0    | 1    | 2    | 1             | 4           | 33*               |

3.4. Biochemical Responses

The data of the biomarkers evaluated in the hepatopancreas are given in Table 3, those in the gill in Table 4, and those in the muscle in Table 5.
3. 4. 1. Cellular Redox Status

In GST activity, there were significant increases in all Thmx concentrations, not dependent on Thmx concentration increase compared to the control group. The highest increase in GST activity was seen in the group in which the LC_{50}/4 dose was applied. The GST activity value at the LC_{50} concentration was close to that of the LC_{50}/4 concentration. Contrary to our study, Han et al. (2016)\(^{43}\) observed a significant increase in GST activity in the liver of zebra fish treated with azoxystrobin for 4 weeks and attributed this increase to the free radical scavenging effect of GST. Husak et al. (2017)\(^{44}\) found that when they applied fenconazole to goldfish, the GST activity in their livers was significantly higher than the control group. Similarly, Korkmaz et al. (2018)\(^{37}\) observed GST was induced by phosalone-based (PBP) and cypermethrin-based (CBP) pesticides in zebrafish (Danio rerio) after 96 h exposure. Liu et al. (2015)\(^{45}\) suggested that when azoxystrobin was applied to green algae Chlorrella vulgaris, GSH level decreased and GST activity increased due to excessive ROS production, thus scavenging free radicals. GST catalyzes the conjugation of xenobiotics with GSH, allowing them to be removed from the organism\(^{46}\) thus, GST induction is used as a biomarker of cellular damage caused by xenobiotics.\(^{47}\) There are many studies in the literature revealing that GST activity increases in aquatic organisms treated with pesticide.\(^{48–52}\)

GR activity decreased significantly in the Thmx applied groups compared to the control. The greatest increase in inhibition was seen at the LC_{50} dose, with a rate of approximately 84% compared to the control. Although all inhibitions were statistically significant, the least inhibition was seen at LC_{50}/8 dose with 76% difference from the control. GR is an enzyme that indirectly acts as an antioxidant by converting oxidized glutathione (GSSG) formed during reactions catalyzed by glutathione peroxidase (GPx) and glutathione S-transferase (GST) into reduced glutathione (GSH).\(^{52}\) In this study, observation of significant decreases in GR activity in all groups may be due to extracellular transport of GSSG rather than GSH to inhibit the cytotoxic effects of Thmx.\(^{54}\)

CarE activity was significantly inhibited in all Thmx concentrations compared to the control. At the highest Thmx concentration (LC_{50}) the greatest inhibition (approximately 55% increase over control) was observed. CarEs are members of the esterase family that catalyze the hydrolysis of substrates such as carboxylic esters, thioesters, amides and carbamates, and various xenobiotics.\(^{55}\) CarEs are involved in important physiological processes such as lipid metabolism,\(^{56}\) pro-drug activation,\(^{57}\) and hydrolysis of phthalates.\(^{59}\) In agreement with our results, Denton et al. (2003)\(^{60}\) reported that CarE activity was inhibited by 50% in fathead minnows compared to the unexposed group due to diazinox exposure. Wheelock et al. (2005)\(^{61}\) observed that after applying chlorpyrifos to Chinook salmon (Oncorhynchus tshawytscha) for 96 hour, CarE activity decreased significantly compared to control. Uçkun and Öz (2020a),\(^{51}\) who first demonstrated that CarE was inhibited as a result of acute application (96 h) of pesticide fenconazole to crayfish, suggested that CarE is a sensitive biomarker of pesticide toxicity in crayfish hepatopancreas. In our study, data on CarE inhibition due to Thmx administration also support this view.

In GPx activity, there were significant increases in all Thmx concentrations. These increases in GPx activity were not dependent on dose increase. The greatest increase was seen at the LC_{50}/8 dose, approximately 44% difference from the control. The main function of GPx is to reduce the lipid hydroperoxides formed in the cell due to xenobiotic exposure to their end product alcohols and to reduce free hydrogen peroxide.\(^{52,61}\) Inhibition in the GPx enzyme may reflect the failure of the antioxidant system to prevent the destructive effect of the pesticide,\(^{64}\) or it may be related to the direct effect of reactive oxygen species formed in cells on the synthesis of this enzyme.\(^{65}\) From this perspective, the GPx increase observed in this study may reflect the protective role of GPx against the oxidative damage induced by Thmx in the cell. In parallel with our findings, Blahova et al.\(^{66}\) found that when they subchronically applied atrazine to zebrafish, GPx activity was significantly increased.

There was a decrease in SOD activity at the LC_{50}/8 dose, and an increase in the other doses compared to the control depending on the dose. Only the increase in the LC_{50} administration dose was statistically significant from the control (p < 0.05). SOD is an important antioxidant enzyme that catalyzes the conversion of superoxide radicals to \( \text{H}_2\text{O}_2 \) and \( \text{O}_2^- \) in organisms and forms the first defense against free oxygen radicals formed in cells.\(^{67}\) When an organism is exposed to a xenobiotic, a decrease in the antioxidant system may be followed by an increase, which may reflect that the organism is adapting.\(^{68,69}\) The increase in SOD activity at high Thmx concentrations indicates that SOD scavenges the overproduction of superoxide ions under the oxidative stress created by Thmx. Many studies have shown that SOD activity is increased in organisms exposed to pesticides.\(^{66,70,71}\)

GSH level decreased significantly in all groups treated with Thmx compared to control. The greatest reduction was seen at the LC_{50} dose, with a rate of 45%. GSH is an essential endogenous tripeptide, which prevents the cell from oxidative injury. GSH acts as a cofactor for GST,\(^{72}\) which is responsible for detoxification of xenobiotics, so an increase or decrease in GSH level can be an important indicator of the detoxification ability of the organism.\(^{73}\) Our findings are in line with many studies in the literature that GSH level decreased as a result of pesticide application to aquatic organisms.\(^{74–78}\) A decrease in GSH may mean that the antioxidant defense system is activated against the oxidative damage caused by ROS in the cell, as this reduction is an indication that GSH is spent converting to oxidized glutathione or regenerating GSH.\(^{79}\) Also, a
There was an increase in the MDA level at all Thmx concentrations and these increases were in a dose-dependent manner. Differences in all concentrations were statistically significant except for the LC50/8 concentration. The highest increase in MDA level was at the LC50 concentration, approximately 43% compared to the control. Lipid peroxidation is the first indicator of cell membrane damage caused by exposure of organisms to pesticides, metals and various xenobiotics. The reason for the high level of MDA in our study may be the peroxidation of unsaturated fatty acids in the cell membranes, as Thmx exposure causes oxidative damage in the cell and increases ROS production. It has been reported that the level of MDA increased significantly in various aquatic organisms exposed to different pesticides compared to the groups not treated with pesticides.

### 3.4.2. Neurotoxicity (AChE)

There was a significant decrease in AChE activity in the Thmx applied groups compared to the control. The reductions in all Thmx concentrations relative to control were not dose dependent. The highest AChE inhibition was observed in the LC50/2 group with an approximately 77% difference from the control. The inhibition in the LC50 application was approximately 71% compared to the control. When AChE is inhibited by xenobiotics, acetylcholine accumulates in the synaptic space and the receptors are highly stimulated. Activation of muscarinic ACh receptors is relatively slow (milliseconds to seconds) and, depending on the subtypes present, they directly alter cellular homeostasis. Unlike muscarinic receptors, the nicotinic receptors are inactivated due to sustained increase in ACh concentrations, which ultimately results in paralysis. Therefore, AChE is used as a biomarker of pesticides that target it directly or indirectly by altering the cholinergic neurotransmission. In our study, significant AChE inhibition due to Thmx administration indicates that Thmx has neurotoxic effects in crayfish at the doses applied. Similar to our findings, AChE inhibition was observed after 96 hours of Thmx application to the midge Chironomus riparius. Many researches reported that AChE is inhibited by neonicotinoid pesticides in various aquatic organisms.

### 3.4.3. Ion Transport

ATPases are responsible for ion homeostasis in cell membranes, play a central role in the physiological functions of the cell by providing energy conversion in chemical reactions, so they are considered a good indicator in toxicological studies. In our study, significant inhibitions at all Thmx doses were significant (p<0.05). Na+K+ATPase inhibition rates in the gill were 25%, 49%, 50% and 71%, respectively, based on the applied Thmx concentrations. In muscle tissue, all Na+K+ATPase inhibitions were statistically significant except for LC50/8 (p<0.05). Na+K+ATPase inhibition rates relative to control in muscle were 6%, 17%, 38% and 42%, respectively. Na+K+ATPase has a vital function in maintaining the cell membrane potential difference by keeping Na+ outside the cell and K+ inside the cell. Inhibitions in Na+K+ATPase activity indicates the destruction of cellular ion regulation in the tissues of fish. The researcher reported that this degradation may also be due to the effect of pesticide on the passive movement of ions, namely its permeability properties. Cirrhinus mrigala, which is exposed to the lethal and sublethal effects of deltamethrin, has been found to decrease Na+K+ATPase activity in gill, liver and muscle tissue. It has been determined that the gill tissue Na+K+ATPase activity of Cyprinus carpio, which is exposed to cypermethrin sub-lethal effect for different periods, shows a decrease depending on the time. Similar observations were reported by Begum (2011) in the fish C. batrachus exposed to carbofuran. In a study conducted by Temiz et al. (2018), it was determined that under the effect of chlorantraniliprole (CHL), the decrease in

### Table 3. Biochemical responses of 96-h Thmx exposure in hepatopancreas. Total protein amount expressed as mg ml⁻¹, and enzyme activities expressed as nmol min⁻¹ mg protein⁻¹ ± mean standard error. GSH and MDA levels expressed as nmol GSH mg protein⁻¹ ± mean standard error and nmol MDA mg protein⁻¹ ± mean standard error, respectively:

| Dose     | N | Total Protein | GST | GR  | AChE | CarE | GPx | SOD | GSH  | MDA  |
|----------|---|---------------|-----|-----|------|------|-----|-----|------|------|
| Control  | 12| 160.86 ± 7.03 | 36.91 ± 1.70 | 6.42 ± 0.36 | 5442.60 ± 278.80 | 8.01 ± 0.33 | 4.47 ± 0.21 | 0.20 ± 0.02 | 3.00 ± 0.17 |
| LC50/8   | 12| 282.77 ± 3.72 | 8.98 ± 0.20 | 2.91 ± 0.10 | 2884.40 ± 39.03 | 14.24 ± 0.51 | 4.04 ± 0.11 | 0.13 ± 0.01 | 3.07 ± 0.04 |
| LC50/4   | 12| 395.32 ± 9.74 | 8.49 ± 0.27 | 3.23 ± 0.11 | 2968.20 ± 88.25 | 14.23 ± 0.51 | 4.61 ± 0.15 | 0.14 ± 0.03 | 4.23 ± 0.21 |
| LC50/2   | 9 | 277.38 ± 10.3 | 8.84 ± 0.31 | 1.50 ± 0.10 | 3075.10 ± 110.50 | 11.22 ± 0.17 | 6.77 ± 0.22 | 0.14 ± 0.01 | 4.58 ± 0.12 |
| LC50    | 8 | 385.33 ± 5.01 | 6.09 ± 0.33 | 1.84 ± 0.07 | 2455.50 ± 61.82 | 13.48 ± 1.13 | 7.92 ± 0.28 | 0.11 ± 0.01 | 5.29 ± 0.30 |

N: Number of animals that survived after the 96-h acute toxicity test. *: p < 0.05 showed statistical importance compared with control group.
Na⁺K⁺-ATPase activity of *O. niloticus* gill tissue increased due to the prolongation of the time. The observed decrease in the activities of Na⁺K⁺-ATPase may be due to the change in ionic homeostasis and may also be due to ATP depletion.⁹²

In both gill and muscle tissues, Mg²⁺-ATPase activity decreased as the applied Thmx concentration increased. The highest reduction was observed in the groups where the highest Thmx concentration (LC⁵₀) was applied. Mg²⁺-ATPase inhibition rates in the gill were 29%, 44%, 47%, 47%; in muscle, it was 15%, 32%, 57% and 63% compared to control depending on the increase in Thmx concentration. Mg²⁺-ATPase is an enzyme that ensures the integrity of the cell membrane by transepithelial regulation of Mg²⁺ ions and is associated with the synthesis of ATP through oxidative phosphorylation in mitochondria.⁹¹ Inhibition of Mg²⁺-ATPase in the present study may have caused a disruption in the transport of ions across the cell membrane and a decrease in ATP production.⁹²,⁹⁶

Ca²⁺-ATPase was inhibited increasingly as Thmx concentration increased in both gill and muscle tissues. The highest inhibitions in the gill and muscle were seen at the LC⁵₀ dose with rates of 57% and 58%, and the lowest were at the LC⁵₀/8 dose with rates of 27% and 13%, respectively. All of these inhibition of Ca²⁺-ATPase activity were statistically significant (p < 0.05). Ca²⁺-ATPase is an enzyme that serves to remove calcium (Ca²⁺) from the cell and is vital in regulating the amount of Ca²⁺ within cells.⁹⁷ Inhibition of Ca²⁺-ATPase activity in gill and muscle tissues may be associated with the disruption of the osmoregulation mechanism due to the blockage of the active transport system by Thmx.⁹⁸ Additionally, Thmx may have caused inhibition of membrane bound enzymes due to degradation products of lipid peroxidation in the cell membrane by inducing oxidative stress.⁹⁹ This may result in disruption of the active transport mechanism due to altered membrane permeability and impaired Ca²⁺-ATPase homeostasis.⁹⁸ Similar to our findings, Uçkun and Öz (2020a, 2020b)⁵¹,⁵² observed that ATPase activities (Na⁺K⁺-ATPase, Mg²⁺-ATPase, Ca²⁺-ATPase) in gill and muscle tissues decreased significantly in a dose-dependent manner in two separate studies in which *A. leptodactylus* applied the fungicides penconazole and azoxystrobin for 96 hours. In our study, the ATPase inhibition rates in the gill were found to be higher than those in the muscle. This decrease is thought to be the result of impairment of ion balance and gill permeability, since it is the first tissue in contact with the pesticide in the aquatic environment. In fish, various toxic substances and ions enter the body by absorption and adsorption by the gill surface, followed by diffusion. Interaction with the membrane may impair the osmotic and ionic regulation of gill tissue by affecting membrane permeability.⁹³ The reason that responses to biomarkers vary according to the organ is related to the defense capacities of the organs as well as their anatomical location that determines the path and distribution of xenobiotic exposure.⁹²

| Dose    | N  | Total protein | Na⁺/K⁺-ATPase | Mg²⁺-ATPase | Ca²⁺-ATPase |
|---------|----|---------------|---------------|-------------|-------------|
| Control | 12 | 12.26         | 40.74 ± 1.58  | 48.72 ± 0.95 | 89.46 ± 2.11 |
| LC⁵₀/8  | 12 | 9.31          | 30.58 ± 0.91  | 34.37 ± 0.39 | 64.94 ± 1.12 |
| LC⁵₀/4  | 12 | 11.49         | 20.82 ± 0.62  | 27.09 ± 0.51 | 48.01 ± 0.62 |
| LC⁵₀/2  | 9  | 10.77         | 20.17 ± 1.03  | 25.80 ± 0.60 | 45.97 ± 0.55 |
| LC⁵₀    | 8  | 11.91         | 11.97 ± 0.37  | 26.07 ± 0.40 | 38.03 ± 0.29 |

N: Number of animals that survived after the 96-h acute toxicity test.
*: p < 0.05 showed statistical importance compared with control group

| Dose    | N  | Total protein | Na⁺/K⁺-ATPase | Mg²⁺-ATPase | Ca²⁺-ATPase |
|---------|----|---------------|---------------|-------------|-------------|
| Control | 12 | 15.13         | 21.18 ± 0.91  | 72.03 ± 1.32 | 93.21 ± 1.32 |
| LC⁵₀/8  | 12 | 11.40         | 19.97 ± 0.44  | 61.51 ± 0.98 | 81.48 ± 1.13 |
| LC⁵₀/4  | 12 | 12.19         | 17.55 ± 0.66  | 49.15 ± 1.04 | 66.70 ± 0.69 |
| LC⁵₀/2  | 9  | 12.02         | 13.18 ± 0.17  | 31.15 ± 0.06 | 44.33 ± 0.18 |
| LC⁵₀    | 8  | 11.37         | 12.32 ± 0.23  | 26.60 ± 1.25 | 38.92 ± 1.25 |

N: Number of animals that survived after the 96-h acute toxicity test.
*: p < 0.05 showed statistical importance compared with control group
When evaluating the responses of biomarkers, we used IBR analysis to allow combining all parameters into one general stress index (Figure 1). IBR analysis is a useful method that provides a brief information in comparing multiple biomarkers. The IBR index expressing the toxicity caused by Thmx in the hepatopancreas was determined to be the highest at the LC50 dose. At the LC50/2 and LC50/4 doses, the IBR index was found to be close to each other and lower than the LC50 dose. Compared to other doses, the lowest IBR index was determined at the LC50/8 dose. As can be seen, although hepatopancreas IBR index rose with increasing Thmx dose, it was suppressed compared to control. This may be because the hepatopancreas plays a role in detoxification. In gill and muscle tissues, IBR index was inhibited compared to the control due to increasing Thmx dose. The IBR index was completely suppressed at the LC50 dose in both tissues because ATPase inhibitions were highest at this dose. The findings of our study are in line with various studies using the IBR index in the assessment of the effects of environmental pollutants on macroinvertebrate, mussel and fish.

Compliance with Ethical Standards
All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. All procedures performed in studies involving animals were in accordance with the ethical standards of the institution or practice at which the studies were conducted.

Conflict of Interest
The authors declare that they have no conflict of interest.

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Povzetek
Tiametoksam (Thmx) je globalno razširjen neonikotinoidni pesticid, ki onesnažuje sladkovodne ekosisteme in katerega ostanke so zaznali v ribiških proizvodih. Astacus leptodactylus je priljubljen sladkovodni rak, ki ga gojijo in izvažajo v mnogih državah. V okviru raziskave smo preučevali akutne toksične učinke Thmx na A. leptodactylus z uporabo različnih biomarkerjev (acetilholinesteraza, karboksilesteraza, glutation S-transferaza, glutation, superoksidna dismutaza, glutation peroksidaza, glutation reduktaza in adenozintrifosfataze). 96-urna vrednost LC₅₀ Thmx je bila izračunana kot 8.95 mg aktivne učinkovine L⁻¹. Ko se je odmerek Thx povečeval, je oksidativni stres povzročil inhibicijo/aktivacijo antioksidativnih encimov, medtem ko so bile aktivnosti acetilholinesteraze, karboksilesterase in adenozintrifosfataz inhibirane. Posledično lahko rečemo, da Thmx izkazuje močno toksične učinke na rake, zato so ti na območjih, kjer se ta pesticid uporablja, ogroženi.