Imidacloprid-induced pathophysiological damage in the midgut of *Locusta migratoria* (Orthoptera: Acrididae) in the field

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**Abstract**

Neonicotinoids are modern insecticides widely used in agriculture worldwide. Their impact on target (nervous system) and non-target (midgut) tissues has been well studied in beneficial insects including honeybees under controlled conditions. However, their detailed effects on pest insects on the field are missing to date. Here, we have studied the effects of the neonicotinoid imidacloprid on the midgut of the pest insect *Locusta migratoria* caught in the field. We found that in the midgut of imidacloprid-exposed locusts the activity of enzymes involved in reactive oxygen metabolism was perturbed. By contrast, the activity of P450 enzymes that have been shown to be activated in a detoxification response and that were also reported to produce reactive oxygen species was elevated. Probably as a consequence, markers of oxidative stress including protein carbonylation and lipid peroxidation accumulated in midgut samples of these locusts. Histological analyses revealed that their midgut epithelium is disorganized and that the brush border of the epithelial cells is markedly reduced. Indeed, microvilli are significantly shorter, misshapen and possibly non-functional in imidacloprid-treated locusts. We hypothesize that imidacloprid induces oxidative stress in the locust midgut, thereby changing the shape of midgut epithelial cells and probably in turn compromising their physiological function. Presumably, these effects reduce the survival rate of imidacloprid-treated locusts and the damage they cause in the field.

**Keywords** Insect · Locust · Neonicotinoid · Detoxification · Oxidative stress · ROS · Midgut

**Background**

Because of its devastating impact on agriculture, the locust *Locusta migratoria migratorioides* is considered the most economic important pest in Assiut governorate in Upper Egypt. The existing control methods against this pest rely on chemical insecticides such as imidacloprid, chlorpyriphos, malathion and others (El-Saad et al. 2017), which are harmful to the environment and potentially also to humans.

Since their introduction in the nineties of the last century, neonicotinoids have become the most widely used chemical class of insecticide in many countries (Casida and Durkin 2013, Jeschke and Nauen 2008, Jeschke et al. 2011, Tomizawa and Casida 2005). Due to their specificity to insects concomitant with low acute toxicity to non-insects and their versatility in application methods, neonicotinoids constitute an attractive tool in modern agriculture (Elbert et al. 2008; Thompson et al. 2020). However, concerns over neonicotinoids are emerging, and in the European Union, for instance, the use of several neonicotinoids has been widely restricted.
because of their adverse effects on honeybees ([https://ec.europa.eu/food/plants/pesticides/approval-active-substances/renewal-approval/neonicotinoids_en](https://ec.europa.eu/food/plants/pesticides/approval-active-substances/renewal-approval/neonicotinoids_en)) (Thompson et al. 2020, Wood and Goulson 2017). Hence, for the near future, it is important to assess the impact of neonicotinoids and to understand their effects on insects at the molecular and cellular levels.

Neonicotinoids target primarily the nervous systems of insects as agonists of postsynaptic acetylcholine receptors by binding to the α subunits of the receptor (Buckingham et al. 1997; Matsuda et al. 2020; Suchail et al. 2004). Unlike the natural ligand of this receptor acetylcholine, neonicotinoids are not deactivated by hydrolyzation by acetylcholinesterase. Therefore, in the presence of neonicotinoids, transmission of nerve impulses does not cease resulting in hyperexcitation of the nervous system. This neuronal dysfunction especially affects visual processing and in turn collision avoidance in, for instance, *Locusta migratoria* (Parkinson et al. 2017, 2020).

Commonly, exposure to toxic xenobiotics including pollutants can be monitored through their negative effects on biochemical processes in cells and tissues. Indeed, pollutants may cause variations in the activity of specific enzymes and associated cellular damage, in turn indicating the presence of these toxic pollutants (Johnston 1995; Lu et al. 2021). For instance, the upregulation of superoxide dismutase (SOD), catalase (CAT), ascorbate peroxidase (APOX), glutathione-S-transferase (GST) and lipid peroxidation has been used in ecotoxicological studies as powerful biomarkers (Scandalias 2005; Zhang et al. 2019). These biomarkers are directly or indirectly involved in reactive oxygen species (ROS) metabolism, which may be perturbed following exposure to xenobiotics such as insecticides or metals in insects (El-Gendy et al. 2020). In *D. melanogaster*, low doses of the neonicotinoid imidacloprid cause mitochondrial dysfunction in glia cells that is associated with oxidative stress triggered by ROS (Martelli et al. 2020; Wang et al. 2018). Moreover, in insects, heat-shock proteins (HSPs) are important players in response to abiotic stressors preventing induction of cell death, for example (King and MacRae 2015), and are therefore highly useful as biomarkers for environmental pollution (Bierkens 2000; El-Saad et al. 2017; Lewis et al. 1999). Besides these enzymatic biomarkers, the use of cellular markers provides an additional means of assessing the toxicity of pollutants inside the insect body. One of these markers is the fragmentation of DNA in necrotic or apoptotic cells after exposure to an environmental stressor.

The cellular consequences of neonicotinoid application in insects have been largely assessed in the target tissue, i.e. the nervous system. A few work described the consequences of controlled application of neonicotinoids on the midgut of the silk moth *Bombyx mori* or the honeybee *Apis mellifera* kept in the laboratory or a mesocosm including detoxification (Carneiro et al. 2022; Wang et al. 2020). The aim of the present study was to evaluate the level of protein carbonyls, lipid peroxides and antioxidant enzyme activity (SOD, CAT, APOX, GR and GPx) in the gut tissue of locusts (*Locusta migratoria*) captured in the field, i.e. on sites along a gradient of imidacloprid pollution near Alexandria, Egypt. At the cytological level, apoptosis, which is the possible consequence of perturbed ROS metabolism, was detected through flow cytometry (annexin-V assay) and by transmission electron microscopy (TEM).

**Materials and methods**

**Insect sampling**

The study site was in Assiut governorate, the Upper Egypt (27° 10′ 48″ N and 31° 11′ 21″ E). The study site received 2–4 applications of 250 mL imidacloprid per feddan (4200 m²) by spraying. There is little available data in the literature regarding the existence of other contaminants in this area. The selected sprayed location is considered a highly active agricultural area for maize crop.

Adult locusts (*Locusta migratoria*) used in the present study were collected during the summer season 2020 and placed in jars. They were kept in our laboratory cages (40×40×70 cm) for 1 day under natural photoperiod conditions at relative humidity of 50–65% at 28 °C and fed with leaves of maize sprouts harvested from the same field. The insects were identified for their species as *L. migratoria* at the Faculty of Agriculture, Alexandria University, Egypt. *L. migratoria* collected from the contaminated area were compared to those reared free of pesticide applications in the laboratory under controlled conditions and were thus considered as the control ones.

In general, ten fresh adult specimens of each control (around 5 days after emergence) and contaminated *L. migratoria* were washed with distilled water and sacrificed after anesthesia with absolute ethanol (95%). The midguts were rapidly dissected out. Half of the sample size of the midgut tissue (20 g) was stored in vials and kept at −80 °C until used in biochemical (except for acetylcholinesterase assays, see below) and neonicotinoid residue analyses, while the other half was used for histological and ultrastructural studies. The technique for collecting the whole midgut tissue is based on the method by Liu (Liu 1984).

**Determination of neonicotinoid pollutant residues**

Twenty grams of midgut tissue from control and contaminated insects was mixed with 10 g of anhydrous sodium sulfate and homogenized in 100 mL of chloroform and acetone. The homogenate was acidified with 10 drops of...
concentrated acetic acid and filtered. The filtrate was dried by evaporation at 35 °C. The residue was redissolved in 1 mL of N-hexane and acetone and cleaned up using 10 g of 1% deactivated Florisil (Diaz et al. 2012). The samples were concentrated using a rotary evaporator to 5 mL and completely dried under a stream of N₂. The residues were dissolved in 0.5 mL of N-hexane. Recovery experiments of the neonicotinoid imidacloprid were carried out by using fortified samples through addition of 25 μg of imidacloprid to specimens of tissue samples. The samples were extracted, cleaned up and determined as described before.

Then, concentrations were corrected for 100% recovery, and procedural recovery trials were undertaken which were above 85%. The analysis was performed on Hewlett-Packard HP-6890 series Gas Chromatograph. The injection pore and flame photometric detector, which is very sensitive to halogenated neonicotinoid, imidacloprid, were operated at 245 and 250 °C, respectively. The data were calculated to final values in parts per billion. The detection limits for the examined compounds were determined. The compound identities were accomplished by comparing retention times against pure standard of imidacloprid at the same conditions. The retention times for imidacloprid were on average 12.52 min.

Biochemical assays

Antioxidant enzyme assays

The midguts of *L. migratoria* collected from control and contaminated areas as described above were rinsed with distilled water and blotted using a filter paper. The tissues were weighed and homogenized for 30 s with 10 volumes (w/v) of ice-cold saline solution (0.9%) and 40 mM sodium phosphate with a homogenizer (Tekmar tissumizer). The homogenates were centrifuged at 6500 rpm for 30 min at 4 °C using IEC-CRU5000 centrifuge. The resulting supernatants were split and frozen at −20 °C for subsequent analysis. Glutathione (GSH) levels were assayed spectrophotometrically using the method published by Anderson (Anderson 1989). We measured the activity of superoxide dismutase (SOD) following the procedure described by Nebot et al. (Nebot et al. 1993). Catalase (CAT) activity was measured following concentration decrease of hydrogen peroxide (H₂O₂) at 240 nm (Aebi 1984). To detect the activity of glutathione peroxidase (GPx), we monitored continuous decrease in NADPH amounts according to the method described by Chu et al. (Chu et al. 1993). GST activity was quantified by the method modified by Carmagnol et al. (1981). Glutathione reductase (GR) was measured according to Smith et al. (Smith et al. 1988). The activity was expressed in U/mg protein by dividing the enzyme activity per milliliter of the sample by the protein concentration in the same sample. The activity of ascorbate peroxidase (APOX) was assessed via the method reported by Levine et al. (Levine et al. 1990).

Acetylcholineesterase assay

Acetylcholineesterase (AChE) activity was assessed in the midguts of *L. migratoria*. Values were measured in pools of three samples in two replicates according to Ellman et al. (Ellman et al. 1961). In brief, in tissue homogenates, AChE catalyzed the conversion of acetylcholine to thiocholine and acetate; thiocholine interacted subsequently with dithiobis-nitrobenzoate ions resulting in a yellow color that was quantified by colorimetry.

Protein carbonyls and lipid peroxidase assays

We isolated the midguts of all locusts from the body and homogenized them in 5 mL of an ice-cold phosphate buffer with additives (60 mL of 50 mM phosphate buffer, 10 mL of 0.1% Triton × 100, 5 mL of 0.05 mM CaCl₂; after adjusting pH to 7.0 with HCl or NaOH, the mixture was filled with distilled water to a volume of 100 mL). After homogenization (mortar, 10 strokes/30 s), the samples were centrifuged at 2000 × g for 10 min at 4 °C. Then, 800 μL aliquots of the supernatant were transferred to a new microtube with 800 μL of 30% trichloroacetic acid (TCA). The samples were incubated for 30 min at room temperature and then centrifuged at 5000 × g for 10 min at 4 °C. The assay of protein carbonyls was conducted on precipitated pellets, and the assay of lipid peroxides was conducted on the supernatant. The concentrations of lipid peroxides (malondialdehyde, MDA) and protein carbonyls were determined using the method of Hermes-Lima et al. (Hermes-Lima et al. 1995) and Levine et al. (1990), respectively. The measurements were done in three replicates; each replicate was a pool of ten insects.

Cytochrome P450 monooxygenase assay

For the determination of cytochrome P450 activity, we used insect homogenates following the protocol of Brogdon and Chan (Brogdon and Chan 2010) introducing minor modifications described in the Anopheles research of US CDC. To prepare the NaOAc buffer, we used 0.25 M sodium acetate (C₂H₃NaO₂) dissolved in 800 mL purified water; the pH of the solution was to 5.0 with acetic acid. Next, 20 mg of TMBZ was dissolved in 25 mL methanol before adding 75 mL of 0.25 M NaOAc buffer (pH 5.0). For the actual assay, to obtain the positive control stock solution, 10 mg of cytochrome-C was added to 100 mL of 0.25 M NaOAc buffer (pH 5.0). One hundred microliters of KPO₄ was pipetted to the plate with the insect homogenates in both the negative and positive sample wells. For the positive controls, 100 μL of the cytochrome-C control solution was
piped into three wells; for the actual assay, we applied 200 μL of TMBZ solution to each test well. The reactions were started by adding 25 μL of 3% hydrogen peroxide (H₂O₂). They were incubated for 5 min at room temperature, and the OD values were recorded at 620 nm for determination of the oxidase activity. Cytochrome P450 (general oxidase) activity obtained from plate reading was expressed as mUnits (mU) of cytochrome P450 per milligram of protein.

**Annexin V-FITC/propidium iodide assay**

Flow cytometric analysis was performed for detecting apoptosis in the midguts tissue samples of control versus contaminated insects using a FACS Calibur flow cytometer (BD-Becton Dickinson, USA), and data were analyzed using Cell Quest software (Becton Dickinson, San Jose, CA, USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA). Apoptosis was assessed with flow cytometry using an annexin V-FITC/propidium iodide (PI) staining kit (TASC USA).

**Midgut histopathological examination and estimation of the mean villus length**

For histological preparations, fresh midgut tissues from control and contaminated locusts were collected and fixed in 10% buffered formalin for 24 h. The specimens were dehydrated in increasing ethyl-alcohol concentrations (70%, 80%, 90% and 100%), double cleared in xylene for 1 h each and embedded in paraffin. Four-micrometer-thick tissue sections were prepared and stained with hematoxylin and eosin (H&E). They were examined by light microscopy and photographed. To calculate mean microvillus length for each group, ten microvilli per section were measured, and ten boundaries/slide/-locust were examined.

**Scanning electron microscopic study of midgut (SEM)**

After dissection, small pieces of fresh midgut tissues were incubated in cold 4F1G for 3 h for fixation; they were post-fixed with 2% osmium tetroxide for 2 h at 4 °C. These samples were washed and dehydrated in an increasing series of ethanol concentration, mounted using carbon paste on an aluminum stub and coated with gold–palladium in a sputter coating unit (JFC-1100 E). Specimens were analyzed using a SEM JSM-5300 (Tahmasebi et al. 2015).

**Transmission electron microscopy study (TEM)**

Midguts of the control and insecticide-treated insect groups were isolated and fixed by incubation in 4% formaldehyde and 1% glutaraldehyde (4F1G) in phosphate-buffer solution (pH 7.2) at 40 °C for 3 h. After post-fixation in 2% osmium tetroxide (OsO₄) in the same buffer for 2 h, samples were rinsed with the buffer. Next, samples were dehydrated in a series of increasing ethanol concentration at 40 °C. Thereafter, they were embedded in a mixture of Epon–Araldite in labeled capsules. Ultra-thin sections (0.06–0.1 μm thick) were produced from the guts of both insect groups for examination with a transmission electron microscope (TEM). Semithin sections were prepared by using LKB ultramicrotome (1 μm thick). For quality check, determination of the orientation and structural features by light microscopy, sections were deposited on a glass slide and stained with toluidine blue. For optimal comparability, ultra-thin sections with either pale gold or silver interference color representing sections with 0.1 μm or 0.06-μm thickness, respectively (Peachey 1958), were chosen for further analyses. They were transferred on 200-mesh naked copper grids. Sections were contrasted with uranyl acetate for half an hour and lead citrate for 25–30 min (Reynolds 1963). Electron micrographs were taken at several magnifications. Observation and photographing the grids were performed on a JEOL JEM-1400 plus Electron Microscope (Jeol, Tokyo, Japan) at 80-kV accelerating voltage, Faculty of Science, Alexandria University, Egypt.

**Assessment of HSP70 and HSP90 by quantitative real-time PCR**

Total RNA was isolated from locust tissue samples using TRIzol reagent (Thermo Fisher Scientific, USA). Sample RNA concentrations were evaluated using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, USA) at 260 nm. After that, total RNA (1.0 μg per sample) was treated with DNase I (Fermentas, USA) to remove potential genomic DNA contamination; first-strand cDNA was synthesized in a 20-μL reaction system using a High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific, USA). The relative expression levels of the HSP70 and HSP90 in locust tissues were analyzed using qRT-PCR. One microgram RNA was used as the template. Specific primers for HSP70, HSP90 and the housekeeping gene β-actin used in the assay are listed in below (Wang et al. 2006, 2007). qRT-PCR reactions were executed using Qiagen Rotor-Gene SYBR Green PCR Kit in a 25 μL mixture containing 1 μL of cDNA, 12.5 μL of SYBR Green, 2.5 μL of each primer and 9 μL of H₂O. The qRT-PCR program consisted of an initial step at 95 °C for 5 min, followed by 40 cycles of 95 °C for 15 s and 60 °C for 10 s. The assays were run in the Rotor-Gene Q using Rotor-Gene
Q-Pure Detection version 2.1.0 (Qiagen, USA). Quantification of the transcript level of HSP70 and HSP90 mRNA was completed using the comparative 2-ΔΔCT method (Livak and Schmittgen 2001). The identification was performed using primers 5′-AAA ATG AAA GAA ACG GCA GAG G-3' (forward), 5′-TAA TAC GCA GCA CAT TGA GAC C-3' (reverse) for HSP70, 5′-GAT ACA TCC ACA ATG GG C-3' (reverse) for HSP90 and 5′-AAT TAC CAT TGG TAA CGA GCC ATT-3' (forward), 5′-TGC TTC CAT ACC CAG GAA TGA-3' (reverse) for β-actin.

**Statistical analysis**

Enzymatic activities and gene expression were measured in biological triplicate. Student tests were performed after checking homoscedasticity with Fisher exact tests between treatments and controls. Data were analyzed using XLstat (c) 2010 from Addinsoft.

**Results**

**Neonicotinoids are present in the locust midgut**

Locusts caught in the field have been exposed to the neonicotinoid imidacloprid (Fig. 1). Our investigations failed to identify the presence of other insecticides at the site of sample collection. In a strict sense, however, we cannot exclude the exposure of our samples to other toxic xenobiotics including insecticides. In other insects, it has been proposed that complex changed acetylcholineesterase (AChE) may be a biomarker for neonicotinoid exposure (Morakchi et al. 2005). To evaluate imidacloprid exposure of our field locusts, we therefore determined the activity of AChE. In a specific enzyme activity assay, we found that AChE performance is significantly reduced by 40% (Fig. 2). As caught locusts were viable, this result is consistent with a sub-lethal exposure of these locusts to imidacloprid.

**Imidacloprid deactivates ROS removal in the midgut**

Neonicotinoid processing involves ROS production (Martelli et al. 2020). We analyzed ROS metabolism in our field locust population by measuring the activity of a series of reactions catalyzed by enzymes involved in the oxidative stress response also in insects including catalase (CAT), superoxide reductase (SOD), glutathione reductase (GR), glutathione S-transferase (GST), guaiacol peroxidase (GPX) and ascorbate peroxidase (APOX) (Fig. 2) (Cui et al. 2020). In addition, we determined the amounts of reduced glutathione (GSH), protein carbonylation (PC) and malondialdehyde (MDA) as markers of oxidative stress (Dalle-Donne et al. 2003). The activities of CAT, SOD, GR, GPX and APOX were substantially reduced in imidacloprid-treated animals. These results suggest that imidacloprid induces oxidative stress. Consistently, GST activities and MDA, PC and GSH amounts were increased in these locusts.

Detoxification upon exposure to xenobiotics including insecticides like imidacloprid is to a large extent managed by P450 cytochromes that catalyze substrate oxidation (Lu et al. 2021). We found that P450 cytochrome activity increased in the gut (Fig. 2). We conclude that imidacloprid triggers detoxification via enhancing the activity of P450 cytochromes.

Next, we monitored the expression of HSP70 and HSP90, two genes coding for chaperones that assist protein folding under stress conditions among others in the nervous system (Lackie et al. 2017). The expression of both genes is elevated suggesting that imidacloprid-treated animals launch
Fig. 2 Biochemical analyses of ROS metabolism. Data are represented as scattergrams. Red crosses indicate the mean. Data were analyzed with Student tests whose results are reported here (Catalase: $t(4) = 5.831$, $p = 0.004$; SOD: $t(4) = 10.119$, $p = 0.001$; GR: $t(4) = 10.423$, $p = 0.0005$; GSH: $t(4) = 12.095$, $p = 0.0002$; MDA: $t(4) = 10.955$, $p = 0.0004$; GPx: $t(4) = 10.094$, $p = 0.001$; GST: $t(4) = 9.255$, $p = 0.001$; APOX: $t(4) = 12.965$, $p = 0.0002$; protein carbonyl: $t(4) = 74.097$, $p < 0.0001$; acetylcholinesterase: $t(4) = 4.315$, $p = 0.013$; HSP70: $t(4) = 15.02$, $p = 0.0001$; HSP90: $t(4) = 22.612$, $p = 0.002$; cytochrome P450 in gut: $t(4) = 12.294$, $p = 0.0002$). All experiments were performed in triplicates.

Fig. 3 The gut tissues of locust showing DNA damage using the comet assay from control (a) and polluted area (b and c) and flow cytometric analysis of annexin V-FITC and PI-staining from control (d) and polluted area (e). Q1: necrotic cells, Q2: late apoptotic cells, Q3: living cells, Q4: early apoptotic cells. The Y-axis represents the PI-labeled population, whereas the X-axis represents the FITC-labeled Annexin V positive cells.
a program to alleviate the effects of the insecticide (Fig. 2). However, together, these results indicate that despite an activated detoxification response, imidacloprid causes oxidative stress.

A consequence of oxidative stress is apoptosis. By flow cytometry (comet assay), we detected apoptosis by marking midgut cells of control and imidacloprid-exposed locusts with annexin-FITC and propidium iodide (Fig. 3). The ratio of apoptotic to non-apoptotic cells was significantly increased in imidacloprid-treated animals. This finding indicates that imidacloprid induces apoptosis in midgut cells of field locusts.

**Microvilli structure and density are compromised upon imidacloprid application**

In various insect species including the honeybee *Apis mellifera* and the silkworm *Bombyx mori*, neonicotinoids have been shown to affect midgut function (Arthidoro de Castro et al. 2020; Wang et al. 2020). To evaluate the impact of imidacloprid on the locust midgut, we examined the ultrastructure of midgut cells by light and electron microscopy.

Sections of control adult desert locust midgut are illustrated in Fig. 4A–D. Images from the control group revealed a cylindrical simple tube of regular thickness with a single layer of epithelial cells. Three types of epithelial cells were found throughout the epithelial layer of the midgut: (1) Columnar digestive cells, the most common cell type that can be recognized by their light staining; these cells produce regular microvilli at their apical site forming the brush border that faces the gut lumen. Their nuclei are oval containing electron-dense chromatin. This layer is supported by the basal lamina and enveloped by a thin bipartite layer of visceral muscles consisting of an inner circular muscle and an outer longitudinal muscle. (2) Regenerative cells with a basophilic cytoplasm and a spherical nucleus cluster together forming distinct nests (nidi) distributed regularly in the basal region of the digestive and secretory cells. Basally, they are delimited by the basement membrane and do not participate at the formation of the luminal surface. (3) The cylindrical or drop-shaped endocrine cells contact the basal lamina via a long, thin cytoplasmic protrusion, while their apical membrane does not join the midgut luminal surface. They insert individually between the digestive cells (nidi). Their nuclei usually localize to the periphery of the niches.

Transverse sections of treated adult desert locust midgut are shown in Fig. 4a–d. Partly, midgut epithelial cells are organized normally, while other parts of the midgut display severe alterations including disrupted epithelial cells, whose debris was found in the midgut lumen. Digestive cells, usually columnar, were rather short containing higher amounts of vacuoles in the cytoplasm; often, their nuclei localized to the midgut lumen. Some cells formed irregularly structured microvilli with reduced tips; occasionally microvillar material budded into the organ lumen due to cell swelling and vacuolization; in some regions, the brush border was completely absent. The lumen shape was irregular because of longitudinal infoldings. The basal lamina and the visceral muscle layer were both thin and, in some parts of the midgut, separated from the epithelium. Regenerative cells
were deformed and occasionally surpassed the apical surface of the epithelium; nidi were disorganized and difficult to identify.

In SEM micrographs of the control midgut, the brush border with the regularly organized microvilli formed a smooth lumen surface (Fig. 5). By contrast, the apical cell surfaces of the midgut of imidacloprid-treated locusts were irregular. Moreover, determination of microvillus length (μm ± SE) revealed a significant decrease (P < 0.05) in the length of microvilli in the imidacloprid-contaminated locust midgut compared to the control (Fig. 6).

To gain a more detailed image of the midgut of control and imidacloprid-treated locusts, we analyzed their ultrastructure by TEM. In these micrographs, the midgut is composed of a single layered epithelium supported by a thin and continuous basal lamina or basement membrane. The basal part of the cells forms the basal labyrinth of in-folding of the basal cell membrane. The apical membrane builds well-organized microvilli with regular thickness and lengths that line the lumen of the gut. At their tips, microvilli may possess vesicle-like or bleb-like structures, which reflect secretory activity. The nuclei generally lie in the middle of digestive cells, but occasionally, they may be shifted basally or apically. Their shape is ovoid, and they contain distinct heterochromatin (Fig. 7). Numerous elongated mitochondria with outlined membranes encasing the dense matrix with cristae occupy the middle and apical cytoplasm. Cisternae of the rough endoplasmic reticulum and Golgi complexes are found randomly in the cytoplasm. We observed numerous small vesicles with electron-dense material and a few lysosomes in the cytoplasm (Fig. 7). The regenerative cells with irregular contours and round or ovoid nuclei characterized by disperse euchromatin and at least one electron-dense nucleolus form the nidi. Finally, there are the endocrine cells
in-between the digestive cells that are slim contacting the basement membrane and the gut lumen. They contain a deformed nucleus, a few mitochondria, rough endoplasmic reticulum and a Golgi apparatus.

In the midgut epithelium of imidacloprid-treated adult desert locusts, the columnar digestive cells display a comparably high vacuolization of the basal cytoplasm and have an irregularly shaped nucleus with a nucleolus and fragmented heterochromatin. The brush border of these cells is disorganized; the microvilli are irregularly thick and distorted. Mitochondria are not uniformly shaped and have altered cristae; they do accumulate at the apical periphery but are distributed throughout the cell (Fig. 7). Membranes representing rough endoplasmic reticulum were bloated and short; fragmented cisternae were observed in the apical portion of the cell. In addition to these structures, we found a few lipid droplets and some lysosomes in these cells. Interestingly, we identified many autophagic vacuoles in imidacloprid-treated locusts. Regenerative cells were deformed and protruded from the surface into the lumen; the nests of regenerative cells were disorganized, reduced in size and number. Closer examinations of the regenerative cells revealed features of necrosis, which is indicated by nuclei with corrugated nuclear membranes, formation of myelinic structures, a few mitochondria, smooth endoplasmic reticulum, a few lipid droplets and various-sized lysosomes (Fig. 7). The ultrastructural changes of the midgut epithelial cells in control and imidacloprid-treated adult L. migratoria are summarized in Table 1.

**Discussion**

**Field locusts are exposed to sub-lethal concentrations of imidacloprid**

Upon uptake by feeding, xenobiotics travel through the digestive tract passing different types of cells. In this work, we show that field captured locusts display a physiological and morphological response to imidacloprid exposure.

Locusts collected on the agricultural site have a lowered AChE activity. To our best knowledge, imidacloprid is the only insecticide applied on the sample collection site. We cautiously conclude that inhibition of AChE activity is due to the presence of imidacloprid. As neonicotinoids cause an immediate paralysis and death of the insect (Bass and Field 2018), we conclude that imidacloprid concentrations in our field caught animals were sub-lethal. The sub-lethal effect of neonicotinoids including imidacloprid has been reported for a number of insect species especially in the honeybees *Apis mellifera* or *cerana* and the model insect *Drosophila melanogaster* (Delkash-Roudsari et al. 2020; Martelli et al. 2020). In the honeybee, chronic exposure

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**Fig. 7** Transmission electron microscopy of control (A–D) and treated (E–H) locust midgut epithelium: (A–C) showing basal and supranuclear region of the digestive cells. (D) showing nidi of regenerative cells with normal nucleus and regular nuclear envelope. Notice: The well-defined basement membrane (BM); junctional complex (JC); circular muscle (CM); nucleus (N) with well-developed nucleoli; vesicles containing granular material (V); rough endoplasmic reticulum (RER); mitochondria (M); secretory vesicles (SV); abundant microvilli (Mv) extending into the lumen (L); serosal barrier (SB); tracheole (T). (E and G) Digestive cells (DC) with disrupted microvilli (Mv), swollen shape and distribution of the mitochondria (M), multivesicular bodies (Mb). (F and H) Regenerative cells with nucleus (N); eccentric and segre gated nucleolus (Nu) and heterochromatin distributed within the nucleoplasm and along the irregular nuclear envelope (Ne). notice: dilated rough endoplasmic reticulum (RER); Golgi apparatus (G) secretory granules (Sg); myelinated fibers (curved arrow)
to low imidacloprid concentration interferes with learning and locomotion (Delkash-Roudsari et al. 2020; Gao et al. 2020). In D. melanogaster, low imidacloprid concentrations induce production of ROS that in turn causes neurological and metabolic defects (Martelli et al. 2020).

The midgut is a target of imidacloprid

An organ recurrently affected by neonicotinoids including imidacloprid is the locust midgut. In various insect species such as Apis mellifera (honeybee), Melipona quadrifasciata (a wild bee), Podisus nigrispinus (a predatory bug) and Bombyx mori (silkworm) especially low concentrations of imidacloprid or acetamiprid cause, among others, reduction and malformation of the microvilli. Defective microvilli may, in turn, impede food processing and thereby reduce insect fitness. The midgut phenotype seems not to be specific to imidacloprid or neonicotinoids in general. Application of chlorantraniliprole, for instance, that does not belong to the neonicotinoid class of insecticides and acts on the insect ryanodine receptor, causes similar defects in the midgut of Anticarsia gemmatalis, a moth (Castro et al. 2019). The pyrethroid insecticide lambda-cyhalothrin also impacts, among other organs, on the midgut of honeybee workers (Arthidoro de Castro et al. 2020). Thus, the non-primary target organ midgut seems to be sensitive to the function of some unrelated insecticides. What could be the mechanism? The central nervous system is the primary target organ of most insecticides including neonicotinoids, pyrethroid and chlorantraniliprole. As a composite organ, the midgut is surrounded by muscles that are enervated by neurons and communicates with the central nervous system via peptides and hormones (Wu et al. 2020). In a simple and speculative scenario, hence, perturbed communication of the defective central nervous system with the midgut may be the reason for midgut cell deformations and malfunction. Alternatively, direct physiological consequences of insecticide function including ROS production may explain the defects observed in the midgut.

Imidacloprid perturbs ROS metabolism in field locusts

Indeed, we found that imidacloprid caused a change in the activity of enzymes that are involved in ROS metabolism. Enzymes needed to eliminate ROS showed lower activity, while molecular responses to elevated ROS levels such as protein carbonylation were enhanced. What is the underlying molecular mechanism? In insects, xenobiotics including insecticides have been repeatedly shown to induce ROS production (Palli 2020). To some extent, hence, the adverse effects of insecticides are not only due to their inhibitory function on their direct target, but also on their capability to promote ROS production that is accompanied by the detoxification program triggered by the contact of the organism with the insecticide. Several pathways are induced in this process (Amezian et al. 2021). Commonly, they regulate the expression of genes coding for detoxification enzymes including P450s that are involved in redox reactions. Consistently, in our field locusts exposed to imidacloprid the activity of P450 enzymes is elevated in the intestines. P450s are known to generate ROS (Li et al. 2021); together with the observation that the activity of those enzymes eliminating ROS declined in our field animals, we believe that due to hyperactivity of P450 enzymes our locusts suffer massive oxidative stress that in turn may be responsible for cell degeneration in the midgut. Indeed, we detected higher levels of apoptotic cells in locusts exposed to imidacloprid than in control locusts. Our data underline that these defects are not only accessible to experimental setups in the laboratory or mesocosm but also in the field where

| Table 1 Summary of ultrastructural changes of the midgut epithelial cells in control and treated old adult L. migratoria | Control | Treated |
|---------------------------------------------------------------|---------|---------|
| Shape of epithelial cells | Columnar | Swollen |
| Microvilli | Well-developed and continuous throughout the apex of the cell | Packed, missing in some regions |
| Cytoplasm | Dense and granular | Less granular |
| Nucleus | Oval | Round, elongated |
| State of mitochondria | Normal with cristae well developed | Swollen, many in process of lysis |
| Distribution of mitochondria | Lying mostly in the apical and basal regions of the cell | Scattered throughout the cell |
| Endoplasmic reticulum | Organized with ribosomes | Less dense and dilated |
| Golgi complex | Well-developed | Present |
| Lysozyme | Present | Present |
| Lipid bodies | Present (few) | Present |
| Autophagic vacuoles | Present | Many in numbers |
the actual insecticide-insect interaction takes place during pest management.

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Data availability Raw data and all materials are available upon request.

Declarations

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