Screening of Carbamate and Organophosphate Pesticides in Food Matrices Using an Affordable and Simple Spectrophotometric Acetylcholinesterase Assay

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Abstract: Carbamates (CMs) and organophosphates (OPs) are widely used pesticides with known neurotoxicity arising from the inhibition of acetylcholinesterase (AChE). When AChE is active, in vitro, it can hydrolyze certain substrates to colored products while in the presence of an inhibitor this color development is decreased. Based on this principle, an AChE assay for CM and OP compounds was optimized and validated for carbofuran, carbofuran-3-hydroxy and dichlorvos in lettuce and strawberry extracts. The analytical performance of the assay was confirmed by an accredited liquid chromatography tandem mass spectrometry (LC–MS/MS) method. The developed AChE assay achieved low limits of detection (LODs) at the part per billion (ppb) level, depending the analyte inhibitory strength, recovery rates higher than 70% and good repeatability. Moreover, the toxic unit (TU) approach was applied, for extracts containing the validated analytes, and antagonism was noticed in all cases. Overall, the developed method is rapid, simple, cost-effective and may find application as a low-cost pre-screening tool of AChE inhibitors presence. Last but not least, this study can be considered a guide on development, validation and benchmarking of bioassays in food safety, a topic, which is commonly misrepresented in the available literature.

Keywords: carbamates; organophosphates; pesticides; acetylcholinesterase; food; screening method; microwell-plate assay; liquid chromatography tandem mass spectrometry

1. Introduction

The ever-increasing demand for food production requires a widespread use of pesticides to prevent pests and boost agricultural productivity. Pesticides encompass a numerous and diverse group of chemical compounds that can be used to combat a wide variety of pests. Regarding insecticides, carbamates (CMs) and organophosphates (OPs) are commonly utilized because, compared to old-school organochlorines (OCs), they do not usually release toxic metabolites and are less persistent in the environment [1]. However, CMs and OPs show relatively high acute toxicity and their residues in food may result in chronic toxicity incidents [2].

The toxicity of CMs and OPs arises from the inhibition of acetylcholinesterase (AChE), a vital enzyme in the neural system of insects and mammals including humans. Typically, AChE hydrolyzes the neurotransmitter acetylcholine into choline and acetic acid (Figure 1a), an essential reaction that enables the cholinergic neuron to return to its resting state after activation. However, AChE activity is reduced in the presence of CMs or OPs, due to carbamylation or phosphorylation of serine hydroxyl group in the enzyme active cite [3], respectively. This results in acetylcholine accumulation, which can
lead, in case of higher exposure, to health problems, including bradycardia, salivation, diarrhea or even coma [4]. A complete overview on AChE physiological function and inhibition mechanisms published by Miao et al. [5] may provide a deeper understanding of the issue. Clearly, there is an urgent need to develop sensitive and accurate analytical methods for OPs and CMs detection considering their impact on health and the potential of dietary exposure as this was reflected in the latest European Food Safety Authority (EFSA) report on pesticides [6] and the EFSA pesticide residue intake model (PRIMO) [7].

Liquid and/or gas chromatography tandem mass spectrometry (LC–MS/MS and/or GC–MS/MS) are commonly applied for pesticide residues monitoring in food matrices [8]. Although these methods provide selectivity, sensitivity with high-throughput capabilities, LC/MS and GC/MS are also costly and time-consuming methodologies that require skilled personnel [9]. To counter these limitations, an ever-increased use of sensor-based methods has been noticed [10]. Various immunoassays were developed against specific CMs and OPs [11,12]. Although this approach may be even more sensitive than chromatographic methods, it detects one or only a limited number of pesticides. Regarding electrochemical (EC) sensors, there were several studies using both enzymatic [13] and non-enzymatic pesticide recognition [14] employing different EC principles, such as amperometry [15] and voltammetry [16]. Despite reaching low limits of detection (LODs), mostly in buffers, these methods suffered from complicated manufacturing and problematic storage stability period [17]. Considering optical detection, surface-enhanced Raman spectroscopy (SERS), surface plasmon resonance (SPR) and spectroscopic methods have been used. SERS has been emerged as an attractive analytical choice due to its portability potential. However, pesticides detection only on fruit peels [18] was a major weakness limiting its application. Although SPR and fluorescent methods showed promising results, especially when combined with nanomaterials (NMs) [19], their sophisticated analytical configuration comes in contrast to the need for simplified alternative methods.

AChE assays may be considered as a potential solution for the aforementioned analytical needs. These assays are based on the principle that the enzyme, in vitro, can produce a colored product (Figure 1b). Under relevant conditions, in the presence of an analyte the residual AChE activity (meaning less color production) can be correlated to an inhibitor concentration by comparing this response with a control sample. The simplest way to monitor this color development is by using an absorption spectroscopy reader, which is a cost efficient, reliable and semi-portable (it fits into a backpack) detector. This principle can be applied for the rapid monitoring of CMs and OPs in food

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**Figure 1.** (a) In vivo, the neurotransmitter acetylcholine is hydrolyzed to choline and acetic acid by AChE; (b) in vitro hydrolysis of various substrates (acetylthiocholine iodide (AThi), Ellman’s assay substrate), indoxyl acetate (IDA) and α-naphthyl acetate (α-NAc) by AChE producing different colored products.
extracts providing high throughput, when using a 96-microwell plate, minimum reagents consumption and LODs at the ppb level, depending on the matrix [20,21].

In this study, the development and validation of an AChE assay for CM and OP pesticides is presented. During assay optimization, various parameters were investigated including a critical comparison of commonly applied AChE substrates for the first time. Following our recently published paper, which addressed the issues related to the validation and benchmarking of biosensors in food safety [8], comprehensive validation datasets for carbofuran, carbofuran-3-hydroxy and dichlorvos in strawberry and lettuce extracts are presented. The optimized AChE assay was benchmarked towards an accredited LC–MS/MS method, following the regulatory requirement (Decision 2002/657/EC) of confirming bioassays results with an instrumental method.

2. Materials and Methods

2.1. Chemicals

Phosphate buffer saline (PBS) tablets, AChE from *Electrophorus electricus*, acetylthiocholine iodide (AThI, purity >99%), 5,5′-dithio bis-2-nitrobenzoic acid (DTNB, purity >99%) indoxyl acetate (IDA, purity >95%), α-naphthyl acetate (α-NAc, purity >98%), fast blue B salt (FBBS, purity 95%), z-sep sorbent, 0.22 µm syringe filters, 5 mL syringes were from Sigma-Aldrich (Prague, Czech Republic). 96-microwell plates were supplied by Gama Group (Ceske Budejovice, Czech Republic). Acetylcholinesterase stock solution in PBS (2 U µL⁻¹) was stored in the fridge at 4 °C. Ethanol (EtOH), methanol, dimethyl sulfoxide, acetonitrile and ethyl acetate were supplied as high purity solvents from Merck (Darmstadt, Germany) and Supelco (Bellefonte, PA, USA). Triphenyl phosphate (TPP), ammonium formate (>99%), and formic acid were from Sigma–Aldrich (Taufkirchen, Germany). Anhydrous magnesium sulphate (MgSO4) was obtained from Fluka (Buchs, Germany), and sodium chloride (NaCl) from Penta (Chrudim, Czech Republic). Sorbent primary–secondary amine (PSA, Bondesil, 40 µm) was purchased from Agilent Technologies (USA). Deionized water was purified using a Milli-Q system (Millipore; Bedford, MA, USA). Aldicarb, carbofuran, carbofuran-3-hydroxy, carbaryl, dichlorvos, chlorpyrifos and paraoxon standards were of analytical standard grade and purchased by Sigma Aldrich (Taufkirchen, Germany). Malathion (purity 99%), malaoxon (purity 99%) and parathion (purity 99%) were bought by Dr. Ehrenstorfer GmbH (Augsburg, Germany). Chlorpyrifos-oxon was of analytical standard grade and supplied by Labicom (Olomouc, Czech Republic). Pesticides stock solutions (3.0 mg mL⁻¹) were prepared in acetonitrile and kept in a freezer at −20 °C protected from ambient light. Pesticides working solutions were daily prepared in PBS-EtOH (7:3, v/v).

2.2. In Vitro AChE Assays

Ellman’s [22], IDA [23] and α-NAc [24] assays were tested to choose which one fits better for the analytical purpose. In all cases, the experimental protocols were modified and transmitted the assay to a microwell plate format. Briefly, in all assays the enzyme was incubated with an inhibitor for 15 min, followed by a substrate addition and absorbance measurement in an Epoch Biotek Reader (Vermont, USA). Assay protocols are further explained in the Supplementary Materials.

The experiments were performed in three independent days, in triplicate and the data were pooled (n = 9 for each calibration point). To obtain the 20% of the maximal and half maximal inhibitory concentration (IC₂₀ and IC₅₀, respectively), a pesticide was prepared in seven different concentration levels (Table S1) and compared to a control sample. Dose-response with variable slope analysis was performed using GraphPad prism 5.0 software. More information on the four-parameter dose-response model can be found in [25].

2.3. AChE Assay Specificity

CMs and OPs have the same mode of action towards AChE, thus, it is important to evaluate AChE specificity towards the tested inhibitors. Assay specificity was evaluated in terms of cross-reactivity
towards carbofuran, which was the strongest AChE inhibitor (see Section 3.1.2). The cross-reactivity was calculated as follows:

Cross-reactivity (%) = \( \frac{IC_{50, \text{carbofuran}}}{IC_{50, \text{pesticide}}} \times 100 \), carbofuran cross-reactivity was set as 100%.

2.4. Food Samples

Strawberry and lettuce bio-samples were purchased by the local market. A representative portion was homogenized with a PM-100 Retsch Planetary Ball Mill (Haan, Germany) and the samples were kept at \(-20^\circ\text{C}\). Prior to the enzyme assay, the samples were analyzed by an LC–MS/MS method to verify that they did not contain detectable levels of OPs and CMs.

2.5. Food Samples Extraction

A modified ethyl acetate extraction [26] followed by a dispersive clean up-step using a zirconium based sorbent (z-sep) was developed to process strawberry and lettuce samples. In detail, 10 g of homogenized samples were weighed into a 50 mL polypropylene tube. When needed, samples were spiked with target analytes from a freshly prepared solution (20 \( \mu \text{g mL}^{-1} \)) to achieve the final concentration and left for at least 30 min at room temperature. Afterwards, 10 mL of ethyl acetate were added and tubes were shaken vigorously for 2 min with hands, followed by the addition of 10 g of Na\(_2\)SO\(_4\) and the tubes were shaken for 30 s. Then, 2 mL from the supernatant were transferred to a 15 mL polypropylene tube containing 65 mg z-sep and 300 mg Na\(_2\)SO\(_4\) and the tubes were shaken vigorously for 1 min. Then, the supernatant was filtered using 0.22 \( \mu \text{m} \) syringe filters, and 500 \( \mu \text{L} \) of the filtrate were transferred to a vial. The filtrate was evaporated close to dryness under a gentle nitrogen steam, reconstituted to 200 \( \mu \text{L} \) PBS:EtOH (7:3, v/v) and subjected to the enzyme assay. The same samples were also analyzed using an accredited LC–MS/MS method [27] as a reference to the enzyme assay.

2.6. Liquid Chromatography-Triple Quadrupole Mass Spectrometry Analysis

A 1290 Infinity II LC system (Agilent Technologies, Santa Clara, CA, USA) coupled to a Triple Quadrupole G6495 (Agilent Technologies) mass spectrometer with electrospray ionization in positive ion mode (ESI+) was used for pesticide residue analysis. Dynamic multiple reaction monitoring (dMRM) mode was selected as operation mode. The separation took place in an Acquity UPLC HSS T3 analytical column (100 mm \( \times \) 2.1 mm i.d. \( \times \) 1.8 \( \mu \text{m} \) particle size, Waters, Milford Massachusetts, USA) with controlled temperature at 40 \( ^\circ\text{C} \), using water and MeOH (with the addition of ammonium formate and formic acid) as a mobile phase and a gradient described in [28]. The obtained data were processed using MassHunter software version B.07.00.

2.7. Methods Validation

The analytical performance of the AChE assay was fully evaluated in terms of repeatability, trueness and limits of detection (LODs). Strawberry and lettuce samples were spiked with carbofuran or carbofuran-3-hydroxy or dichlorvos. In the case of carbofuran, repeatability and trueness testing were performed in six-fold at two concentration levels (0.020 and 0.100 mg Kg\(^{-1}\)). Aliquots of the control extracts were used to prepare matrix matched calibration standards at 0.005, 0.010, 0.020, 0.050, 0.100, 0.250 and 0.500 mg Kg\(^{-1}\) for the enzyme assay. In terms of carbofuran-3-hydroxy and dichlorvos, repeatability and trueness testing were performed at 0.300 and 1.20 mg Kg\(^{-1}\) as they were weaker inhibitors compared to carbofuran. The calibration range was also different for these two compounds, in detail 0.078, 0.150, 0.300, 0.600, 1.20, 2.5 and 5.0 mg Kg\(^{-1}\). The LODs were calculated using the formula

\[
\text{LOD enzyme assay} = \frac{\text{meanblank}}{3.3 \times \text{SDblank}}, \quad n = 6 \text{ blank samples}
\]
To investigate the assay quantification capability in the presence of multiple residues, as well as to monitor their toxicity potential, the widely accepted toxic unit (TU) approach was applied in matrix matched calibration curves containing two analytes [29]. The calibration range for each analyte was based on the IC$_{50}$ obtained in the matrix matched calibration curve. In detail:

- Three points below the IC$_{50}$ value, IC$_{50}$/8, IC$_{50}$/4, IC$_{50}$/2;
- One at the IC$_{50}$;
- Three points above the IC$_{50}$, 2 × IC$_{50}$, 4 × IC$_{50}$, 8 × IC$_{50}$.

Moreover, a TU is the sum of the toxic contributions of each component in the mixture. The calculation formula for TU is the following:

$$\text{TU} = \frac{\text{IC}_{50} \text{A (mix)}}{\text{IC}_{50} \text{A (individual)}} + \frac{\text{IC}_{50} \text{B (mix)}}{\text{IC}_{50} \text{B (individual)}}$$

where A and B are pesticides, IC$_{50}$ (mix) is the effect of each component in the binary mixture and IC$_{50}$ (individual) is the individually induced inhibitory effect, at the IC$_{50}$ level. The TU model can inform about the joint inhibitory effect type, with TU equals to one meaning additive effect, in other words 0.5 TUA + 0.5 TUB = 1 TUA + B. Alternatively, TU < 1 indicates synergistic effect whilst TU >1 shows antagonism between the inhibitors.

Finally, an accredited LC–MS/MS method was also applied for the same samples to compare the assay performance. Repeatability and trueness were also tested at the same conditions as for the enzyme assay and matrix matched calibration standards were prepared at 0.001, 0.002, 0.005, 0.010, 0.020, 0.050 and 0.100 mg Kg$^{-1}$. The LODs were calculated using the formula.

$$\text{LOD LC-MS/MS} = \frac{\text{SD regression}}{\text{slope}} \times 3.3.$$

3. Results and Discussion

As mentioned in the Introduction, the potential dietary intake of CM and/or OP residues poses a threat for human health due to their neurotoxicity. However, unless sophisticated laboratory instrumentation is available, then controlling residues is in most cases a rather difficult task. On this account, we focused on the development of a rapid, simple and cost-effective method capable to detect CMs and OPs in food matrices. In this study, critical assay parameters such as enzyme tolerance towards organic solvents (see Supplementary Materials), enzyme concentration and the enzyme substrate were optimized. In addition, AChE specificity towards 11 widely applied OP and CM pesticides was investigated since all these compounds have the same inhibitory mode of action. Based on this information, the assay was validated for carbofuran (a CM compound), carbofuran-3-hydroxy (a carbofuran toxic metabolite) and dichlorvos (an OP compound) in strawberry and lettuce using a fast and simple ethyl acetate extraction. Additionally, the AChE assay results were confirmed by an accredited LC–MS/MS method. Finally, the TU approach was applied to investigate the effect of mixtures containing two pesticide residues, as a simplified analogue towards real life conditions, since samples may be contaminated with multiple residues.

3.1. AChE Assay Optimization

3.1.1. AChE Concentration and Substrates

The suitability of conceivable substrates yielding color products [22–24] was considered. Comparing AThI:DTNB (Ellman’s reagent), IDA and α-NAc for AChE activity measurement, the lower the enzyme concentration the lower the obtained IC$_{50}$; this shows the clear connection between enzyme activity and assay sensitivity. Importantly, Ellman’s assay needed about 80 and 120 times less enzyme per well (compared to IDA and α-NAc assays), indicating the higher affinity of AChE towards AThI (Ellman’s assay substrate; Table 1). Regarding the total assay duration, it lasted from 15 (Ellman’s and α-NAc assays) to 45 min (IDA assay), depending on the assay used. Significantly, all assays were quite
affordable, with the cost of the Ellman’s assay being surprisingly low at about 0.16 euro per plate. Therefore, Ellman’s assay was selected as it combined the best sensitivity with the least cost.

Table 1. Critical comparison of Ellman’s, IDA and α-NAc assays.

| Assay Characteristic | Ellman’s | IDA | α-NAc |
|----------------------|----------|-----|-------|
| product color        | yellow (412 nm) | blue (670 nm) | purple (525 nm) |
| AChE activity, U well$^{-1}$ | 0.009 | 0.75 | 1.1 |
| total assay time     | 15 min | 45 min | 15 min |
| $IC_{50}$ for dichlorvos, µM, n = 9 $^1$ | 3.5 (1.7–7.0) | 13 (8.3–22) | 11 (7.4–19) |
| $IC_{50}$ for carbofuran, µM, n = 9 $^1$ | 0.099 (0.092–0.11) | 0.27 (0.22–0.36) | 0.23 (0.18–0.31) |
| cost, euro/plate     | 0.16 | 14 | 22 |

$^1$: 95% confidence intervals in brackets.

3.1.2. Studied CMs and OPs Inhibitory Strength and AChE Assay Specificity

The inhibition potency of the studied pesticides largely differed, with carbofuran being the strongest inhibitor followed by chlorpyrifos-oxon and malaoxon (Table 2). Worthy to notice that the tested oxon forms (OPs with a P=O moiety) were much stronger inhibitors compared to the thiono analogues (OPs with a P=S moiety). This fact was expected due to the low sulfur electronegativity of the P=S moiety resulting in low reactivity with the serine hydroxyl group at the active site of AChE. On the other hand, oxon forms (P=O) have relatively higher affinity for AChE active site compared to acetythiocholine (Ellman’s substrate), thus they undergo nucleophilic substitution reaction by serine hydroxyl group resulting in a covalent binding of a pesticide to AChE [30]. Regarding the oxidation, this physiologically takes part during the metabolism, probably by the mixed function oxidases (MFO), resulting in a stronger AChE inhibitor [31]. Except the oxidation, which can occur at the P=S bond, also hydrolysis is possible [32], for example under acidic induced conditions resulting in phosphate ester hydrolysis and phosphoryl transfer [33]. Enzymatic hydrolysis of organophosphates also acts as a detoxification process exploiting various esterases and amidases depending the organism [34].

To assess AChE specificity towards carbofuran (the strongest AChE inhibitor), the cross-reactivity was calculated (Table 2). In the case of CM compounds, the cross-reactivity with aldicarb and carbaryl was less than 2% whilst with carbofuran-3-hydroxy (a carbofuran metabolite) was about 14% meaning that about 10-times higher concentration of carbofuran-3-hydroxy is necessary to achieve the same inhibition as in the case of carbofuran. Regarding chlorpyrifos, a widely used insecticide occasionally exceeding the EU maximum residue limits (MRLs) in various matrices, a very low cross-reactivity was calculated (about 0.024%). Nevertheless, chlorpyrifos-oxon and malaoxon showed a significant cross-reactivity (about 45% and 25%, respectively), which cannot be considered a problem for carbofuran detection in real food matrices because the P=S bond is metabolically oxidized to P=O. In other words, chlorpyrifos-oxon and malaoxon forms cannot occur in plant crops. In terms of dichlorvos, paraoxon, malathion and parathion, the cross-reactivity was negligible (<3% in all cases). All in all, a minor cross reactivity was noticed only for carbofuran-3-hydroxy.
Table 2. Inhibitory strength of tested CMs and OPs in terms of IC$_{20}$ and IC$_{50}$ and AChE specificity towards carbofuran in terms of cross-reactivity %.

| Pesticide            | Structure | IC$_{20}$, µM$^1$ | IC$_{50}$, µM$^1$ | Cross-Reactivity % |
|----------------------|-----------|-------------------|-------------------|--------------------|
| carbofuran           | ![Carbofuran](image) | 0.037 (0.033–0.041) | 0.099 (0.092–0.11) | 100                |
| aldicarb             | ![Aldicarb](image) | 2.1 (1.1–4.0)     | 6.1 (4.4–8.5)     | 1.6                |
| carbofuran-3-hyrdoxy | ![Carbofuran-3-Hydroxy](image) | 0.25 (0.20–0.30) | 0.72 (0.62–0.83) | 14                 |
| carbaryl             | ![Carbaryl](image) | 2.1 (1.1–3.7)     | 6.2 (4.5–8.5)     | 1.6                |
| chlorpyrifos         | ![Chlorpyrifos](image) | 251 (157–401)     | 420 (298–594)     | 0.024              |
| chlorpyrifos-oxon    | ![Chlorpyrifos-Oxon](image) | 0.081 (0.050–0.13) | 0.22 (0.16–0.30) | 45                 |
| dichlorvos           | ![Dichlorvos](image) | 1.2 (0.80–1.9)    | 3.5 (1.7–7.0)     | 2.8                |
| malaoxon             | ![Malaoxon](image) | 0.92 (0.49–1.8)   | 0.40 (0.25–0.62)  | 25                 |
| malathion            | ![Malathion](image) | 85 (72–96)        | 294 (220–305)     | 0.034              |
| paraoxon             | ![Paraoxon](image) | 0.92 (0.49–1.8)   | 2.9 (1.7–4.8)     | 3.4                |
| parathion            | ![Parathion](image) | 125 (102–145)     | 357 (330–395)     | 0.028              |

$^1$: 95% confidence intervals in brackets.
3.2. Optimized AChE Assay Application in Lettuce and Strawberry

As carbofuran was the strongest AChE inhibitor tested, it was chosen for assay performance demonstration in real life samples. Carbofuran screening is also important because despite being an unauthorized pesticide in the EU, it still remains a problem, which can potentially impact consumer’s health. For example, carbofuran detection in goji berries from China resulted in the 2018 amendments of Regulation (EC) No 669/2009, which prescribes more frequent official controls at the point of entry into the EU territory [6]. During the latest European Pesticide Residue Workshop (EPRW) in 2018, carbofuran was detected in the so called “superfoods” such as goji berries and matcha tea [35], a group of “nutrient-rich” trending foodstuffs without a clear legal definition. Usually, these products are imported from countries where legislative regulatory mechanisms, requirements and official control systems are different than those in the European countries.

The assay was validated also for the detection of carbofuran-3-hydroxy (a carbofuran toxic metabolite, which is included also in carbofuran residue definition, https://ec.europa.eu/food/plant/pesticides/eu-pesticides-database/public/?event=activesubstance.detail&language=EN&selectedID=1082, last accessed 03/12/2019) and dichlorvos (an OP) since they share a common inhibitory effect against AChE (see Supplementary Materials). To investigate the cumulative inhibitory effect in food extracts, the TU approach was applied and matrix matched calibration curves containing two of the analytes were measured. Regarding the tested matrices, lettuce and strawberry were selected since they are crops in which carbofuran and dichlorvos were earlier reported to reach the toxicological reference value [36]. Another reason for selecting these matrices is their different composition mainly meaning different natural pigments transferred into extracts, which can potentially affect the AChE assay performance.

3.2.1. Carbofuran Validation and Benchmarking

Carbofuran analysis was successfully validated using the optimized AChE assay in lettuce and strawberry (Table 3). The obtained recoveries (n = 6) at two spiking levels, 0.020 mg Kg⁻¹ and 0.100 mg Kg⁻¹, were higher than 70% in both matrices. Better repeatability was noticed in lettuce compared to strawberry and a LOD equal to 0.013 mg Kg⁻¹ was achieved for lettuce and 0.012 mg Kg⁻¹ for strawberry. Worthy to notice is that LODs do not allow us to determine the measurement of uncertainty and this is a reason why screening methods results need to be verified by instrumental methods. Other spectrophotometric AChE assays have also achieved LODs at the ppb level [20,21], however, insufficient validation was reported (solely focused on LODs). Comparing to EC methods, an AChE based EC sensor was utilized for carbofuran detection in cucumber achieving worse LODs than our study (0.22 µM meaning 0.048 mg Kg⁻¹) [37]. In another paper, a non-enzymatic EC sensor was used and lower LOD was reported for carbofuran residues in grape, orange, tomato and cabbage matrices [14]. However, such a sensor is not available at the market probably due to a complicated manufacture.

Although it was not feasible to achieve LODs lower than EU MRLs (0.002 mg Kg⁻¹ and 0.005 mg Kg⁻¹ for lettuce and strawberry, respectively), the developed assay was able to detect carbofuran about two times lower than the Chinese MRL (0.020 mg Kg⁻¹ for fruits and vegetables) [38] and ten times lower than the Indian MRL (0.100 mg Kg⁻¹ for fruits and vegetables) [39]. Our method can enhance food testing in these countries, as carbofuran is commonly detected in foodstuffs originating from China [40] or India [41]. Additionally, the AChE assay can be used, in the EU, as a cost-effective pre-screening tool for imported food. Moreover, the developed method can be beneficial also for checking the EU acute reference dose (ARfD) exceedance. In detail, carbofuran ARfD is equal to 0.00015 mg Kg⁻¹ Bw⁻¹ and the AChE assay could enable the concentration control that might, under certain consumption conditions, approach or even exceed it. Considering, e.g., an individual with a 70 kg body weight, the ARfD is 0.0105 mg, and then this value would be achieved when eating 150 g of a contaminated foodstuff containing 0.070 mg Kg⁻¹ of carbofuran residues. In fact, such cases have been recently reported in the EU Rapid Alert System for Food and Feed (RASFF) portal (shorturl.at/quxKO,
last accessed 10/01/2020). For example, carbofuran was detected at concentrations higher than 0.070 mg Kg$^{-1}$ both in eggplants and goji berries in Germany, Czech Republic and Switzerland the last two years.

To benchmark AChE assay analytical performance, the same strawberry and lettuce samples were analyzed by an accredited LC-MS/MS reference method (Figure 2). Recoveries ($n=6$) at two spiking levels, 0.020 mg Kg$^{-1}$ and 0.100 mg Kg$^{-1}$, were higher than 80% in both matrices by using the Quick, Easy, Cheap, Effective, Rugged and Safe (QuEChERS) extraction. Although the QuEChERS extraction has been addressed as the golden standard for multiple residues extraction, an ethyl acetate extraction was modified for the AChE assay. In contrast to QuEChERS extraction, which includes several analytical steps, the developed ethyl acetate extraction, did not include any centrifugation step resulting in a faster extraction protocol. Another advantage was that ethyl acetate can be evaporated considerably faster compared to acetonitrile (QuEChERS solvent) due to a lower boiling point and thus a reduced extraction time was achieved. Furthermore, acetonitrile is related to difficulties associated with the removal of water molecules, therefore, ethyl acetate (a less polar solvent) fits better to the sample preparation simplification concept. Considering the LC–MS/MS method repeatability, it was significantly lower, less than 2%, while LOD was equal to 0.0014 mg Kg$^{-1}$. Comparing to the AChE assay, LC–MS/MS precision was fairly better and the LOD lower about one order of magnitude due to the highly selective and sensitive tandem mass detector.

| Spiking Level (mg Kg$^{-1}$) | Lettuce | Strawberry | Lettuce | Strawberry |
|-------------------------------|---------|------------|---------|------------|
| 0.020                         | 91      | 0.020      | 79      | 0.013      |
| 0.100                         | 94      | 0.020      | 91      | 0.012      |

1 R%: recovery %. 2 RSDr%: Relative standard deviation under repeatability conditions %.

Table 3. Carbofuran validation results in strawberry and lettuce for AChE assay and LC–MS/MS method.

Figure 2. Cont.
at concentrations higher than 0.070 mg Kg$^{-1}$ both in eggplants and goji berries in Germany, Czech Republic and Switzerland the last two years.

Table 3. Carbofuran validation results in strawberry and lettuce for AChE assay and LC–MS/MS method.

| Method      | AChE Assay | LC–MS/MS |
|-------------|------------|----------|
| Matrix      | Lettuce    | Strawberry |
| Spiking Level (mg Kg$^{-1}$) | 0.020 | 0.020 | 0.100 | 0.100 |
| R%          | 91         | 71       | 79     | 78     |
| RSDr%       | 5.2        | 19       | 1.9    | 1.2    |
| LOD (mg Kg$^{-1}$) | 0.013 | 0.0014 | 0.012 | 0.0012 |

**Figure 2.** Extracted ion chromatograms (XIC) of carbofuran, carbofuran-3-hydroxy and dichlorvos at 0.020 mg Kg$^{-1}$ in (a) lettuce and (b) strawberry matrix matched solutions using the LC–MS/MS system.

3.2.2. TU Approach in Lettuce and Strawberry Extracts

The co-occurrence of multiple pesticide residues in a single sample is not unusual indicating the need to evaluate their potential cumulative effect on the AChE assay. Actually, this topic has drawn EFSA interest, which recently published an opinion on cumulative risk assessment to pesticides affecting the nervous system [42]. In this way, mixtures containing two analytes (carbofuran, carbofuran-3-hydroxy or dichlorvos) were prepared in lettuce and strawberry blank extracts. The TU approach was applied and antagonism was noticed for all the binary pesticide mixtures in both matrices (Table 4). Important to notice is that in previous studies the cumulative inhibitory effect was evaluated only in buffers [29,43]. The presence of food matrix components in extracts may alternate the results of TU approach as synergistic effect was reported for a carbofuran-malathion mixture whilst mild antagonism for a carbofuran-chlorpyrifos mixture [29]. The cumulative effect may also vary in vivo, for example additive or synergistic effect was reported for salmon exposed to binary CM and/or OP mixtures [44]. In any case, implications related to pesticide cocktail effects should be considered carefully [45].
Table 4. IC_{50} for the tested binary mixtures in lettuce and strawberry extracts. TU were calculated as explained in Section 2.7. TU < 1, TU = 1, TU > 1 shows synergism, dose addition and antagonism, respectively.

| Mixture                        | Matrix          | Analyte                  | Lettuce |                           |                           |                           |                           |                           |                           |
|--------------------------------|-----------------|--------------------------|---------|---------------------------|---------------------------|---------------------------|---------------------------|---------------------------|---------------------------|
| carbofuran + carbofuran-3-hydroxy | carbofuran      | carbofuran-3-hydroxy     | 0.055 (0.038–0.078) | 0.041 (0.021–0.080)  | 1.70 (antagonism)        | 0.050 (0.032–0.065)  | 0.043 (0.022–0.098)  | 1.74 (antagonism)        |
| Carbofuran + dichlorvos        | carbofuran      | dichlorvos               | 0.055 (0.038–0.078) | 0.041 (0.021–0.080)  | 1.50 (antagonism)        | 0.050 (0.032–0.065)  | 0.037 (0.018–0.095)  | 1.52 (antagonism)        |
| carbofuran-3-hydroxy +        | carbofuran-3-hydroxy | dichlorvos               | 0.055 (0.038–0.078) | 0.041 (0.021–0.080)  | 1.50 (antagonism)        | 0.050 (0.032–0.065)  | 0.037 (0.018–0.095)  | 1.52 (antagonism)        |
| dichlorvos                     |                 |                          | 0.78 (0.41–1.5)     | 2.8 (2.2–3.5)        | 2.5 (antagonism)         | 0.85 (0.55–1.8)     | 0.95 (0.75–3.0)      | 2.14 (antagonism)         |

1: 95% confidence intervals in brackets.
4. Conclusions

A quantitative spectrophotometric AChE assay for the studied CM and OP pesticides monitoring in lettuce and strawberry extracts was presented. Importantly, the AChE assay was validated in the food matrices and its performance was confirmed by an accredited LC–MS/MS, a practice commonly missing or misrepresented in the available literature. The short analysis time (20 min for sample preparation/ 6 samples and 15 min for AChE assay) combined to the affordable cost (0.16 euro/96-microwell plate) showed that the AChE assay can be applied by laboratories with limited resources as a pre-screening tool. In contrast to chromatographic methods, which are serial meaning that only one sample is analyzed per run, up to 96 samples could be analyzed using the developed method achieving high-throughput. All in all, our contribution can be extremely useful for monitoring carbofuran in countries where this pesticide is still used or testing imports originating from those countries and a rapid answer is needed. Future work is under way to expand target analytes-food matrices portfolio and further improve LODs.

Supplementary Materials: The following are available online at http://www.mdpi.com/2076-3417/10/2/565/s1, Figure S1: AChE tolerance towards various organic solvents after 30 min of incubation with ethanol, methanol, acetonitrile and dimethyl sulfoxide (n = 3). The 100% AChE activity was set for 0% organic solvent in PBS., Table S1: Calibration ranges for each pesticide investigated for potential cross-reaction with carbofuran. (n = 7 calibration points and one blank sample), Table S2: Carbofuran-3-hydroxy and dichlorvos validation results in strawberry and lettuce for AChE assay, Table S3: Carbofuran-3-hydroxy and dichlorvos validation results in strawberry and lettuce for LC-MS/MS method.

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