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ONE-STEP EXTRACTION VERSUS QUECHERS FOR PESTICIDE ANALYSIS IN SELECTED FRUITS AND VEGETABLES

Darko Andjelković¹, Milica Branković^{2*}

¹University of Niš, Faculty of Agriculture, Kosancićeva 4, 37 000 Kruševac, Serbia ²University of Niš, Faculty of Science and Mathematics, Višegradska 33, 18 000 Niš, Serbia milica.chem@outlook.com

This research was focused on the performance evaluation of a simple sample preparation involving acetonitrile extraction followed by liquid chromatography/mass spectrometry (LC/MS) analysis. Simplified method validation parameters, along with several other features, were compared to those of the citrate QuEChERS for 19 pesticides analyzed in four representative fruits and vegetables. The results showed comparable performances of the two methods for 5 of the 6 investigated validation parameters. The simplified method had better performance regarding the selectivity, since three analytes experienced a selectivity issue in one of four QuEChERS-treated matrices. Overall, results lead to an assumption that acetonitrile extraction could be reasonably implemented in certain cases of pesticide analysis, as an efficient and economical alternative to the official method. Since the research provides an insight into acetonitrile extraction capabilities in the domain of pesticide analysis in complex matrices, scientists, researchers or analytical practitioners can determine which method is most beneficial for a particular analysis.

Keywords: acetonitrile; EN 15662; HPLC; sample preparation

ЕКСТРАКЦИЈА ВО ЕДЕН ЧЕКОР НАСПРОТИ QUEChERS ЗА АНАЛИЗА НА ПЕСТИЦИДИ ВО ИЗБРАНО ОВОШЈЕ И ЗЕЛЕНЧУК

Ова истражување беше фокусирано на евалуација на перформансите на едноставна подготовка на примерок која вклучува екстракција со ацетонитрил проследена со анализа со течна хроматографија/масена спектрометрија (LC/MS). Параметрите за валидација на поедноставениот метод, заедно со неколку други карактеристики, беа споредени со оние на цитратно пуфериран QuEChERS за 19 пестициди анализирани во четири репрезентативни видови овошје и зеленчук. Резултатите покажаа споредливи перформанси на двата метода за 5 од 6-те параметри истражени за валидација. Поедноставениот метод имаше подобри перформанси во однос на селективноста, бидејќи три аналити покажаа проблем со селективноста во една од четирите матрици третирани со QuEChERS. Општо земено, резултатите водат до претпоставка дека екстракцијата со ацетонитрил може разумно да се спроведе во одредени случаи на анализа на пестициди, како ефикасна и економична алтернатива на официјалниот метод. Бидејќи истражувањето дава увид во способностите за екстракција со ацетонитрил во доменот на анализа на пестициди во сложени матрици, научниците, истражувачите односно аналитичките практичари можат да одредат кој метод е најкорисен за одредена анализа.

Клучни зборови: ацетонитрил; EN 15662; HPLC; подготовка на примероци

1. INTRODUCTION

The diversity and complexity of fruit and vegetable contents remain the main challenge in pesticide monitoring in these matrices. Consequently, various sample preparation methods have been developed, each one with the same goal, to generate consistent high-quality results of multiresidue analysis in many plant matrices at the lowest possible pesticide concentrations. At this point, the established sample preparation method is QuEChERS. Owing to the simple steps, QuEChERS is time effective and less prone to errors, thus it represents a streamlined approach to assess multiple pesticide residues in food.

A recent literature overview shows that the established QuEChERS method and its versions are dominantly employed for pesticide analysis in fruits and vegetables. 1-4 The non-QuEChERS sample preparation methods, 5,6 simple solvent extraction methods^{7,8} and methods comprising minimal sample preparation⁹⁻¹² are less frequently reviewed. A gas-liquid microextraction technique (GLME), based on the distinctive boiling points between the analytes and the interferences to achieve effective separation, has been coupled to a dispersive solidphase extraction (dSPE) in a one-step sample pretreatment approach by Jin and co-workers.⁶ The main feature of such a method is that the integrated extraction and clean-up can be performed in several minutes. This method was evaluated for 35 pesticides in apple, leek, orange and honey matrices. Guo and co-workers⁵ implemented the reverse approach, involving carbamate pesticide extraction from tomato and apple samples onto columnpacked covalent organic frameworks with acrylamide sites, eluted with acetonitrile and subjected to the liquid chromatography/ultraviolet-visible analysis (LC/UV-Vis). Solid-liquid extraction with acetonitrile was employed for the analysis of chlorpyrifos and acetamiprid in tomato peel.^{7,8}

Methods comprising minimal sample preparation depend on the technique used for instrumental analysis. A minimal sample preparation was utilized in the screening of 6 pesticides on lemon surface by the paper-spray ionization mass spectrometry. Pesticide detection can be achieved by rubbing the fruit surface with some tool, then exposing it to the mass spectrometer inlet. Another technique involving minimal sample preparation is the surface-enhanced Raman spectroscopy (SERS), an optimized version of Raman spectroscopy, that involves an application of gold- and/or silver-based nanoparticles onto the analytical surface to enhance the Raman signal. The technique was developed for the analysis of thiram in apples, ¹¹ triazo-

phos in apples and cherry tomatoes, ¹⁰ and chlorpyrifos on tomato and grape surfaces. ¹²

The objective of this study was to evaluate the performance of solvent extraction as the simplest possible sample preparation method. Evaluation included 19 pesticides, acetonitrile as an extraction solvent and four commodities (cucumber, lettuce, tomato and lemon) chosen for diverse texture, pigmentation and acidity.

The performances of pesticide analysis in the selected fruits and vegetables were compared to those of the citrate-buffered QuEChERS followed by dispersive solid phase extraction (dSPE) with PSA. Contrary to the practice of analysis in multiple reaction monitoring (MRM) mode, analytes were detected and quantified in the full scan MS¹ mode. Since the goal was to evaluate and compare the performances of the two methods, the simplification in mass analysis was intentionally implemented. Furthermore, if the screening itself is a purpose, recording a sample's spectrum in full scan mode will offer a possibility for the retroactive analysis of some other analytes, not covered by the current plan of analysis.

2. MATERIALS AND METHODS

2.1. Chemicals and consumables

Formic acid (FA) (98 %) and high purity pesticide standards (acetamiprid, azoxystrobin, boscalid, buprofezin, chlorpyrifos, cyprodinil, difenoconazole, fenhexamid, imidacloprid, kresoximmethyl, metsulfuron-methyl, propiconazole, pyraclostrobin, pyrimethanil, pyriproxyfen, tebuconazole, thiacloprid, thiamethoxam, trifloxystrobin) were produced by Sigma-Aldrich® (St. Louis, Missouri, USA). Ammonium formate (AMF) (98 %), deionized water and high-performance liquid chromatography (HPLC) grade ethanol were produced by Carlo Erba (Emmendingen, Germany). HPLC grade methanol (MeOH) and HPLC grade acetonitrile (AcN) were produced by J.T. Baker, Thermo Fisher Scientific (Waltham, Massachusetts, USA). For the sample preparation, procedure prepacked Hillium QuEChERS extraction pouches (1 g of NaCl, 4 g of MgSO₄, 1 g of trisodium citrate dihydrate and 0.5 g disodium hydrogen citrate) and Hillium OuEChERS dispersion kits (25 mg PSA and 150 mg MgSO₄) were used. Syringe microfilters (Nylon Hydrophilic 0.22 µm) were produced by Membrane Solutions (Auburn, Washington, USA).

Fresh fruits and vegetables (tomato, cucumber, lettuce, and lemon) were purchased in a local supermarket.

2.2. Instruments and instrumental parameters

Appliances. For high purity standards weighing procedures, the analytical balance Sartorius BP110S (Göttingen, Germany) was used. The sample preparation procedure used the following appliances: blender 0.9 l BL142A by TEFAL (Rumilly, Haute-Savoie, France); balance (acc. ± 0.01 g) KB 2000-2N by KERN & SOHN GmbH (Balingen, Germany); and centrifuge Jouan C4i by Thermo Fisher Scientific. To facilitate the extraction, Digital Vortex-Genie 2 by Scientific Industries (Bohemia, New York, USA) was used. Nitrogen (99 %) was supplied by a nitrogen generator by PEAK Scientific (Glasgow, Scotland, UK).

Analytical instruments. Instrumental analysis was performed on an LC/MS system including a Surveyor autosampler by Thermo Finnigan LLC (San Jose, California, USA), and an Accela MS pump and LTQ XL mass spectrometer with linear ion trap analyzer by Thermo Fischer Scientific. Analytes were separated on a Hypersil GOLD column (C_{18} , 150 mm \times 2.1 mm, particle size 3 μ m), ionized in ESI+ ionization mode and monitored in MS¹ full scan mode (scan range m/z 150–600). Data was acquired and analyzed by Thermo XcaliburTM software, version 2.1.0, SP1.1160.

Instrumental parameters. Ten microliters of sample were loaded on to a column in a partial loop injection mode and eluted with a mixture containing eluent A (buffer solution -0.1 % of FA and 0.03 % of AMF in water) and eluent B (MeOH), following the gradient: 0 min (90 % A), 0–2 min (90 % A), 2– 7 min (30 % A), 7–30 min (30 % A), 30–35 min (90 % A) and 35-40 min (90 % A) with a flow rate equal to 300 μl min⁻¹. The chromatographic column was kept at thermostatic conditions at 25 °C. ESI source parameters were: sheath gas = 21 arbitrary units; auxiliary gas = 18 arb; I (spray voltage) = 5kV; capillary T = 275 °C. Protonated molecular ions of tested analytes cover the m/z range from 200 to 409 (Table S1), therefore, the ion optics were optimized according to the pesticides' ions from the lower, middle and higher part of the m/z range, that is, according to the ions of pyrimethanil, buprofezin and trifloxystrobin. Tuned parameters meeting optimal detectability for all analytes were chosen.

2.3. Procedures

Stock preparation. Single-pesticide stock solutions (1 mg ml⁻¹ each) were prepared by dissolving high purity pesticide standards in ethanol. Multi-pesticide solutions were prepared by mixing and diluting single stocks in ethanol.

Sample homogenization. Lemon, tomato, cucumber and lettuce, one kilogram of each, were cut and homogenized by blending for 5 min. All parts (flesh and peel) of the tomato, lettuce and cucumber were included in analysis. For lemon samples, only the flesh was used.

Sample preparation – extraction with AcN. Ten grams of homogenate were extracted with 10 ml of AcN. Extracts were centrifuged (10 min/3000 rpm) and supernatants were microfiltered prior to instrumental analysis. In the case of procedural standards preparation, the homogenate portion was spiked prior to the solvent extraction.

Sample preparation – citrate-buffered QuEChERS (EN 15662). Ten grams of homogenate were extracted with 10 ml of AcN, after which an extraction pouch was added. The mixture was immediately vortexed for one minute and centrifugated (10 min/3000 rpm). A supernatant aliquot was subjected to a dispersive extraction by the addition of one dispersion kit per ml of supernatant. The mixture was vortexed for one minute and centrifugated (10 min/3000 rpm). Supernatant was microfiltered prior to instrumental analysis. In the case of procedural standards preparation, the homogenate portion was spiked prior to the solvent extraction.

Validation study. Samples obtained from the market were initially screened for tested pesticides residues. Since none of the targeted pesticides was detected, the samples were considered blank samples and used for validation procedures. The matrix effect (ME) was evaluated by comparing the slopes of the calibration curves (0.01–15.00 µg ml⁻¹) of solvent-based and matrix-based multi-pesticide standards. The linear range of methods was evaluated with 7 procedural standards in the concentration range 0.00-15.00 mg kg⁻¹. Chromatographic repeatability was evaluated from the successive injections of ten procedural-based (5.00 mg kg⁻¹) and ten solvent-based standards (5.00 µg ml⁻¹). Trueness and precision were evaluated at three concentration levels (0.50, 5.00 and 15.00 mg kg⁻¹), each level at 5 replicates. Detection limits (DLs) were evaluated by the S/N criterion. The concentration of spiked sample that produced an analyte peak with S/N = 3 is established as the method's DL. In cases where the spiked samples produced an S/N ratio other than 3, the DL was estimated by a proportion involving the procedural standard with S/N ratio closest to 3.

3. RESULTS AND DISCUSSION

A prerequisite for a successful analysis is chromatographic or mass separation of analytes. Among the targeted analytes, boscalid and propiconazole would express a selectivity issue if they were not chromatographically separated, due to the closeness of MH⁺ *m*/*z* values (Table S1).

Both one-step solvent extraction and the QuEChERS method expressed similar performances in terms of analyte recovery. Recoveries of the targeted analytes from tomato and lemon treated with the QuEChERS were within the limits (70–120 %) (Figs. S1 and S4). In lettuce and cucumber (Figs S2 and S3), at 0.50 mg kg⁻¹ spike level, the recovery of imidacloprid was much below 70 %. Recoveries of the targeted analytes from each matrix treated with the simplified method (AcN extraction) were within the limits (70–120 %) (Figs. S1–S4). Considering the average recovery, both methods demonstrated satisfactory performance for each analyte (Fig. 1). The average recovery of thi-

amethoxam, after AcN extraction from lemon samples was less than 70 % (Fig. 1), however, out-of-limits recovery was 68 %. According to SAN-TE/11312/2021, 13 recoveries out of the 70–120 % range are acceptable if they are consistent (RSD < 20 %), but the mean recovery must not be lower than 30 % or higher than 140 %.

The detection limits of the two methods were comparable, and in most cases below or equal to the pesticide maximum residue limit (MRL) (Fig. 1). In three cases, the DLs were higher than the MRLs. The DLs of both methods were slightly higher than fenhexamid MRL in lemon; the DL of the QuEChERS was higher than cyprodinil MRL in lemon (Fig. 1).

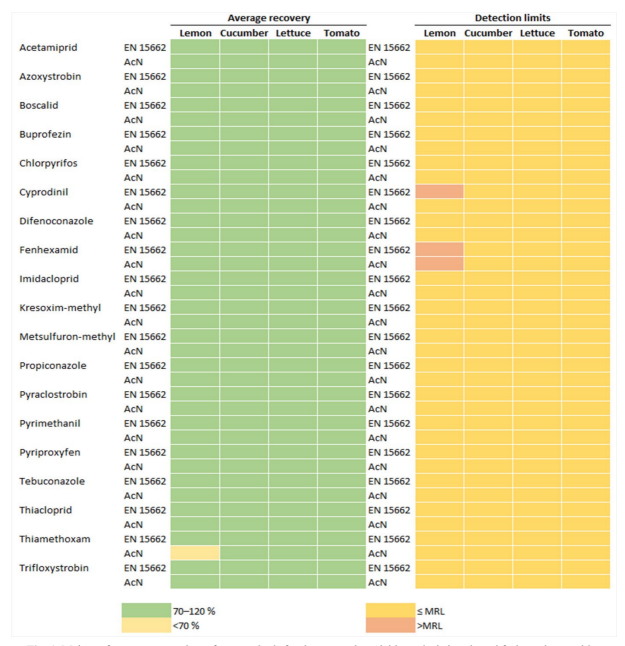


Fig. 1. Main performances overview of two methods for the targeted pesticide analysis in selected fruits and vegetables

Both methods demonstrated similar performances regarding the matrix effect and linearity range. Strong signal suppression was observed for each analyte in each tested matrix, regardless of the sample preparation procedure (Figures S1-S4). Analyte signals were suppressed by at least 37 %. Methods were linear in the tested analytical range $(0.00-15.00~{\rm mg~kg^{-1}})$, with back-calculated concentration deviations < 20 % and correlation coefficients > 0.90 (Table 1).

Significant pesticide retention time shifts were observed in sample extracts, regardless of the sample preparation procedure (Figure S5). The shift ranged from 0.12 to 1.11 min. The tolerable limit of \pm 0.10 min was exceeded by 8 analytes in tomato, 4 in lettuce and cucumber and 12 analytes in lemon, all treated with the QuEChERS. The exceedance rate in samples treated with the simplified method was similar i.e., the retention time shift of 3 analytes in tomato, 5 in lettuce, 12 in cucumber and 11 in lemon exceeded the limits. The strongest shift was noticed in the lemon matrix, which stands out with the lowest pH value of the final extracts.

In terms of method selectivity, lemon was identified as a difficult matrix. A selectivity issue was observed for pyraclostrobin, tebuconazole and fenhexamid in the QuEChERS-treated lemon (Figure S6) and for fenhexamid in AcN-treated lemon (Figure S7). The issue for pyraclostrobin and fenhexamid was related to the overlay of analyte chromatographic peaks with the peaks of interferences. The issue for tebuconazole was related to the overlay of its peak with the peak of buprofezin, due to the heavy retention time shift of tebuconazole in lemon. Interestingly, the selectivity issue for pyraclostrobin and tebuconazole could be exclusively observed in the QuEChERS extracts.

Analytical data for herein investigated methods and for similar methods found in the literature are generally in mutual agreement. Melton and Taylor⁴ implemented the same QuEChERS procedure for the analysis of buprofezin and chlorpyrifos in lemon by gas chromatography/mass spectrometry (GC/MS), which resulted in more than 94 % of recovered analytes and a DL of 0.01 mg kg⁻¹. Martinez Bueno et al.³ also implemented the citrate QuEChERS for multi-pesticide LC/MS analysis in lettuce, but the *d*SPE step was performed with C₁₈. Percentages of recovered analytes

ranged from 77 for azoxystrobin to 91 for thiacloprid. Another variation of the citrate QuEChERS followed by the dSPE with PSA and ENVI-Carb was implemented by Fearracane et al.1 for the analysis of chlorpyrifos in tomatoes, cucumbers and lettuce by flow-modulated GC/MS. The recovery of chlorpyrifos was higher than 99 % in each matrix. The lowest DL of 1.8 µg kg⁻¹ was established for cucumbers. Mahdavi et al.² implemented the dispersive solid-phase extraction with primarysecondary amine (PSA) for pesticide LC/MS analysis in cucumbers, but as a part of the acetatebuffered QuEChERS. Ten of more than 50 investigated pesticides matched our study. The recovery ranged from 74 % for kresoxim-methyl to 107 % for propiconazole. The lowest and the highest DL of 0.002 and 0.01 mg kg⁻¹ were established for kresoxim-methyl and imidacloprid, respectively. Çatak and Tiryaki¹⁴ implemented the same acetatebuffered QuEChERS for GC/MS analysis of chlorpyrifos and acetamiprid in cucumbers. A recovery higher than 80 % and LOQs of 2 and 10 µg kg⁻¹ for acetamiprid and chlorpyrifos, respectively, were established. Dashtbozorgi et al. 15 applied a dispersive liquid-liquid microextraction technique for the extraction and pre-concentration of acetamiprid, imidacloprid, azoxystrobin and 17 other pesticide residues from QuEChERS extracts of tomato and cucumber. This combined procedure resulted in recovery of around 100 % and DLs ranging from 3.9 to 9.4 μ g kg⁻¹.

In addition to our study, acetonitrile-based solvent extractions were implemented by Moura et al.8 and Hegazy et al.7 for the analysis of chlorpyrifos and acetamiprid in tomatoes. The detection techniques, however, were the paper-spray ionization mass spectrometry (chlorpyrifos) and the LC/UV-Vis (acetamiprid), providing the DL of 0.01 ppm and 0.03 µg ml⁻¹, respectively. In each case, pesticide recovery was higher than 94 %. A different solvent extraction procedure was implemented by Mohamed et al.16 for the analysis of chlorpyrifos and five other pesticides in tomatoes and cucumbers by gas chromatography/flame ionization detection (GC/FID). Pesticides were extracted from samples in a successive extraction with acetone and dichloromethane, after which the extract was cleaned-up on a Florisil® stationary phase. Achieved DLs ranged from 0.001 (chlorpyrifos) to 0.20 mg kg⁻¹ (profenofos).

Table 1

Linear regression parameters $(y = ax + b, conc. range 0.00 - 15.00 \text{ mg kg}^{-1}, 7 \text{ points})$ for EN 15662 and AcN method

Amount			Tomato			Cucumber			Lettuce			Lemon		
Thimmetry Final 15662 49,942 607 999 39,064 4.578 0.997 1 4,5668 -5,6647 0.999 0.23,806 8.495 0.998 0.808 0.908 0.			а	b	\mathbb{R}^2	a	b	\mathbb{R}^2	a	b	\mathbb{R}^2	а	b	\mathbb{R}^2
Acctamiprice Acctam	Thiametox-	EN	49,942	607		39,064	4,578	0.997	45,668	-5,647		23,806	8,495	
Acetamiprid Fine 486,466 248,738 43 379,027 278,482 0.78 425,279 587,260 0.818 16.010 165.89 0.935	am		29,397	-4,759	0.998	22,034	-1,084		25,274	4,890	0.984	21,210	7,403	
Thing Thin	Acetamiprid		486.466	248.738	0.983	379.027	278.482	0.981	425.279	587.260	0.981	165.015		_
This cloppid FN			ŕ		-	,	ŕ			ŕ		ŕ	•	
	Thiacloprid		ŕ		_	,	ŕ	,	ŕ	ŕ			ŕ	
Metsulfuron EN 2,504 2,936 0	F		ŕ		2	ŕ	ŕ	0	ŕ	ŕ	2	51,924	20,519	1
15662 21.00 29.95 1 12.00 7.91 7 14.239 15.270 0 7.042 0.994 0.996	T '1 1 '1		76,465	9,992	5	49,158	16,813	8	63,079	11,697	9	29,673	11,311	9
Metsulfuro	Imidacloprid		21,500	29,936	1	12,056	7,919	7	14,559	18,370	0	7,042	6,694	2
methyl 15662 38.308 14/47/ 4 22.441 99.89 2 22.97/4 123.488 0 15.044 63.143 5 5 6.048 64.143 103.910 0.979 18.556 70.425 0.955 17.556 0.932 0.932 0.937 83.970 0.995 8 9.071 15.008 10.0		AcN	12,383	11,748		7,176	4,941		13,209	27,382		5,332	2,489	_
Pyrimethania	Metsulfuron methyl		38,308	147,477		22,447	99,689		22,974	123,848		15,644	63,143	_
Pyrimethanian EN 1.70.5 926.818 0.990 742.540 756.042 0.991 993.453 223.656 0.999 403.327 23.229 0.999 99.99 1.5662 0.66 6.6 6.6 6.6 6.6 0.999 0	,		25,804	103,910	0.979	18,556	70,425		17,295	175,560		9,075	83,970	0.995
Aco	Pyrimethanil			926,818	0.990	742,540	756,042	0.991	993,453	223,656	0.999	403,327	23,229	0.999
Azoxystrobin EN 1,566,2 1,241,2 0,966 982,868 754,357 7 63 00 00 624,781 0 0 3 3 0 0 3 3 0 0						353.311	73.690	0.999	527.774	136.482	0.998	273.318		0.999
No. 1,047,7 641,847 0,985 0,944 1,047,6 641,847 0,985 642,729 403,148 0,976 926,415 668,696 0,976 3 596,502 380,28 0,977 780,10 0,989 1,5662 60 0,54 60 0,91 7 7 88 9 0,5 7 7 2 2 2 2 2 2 2 2	Azoxystrobin			ŕ		ŕ	,		-	ŕ		ŕ		8 0.975
Cyprodini	•				-		ŕ						0 380.28	
15662 60 05 4 60 91 7 17 7 88 9 05 7 22 24 24 26 25 25 25 25 25 25 25	Cymrodinil		86	ŕ	0	The state of the s	,	5	-	ŕ	3		6	1
Boscalid	Сургосиии		60	05	4	60		7	17	88	9		7	2
15662 164,078 568,662 2 105,761 509,623 9 158,072 417,802 2 73,347 0 0 9 AcN 102,244 526,804 6 6 47,261 672,624 0.969 1 74,009 595,158 3 37,466 584,88 0.966 Fenhexamid EN 15662 332,842 55,619 0.998 4 200,520 29,784 9 288,272 27,730 0.999 61,730 8,002 1.000 AcN 201,886 -2,966 0.999 108,899 12,473 0.999 157,094 31,708 0.998 106,134 16,633 0.999 Methyl 15662 115,839 26,132 5 5 81,169 7,792 0.997 4 112,297 -35,606 0.999 55,463 3,860 0.999 methyl 15662 456,056 44,800 5 335,944 46,944 5 405,717 15,865 0.999 49,270 4,971 0.999 Propiconazole EN 15662 506,139 55,230 0.999 335,944 46,944 5 9 212,257 42,837 0.998 161,520 349 0.999 Propiconazole EN 256,244 257,248 257,448 257		AcN					945,764				5	857,576		1
Fenhexamid	Boscalid		164,078	568,662		105,761	509,623		158,072	417,802		73,347		
Fenhexamid EN 15662 332,842 55,619 0.998 4 200,520 29,784 0.998 288,272 27,730 0.999 61,730 8,002 1.000 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		AcN	102,244	526,804		47,261	672,624	0.969	74,009	595,158		37,466		0.966
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Fenhexamid		332,842	55,619	0.998	200,520	29,784		288,272	27,730	0.999	61,730	•	
Kresoxim methyl EN 15662 115,839 26,132 0.997 5 5 5 81,169 7,792 0.997 4 112,297 -35,606 0.999 5 55,463 3,860 0.999 7 7 0.999 7 0.999 AcN 67,531 -9,810 0.999 2 44,782 -5,484 0.999 2 55,713 -1,362 0.996 7 7 7 0.996 44,333 27,402 0.996 7 7 0.999 Tebuconazole EN 15662 456,056 44,800 0.999 3 335,944 46,944 0.999 9 0.999 405,717 15,865 0.999 9 0.999 9 0.999 49,270 4,971 0.999 0.999 Propiconazole EN 15662 506,139 55,230 0.999 0.999 0.999 39,303 0.999 0.999 0.999 212,257 42,837 0.999 0.999 0.999 0.999 249,045 24,293 0.999 0.999 0.999 Pyracolostrobin 15662 506,139 55,230 0.999 0.999 0.999 198,263 9,269 0.999 0.999 0.999 241,819 108,716 0.993 0.999 0.999 178,305 68,814 0.995 0.995 0.999 241,819 108,716 0.993 0.999 0.999 112,019 53,701 0.990 0.999 0.992 0.999 241,819 108,71			201.886	-2.966	0.999	108.899	12,473	0.999	157.094	31,708	0.998	106,134	16,633	0.999
AcN 67,531 -9,810 0.999 44,782 -5,484 0.999 55,713 -1,362 0.996 7 44,333 27,402 0.996 5 5 5 5 5 5 5 5 5	Kresoxim	EN	ŕ		0.997		ŕ	-	ŕ	ŕ	0.999	ŕ	ĺ	
Tebucona- zole	methyl		ŕ	ŕ	-	,	ŕ	-	ŕ	ŕ	_	ŕ	ŕ	7 0.996
zole 15662 436,036 44,000 5 333,944 40,944 5 403,717 13,603 9 49,270 4,971 6 AcN 264,252 -15,632 0.999 3 180,366 4,045 0.999 212,257 42,837 0.998 161,520 349 0.999 212,257 42,837 0.998 161,520 349 0.999 249,045 24,293 0.999 249,045 24,293 0.999 241,819 108,716 0.993 178,305 68,814 0.995 0.999 241,819 108,716 0.993 178,305 68,814 0.995 0.999 241,819 108,716 0.993 178,305 68,814 0.995 0.999 15662 0.989 240,437 81,509 0.995 241,819 108,716 0.997 112,019 53,701 0.992 0.998 15662 0.999 140,144 0.997 112,019 53,701 0.992 0.998 140,144 0.997 15662 0.998 140,144 0.998 183,003 105,754 0.995 0.995 0.998 15662 0.998 140,144 0.998 0.998 183,003 105,754 0.995 0.995 0.998 15662 0.998 0.998 0.998 0.998 0.999	Tebucona-				_	,	ŕ	_		ĺ	7 0.999	,	ŕ	
Propicona- zole			456,056		5	335,944	46,944	5	405,717		9		4,971	6
zole 15662 306,139 33,230 5 339,103 39,303 6 462,719 23,161 9 249,043 24,293 7 AcN 298,162 -2,832 0.999 198,263 9,269 0.999 241,819 108,716 0.993 178,305 68,814 0.995 Pyra- clostrobin 15662 386,406 200,652 5 240,437 81,509 2 353,274 170,541 2 112,019 53,701 6 AcN 230,295 45,069 8 140,144 32,557 8 183,003 105,754 1 92,956 61,009 6 Buprofezin EN 2,104,0 1,128,6 0.982 1,408,6 29 621,707 0.989 2,017,3 702,916 0.992 881,980 275,50 0.994 15662 04 57 6 29 621,707 3 41 702,916 0.992 881,980 275,50 0.994 1,186,9 40,000 1,295,5 361,312 0.996 860,423 232,692 0.994 1,186,9 410,927 0.991 705,379 256,71 0.991 705,379 256,7	.		264,252	-15,632	3	180,366	4,045	9	212,257	42,837	1	161,520	349	9
Pyraclostrobin			506,139	55,230	5	359,105	39,303	6	462,719	25,161	9	249,045	24,293	7
Clostrobin 15662 386,406 200,652 5 240,437 81,509 2 353,274 170,541 2 112,019 53,701 6 AcN 230,295 45,069 0.997 8 140,144 32,557 0.997 8 183,003 105,754 0.995 92,956 61,009 0.978 8 183,003 105,754 1 92,956 61,009 0.978 15662 0.904 57 6 29 621,707 3 41 702,916 1 881,980 275,50 0.994 1,186,9 410,927 2 361,312 0.996 860,423 232,692 0.994 1,186,9 410,927 2 2 56,71 0.991 5 4 1,186,9 15662 0.995 15662 0.998 201,000 0.998 201,000 0.999 201		AcN	298,162	-2,832	0.999 7	198,263	9,269		241,819	108,716		178,305	68,814	0.995 5
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$			386,406	200,652	0.989	240,437	81,509		353,274	170,541		112,019	53,701	0.992
Buprofezin EN 2,104,0 1,128,6 0.982 1,408,6 29 621,707 0.989 2,017,3 702,916 0.992 881,980 275,50 0.994 $\frac{1}{3}$ AcN 1,295,5 $\frac{1}{2}$ 361,312 0.996 860,423 232,692 0.994 1,186,9 $\frac{1}{2}$ 410,927 0.991 705,379 256,71 0.991 $\frac{1}{2}$ Difenocona-zole EN 706,205 -43,422 0.998 2664,815 -75,625 0.999 664,133 0.999 232,682 0.999 323,682 0.995 0.998 0.995 0.998	Closuloom		230,295	45,069		140,144	32,557	0.997	183,003	105,754	0.995	92,956	61,009	
13602 04 37 6 29 3 41 1 3 9 41 1 3 9 41 1 1 3 9 41 1 1 3 9 41 1 1 3 9 41 1 1 1 3 9 41 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Buprofezin				0.982			0.989		702,916	0.992	881,980	_	
Difenocona- EN 706,205 -43,422 0.998 204,815 -75,625 0.999 21 2 0.999 329,484 9,350 1.000 0 0.998 204,21			1,295,5							ŕ				9 0.991
zole 15662 /06,205 -43,422 2 504,815 -/5,625 2 664,135 216,835 3 329,484 9,350 0	Difenocona-									-10,327				4 1.000
AcN 408,421 135.632 6 266,223 110,243 9 323,682 282,868 0.932 213,260 24,047 8		15662			2		-75,625	2			3		9,350	0
	T. :			135,632	6		110,243	9		282,868	2	213,260	24,047	8
$\begin{array}{cccccccccccccccccccccccccccccccccccc$				366,819			311,321			265,710		802,219		

			Tomato		Cucumber			Lettuce			Lemon		
	AcN	1,015,2	65,089	0.999	590,471	255,918	0.998	822,058	314,827	0.996	541,449	159,25	0.996
	71011	59	05,007	9	570,171	255,710	7	022,030	311,027	7	311,117	8	8
Pyriproxyfen	EN	1,716,3	539,341	0.993	1,117,8	303,442	0.997	1,574,5	351,082	0.998	767,966	110,02	0.999
	15662	52		7	34		3	90		6		8	7
	ACN	1,019,9	69 890	0.999	616,235	106,065	0.998	852,834	232,179	0.995	546,597	134,72	0.996
		48		8			1			3		3	2
Chorpyrifos	EN	270 (00	12.020	0.996	224 162	24.077	0.998	210.002	01 222	0.994	170 ((0	1.050	0.999
	15662	370,699	13,928	7	234,163	34,077	9	310,092	91,233	1	178,660	-1,059	9
		100 140	11.644	0.993	114 122	22 707	0.994	1.40.077	62.707	0.989	114 402	17.045	0.998
	AcN	182,143	11,644	3	114,122	32,707	5	142,877	63,797	1	114,403	17,945	4

4. CONCLUSION

Performances of the first step of the QuEChERS sample preparation procedure, which is the acetonitrile extraction, were evaluated for 19 pesticides in four representatives of fruits and vegetables and were compared to the performances of the citrate-buffered QuEChERS. Both methods expressed comparable performances regarding most of the validated parameters, including the matrix effect, chromatographic repeatability, recovery, detection limits and linearity. Better performances regarding the selectivity were expressed in the acetonitrile extraction method, since 3 anaexperienced a selectivity QuEChERS-treated lemon. In the end, the comparable performances of the simplified method and QuEChERS reasonably qualify the simplified method for implementation in certain cases of pesticide analysis, as an efficient and economical alternative to the official method.

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