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# Multi-residue method of pesticides by UPLC-MS/MS in bivalve mollusks samples as a tool for food quality and safety

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## Abstract

Bivalve molluscs (BM) are filter animals and bioaccumulators of substances from the environment. This characteristic allows a great absorption of nutrients that makes them a source of protein-rich foods. On the other hand, if there are toxic contaminants in the environment, their absorption by animals may occur. This happens with pesticides coming from agricultural and livestock production systems that can migrate to areas of BM crops. Considering the high nutritional value of bivalve molluscs and their positive impact on the human diet, these products must be carefully evaluated for the possible presence of toxic substances in order to guarantee their safety.

Thus, the aim of this work was to implement and validate a multi-residue method using tandem mass spectrometry to evaluate pesticide residues commonly used in agricultural production systems present in these matrices. Extraction and cleaning steps were optimized and the method proved to be adequate to quantify 322 pesticides. The samples come from five different areas of culture of bivalve molluscs in the southeast, north and northeast regions of Brazil. The analysis of the mollusc samples showed the presence at the trace level of seven different pesticide residues in four of the five evaluated samples.

Keywords: mollusk bivalves; pesticides; UPLC-MS/MS; oyster; scallop.

**Practical Application:** Evaluation of the contamination by pesticides in samples of bivalve molluscs to guarantee food safety; to point out the presence of pesticides coming from areas far from the shellfish cultivations that may represent possible contamination for of the environment.

## **1** Introduction

*In* recent years, aquaculture has become one of the most developed food sources (Food and Agriculture Organization of the United Nations, 2018). The expansion of this type of activity is one of the most viable and sustainable alternatives for the production of high-protein foods for human consumption. In the future, this branch of activity may assume an even more important role capable of meeting global needs in terms of food production and nutrition.

The increase in fish consumption favored, at the same time, the development of oyster crops, which are considered a delicacy appreciated in several continents and have become an excellent option for genuinely protein food. Oysters are also an important source of minerals, amino acids, glycogen and essential fatty acids (Asha et al., 2014). This composition, rich in nutritional components, is essential to define its quality and commercial value.

On the other hand, the intensification of agricultural activities to increase food production and meet world demand has resulted in the intense use of pesticides in plantations to control diseases and parasites. The drift of pesticides to non-agricultural areas is a common problem in regions of intensive agriculture (Cech et al., 2022). Cui et al. (2020) showed that the pollution load of diffuse origin from agricultural activities depends on the amounts and frequency of irrigation and fertilization in the plantations. These pesticide residues are not restricted to the application site, but can be diverted to surrounding areas due to droplets that evaporate before reaching the target and travel with fine particles of spray, air movement and volatilization during application (European Food Safety Authority, 2008), or later by leaching and soil erosion (Linhart et al., 2019; Zivan et al., 2017). Burket et al. (2018) observed in samples of mussels and oysters collected in the outskirts of Hong Kong, the presence of pesticide substances at low levels ( $\mu$ g/kg). The presence of potentially toxic substances started to be commonly observed in surface waters as well as their accumulation in animals that make use of it. Its monitoring can point out possible contamination in food by pesticides that originate from their intense application in adjacent agricultural areas (Iliff et al., 2019; Bringer et al., 2021).

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The place where bivalve mollusks are cultivated is a very important factor that directly affects the quality of the product and also its characteristics as food. The risk associated with human exposure to pesticides occurs notably through ingestion. The diet of the population constitutes the main route of human exposure to pesticide residues, contributing with more than 90% of the total exposure (Fazal et al., 2022; Riaz et al., 2018). It should be noted that oysters are filtering animals and, admittedly, bioaccumulators of substances present in the environment in which they live, such as nutrients, minerals, microorganisms and also chemical contaminants. When these animals develop in polluted waters they can become vehicles of contamination for humans through the food route (Petrarca et al., 2022; Hussein et al., 2022). The health of these foods is essential and, as the cultivation of bivalve mollusks is intended to compose the human diet, these products must be carefully evaluated for the presence of contaminants in order to guarantee their safety and health (Onac et al., 2022). In order to carry out monitoring for the evaluation of contaminants in mollusks, it is essential to implement a validated analytical method for the matrices of interest capable of answering questions about the presence of toxic substances potentially harmful to human health.

The objective of this work was to validate a multiresidue method in order to investigate the occurrence of pesticide residues in samples of bivalve molluscs. For this, samples of molluscs from different regions of Brazil were evaluated. Four samples of *Crassostrea gasar* oysters and one sample of scallops (*Nodipecten nodosus*) were analyzed. The multiresidues method was implemented and validated for these matrices. The extraction and clean up conditions were optimized for the quantification of 322 active ingredients from pesticide residues of different chemical classes commonly used in agricultural production systems.

Residue analyzes were performed using the extraction method called QuEChERS (acronym for Quick, Easy, Cheap, Effective, Rugged, Safe) (Anastassiades et al., 2003). The complexity of the study matrix makes it difficult to quantify a large number of analytes, requiring a purification step (clean-up) of the extract and quantification of the active ingredients of pesticides using chromatographic methods coupled with sequential mass spectrometry (Lehotay et al., 2007).

# 2 Material and methods

#### 2.1 Chemicals and reagents

Acetonitrile and methanol, LC-MS grade, were purchased from BIO-GRADE Chem (San Francisco, CA, USA); formic acid was obtained from Merck (Darmstadt, Germany) and ammonium formate was purchased from Fluka (Steinhein, Swirtzerland). MgSO<sub>4</sub> was obtained from Sigma-Aldrich (St. Louis, MO, USA); NaCl ACS grade were purchased from F. Maia (São Paulo, Brazil); citrate tribasic sodium dehydrate purchased from Vetec (Rio de Janeiro, Brazil); sodium hydrogen citrate sesquihydrate ACS grade was purchased from Sigma-Aldrich (St. Louis, MO, USA);Bondesil\* PSA from Agilent Technologies (Santa Clara, CA, USA); ultrapure water was obtained from Milli-Q Advantage A10 Millipore System at 18.2 M $\Omega$  cm-1 (Molsheim, France); all the 346 pesticide reference standards were purchased

from Dr. Ehrenstorfer Laboratory (Augsburg, Germany) and manufactured under ISO 17034.

## 2.2 Standards solutions

The standard solutions used in this study were prepared and stored at concentrations ranged from 1000 to 430  $\mu$ g/mL and stored in a freezer at -25 °C. Calibration curves were constructed with a minimum of six points with concentration levels ranging from 0.005 to 0.08  $\mu$ g/mL.

#### 2.3 Bivalve samples

The culture of molluscs that generated the samples used in this work was structured after several stages that began with the domestication of native species such as *C. gasar*. The entire production process has been optimized from seed cultivation, animal handling and harvesting to obtain individuals in commercial size to add the correct value to the product.

Bivalve mollusc samples were acquired from five mollusc farms. A total of 5 samples were collected from four oyster farms and one scallop farm. Sample collection was carried out in the same way in the five regions. Each sample actually represents a total of 2.5-3.0 kg of oyster or scallop meat. This makes a considerable number of individuals.

The oysters came from four different regions located in the north/northeast of Brazil and a single sample of scallops (Nodipecten nodosus) grown on a farm in the southeastern region. These 5 samples were named A, B, C, D and E, respectively.

## 2.4 Fortified samples

The fortified oyster sample used in the recovery trials was prepared by mixing samples from each of the four cultures from the four different states in equal proportions. To this composite sample, 1.0 or 0.5 mL of a standard pesticide solution (MIX) of 346 pesticides of different functional classes was added to carry out the tests at two levels of fortification - 0.02 or 0.01 mg.kg<sup>-1</sup>, respectively.

The fortified scallop sample was obtained by mixing the meat of all the animals collected and sent for analysis in the laboratory. In a sample containing about 10 g of this homogenized material, the Mix of 346 certified pesticide standards was added for the recovery tests at the same two levels of fortification.

#### 2.5 Sample preparation

All samples were collected in a representative way in the different farms, properly packaged and kept at 4 °C until their arrival at the Residues and Contaminants Laboratory until the beginning of the analyses. Sample preparation consists of homogenizing the mollusk muscle tissue. The samples were properly ground in a food processor and stored in flasks in a freezer kept at -20 °C until the moment of analysis.

#### 2.6 Extraction and clean-up method

The QuEChERS method was used (Anastassiades et al., 2003; Lehotay et al., 2007) as described below. About  $10 \pm 0.5$  g

homogenized samples were weighed in a 50 mL polypropylene tube, 10 mL of extraction solvent (ACN) and 1 mL of ultrapure water were added. After vortexing for 1 min, 1 g NaCl, 4 g MgSO4, 1 g sodium citrate and 0,5 g sodium citrate sesquihydrate were added. The mixture was vortexed for another 1 min, subjected to ultrasound for 20 minutes and centrifuged at 4000 rpm for 5 min.

A 7 mL aliquot of the supernatant was transferred to a 10 mL test tube and left in the freezer at -20 °C for 2 hours so that the fat layer could decant. After centrifugation at 4000RPM for 5 min, a 5 mL aliquot of supernatant was transferred to another 15 mL test tube, containing 125 mg of Bondesil - PSA and 750 mg of MgSO4. After homogenation using vortex for 1 min, the tubes were centrifuged at 4000 rpm for 5 min. Finally, 2.0 mL of supernatant were transferred to a vial for UPLC/MS-MS analysis. All the analytical conditions of the method used in this work are summarized in Figure 1.

#### 2.7 UPLC-ESI-MS/MS analysis

All pesticide residue analyzes were performed using an ultra-performance liquid chromatography system coupled with tandem mass spectrometry (UPLC-MS/MS).

The ultra-pure water was produced in the RiOs-Advantage A10 model water deionizer - Milli-Q<sup>®</sup> (Merck Millipore); The Waters ultra-efficiency liquid chromatograph, model Acquity UPLC<sup>®</sup> (Milford, USA) coupled to the Waters sequential mass spectrometer model Quattro Premier XE<sup>®</sup> (Milford, USA). The Waters Acquity UPLC<sup>®</sup> chromatograph has a binary pump system, automatic injector, degasser and column oven.

An elution gradient was used starting with mobile phase A (5 mmol  $L^{-1}$  ammonium formate in water with 10% methanol) with 82.5% (v/v) with a linear ramp until reaching 5.5% of the same. linear curve phase in 25 minutes.

The Quattro Premier XE<sup>\*</sup> mass spectrometer was operated in electrospray ionization (ESI) and multiple reaction monitoring (MRM) in positive and negative modes. The collision gas was argon and the desolvation gas was nitrogen. Table 1 presents the chromatographic parameters of the UPLC-MS/MS system.

Table 1. Conditions of multi-residue analysis of pesticides by UPLC-MS/MS.

|                        | UPLC   |  |  |  |
|------------------------|--|--|--|--|
| Analytical column      | BEH C <sub>18</sub> (1.7 $\mu$ m, 100 × 2,1 mm)                              |  |  |  |
| Pre-column             | VanGuard <sup>®</sup> BEH C18 (Waters, USA)                                  |  |  |  |
| Column temperature     | 35 °C  |  |  |  |
| Mobile fase            | A - 5 mmol L <sup>-1</sup> ammonium formate<br>in methanol 10%; B - Methanol |  |  |  |
| Injection volume       | 5 µL   |  |  |  |
| Flow rate              | 0.3 mL min <sup>-1</sup>   |  |  |  |
| MS/MS                  |  |  |  |  |
| Electronspray Source   | ESI <sup>+</sup> ;ESI <sup>-</sup>   |  |  |  |
| Capillar voltage       | 0.98 kV  |  |  |  |
| Source temperature     | 110 °C   |  |  |  |
| Interface              | Electrospray (Z-Spray)   |  |  |  |
| Cone gas flow          | Nitrogen (50 Lh <sup>-1</sup> )  |  |  |  |
| Heated desolvation gas | Nitrogen; 400 °C   |  |  |  |
| Colision gas           | Argon (3.5 10 <sup>-3</sup> mbar)  |  |  |  |
|                        |  |  |  |  |



Figure 1. Analytical protocol used in the multi-residue monitoring of 320 pesticides in this work.

## 2.8 Method validation

The use of an analytical method to quantify pesticide residues in a matrix requires careful validation of all parameters involved and also the scope that must be achieved. The validation of the multi-residue method carried out in this study is suitable to the criteria described in SANTE (European Union, 2020) which specifically addresses the performance of methods for the analysis of contaminants in food.

The identity criteria specifically used in the sequential mass spectrometer to confirm the pesticide peaks in the samples were (1) Signal-to-noise ratio greater than or equal to 3; (2) Signals from the transitions between the precursor ion and its fragments (quantitation transition and confirmation transition) totally superimposed; (3) Sample retention time meeting a tolerance criterion of  $\pm$  0.1 minute in relation to the standard retention time; (4) Relative intensity of the ionic transitions detected in the evaluated sample and in the standard, is expressed as the ratio of the intensity of the most abundant transition and the corresponding transition of the standard. Assays are performed using the same concentrations and analytical conditions, meeting a tolerance criterion of up to 30% (European Union, 2020).

#### 2.9 Pesticides monitored in this study

The parameters related to the mass spectrometer of all tested pesticides were individually determined in the UPLC-MS/MS system. The transitions used for the monitoring of pesticides are shown in Table 2, which contains only the active ingredients that were considered tested and approved by the validation of this method.

## 3 Results and discussion

# 3.1 Method validation

The QuEChERS method was used (Anastassiades et al., 2003; Lehotay et al., 2007) to evaluate the pesticide residues in all mollusk bivalve samples. The validation of the multiresidue method constitutes the most important and critical part of this work and was carried out for the two matrices studied - oysters and scallops. All steps of the proposed method must meet the quality criteria to be considered valid.

Each step of the analytical procedure was separately optimized for all 346 pesticide standards surveyed in this study. The method validation was performed using the following parameters: accuracy (expressed as recovery), precision (expressed as RSD), linearity (expressed as R<sup>2</sup>), limit of quantification and detection in accordance with the European Union (2020).

#### 3.2 Optimization of extraction / clean-up

QuEChERS extraction parameters such as organic solvent (MeOH and/or ACN), amount of water and added acid were tested considering the response of samples fortified with Mix of standards in the UPLC-MS/MS system. Both organic solvents had a good response. However, the mixture of ACN: $H_2O$  (10:1 v/v) presented better performance than the others.

The cleaning step using only PSA and MgSO4 produced extracts free of impurities for analysis. These results obtained with ACN and cleaning salts are compatible with previous studies using the QuEChERS method with fish samples.

## 3.3 Recovery and precision

One of the most important quality assessment parameters is accuracy. The recovery tests of all analytes in each matrix, native oyster Crassostrea gasar and scallop Nodipecten nodosus, were carried out at two recovery levels: 0.02 and 0.01 mg.kg<sup>-1</sup> of sample. The first recovery level corresponds to the first point on the curve and the second level to half of this value. These recovery tests were carried out on purpose at these low levels of concentration as we expected to find residues at really very low concentrations as is usually observed in the literature.

The study of the recovery and precision of all these 346 active principles was done through recovery tests in the matrix fortified with the MIX of standards as indicated in item 2.4.

Considering the linearity parameters (curves and the R<sup>2</sup> correlation coefficients), the accuracy and precision obtained for the 346 standards tested in this work, allowed proving that the vast majority (322 pesticides) among all the tested standards presented recovery in the range of 70 to 120%, recommended by the EU (European Union, 2020). In this specific case, the recovery was in the range of 76 to 106% and the accuracy was in the range of 3.5 to 13.6%. These values are fully consistent and within the criteria established by the EC. These results are very good and positively attest to the application of this method for the evaluation of pesticide residues in oyster and scallop meat.

#### 3.4 Linearity, LOD and LOQ

The determination of the working ranges as well as the linearity of the calibration curves were evaluated by the correlation coefficients ( $R^2$ ) of the 322 standards considered approved by the recovery tests. In addition, all calibration curves of pesticide standards provided  $R^2$  values greater than 0.95.

The limits of detection and quantification (LOD and LOQ) in the UPLC-MS/MS system were calculated using the signal-tonoise (s/r) ratio. The default values in this s/r ratio correspond to 3 for the limit of detection (LOD, s/r = 3) and 10 for the limit of quantification (LOQ, s/r = 10). In our case, the concentration of the first point of the curves corresponds to the LOQ. Therefore the values of detection and quantification limits for almost all analytes correspond to LOD=2 ng/mL and LOQ=6 ng/mL or LOD=0.2 and LOQ=0.6 ng/g of the sample.

#### 3.5 Analysis of oyster and scallop samples

The application of this multi-residue method of pesticides, validated for the two matrices in question, was carried out for the four samples of oysters and one of scallops produced in different cultures in the north / northeast / southeast of Brazil, being called A, B, C, D and E. All samples were fully evaluated following the analytical procedure described in the Material and Methods item.

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| Table 2 | . Mon | itored | transitions | of eva | luated | pesticides | s (to | be cont | inued). |
|---------|-------|--------|-------------|--------|--------|------------|-------|---------|---------|
|---------|-------|--------|-------------|--------|--------|------------|-------|---------|---------|

| Docticido                | MS/MS transitions $(m/z)$         | Docticido              | MS/MS transitions (m/z)                        |
|--------------------------|-----------------------------------|------------------------|--|
| Pesticide                |                                   | Preserve               |  |
| 2,4-D                    | 219 > 161   221 > 163             | Bromopnos metnyi       | 366 > 125   369 > 125                          |
| 2,4-DB                   | 247 > 161   247 > 125             | Bromuconazole          | 376 > 159   376 > 70                           |
| 2,6-Dichlorobenzamide    | 190 > 109   190 > 145             | Bupirimate             | 317 > 108   317 > 272                          |
| 3-Hydroxycarbofuran      | 238 > 163   238 > 181             | Buprofezin             | 306 > 201   306 > 116                          |
| Abamectin                | 891 > 305   891 > 567             | Butachlor              | 312 > 238   312 > 162                          |
| Acephate                 | 184 > 143   184 > 95              | Butocarboxim           | 213 > 75   213 > 116                           |
| Acetamiprid              | 223 > 126   223 > 90              | Butocarboxim-sulfoxide | 207 > 132   207 > 75                           |
| Acetochlor               | 270 > 224   270 > 148             | Cadusafos              | 271 > 159   271 > 215                          |
| Acibenzolar-S-methyl     | 211 > 136   211 > 140             | Carbaryl               | 219 > 145   219 > 127                          |
| Alachlor                 | 270 > 238 270 > 162               | Carbendazim            | 192 > 160   192 > 132                          |
| Alanycarb                | 400 > 238   400 > 91              | Carbetamide            | 237 > 192 237 > 118                            |
| Aldicarb                 | 191 > 116   191 > 89              | Carbofuran             | 227 > 165   227 > 123                          |
| Aldicarb sulfone         | 223 \ 86 223 \ 76                 | Carbosulfan            | $222 \times 103   222 \times 123$              |
| Aldiersk sulfáside       | 225 > 00   225 > 70               | Carbosulai             | $226 \times 142   226 \times 97$               |
| Aldicard suitoxide       | 207 > 132   207 > 89              | Carboxin               | 256 > 145   256 > 87                           |
| Ametryn                  | 228 > 186   228 > 96              | Carfentrazone-ethyl    | 412 > 346   412 > 266                          |
| Amicarbazone             | 242 > 143   242 > 85              | Carpropamid            | 334 > 139   334 > 196                          |
| Aminocarb                | 209 > 137   209 > 152             | Cartap                 | 238 > 73   238 > 150                           |
| Atrazine                 | 216 > 174   216 > 96              | Chlorantraniliprole    | 484 > 453   484 > 286                          |
| Azaconazole              | 300 > 159   300 > 231             | Chlorbromuron          | 294 > 206   294 > 182                          |
| Azadirachtin             | 719 > 687   719 > 491             | Chlordimeform          | 197 > 46   197 > 117                           |
| Azamethiphos             | 325 > 112   325 > 139             | Chlorfluazuron         | 540 > 383   540 > 158                          |
| Azinphos-ethyl           | 345 > 132   345 > 160             | Chlorimuron-ethyl      | 415 > 186   415 > 83                           |
| Azinphos-methyl          | 318 > 132   318 > 104             | Chloroxuron            | 291 > 72   291 > 164                           |
| Azocyclotin              | 369 > 205   369 > 287             | Chlorpyrifos           | 350 > 98   350 > 97                            |
| Azoxystrobin             | 404 > 372   404 > 329             | Chlorpyrifos-methyl    | 322 > 125   322 > 290                          |
| Benalaxyl                | 326 > 148   326 > 294             | Clethodim *            | 358 > 238   358 > 268                          |
| Bendiocarb               | 224 > 167   224 > 109             | Clofentezine           | 303 > 138 303 > 102                            |
| Benfuracarb              | 411 > 252 411 > 158               | Clomazone              | 240 > 125   240 > 89                           |
| Bentazone *              | 239 > 132   239 > 197             | Clorfenvinnhos         | 359 > 99   359 > 127                           |
| Bifenazate               | $200 \times 100   200 \times 100$ | Clothianidin           | 250 > 169   250 > 132                          |
| Bitertanol               | 338 \ 00   338 \ 70               | Coumanhas              | 250 > 105   250 > 152                          |
| Ditertation<br>Described | 242 > 207 242 > 271               | Countaprios            | 303 > 307   303 > 203                          |
| Boscalid                 | 543 > 507   543 > 271             | Desticite              | 303 > 185   303 > 125                          |
| Pesticide                | ms/ms transitions $(m/z)$         | Pesticide              | MS/MS transitions $(m/z)$                      |
| Cyazofamid               | 325 > 108   325 > 261             | DMSA                   | 201 > 92   201 > 137                           |
| Cycloxydim               | 326 > 280   326 > 180             | DMST                   | 215 > 106   215 > 79                           |
| Cyfçufenamid             | 413 > 203   413 > 295             | Dodemorph              | 282 > 116   282 > 98                           |
| Cyfluthrin               | 451 > 191   451 > 127             | Dodine                 | 228 > 57   228 > 60                            |
| Cyhexatin                | 369 > 205   369 > 287             | Doramectin             | 917 > 331   917 > 593                          |
| Cymoxanil                | 199 > 128   199 > 111             | Emamectin benzoate     | 886 > 126   886 > 302                          |
| Cypermethrin             | 433 > 191   433 > 416             | Epoxiconazole          | 330 > 121   330 > 123                          |
| Cyproconazole            | 292 > 70   292 > 125              | Eprinomectin           | 915 > 186   915 > 144                          |
| Cyprodinil               | 226 > 93   226 > 108              | EPTC                   | 190 > 128   190 > 86                           |
| Cyromazine               | 167 > 60   167 > 125              | Esfenvalerate          | 437 > 167   439 > 169                          |
| Daimuron                 | 269 > 151   269 > 91              | Esprocarb              | 266 > 91   266 > 71                            |
| Deltamethrin             | 523 > 281   523 > 506             | Ethidimuron            | 265 > 208   265 > 114                          |
| Demeton-S-methyl         | 231 > 89   231 > 61               | Ethiofencarb           | 226 > 107   226 > 169                          |
| Desmedinham              | 318 > 182   318 > 136             | Ethiofencarb-sulfone   | 275 > 107   275 > 201                          |
| Diafenthiuron            | 385 > 329   385 > 278             | Fthiofencarb-sulfoxide | 242 > 107   242 > 185                          |
| Diazinon                 | 305 > 169   305 > 97              | Ethion                 | 212 > 103   212 > 103<br>385 > 199   385 > 143 |
| Dichlomos                | $305 \times 100   305 \times 57$  | Ethicimol              | 210 > 140   210 > 08                           |
| Dialefront               | 221 > 107   221 > 127             |                        | 210 > 140   210 > 90                           |
| Diciotuanid              | 350 > 123   350 > 224             | Ethorumesate           | 28/ > 121   28/ > 259                          |
| Dicrotophos              | 238 > 112   238 > 72              | Ethoprophos            | 243 > 131   243 > 97                           |
| Diethotencarb            | 268 > 226   268 > 124             | Etiprole               | 414 > 351   414 > 255                          |
| Difenoconazole           | 406 > 251   406 > 188             | Etobenzanid            | 340 > 179   340 > 149                          |
| Difenoxuron              | 287 > 122   287 > 71              | Etofenprox             | 394 > 177   394 > 107                          |
| Diflubenzuron            | 311 > 158   311 > 113             | Etoxazole              | 360 > 141   360 > 57                           |
| Dimethenamid             | 276 > 244   276 > 168             | Etrimfos               | 293 > 125   293 > 265                          |
| Dimethoate               | 230 > 199   230 > 125             | Famoxadone             | 392 > 331   392 > 238                          |
| Dimethomorph             | 388 > 301   388 > 165             | Fenamidone             | 312 > 92   312 > 236                           |

Caption: pesticides with \* in bold are analyzed in ESI- mode; the others are analyzed in ESI+ mode.

# Table 2. Continued...

| Pesticide            | MS/MS transitions $(m/z)$                      | Pesticide          | MS/MS transitions $(m/z)$                      |
|----------------------|--|--------------------|--|
| Dimoxystrobin        | 327 > 116   327 > 89                           | Fenamiphos         | 304 > 217   304 > 202                          |
| Diniconazol          | 326 > 70   326 > 159                           | Fenarimol          | 331 > 268   331 > 81                           |
| Dinotefuran          | 203 > 129   203 > 123                          | Fenazaquin         | 307 > 57   307 > 161                           |
| Dioxacarb            | 224 > 167   224 > 123                          | Fenbuconazole      | 337 > 125   337 > 70                           |
| Disulfoton           | 275 > 89   275 > 61                            | Fenhexamid         | 302 > 97   302 > 55                            |
| Diuron               | 233 > 72   233 > 160                           | Fenitrothion       | 278 >184   278 > 125                           |
| Pesticide            | MS/MS transitions $(m/z)$                      | Pesticide          | MS/MS transitions $(m/z)$                      |
| Fenobucarb           | 208 > 95   208 > 152                           | Heptenophos        | 251 > 127   251 > 109                          |
| Fenoxycarb           | 302 > 88   302 > 116                           | Hexaconazole       | 314 > 70   314 > 159                           |
| Fenpropathrin        | 367 > 125   367 > 250                          | Hexythiazox        | 353 > 228   353 > 168                          |
| Fenpropidin          | 274 > 147   274 > 86                           | Imazalil           | 297 > 159   297 > 69                           |
| Fenpropimorph        | 304 > 147   304 > 130                          | Imazapic           | 276 > 231   276 > 163                          |
| Fenpyroximate        | 422 > 366   422 > 138                          | Imazapyr           | 262 > 69   262 > 86                            |
| Fenthion             | 279 > 169   279 >105                           | Imazaquin          | 312 > 266   312 > 86                           |
| Fenthion-sulfoxide   | 295 > 109   295 > 79                           | Imazethapyr        | 290 > 245   290 > 86                           |
| Fenuron              | 165 > 72   165 > 46                            | Imazosulfuron      | 413 > 153   413 > 156                          |
| Fenvalerate          | 437 > 167   439 > 169                          | Imibenconazole     | 411 > 125   411 > 171                          |
| Fipronil             | 435 > 330   435 > 250                          | Imidacloprid       | 256 > 175   256 > 209                          |
| Flonicamid           | 230 > 203   230 > 148                          | Indoxacarb         | 528 > 203   528 > 218                          |
| Fluazifop-p-butyl    | 384 > 282   384 > 328                          | Ioxynil            | 370 > 127   370 > 243                          |
| Fluazinam *          | 463 > 416   463 > 398                          | Iprovalicarb       | 321 > 119   321 > 203                          |
| Flubendiamide        | 683 > 274   683 > 408                          | Isocarbamid        | 186 > 87   186 > 130                           |
| Flufenacet           | 364 > 194   364 >152                           | Isocarbophos       | 291 > 231   291 > 121                          |
| Flutenoxuron         | 489 > 158   489 > 141                          | Isotenphos         | 346 > 245   346 > 217                          |
| Fluoxastrobin        | 459 > 427   459 > 188                          | Isoprocarb         | 194 > 95   194 > 137                           |
| Fluquinconazole      | 376 > 349   376 > 108                          | Isoprothiolane     | 291 > 231   291 > 189                          |
| Flusilasole          | 316 > 24/   316 > 165                          | Isoproturon        | 20/ > /2   20/ > 46                            |
| Flusuitamide         | 413 > 1/1   413 > 1/9                          | Isoxaflutole       | 359 > 251   359 > 220                          |
| Flutniacet-metnyi    | $404 > 2/4 \mid 404 > 215$                     | Isoxathion         | 314 > 105   314 > 286                          |
| Flutoiann            | 324 > 202   324 > 05                           | Verhetilete        | 895 > 307   278 > 124                          |
| Fluthalol            | 502 > 70   502 > 125                           | Karbutilate        | 2/8 > 1/9   2/8 > 134                          |
| Fluxapyroxad         | 382 > 342   382 > 314<br>437 > 195   437 > 286 | Lactofen           | 514 > 110   514 > 207<br>479 > 344   479 > 462 |
| Forchlorfenuron      | 437 > 193   437 > 200<br>248 > 120   248 > 03  | Lambda cybalothrin | 479 > 344   479 > 402                          |
| Formetanate          | 240 > 129   240 > 93                           | Lambua-Cynaiotinin | 407 > 225   407 > 450<br>249 > 160   249 > 182 |
| Fuberidazole         | 185 > 157   185 > 156                          | Lufenuron *        | 509 > 323   509 > 339                          |
| Furalaxyl            | 302 > 95   302 > 242                           | Malathion          | 331 > 127   331 > 99                           |
| Furathiocarb         | 383 > 195   383 > 252                          | Mandipropamid      | 412 > 328   412 > 125                          |
| Halofenozide         | 331 > 275   331 > 105                          | Mefenacet          | 299 > 148   299 > 120                          |
| Pesticide            | MS/MS transitions $(m/z)$                      | Pesticide          | MS/MS transitions $(m/z)$                      |
| Mepanipyrim          | 224 > 106   224 > 77                           | Omethoate          | 214 > 183   214 > 125                          |
| Mephosfolan          | 270 > 140   270 > 196                          | Oxadiargyl         | 341 > 151   341 > 230                          |
| Mepronil             | 270 > 119 270 > 91                             | Oxadixyl           | 279 > 219   279 > 132                          |
| Mesotrione           | 340 > 228   340 > 104                          | Oxamyl             | 237 > 72   237 > 90                            |
| Metalaxyl-M          | 280 > 220   280 > 192                          | Oxamyl Oxime       | 163 > 72   163 > 90                            |
| Metamidophos         | 142 > 94   142 > 125                           | Oxycarboxin        | 268 > 175   268 > 147                          |
| Metconazole          | 320 > 70   320 > 125                           | Paclobutrazol      | 294 > 70   294 > 125                           |
| Methfuroxan          | 230 > 137   230 > 111                          | Penconazole        | 284 > 70   284 > 159                           |
| Methidation          | 303 > 145   303 > 85                           | Pencycuron         | 329 > 125   329 > 218                          |
| Methiocarb           | 226 > 169   226 > 121                          | Pendimethalin      | 282 > 212   282 > 194                          |
| Methiocarb-sulfone   | 275 > 122   275 > 201                          | Permethrin         | 408 > 183   408 > 335                          |
| Methiocarb-sulfoxide | 242 > 185   242 > 122                          | Phenmedipham       | 301 > 168   301 > 136                          |
| Methomyl             | 163 > 88   163 > 106                           | Phentoato          | 321 > 247   321 > 163                          |
| Methoprene           | 311 > 279   311 > 191                          | Phosalone          | 368 > 182   368 > 111                          |
| Methoprotryne        | 272 > 198   272 > 170                          | Phosmet            | 318 > 160   318 > 133                          |
| Metobromuron         | 259 > 170   259 > 148                          | Phosphamidon       | 300 > 174   300 > 127                          |
| Metoxuron            | 229 > 72   229 > 156                           | Phoxim             | 300 > 129   300 > 125                          |
| Metoxyfenozide       | 369 > 149   369 > 313                          | Picoxystrobin      | 368 > 205   368 > 145                          |
| Metrafenone          | 409 > 209   409 > 227                          | Piperonyl butoxide | 356 > 177   356 > 119                          |

Caption: pesticides with \* in bold are analyzed in ESI- mode; the others are analyzed in ESI+ mode.

## Table 2. Continued...

| Pesticide          | MS/MS transitions $(m/z)$ | Pesticide             | MS/MS transitions $(m/z)$ |
|--------------------|---------------------------|-----------------------|---------------------------|
| Metribuzin         | 215 > 131   215 > 89      | Pirimicarb            | 239 > 72   239 > 182      |
| Metsulfuron-methyl | 382 > 167   382 > 199     | Pirimicarb-desmethyl  | 225 > 72   225 > 168      |
| Mevinphos          | 225 > 127   225 > 193     | Pirimiphos-ethyl      | 334 > 198   334 > 182     |
| Molinate           | 188 > 126   188 > 55      | Pirimiphos-methyl     | 306 > 108   306 > 67      |
| Monalide           | 240 > 85   240 > 128      | Prochloraz            | 376 > 308   376 > 266     |
| Monocrotophos      | 224 > 127   224 > 98      | Profenofos            | 375 > 305   375 > 347     |
| Monolinuron        | 215 > 148   215 > 99      | Prometon              | 226 > 184   226 > 86      |
| Moxidectin         | 641 > 528   641 > 498     | Prometryne            | 242 > 158   242 > 200     |
| Myclobutanil       | 289 > 70   289 > 125      | Propanil              | 218 > 162   218 > 127     |
| Neburon            | 275 > 88   275 > 57       | Propargite            | 368 > 231   368 > 175     |
| Nitenpyram         | 271 > 225   271 > 126     | Propazine             | 230 > 146   230 > 188     |
| Norflurazon        | 304 > 284   304 > 160     | Propham               | 180 > 120   180 > 138     |
| Novaluron          | 493 > 158   493 > 141     | Propiconazole         | 342 > 69   342 > 159      |
| Nuarimol           | 315 > 252   315 > 81      | Propoxur              | 210 > 111   210 > 93      |
| Pesticide          | MS/MS transitions $(m/z)$ | Pesticide             | MS/MS transitions (m/z)   |
| Propyzamide        | 256 > 190   256 > 173     | Teflubenzuron *       | 379 > 339   379 > 196     |
| Proquinazid        | 373 > 289   373 > 331     | Temephos              | 467 > 419   467 > 125     |
| Prothioconazole    | 344 > 189   344 > 326     | Tepraloxydim          | 342 > 250   342 > 166     |
| Pymetrozin         | 218 > 105   218 > 78      | Terbufos              | 289 > 103   289 > 57      |
| Pyraclostrobin     | 388 > 194   388 > 163     | Terbumeton            | 226 > 170   226 > 114     |
| Pyrazophos         | 374 > 222   374 > 194     | Terbutryn             | 242 > 186   242 > 91      |
| Pyridaben          | 365 > 147   365 > 309     | Tetraconazole         | 372 > 159   372 > 70      |
| Pyridaphention     | 341 > 189   341 > 92      | Thiacloprid           | 253 > 126   253 > 90      |
| Pyrifenox          | 295 > 93   295 > 66       | Thiobencarb           | 257 > 124   257 > 100     |
| Pyrimethanil       | 200 > 107   200 > 82      | Thiodicarb            | 355 > 88   355 > 108      |
| Pyriproxyfen       | 322 > 96   322 > 185      | Thiofanate-metyl      | 343 > 151   343 > 93      |
| Quinalphos         | 299 > 163   299 > 147     | Thiofanox             | 219 > 57   219 > 76       |
| Quinoxyfen         | 308 > 197   308 > 162     | Thiofanox-sulfone     | 268 > 57   268 > 76       |
| Quizalofop-P-ethyl | 379 > 211   379 > 115     | Thiofanox-sulfoxide   | 252 > 235   252 > 178     |
| Rotenone           | 395 > 213   395 > 192     | Tiabendazole          | 202 > 175   202 > 131     |
| Sebuthylazine      | 230 > 174   230 > 96      | Tiamethoxam           | 292 > 211   292 > 181     |
| Siduron            | 233 > 94   233 > 137      | Tolclofos-metyl       | 301 > 269   301 > 175     |
| Simazine           | 202 > 132   202 > 124     | Tolylfluanide         | 363 > 238   363 > 137     |
| Simetryn           | 214 > 124   214 > 96      | Triadimefon           | 294 > 69   294 > 197      |
| Spinetoram         | 749 > 142   749 > 98      | Triadimenol           | 296 > 70   296 > 99       |
| Spinosad A         | 733 > 142   733 > 98      | Triazophos            | 314 > 162   314 > 119     |
| Spinosad D         | 747 > 142   747 > 98      | Triclhorfon           | 257 > 109   257 > 127     |
| Spirodiclofen      | 411 > 71   411 > 313      | Tricyclazole          | 190 > 162   190 > 136     |
| Spiromesifen       | 371 > 273   371 > 255     | Tridemorph            | 298 > 57   298 > 98       |
| Spirotetramat      | 374 > 330   374 > 302     | Trifloxystrobin       | 409 > 186   409 > 145     |
| Spiroxamine        | 298 > 144   298 > 100     | Triflumizole          | 346 > 278   346 > 73      |
| Sulfentrazone      | 387 > 146   387 > 307     | Triflumuron           | 359 > 156   359 > 139     |
| Tebuconazole       | 308 > 70   308 > 125      | Triflusulfuron-methyl | 493 > 264   493 > 96      |
| Tebufenozide       | 353 > 133   353 > 297     | Triforine             | 435 > 390   435 > 215     |
| Tebufenpyrad       | 334 > 117   334 > 145     | Triticonazole         | 318 > 70   318 > 125      |
| Tebupirimfos       | 319 > 276   319 > 153     | Vamidothion           | 288 > 146   288 > 118     |
| Tebuthiuron        | 229 > 172   229 > 116     | Zoxamide              | 336 > 187   336 > 159     |

Caption: pesticides with \* in bold are analyzed in ESI- mode; the others are analyzed in ESI+ mode.

These analyzes produced very interesting results. The data obtained clearly showed residues of some pesticides at the trace level for four of the five samples evaluated. One of the oyster samples (C) did not show any residue from any of the 322 agrochemicals monitored in this study.

This result, at first, surprised us, but proved total adherence with the fact that this sample comes from an organic crop considered as standard. The other four samples evaluated, A, B, D and E, showed the presence of pesticide residues in very low concentrations, at the trace level, as shown in Table 3.

The application of the pesticide multi-residue method proposed in this study in the evaluation of bivalve molluscs allowed the presence of some pesticides to be evidenced. If only routine analyzes were carried out for these samples, this occurrence would not be pointed out because they are not yet part of the mollusc production

Table 3. Pesticides found in BM samples through multiresidue analysis.

| Samp     | ole | Pesticide          |
|----------|-----|--------------------|
| oysters  | А   | Methyl pyrimiphos  |
|          | В   | Methyl pyrimiphos  |
|          |     | Clomazone          |
|          | С   | -                  |
|          | D   | Sprocarb           |
|          |     | Picoxystrobin      |
|          |     | Promethrin         |
|          |     | Tebupirinphos      |
|          |     | Tricyclazole       |
| scallops | Е   | Piperonyl butoxide |
|          |     | Methyl pyrimiphos  |
|          |     | Thiabendazole      |

process and, therefore, would not be monitored. In the context of food safety, it is important to consider the danger associated with contamination of the environment in which molluscs live.

Pesticide residues from areas adjacent to the cultivations may be present in the muscle of molluscs, indicating that these toxic substances from other regions are contributing to the increase of contamination in areas where shellfish are cultivated. Careful analysis of data on food contamination associated with environmental characteristics around mollusc cultivation areas may generate solutions to this problem through the application of corrective measures.

It should be considered that the quality of a food is intrinsically linked to both its phytosanitary nature and the presence of contaminants. Pesticides affect human health in many ways and the increase in levels of contamination in food can drastically affect its quality and even make it unsuitable for consumption, greatly reducing its commercial value.

Comprehensive and effective control of all possible sources of contamination can provide a complete and more accurate assessment of food quality. In this context, the application of a multi-residue method with 322 analytes, commonly used in agricultural activities as proposed in this work, is extremely valuable.

## **4** Conclusion

The multi-residue pesticide method validated for oyster (*Crassostrea gasar*) and scallop (*Nodipecten nodosus*) matrices according to the requirements recommended by the European Union (2020) proved to be adequate to quantify 322 pesticides listed in Table 2.

The results of the pesticide residue analysis of all five samples of bivalve mollusks in this study clearly showed that they all meet the level of healthiness and fully meet the quality standard of the legislation.

Among the five samples of this work (A, B, C, D and E), three of the four samples of oysters and one of scallops (named A, B, D and E) showed the presence of residues only at the trace level and not causes damage to health. This result attests to the quality of oyster and scallop meat from these crops, guaranteeing its safety as food.

Among the oyster samples, only the sample C from Rio Grande do Norte did not present any pesticide residue, not even at the trace level. This sample was produced in an "organic area" and this fact can be proven because it did not present abiotic stresses.

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