

FOOD CHEMICAL CONTAMINANTS

Quantitative Determination of the Okadaic Acid Toxins Group by a Colorimetric Phosphatase Inhibition Assay: Interlaboratory Study

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An interlaboratory collaborative study to validate a colorimetric phosphatase inhibition assay for quantitative determination of the okadaic acid (OA) toxins group in molluscs, OkaTest, was conducted. Eight test materials, including mussels, scallops, clams, and cockles, were analyzed as blind duplicates. Blank samples and materials containing different OA toxin levels ranging from 98 to 275 µg/kg OA equivalents were included. The study was carried out by a total of 16 laboratories from 11 different countries. Values obtained for repeatability relative standard deviations (RSD_r) ranged from 5.4 to 11.2% (mean 7.5%). Reproducibility RSD (RSD_R) values were between 7.6 and 13.2% (mean 9.9%). The Horwitz ratio (HorRat) values ranged between 0.4 and 0.6. A recovery assay was also carried out using a sample spiked with OA. A mean recovery of 98.0% and an RSD of 14.5% were obtained. The results obtained in this validation study indicate that the colorimetric phosphatase inhibition assay, OkaTest, is suitable for quantitative determination of the OA toxins group. OkaTest could be used as a test that is complementary to the reference method for monitoring the OA toxins group.

characterized by symptoms, such as diarrhea, nausea, vomiting, and abdominal pain. These symptoms may occur in humans shortly after consumption of contaminated bivalve molluscs, such as mussels, clams, scallops, or oysters. Inhibition of serine/threonine phosphoprotein phosphatases (PPs) is assumed to be responsible for these toxic effects. These compounds are also involved in tumor promotion (1). Therefore, these toxins are regulated by European Union law.

Regulation (EC) No. 853/2004 (2) states that live bivalve molluscs placed on the market for human consumption must not contain marine biotoxins in total quantities (measured in the whole body or any part edible separately) that exceed 160 µg of OA equivalents/kg for OA, dinophysistoxins, and pectenotoxins together.

Commission Regulation (EC) No. 15/2011 (3) indicates that in the case of lipophilic toxins including OA toxins, LC/MS/MS is the reference method for routine testing of official controls or any checks done by food operators. This regulation has recently amended the Commission Regulation (EC) No. 2074/2005 (4), in which biological methods (mouse and rat bioassay) were considered the reference. From now on, they will only be used for a transitional period of time (until the end of 2014) or in special circumstances.

Both regulations (No. 2074/2005 and No. 15/2011) contemplate other methods for routine testing of lipophilic toxins, providing they are intralaboratory-validated and successfully tested under a recognized proficiency test scheme. Those methods should detect, either alone or in combination with others, all of the lipophilic toxin analogs (OA, pectenotoxins, yesotoxins, and azaspiracids group toxins). The protein phosphatase inhibition assay (PIIA) is specifically mentioned in these regulations as an alternative or complementary method, considering that the PPs are known to be OA-toxins natural targets (5, 6). In-house PPIAs using different phosphatase sources and colorimetric or fluorometric substrates have been previously developed (7–12). Later improvements to detect all OA derivatives by hydrolysis of samples were also suggested

Okadaic acid (OA) and its analogs dinophysistoxin-1 and -2 (DTX1, DTX2), together with their ester forms, are known as the OA toxins group. These lipophilic and heat stable toxins are produced by dinoflagellates and can be found in various species of shellfish, mainly in filter-feeding bivalve molluscs.

OA toxins causes diarrhetic shellfish poisoning, which is

Received October 28, 2011. Accepted by AP May 23, 2012.

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DOI: 10.5740/jaoacint.11-465

Table 1. Details of matrixes and species origin of test materials used in this study

| Code | Matrix/Species | Origin |
|------|--|--|
| A | Mussel (<i>M. galloprovincialis</i>) | Galicia (NW Spain) |
| D | Clam (<i>V. pullastra</i>) | Food & Agricultural Organization, 37 Mediterranean Sea |
| E | Mussel (<i>M. galloprovincialis</i>) | Galicia (NW Spain) |
| F | Scallop (<i>P. maximus</i>) | FAO 27 NE Atlantic |
| G | Clam (<i>V. decussatus</i>) | Galicia (NW Spain) |
| K | Clam (<i>V. romboides</i>) | Galicia (NW Spain) |
| L | Cockle (<i>C. edulis</i>) | Portugal and Galicia (NW Spain) |
| N | Mussel (<i>M. edulis</i>) | Ireland |
| BM | Scallop (<i>P. maximus</i>) | Scotland |

(13), and a collaborative study was also performed with a fluorometric PPIA (14). However, none of those assays was commercially available for routine analysis, nor were they demonstrated to comply with the legislation requirements.

ZEU-INMUNOTEC (Zaragoza, Spain) has developed a commercial kit (OkaTest, formerly Toxiline-DSP) based on a colorimetric PP2A inhibition assay for quantification of the OA toxins group in molluscs (15).

The PPIA described in this study uses a human PP2A purified by ZEU-INMUNOTEC that has showed higher sensitivity than other commercial and genetic engineering produced enzymes (16). PP2A was stabilized by freeze-drying to obtain a standardized assay with shelf life of up to 12 months at 4°C (15). Colorimetric substrate was chosen over a fluorometric one as the latter is less stable and, therefore, less appropriate for ready-to-use kits. Besides, fluorometric assays require specific equipment not often available in routine testing laboratories; therefore, they are difficult to use for monitoring purposes.

The robustness and performance of OkaTest were evaluated by the manufacturer in a single-laboratory validation according to AOAC and Eurachem guidelines (15). All of the results obtained showed that the OkaTest kit is robust and accurate, and, therefore, suitable for an interlaboratory study.

Interlaboratory Study

A colorimetric PPIA, OkaTest, was interlaboratory-validated for quantification of the OA toxins group. The main purpose of this study was to determine repeatability and between-laboratory reproducibility. A recovery assay was also carried out, and accuracy of the method confirmed.

A validation management team (David Clarke, Elena Domínguez, Katrin Kapp, Panagiota Katikou, and María Luisa Rodríguez) was appointed to supervise, advise on the accomplishment of the study, and ensure its independence. A total of 16 laboratories from 11 different countries in Europe and South America participated in the study.

The study plan including details of the test method, experimental design, preparation of test materials, instructions for participants, key personnel, schedule, and data analysis was prepared and agreed to by the validation management team.

Participants were fully informed of the study design prior to distribution of testing materials.

Eight different test materials, as blind duplicates, were analyzed by each laboratory on 2 different days. Five materials contained different OA toxin levels, all naturally contaminated except for one that was partially spiked. Three of the test materials were blank samples. An additional blank material (BM) was used in the recovery study. The test materials comprised four different genera of molluscs (*Mytilus spp.*, *Pecten spp.*, *Venerupis spp.*, and *Cerastoderma spp.*) and seven different species. Details of the materials used are shown in Table 1. The materials were prepared by the Spanish Association of Seafood Products Manufacturers (ANFACO-CECOPECA; Vigo, Spain) as explained below.

All participants sent back an electronic copy of a tailor-made Excel reporting sheet for each day of analysis with raw data and final results for each test material. The reporting sheets were checked upon receipt for obvious errors in sample codes and calculations.

Participants also completed a questionnaire with details of the equipment used and preparation of reagents and samples, as well as feedback on the assay.

Preparation of Test Materials

Materials A and E (mussel) and D and K (clam) were purchased from the retail market fresh and alive. They were thoroughly cleaned outside and inside with fresh water to remove sand and any other foreign materials. Tissues were removed from the shell, transferred to strainers, and drained for 5 min before homogenization (blender and Ultraturrax[®]; IKA, Staufen, Germany). The homogenate (at least 450 g) was then distributed into plastic containers (5.0 ± 0.1 g), frozen, and stored at -20 ± 2°C until analysis or the day of shipment.

Materials F (scallop) and G (clam) were purchased frozen

Table 2. Total concentration of OA toxins group (µg/kg) determined by OkaTest, and toxins profile by LC/MS/MS

| Test material ^a | Matrix/species | Total OA equivalents, µg/kg ^b | OA toxins content ^c |
|----------------------------|--|--|--------------------------------|
| BM | Scallop (<i>P. maximus</i>) | <LOD | — |
| A | Mussel (<i>M. galloprovincialis</i>) | <LOD | — |
| F | Scallop (<i>P. maximus</i>) | <LOD | — |
| G | Clam (<i>V. decussatus</i>) | <LOD | — |
| E | Mussel (<i>M. galloprovincialis</i>) | 79 ± 5 | OA |
| L | Cockle ^d (<i>C. edulis</i>) | 168 ± 11 | OA, DTX1, and DTX2 |
| D | Clam (<i>V. pullastra</i>) | 240 ± 9 | OA |
| K | Clam (<i>V. romboides</i>) | 250 ± 6 | OA |
| N | Mussel ^e (<i>M. edulis</i>) | 276 ± 6 | OA and DTX2 |

^a Samples presented in increasing order of concentration.

^b Determined by OkaTest; LOD = 44 OA equivalents µg/kg.

^c Determined by LC/MS/MS.

^d Artificially contaminated with DTX1 and mixed with blank material.

^e Mixed with blank material.

Table 3. Results from homogeneity study for test materials for the determination of OA (μg OA total equivalents/kg)

| Test material | Variance of sums, V_s | Analytical variance, | Allowable sampling | Sampling variance, | Critical value, c | Test for homogeneity result |
|---------------|-------------------------|----------------------|----------------------------|--------------------|---------------------|-----------------------------|
| | | s_{an}^2 | variance, σ_{all}^2 | S_{sam}^2 | | |
| D | 166 | 90.7 | 36.8 | 116 | 310 | $S_{sam}^2 < c$ |
| E | 84.7 | 8.09 | 19.8 | 11.1 | 29.1 | $S_{sam}^2 < c$ |
| K | 139 | 19.6 | 32.5 | 126 | 257 | $S_{sam}^2 < c$ |
| L | 356 | 46.9 | 85.7 | 55.6 | 152 | $S_{sam}^2 < c$ |
| N | 124 | 24.2 | 28.4 | 154 | 314 | $S_{sam}^2 < c$ |

from the retail market. They were thawed at room temperature, cleaned, and prepared as described above.

Material L (cockle) was provided cleaned, blended, and frozen by the European Reference Laboratory for Marine Biotoxins (EURLMB, Vigo, Spain). The sample contained OA, DTX2, and traces of DTX1. In order to achieve a suitable toxin profile, the sample was mixed with fresh cockle from the same species (*C. edulis*) without toxin prior to being spiked with DTX1 (Wako Chemicals, Neuss, Germany). The sample was thawed at room temperature, mixed with the cockle blank material (purchased in Porto, Portugal), and spiked. Then, it was distributed into plastic containers (5.0 ± 0.1 g), frozen, and stored at $-20 \pm 2^\circ\text{C}$ until the day of shipment.

Material N (mussel) was provided cleaned, blended, and frozen by the National Reference Laboratory of Ireland, Galway, Ireland. The sample contained a high level of OA toxins, so it was mixed with mussel (*M. edulis*) without toxin (purchased in a retail market in Ireland) to achieve a suitable toxin concentration. The sample was thawed at room temperature, mixed, and distributed into plastic containers (5.0 ± 0.1 g). The material was then frozen and stored at $-20 \pm 2^\circ\text{C}$ until the day of shipment.

The BM (scallop) was provided blended and homogenized by Integrin Advanced Bioscience (Oban, Scotland) and stored frozen at approximately $-20 \pm 2^\circ\text{C}$ until the day of shipment.

Homogeneity and stability of test materials were studied according to the International Harmonized Protocol for the Proficiency Testing of Analytical Chemistry Laboratories (17). Ten containers of 5 g were randomly selected for each material. The content of each container was homogenized and extracted, and two test portions (from the sample extract) were analyzed to estimate the analytical variance. A total of 20 portions/material

were tested under repeatability conditions and in a random order using the OkaTest kit.

To ensure the stability of the materials during shipment to participants and the study duration, aliquots of each material were taken randomly and split into two subsets, each of them containing five samples. One subset was used as control and stored at $-18 \pm 1^\circ\text{C}$. The second was stored under experimental conditions of $9.0 \pm 1^\circ\text{C}$ for 5 days. Samples of both subsets were randomized before testing and analysis simultaneously using the OkaTest kit under repeatability conditions. The test materials were also analyzed by LC/MS/MS (18, 19) to determine the OA toxin profile.

The test materials were blind coded by EURLMB and distributed by ANFACO-CECOPESCA to the participants. The codes were securely kept by EURLMB until statistical analysis was carried out.

The materials were shipped in isothermal boxes with dry ice and were received within the following 2 days by most participants. Materials sent to South American countries were delivered more than a week after the dispatch date, as they have long customs check up procedures. Samples were, however, reported to have been kept frozen while stored at customs. Two laboratories informed that the box containing the samples did not arrive in good conditions, and six reported that samples were cold, but defrosted.

PPIA

Principle

OkaTest is an enzymatic test based on a colorimetric PPIA for quantitative determination of OA and other toxins of the OA group, including DTX1, DTX2, and their ester forms.

Table 4. Results obtained for the stability assays conducted for materials D, E, K, L, and N

| Test material | Storage conditions | | Absolute difference D | Variance F -test | t -test | Test criterion | |
|---|---------------------------|---------------------------|-----------------------|--------------------|-----------|----------------|-------|
| | $-18 \pm 1^\circ\text{C}$ | $9.0 \pm 1^\circ\text{C}$ | | | | C | D < C |
| | Mean | | | | | | |
| Total OA equivalents, $\mu\text{g}/\text{kg}$ | | | | | | | |
| D | 265 ± 10 | 262 ± 15 | 3.02 | 0.54 | 0.71 | 34.5 | Pass |
| E | 84.0 ± 4 | 85.1 ± 3 | -1.19 | 0.45 | 0.62 | 10.9 | Pass |
| K | 255 ± 8 | 257 ± 7 | -1.57 | 0.87 | 0.75 | 33.2 | Pass |
| L | 171 ± 7 | 169 ± 8 | 1.63 | 0.79 | 0.73 | 22.2 | Pass |
| N | 343 ± 24 | 355 ± 32 | -13.0 | 0.58 | 0.49 | 44.6 | Pass |

Table 5. Calibration curve parameters obtained by each laboratory every day of the study

| Lab | R ² | | Slope | | Absorbance 405 nm, lowest standard 0.5 nM | | Absorbance 405 nm, highest standard 2.8 nM | |
|----------------|----------------|-------|-------|-------|--|-------|---|-------|
| | Day 1 | Day 2 | Day 1 | Day 2 | Day 1 | Day 2 | Day 1 | Day 2 |
| A | 0.99 | 0.98 | -0.12 | -0.45 | 0.734 | 1.287 | 0.524 | 0.505 |
| B | 0.99 | 0.99 | -0.50 | -0.65 | 1.157 | 1.425 | 0.334 | 0.339 |
| C | 0.98 | 0.98 | -0.64 | -0.44 | 1.530 | 1.177 | 0.496 | 0.468 |
| D | 0.98 | 0.98 | -0.67 | -0.58 | 1.537 | 1.402 | 0.430 | 0.459 |
| E | 0.97 | 0.98 | -0.51 | -0.48 | 1.222 | 1.221 | 0.409 | 0.436 |
| F | 1.00 | 0.99 | -0.72 | -0.74 | 1.684 | 1.726 | 0.482 | 0.491 |
| G | 0.98 | 1.00 | -0.79 | -0.58 | 1.781 | 1.411 | 0.462 | 0.423 |
| H | 0.99 | 0.99 | -0.78 | -0.73 | 1.644 | 1.609 | 0.366 | 0.414 |
| I | 0.99 | 0.99 | -0.76 | -0.68 | 1.661 | 1.486 | 0.409 | 0.357 |
| J | 0.97 | 0.98 | -0.41 | -0.45 | 1.164 | 1.204 | 0.498 | 0.458 |
| K | 0.99 | 0.98 | -0.77 | -0.74 | 1.712 | 1.690 | 0.438 | 0.485 |
| L ^a | 0.93 | 0.96 | -0.63 | -1.13 | 1.488 | 2.588 | 0.425 | 0.709 |
| M | 0.99 | 0.99 | -0.78 | -0.65 | 1.697 | 1.464 | 0.419 | 0.390 |
| N | 0.99 | 0.98 | -0.54 | -0.65 | 1.273 | 1.497 | 0.384 | 0.444 |
| O | 0.97 | 0.98 | -0.49 | -0.32 | 1.188 | 0.992 | 0.396 | 0.470 |
| P | 0.97 | 0.99 | -0.27 | -0.58 | 1.015 | 1.474 | 0.549 | 0.520 |

^a Standard curve obtained by Laboratory L on Day 1 was rejected as R² criterion was not met. Assay could not be repeated due to time issues.

This method is applicable to shellfish species, such as mussels, clams, cockles, and scallops.

The toxicity of the OA toxins group is directly related to its inhibitory activity against a family of structurally related PPs, in particular PP1 and PP2A. OkaTest uses this strong inhibitory activity to determine the OA content in shellfish using the PP2A with a chromogenic substrate for this enzyme. After the substrate's hydrolysis by the enzyme, the product can be measured at 405 nm by a microplate reader. As the ability of the PPs to hydrolyze the substrate depends on the amount of OA and analogs in the samples, the toxin concentration can be calculated by using a standard curve.

Apparatus

(a) *Micropipets*.—Adjustable 100, 200, and 1000 μ L (Thermo LabSystems, Helsinki, Finland).

(b) *Ultra homogenizer*.

(c) *Block heater or incubator*.—For $30 \pm 2^\circ\text{C}$ (ZEU-INMUNOTEC, Zaragoza, Spain).

(d) *Microwell absorbance reader*.—405 \pm 10 nm wavelength filter (Thermo LabSystems).

(e) *Water bath*.—Set at $76 \pm 2^\circ\text{C}$ (Raypa, Barcelona, Spain).

(f) *Centrifuge tubes*.—Graduated 50 mL.

(g) *Laboratory glassware*.

Reagents

(a) *Extraction solvent*.—Methanol, reagent grade, 100% (v/v; Sharlab, Barcelona, Spain).

(b) *HCl*.—Reagent grade, 37% (v/v; Sharlab).

(c) *NaOH*.—Reagent grade (Sharlab).

(d) *Deionized water*.—Type II, ISO 3696 (Ellix 5; Millipore, Germany).

(e) *OkaTest kit*.—From ZEU-INMUNOTEC containing:

(1) 96-well microtiter plate and plate adhesive film.

(2) Lyophilized PP2A purified from human blood cells.

(3) Ready-to-use OA Standards of 0.5, 0.8, 1.2, 1.8, and 2.8 nM, prepared from the OA reference solution (NRC CRM-OA-c, Institute for Marine Biosciences, Halifax, Canada).

(4) Chromogenic substrate.

(5) Phosphatase dilution buffer.

(6) Stock buffer solution.

(7) OA Spiking solution (2 μ M) prepared from the OA reference solution (NRC CRM-OA-c, Institute for Marine Biosciences).

Spiking Procedure

Due to the limited experience on the homogeneity and stability of spiked samples with OA toxins, each participant prepared a spiked sample on the day of the assay. A BM and an OA solution of known concentration (2 μ M, to prepare a final concentration of 161 μ g/kg) were provided to each participant.

A blank sample was spiked with OA solution for the recovery study as follows:

(a) Mix 500 μ L OA spiking solution (2 μ M) with 5.0 ± 0.1 g homogenous blank sample.

(b) Add 25 mL extraction solvent [methanol, 100% (v/v)] to the mixture and shake for 2 min by vortexing. Proceed with the extraction procedure described below under point (b).

Sample Extraction

(a) Thaw each aliquot with 5.0 ± 0.1 g homogenized mollusc at room temperature ($22 \pm 2^\circ\text{C}$). Add 25 mL extraction solvent [methanol, 100% (v/v)]; then mix for 2 min using an ultra homogenizer.

(b) Centrifuge at 2000 g for 10 min at 4°C. The supernatant is called “methanolic extract.”

(c) Pipet 640 µL methanolic extract into a 50 mL graduated centrifuge tube and add 100 µL 2.5 M NaOH.

(d) Seal the test tube and heat at $76 \pm 2^\circ\text{C}$ for 40 min in a water bath.

(e) Do not cool the sample; add 80 µL 2.5 M HCl immediately.

(f) Add 19.18 mL buffer solution with a glass pipet up to a total volume of 20 mL.

Assay Procedure

(a) Rehydrate the lyophilized phosphatase (PP2A) by adding 2.0 mL phosphatase dilution buffer to the vial and mix gently for 60 ± 5 min at room temperature ($22 \pm 2^\circ\text{C}$) on a roller mixer or a shaker (maximum 60 rpm) (both from JP Selecta, Barcelona, Spain).

(b) Add 50 µL each sample extract or standard to wells. Samples and standards have to be analyzed in duplicate.

(c) Add 70 µL phosphatase solution to each well. Cover the plate with the adhesive film provided in the kit, and mix by gentle tapping on the side.

(d) Incubate at $30 \pm 2^\circ\text{C}$ for 20 ± 0.5 min.

(e) Remove the adhesive film and add 90 µL chromogenic substrate to each well and mix by tapping gently on the side. Incubate at $30 \pm 2^\circ\text{C}$ for 30 ± 0.5 min.

(f) Read the absorbance of samples and standards at 405 ± 10 nm.

Calculations

The results were calculated from a standard curve by plotting the absorbance values on a linear y axis and the concentration of OA on a logarithmic x axis, and using a logarithmic fitting. As an acceptability criterion for the assay, the Pearson correlation coefficient R^2 had to be equal to or greater than 0.96. The OA concentration contained in the sample was then calculated using the following equation:

$$x = \text{EXP}(y - b)/a$$

where x is the OA concentration in the sample (C_s), y the absorbance of the sample, a is the slope, and b is the y-intercept.

The OA toxin concentration in shellfish tissue was calculated as follows:

$$C_t, \mu\text{g}/\text{kg} = [C_s (\text{nM}) \times \text{FD} \times \text{MW} (\text{g}/\text{mol}) \times V_e (\text{L})] / M_t (\text{g})$$

where C_t is the toxin concentration in tissue expressed as equivalents of OA, FD is the methanolic extract dilution factor, MW of OA = 805, V_e is the methanolic extract volume (0.025 L), and M_t is the tissue weight (5 g).

Samples with an OA concentration falling outside the working range (<0.5 nM or >2.8 nM) will be reported as <63 µg/kg (or <0.5 nM) or >352 µg/kg (or >2.8 nM), respectively.

Results were recorded by each participant in a tailor-made Excel spreadsheet with which the results were automatically calculated when the absorbance values were entered. All participants sent back an electronic copy of the reporting sheet for each day of analysis.

Table 6. Individual results (µg OA total equivalents/kg) reported from laboratories A to P for Materials A, D, E, F, G, K, L, and N on Days 1 and 2. Invalid or incorrect results are those in bold type.

| Lab | µg OA total equivalents/kg | | | | | | | | | | | | | | | |
|-----|----------------------------|-----------------|-----|------------|------------------|------------------|-----|---------------|-----------------|-----------------|------------------|------------------|-----|------------|-----|------------|
| | Material | | | | | | | | | | | | | | | |
| | A | | D | | E | | F | | G | | K | | L | | N | |
| | Day | | | | | | | | | | | | | | | |
| | 1 | 2 | 1 | 2 | 1 | 2 | 1 | 2 | 1 | 2 | 1 | 2 | 1 | 2 | 1 | 2 |
| A | <63 | <63 | 186 | 239 | 97 | 102 | <63 | <63 | <63 | <63 | 248 | 281 | 167 | 174 | 210 | 247 |
| B | <63 | <63 | 251 | 266 | 100 | 101 | <63 | <63 | <63 | <63 | 302 | 299 | 177 | 190 | 273 | 277 |
| C | <63 | <63 | 244 | 233 | 96 | 87 | <63 | <63 | <63 | <63 | 279 | 246 | 174 | 160 | 256 | 251 |
| D | <63 | <63 | 264 | 253 | 125 | 100 | <63 | <63 | <63 | <63 | 282 | 277 | 189 | 223 | 269 | 295 |
| E | <63 | <63 | 210 | 233 | 101 | 120 | <63 | <63 | <63 | <63 | 239 | 244 | 156 | 181 | 226 | 219 |
| F | <63 | <63 | 252 | 250 | 113 | 116 | <63 | <63 | <63 | <63 | 287 | 286 | 166 | 165 | 271 | 275 |
| G | <63 | <63 | 246 | 252 | 89 | 100 | <63 | <63 | <63 | <63 | 356 ^a | 269 ^a | 192 | 192 | 274 | 236 |
| H | <63 | <63 | 253 | 250 | 90 | 99 | <63 | <63 | <63 | <63 | 291 | 301 | 175 | 179 | 271 | 270 |
| I | <63 | <63 | 252 | 254 | 95 | 87 | <63 | <63 | <63 | <63 | 284 | 283 | 169 | 161 | 265 | 253 |
| J | 70 ^a | 98 ^a | 238 | 239 | 163 ^a | 102 ^a | <63 | <63 | 78 ^a | 67 ^a | 248 | 268 | 239 | 184 | 246 | 235 |
| K | <63 | <63 | 253 | 264 | 81 | 81 | <63 | <63 | <63 | <63 | 295 | 300 | 152 | 160 | 247 | 266 |
| L | — | <63 | — | 242 | — | 145 | — | <63 | — | — | — | 266 | — | 202 | — | 182 |
| M | <63 | <63 | 257 | 255 | 101 | 104 | <63 | <63 | <63 | <63 | 292 | 274 | 177 | 176 | 271 | 272 |
| N | <63 | <63 | 261 | 251 | 98 | 101 | <63 | <63 | <63 | <63 | 285 | 285 | 161 | 181 | 257 | 250 |
| O | <63 | <63 | 221 | 223 | 91 | 94 | <63 | <63 | <63 | <63 | 270 | 249 | 179 | 184 | 259 | 244 |
| P | <63 | <63 | 192 | 241 | 69 ^a | 153 ^a | <63 | <63 | <63 | <63 | 226 | 278 | 97 | 173 | 206 | 259 |

^a Outlier.

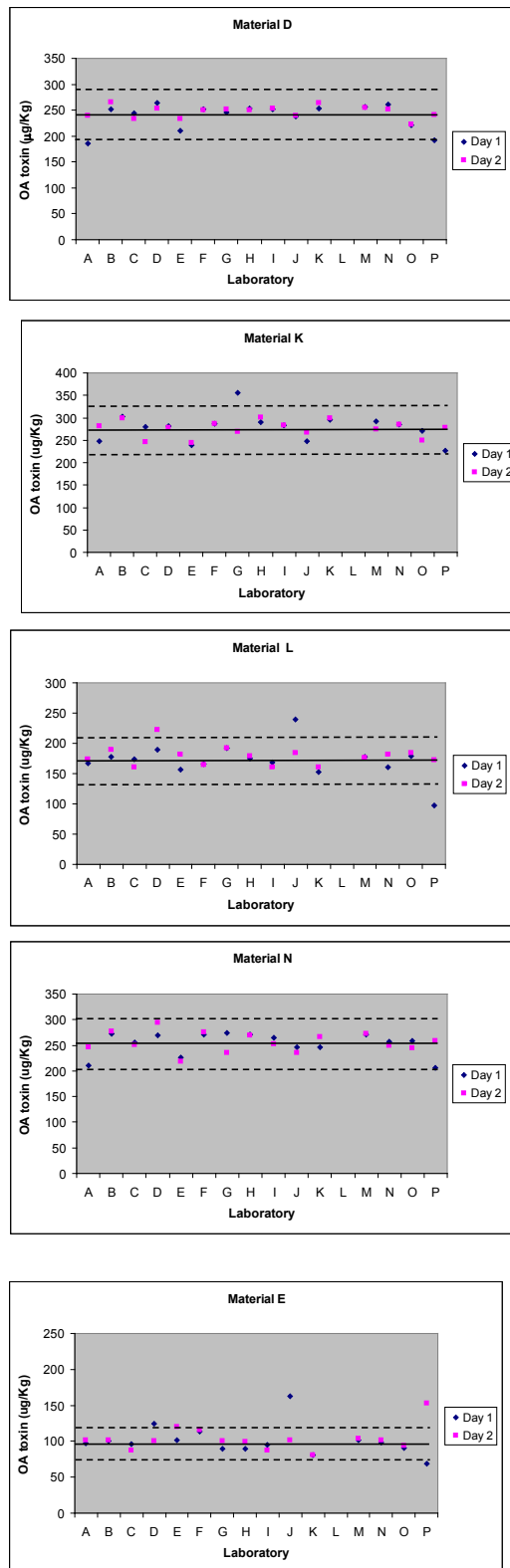


Figure 1. Individual results for each test material obtained per lab and per day of analysis (including outliers). The solid line shows the assigned mean value calculated in this study for each material. The dashed lines indicate the theoretical reproducibility SD determined for each material in this study (PRSD_R).

Statistics

Analysis of Valid Data and Outliers

Statistical data analysis was carried out following the approach described in the AOAC/IUPAC guidelines (17, 20). Submitted results were initially reviewed to remove invalid data. Results from assays with calibration curves with a $R^2 < 0.96$ and results outside the working range or showing deviations from the Standard Operating Procedure were considered invalid.

The valid data were first analyzed for possible outliers applying the Cochran and Grubbs tests. Then, precision parameters, HorRat values, and recovery were calculated.

The Cochran test was applied to remove laboratories showing significantly greater variability among replicate (within-laboratory) analyses than the other laboratories for a given material. A 1-tail test at a probability value of 2.5% was applied (17, 20).

The Grubbs test was used to remove results from laboratories with extreme averages (17, 20). This test was applied to the remaining values from the Cochran test. A single value test (two-tail, $P = 2.5\%$) was first applied, followed by a pair value test (two values at the highest end, two at the lowest end, and one at each end, at an overall $P = 2.5\%$).

Precision

To estimate the precision of the method, the within-laboratory repeatability and between-laboratory reproducibility were determined by calculating s_r (repeatability SD), s_R (reproducibility SD), RSDs (RSD_r and RSD_R), repeatability and reproducibility limits (r and R), and HorRat values. These parameters were calculated following the AOAC guidelines (20).

Recovery

For recovery calculations, the marginal recovery was calculated as follows:

$$\text{Recovery, \%} = 100 (C_f - C_u) / C_A,$$

where C_f is the amount found for the spiked concentration, C_u is the amount present originally for the unspiked concentration, and C_A is the amount added.

Results and Discussion

Test Material Results

The test materials were first analyzed by OkaTest and LC/MS/MS to determine the content and profile of OA toxins. Results obtained by both methods for samples A, F, and G showed concentration for OA toxins below their LOD (44 and 40 µg/kg, respectively). The BM was tested by LC/MS/MS (19) at EURLMB, and no peaks were detected for this group of toxins (LOD for this method is 15 µg/kg). Therefore, materials A, F, G, and BM were considered blank; therefore, no homogeneity or stability studies were carried out.

Analyses by LC/MS/MS were used to identify the toxin profile and to ensure that all toxins belonging to the OA group were present in the materials. Table 2 shows concentration

Table 7. Details of the test materials, number of results submitted, and results after removing outliers, together with performance values of precision (repeatability and reproducibility) obtained for the colorimetric OkaTest^a

| Test material | Matrix | Runs/lab | No. labs submitting results | No. labs after invalid/incorrect results | No. of labs after outliers ^b | Mean (μg total equivalent OA/kg) ^c | Repeatability ^c | | | Reproducibility ^c | | | |
|---------------|---------------------------------------|----------|-----------------------------|--|---|---|----------------------------|----------------|--------------------|---------------------------------|----------------|--------------------|--------------|
| | | | | | | | S_r | r | RSD _r % | μg total equiv.OA/kg | | | |
| | | | | | | | | | | S_R | R | RSD _R % | HorRat |
| A | Mussel <i>M. galloprovincialis</i> | 2 | 16 | 14 | — | <63 | — | — | — | — | — | — | — |
| D | Clam <i>V. pullastra</i> | 2 | 16 | 15 | 15 (0) | 242 | 14.7 | 41.2 | 6.1 | 19.4 | 54.4 | 8.0 | 0.4 |
| E | Mussel <i>M. galloprovincialis</i> | 2 | 16 | 15 | 13 (2) | 98.8 (102) | 7.32 (20.8) | 20.5 (58.4) | 7.4 (20.5) | 10.7 (19.6) | 30.0 (54.8) | 10.7 (19.2) | 0.5 (0.8) |
| F | Scallop <i>P. maximus</i> | 2 | 16 | 15 | — | <63 | — | — | — | — | — | — | — |
| G | Clam <i>V. decussatus</i> | 2 | 16 | 14 | — | <63 | — | — | — | — | — | — | — |
| K | Clam <i>V. rombooides</i> | 2 | 16 | 15 | 14 (1) | 275 (277) | 14.9 (21.4) | 41.8 (60.1) | 5.4 (7.7) | 21.0 (25.0) | 58.7 (70.1) | 7.6 (9.0) | 0.4 (0.5) |
| L | Cockle <i>C. edulis</i> | 2 | 16 | 15 | 15 (0) | 175 | 19.6 | 55.0 | 11.2 | 23.2 | 64.9 | 13.2 | 0.6 |
| N | Mussel <i>M. edulis</i> | 2 | 16 | 15 | 15 (0) | 255 | 15.6 | 43.7 | 6.1 | 20.7 | 58.1 | 8.1 | 0.4 |

^a S_r = Repeatability SD, S_R = reproducibility SD, RSD_r = repeatability RSD, RSD_R = reproducibility RSD, r = repeatability limit, R = reproducibility limit.

^b Number of laboratories remaining after removal of outliers (number of outliers).

^c Mean, repeatability, and reproducibility (values obtained including outliers).

in OA equivalents determined by OkaTest and toxins profile of the different materials used. All test materials were found to be stable for the duration of the study and with sufficient homogeneity (Tables 3 and 4).

Interlaboratory Study Results

All participants who received test materials reported results. The sample concentration was calculated by standard curves obtained by each laboratory every day of analysis. Fit parameters of each standard curve are shown Table 5. Although the slopes show differences depending on the laboratory and day, the calculated samples concentration was not affected. The data obtained by each laboratory per test material and day of analysis are shown in Table 6.

All individual values obtained per material, day and laboratory were also plotted. One graph per material is shown in Figure 1. The solid lines represent the assigned mean value obtained for each material in this study (Table 7). The area between the dashed lines demonstrates the range of deviation from the mean value based on the theoretical reproducibility SD (PRSD_R).

Two laboratories reported one of the assays with $R^2 < 0.96$; one (Laboratory A) repeated the analysis obtaining R^2 within the required criterion. Laboratory L, however, could not repeat the assay on time, and those results were considered invalid and removed for statistical analysis.

Materials A, F, and G were not statistically analyzed, as they were blank samples. However, Laboratory J reported values within the working range of the test for Materials A and G. These values are considered incorrect according to the AOAC

guidelines (20), as they are positive values found for a blank material. All the other laboratories in the study identified the blank materials below the working range of the test.

The valid data from the contaminated test materials (D, E, K, L, and N) were then analyzed for identification of outliers applying Cochran and Grubbs tests (20). Results from Laboratory L could not be included in the statistical analysis, as only one value per material was available.

The Cochran test showed Laboratory G for Material K and Laboratory P for Material E as outliers. This test was applied again after these outliers were removed. Laboratory J for Material E was also excluded in a second round. The Grubbs single and pair values tests were then applied; no further outliers were identified.

The mean values assigned for OA-toxins for the test materials were 98.8, 175.4, 242.8, 255.0, and 275.0 μg total equivalents OA/kg for Materials E, L, D, N, and K, respectively (Table 7).

Values obtained for repeatability SD (S_r) ranged from 7.3 $\mu\text{g}/\text{kg}$ for Material E to 19.6 $\mu\text{g}/\text{kg}$ for Material L, with repeatability RSDs (RSD_r) from 5.4% for Material K to 11.2% for Material L (Table 7). The reproducibility SD (S_R) calculated for the five test materials ranged from 10.7 to 23.2 $\mu\text{g}/\text{kg}$, with reproducibility RSD (RSD_R) values from 7.6 to 13.2% for Materials K and L, respectively (Table 7).

The HorRat values obtained were 0.4 for Materials D, K, and N, 0.5 for Material E, and 0.6 for Material L (Table 7), indicating a very good performance of the method. These values are just at the lower limit of the range considered as normally expected for a good reproducibility of a method ($0.5 < \text{HorRat} \leq 1.5$), according to the AOAC guidelines (20). HorRat values between 0.64 and 2.61 for OA-toxins group (21), 0.3 and 2.0 for paralytic

Table 8. Results from the recovery experiment carried out during Day 2 of the interlaboratory study

| Lab code | µg OA total eq./kg | | | Recovery, % |
|------------------|--------------------|---------------|----------------------|-------------|
| | BM ^a | Spiked concn. | BM + OA ^b | |
| A | — | 161 | 172 | 107.1 |
| B | — | 161 | 162 | 100.7 |
| C | — | 161 | 155 | 96.3 |
| D | — | 161 | 115 | 71.6 |
| E | — | 161 | 124 | 77.3 |
| F | — | 161 | 138 | 85.5 |
| G | — | 161 | 162 | 100.7 |
| H | — | 161 | 131 | 81.1 |
| I | — | 161 | 152 | 94.4 |
| J | — | 161 | 197 | 122.3 |
| K | — | 161 | 152 | 94.4 |
| L | — | 161 | 196 | 121.6 |
| M | — | 161 | 153 | 95.0 |
| N | — | 161 | 174 | 108.3 |
| O | — | 161 | 155 | 96.3 |
| P | — | 161 | 185 | 114.7 |
| Mean recovery, % | | | | 98.0 |
| SD | | | | 14.2 |
| RSD, % | | | | 14.5 |

^a BM = Blank material. No OA toxins were detected; therefore, a concentration of zero was considered for calculation purposes.

^b BM + OA = Concentration of the samples spiked with 161 µg/kg.

shellfish toxins (22) and 1.1 to 2.4 for domoic acid (23) were previously described for other methods.

The statistical analysis was also carried out including outliers (Table 7). Although there were some differences when including outlier values, repeatability and reproducibility remained satisfactory and within the expected values for this type of interlaboratory study.

Although the main objective of the validation study was to determine the repeatability and between-laboratory reproducibility of the OkaTest kit, a recovery assay was also carried out. A scallop blank sample (BM) was spiked with OA by each laboratory, and the recovery of OkaTest calculated. Recovery values from all participants ranged from 71.6 to 122.3%. The mean and RSD were 98.0 and 14.5%, respectively (Table 8). These recoveries met the criteria set in the AOAC Guidelines for Single Laboratory Validation of Chemical Methods for Dietary Supplements and Botanicals (24).

Comments from Participants

Most participants reported that the SOP for the method provided all the information they needed to perform the assay and that they did not have difficulties understanding any part of it. Some comments were made about the phosphatase preparation. Those led to the conclusion that the use of a nonorbital shaker does not always guarantee full dissolution of this reagent. Manual mixing, longer preparation, and a final visual check of the solution should be included in the SOP. Other

minor comments were made, and were answered or resolved by the study director.

Conclusions

The precision and recovery values determined in this study for OkaTest can be considered satisfactory for this type of methodology and the concentration range required. The colorimetric PPIA, OkaTest, could be used as an assay complementary to the reference method for determination of the OA toxins group in molluscs according to the Commission Regulations (EC) No. 2074/2005 and No. 15/2011. Additional methods have to be implemented in a laboratory to analyze all regulated lipophilic marine biotoxins.

Acknowledgments

We thank the following collaborators for their participation, time, effort, and comments towards this study:

Dolores Calvo, ZEU-INMUNOTEC, Spain;

Monica Campàs, IRTA, Spain;

Guntis Cepurnieks, Institute of Food Safety, Animal Health and Environment, Latvia;

David Clarke, Marine Environment and Food Safety Services, Ireland;

Jorge Correa, INTECMAR, Spain;

Lorena A. Delgado, Public Health Institute, Chile;

Alejandra Goya A, SENASA, Argentina;

Katrin Kapp, BfR (Federal Institute for Risk Assessment), Germany;

Panagiota Katikou, Ministry of Rural Development and Food Centre of Veterinary Institutions of Thessaloniki Institute of Food Hygiene, Greece;

Mirslaw Michalski and Kataryna Grazil, National Veterinary Research Institute, Poland;

María José Chapela, ANFACO-CECOPESCA, Spain;

Sonia Piñero, European Reference Laboratory for Marine Toxins, Spain;

Vlad Serafim, The Institute for Diagnosis and Animal Health, Romania;

Ulrich Schwank, Bavaria's policies on health and consumer protection (LGL), Germany;

Andrew Turner and Clothilde Brunet, CEFAS, UK; and

Paulo Vale and Susana Rodrigues, IPIMAR, Portugal.

Note: Collaborative efforts among the European Reference Laboratory for Marine Toxins (EURLMB), NRLs, and ZEU-INMUNOTEC does not amount to an endorsement of the firm's products.

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