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Ultra-trace determination of domoic acid in the Ebro Delta estuary by SPE-HILIC-HRMS†

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Marine phytoplankton, such as diatoms, are responsible for a considerable part of carbon fixation and form the basis of the marine food web. However, different factors, such as eutrophication, can affect their population growth and induce the production of toxins. During the last few years, the presence of domoic acid (DA) has increased worldwide in coastal areas. Its toxic properties and capacity to bioaccumulate through the food chain have reinforced the need for developing monitoring strategies to ensure the safety and quality of marine resources. In this work, a highly sensitive analytical method has been developed to be used as an early detection and quantification tool for DA in seawater. The method is based on a sample pre-treatment step by solid-phase extraction (SPE) and an instrumental analysis by hydrophilic interaction liquid chromatography coupled with high-resolution mass spectrometry (HILIC-HRMS). The method has been validated, showing good results in terms of sensitivity and repeatability. Limits of detection and quantification were 0.25 and 0.75 ng L⁻¹ respectively, and acceptable recovery was in the range of 48–69%. Then, the method was applied to study a real scenario in the Ebro Delta where mariculture activities are relevant for the economy of the area. A total of 34 samples were collected in different campaigns during three seasons covering the two main delta bays (Alfacs and Fangar) and four lagoons (La Tancada, Illa de Buda, l'Encanyissada and Canal Vell). 65% of the samples had concentrations of DA at the ng L⁻¹ level, ranging from 0.90 to 69.6 ng L⁻¹. The highest levels were found in samples from Alfacs bay and the summer campaign. In the case of the lagoons, DA was detected only in La Tancada at 8.78 ng L⁻¹ in the winter campaign. This study proves the prevalence of DA in highly eutrophic areas, during all seasons, often at trace level concentrations.

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1. Introduction

The frequency of harmful algal blooms (HABs) has globally increased due to water eutrophication and global warming. This results in negative consequences for ecosystems because of the release of toxins by such harmful algae. Domoic acid (DA) is a potent neurotoxin produced by diatoms of the genus *Pseudo-nitzschia* (present in all coastal areas). DA has bioaccumulation capacity¹ and is the cause of the Amnesic Shellfish Poisoning (ASP) syndrome, known since the first documented intoxication in 1987, through blue mussel consumption in Canada.² Since this event, many programs have been established to understand the toxin-release mechanism from algae and to predict and prevent contamination incidents. However, the subject still remains poorly understood; in the case of diatoms, not all species of this genus can produce DA, and the toxicity within the

same species can vary even within different strains of the same toxicogenic species.³ Moreover, the presence of other microbial communities can induce the release of DA as a defensive mechanism.⁴ In addition, the production of DA depends on miscellaneous factors; the availability of nutrients, temperature and growth phase of the organisms are also relevant parameters. Due to the difficulties in controlling all these abiotic and biotic factors, the assessment of DA has to be done with more selective methodologies to have reliable results.

Liquid chromatography coupled with mass spectrometry (LC-MS) has been proved to be a sensitive and selective technique in the analysis of marine toxins. The European Commission⁵ established the use of LC-MS techniques as reference methods for the determination of toxins in shellfish. These techniques, in combination with appropriate sample extraction and clean-up procedures, permit the determination of DA in seafood and other matrices such as phytoplankton and seawater.^{6–10} Solid-phase extraction (SPE) is an exceptional technique that provides the isolation of DA from seawater and the elimination of matrix interference, especially the high content of dissolved salts. Moreover, the possibility of concentrating the sample by several orders of magnitude greatly benefits the increase of the overall sensitivity of the method.

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In this study, the development of a sensitive and selective analytical method to determine and quantify DA in seawater is presented. The analytical approach is based on solid-phase extraction (SPE) followed by hydrophilic interaction liquid chromatography (HILIC) coupled with high resolution mass spectrometry (HRMS) using electrospray ionisation (ESI). The analytical approach has been in-house validated and applied to assess DA in the Ebro Delta embayment, which is the major mollusc harvesting area (mussels, oysters and clams) of the Catalan coast. The assurance of the quality and safety of these products is mandatory. Evidence of *Pseudo-nitzschia* spp. has been widely reported in the Ebro Delta bays during the last few years although, to date, the occurrence of *Pseudo-nitzschia* has never been directly related to the presence of DA.^{11–14} The developed method has allowed the direct accurate measurement of DA in this area and, consequently, the possibility of studying its spatial and temporal distribution. Three sample campaigns have been conducted in two bays, Fangar and Alfacs, and four lagoons, La Tancada, Illa de Buda, l'Encanyissada and Canal Vell, during three different seasons. Then, the occurrence of DA has been discussed.

2. Materials and methods

2.1 Chemicals and reagents

Domoic acid and C5'-*epi*-domoic acid (a diastereomer of domoic acid) (DA, $44.5 \pm 3.1 \mu\text{g g}^{-1}$, $\geq 99\%$ purity and reference CRM-03-DA) were purchased from the Cifga laboratory (Lugo, Spain). Auxiliary reagents such as ammonium formate, HPLC-grade methanol (MeOH), ultra-pure water, acetonitrile (ACN) and formic acid (FA) were supplied by Merck (Darmstadt, Germany). Standard solutions of DA were prepared in MeOH.

2.2 Sampling

Three sampling campaigns were carried out in the Ebro Delta wetland (north-east Spain, Mediterranean Sea) during October 2015, February 2016 and June 2016. Surface seawater was collected from two semi-closed bays, Fangar and Alfacs bays, and from four lagoons, La Tancada, Illa de Buda, l'Encanyissada and Canal Vell. Temperature (°C), salinity and pH (Table S2†) were measured with a multiparameter probe Professional Plus YSI 2030 from Yellow Springs Instruments (Ohio, USA) on each sampling date. Sampling point details are shown in Fig. S1,† and detailed information is provided in Table S1 of the ESI.†

Seawater samples were collected at a 1 m depth using a Niskin bottle and were stored in amber glass bottles at $-20\text{ }^\circ\text{C}$ until the analyses.

2.3 Sample treatment

The optimized sample treatment (see optimization details in Results section 3.1) was as follows: 500 mL of seawater was extracted by solid-phase extraction (SPE) employing hydrophilic-lipophilic balance cartridges (OASIS HLB, 200 mg) from Waters (Cerdanyola del Vallès, Spain). Cartridge conditioning and equilibration were accomplished with 6 mL of MeOH and 6 mL of ultrapure water with 0.1% FA. Then, the samples were acidified with 0.1% FA, and cartridge loading was performed at

a flow rate of 1 mL min⁻¹. After loading, the cartridges were cleaned with 6 mL of ultra-pure water acidified with 0.1% FA. Finally, MeOH was evaporated from the extracts under gentle N₂ flow in a TurboVap from Biotage (Uppsala, Sweden). Extracts were reconstituted with 250 μL of acetonitrile–ammonium formate/FA buffer (9 : 1), which is the mobile phase composition at the beginning of the chromatographic run.

2.4 Instrumental analysis by HILIC-HRMS

The optimal instrumental conditions (see optimization details in Results section 3.1) were as follows: chromatographic separation was achieved by ultra-high performance liquid chromatography (UPLC) using the Acquity Waters (Harwich, Massachusetts, USA) system equipped with a HILIC (HILIC LUNA® 150 mm \times 2 mm, 3 μm , 200 Å) column from Phenomenex (Torrance, California, USA). The mobile phase was composed of (A) 90% of acetonitrile and 10% of 50 mM ammonium formate at pH 3.2 and (B) 50% acetonitrile, 40% water and 10% of 50 mM ammonium formate at pH 3. The elution gradient was programmed as follows: 0 min (100% A) – 2 min (100% A) – 8 min (100% B) – 9 min (100% A) – 10 min (100% A). An extra 5 minutes was added between each analysis in order to stabilize the pressure in the column. The flow rate was established to be 0.3 mL min⁻¹. The samples were kept at 10 °C in an auto-sampler, and the injection volume was 20 μL .

Liquid chromatography was coupled with mass spectrometry through a heated electrospray ionization source (HESI-II) from Thermo Fisher Scientific (San Jose, California, USA) operating in positive ion mode. The optimal parameters of the source are as follows: sheath flow gas of 60 a.u., auxiliary gas of 15 a.u., and sweep gas of 2 a.u. Heater and capillary temperatures were set at 350 and 320 °C, respectively. The S-lens RF level was 60% and the spray voltage was 3.5 kV.

Mass spectrometry was performed with a Thermo Scientific Q-Exactive™ mass spectrometer from Thermo Fisher Scientific (San Jose, California, USA) with a hybrid quadrupole-Orbitrap analyser. Data were acquired in full scan mode in the range of 50–800 *m/z* at a resolution of 70 000 [full width at half maximum (FWHM)], and contemporary in the MS/MS spectrum at a resolution of 35 000 FWHM in parallel reaction monitoring (PRM) mode. The normalized collision energy (NCE) was set at 10 a percentage of 30 for the molecular ion $[\text{M} + \text{H}]^+ = 312.1$ of the target toxin.

Data analysis was carried out using Thermo Xcalibur 3.1.667 software from Thermo Fisher Scientific. Identification of DA was achieved with the retention time and exact mass (never exceeding 2 ppm of error between the accuracy mass and the calculated exact mass) of the precursor ion in the full scan mode and the fragment ions. Using the same criteria, quantification was performed by integrating the area of the most intense product ions (peaks with at least seven scan points) and the confirmation was performed with two other fragment ions following the abundance ratio of each one.

2.5 Validation of the method

Method validation was accomplished with the evaluation of the selectivity, linearity, precision, sensibility, accuracy, limits of



detection (LODs) and limits of quantification (LOQs) using standard solutions of DA and fortified seawater.

2.5.1. Selectivity. For identification purposes, the exact mass of the precursor ion in the full-scan mode, the products ions in the fragmentation pattern, and the retention time of DA in the standard and in the spiked seawater blank samples were compared at a tolerance of $\pm 2.5\%$. Moreover, in accordance with the 2002/657/EC decision, the relative ion intensities (each product ion area signal *versus* the base product ion area signal) of the spiked samples were compared with the relative ion intensities of the DA standard solutions, at the same concentration levels used for the construction of the calibration curve.

2.5.2. Linearity. The linearity of the measurements in the instrumentation was estimated in the concentration range of 0.5 to 100 $\mu\text{g L}^{-1}$ of DA. Pearson's correlation coefficient (R^2) and the slope of the calibration curve in solvent and matrix were determined to evaluate the sensitivity.

2.5.3. Limits of detection and quantification. The instrumental LOD (iLOD) was experimentally determined by gradual dilutions of the standard solution of domoic acid, starting at 40 $\mu\text{g L}^{-1}$ concentration.

Because of the lack of noticeable instrumental noise, the lowest concentration at which the signal can be measured (fulfilling the required points per peak and selectivity criteria) was considered the iLOD. Consequently, the instrumental LOQ (iLOQ) was calculated, which was 10/3 times the iLOD.

2.5.4. Recovery. Since no certified reference material was available, recovery experiments were carried out using fortified samples in triplicate at two different levels 20 and 50 ng L^{-1} . In all batches of samples experimental blanks were analysed.

The method LOD (mLOD) for each extraction procedure was determined by the analysis of the standard solution in the matrix resulting from the particulate and the filtrate extracts. LODs were established to be the lowest concentration measurable of the matrix-matched calibration curve. The method LOQs (mLOQs) were estimated to be 10/3 times the mLODs.

2.5.5. Intra-day and inter-day precision. The repeatability of the instrument was evaluated with the measurement of a DA solution on the same day ($n = 6$) and consecutive days ($n = 3$), respectively, under the same instrumental conditions.

2.5.6. Accuracy. It was evaluated with the calculation of the DA recovery during the pre-treatment process.

For this, fortified blanks of seawater were subjected to the pre-treatment process. Values obtained were compared with those from the extracts subjected to the same process but fortified for the LC-MS analysis.

The matrix effect was evaluated to determine a possible signal enhancement or ion suppression during the ionization process at source by interferent substances present in seawater. To assess the effects of the matrix, fortified solutions of DA in seawater extracts and pure solvent were compared, and the percentage of effect was calculated according to the following expression:

$\% \text{ Matrix effects} = ([\text{area}]_{\text{seawater}}/[\text{area}]_{\text{solvent}}) \times 100$, where $[\text{area}]_{\text{seawater}}$ is the integrated area of DA in the seawater extracts and $[\text{area}]_{\text{solvent}}$ is the corresponding area in pure solvent.

3. Results and discussion

3.1 Analytical method optimisation

The high polarity of DA increases the difficulty in isolating it from the aqueous phase during extraction. The molecule possesses three carboxylic acid groups and one amino group, with pK_a s ranging from 1.85, 4.47, and 4.75 to 10.6.¹⁵ To optimise extraction from seawater different stationary phases and procedures have been studied: (i) the hydrophilic-lipophilic balance using OASIS HLB (500 mg, 6 cm³) cartridges from Waters; (ii) carbon adsorption using ENVI CARB graphitized charcoal cartridges (500 mg, 6 cm³), from Merck; and (iii) ionic exchange, in both cation and anion exchange modalities, with Isolute WCX (500 mg, 6 cm³) and Isolute SAX (500 mg, 6 cm³) cartridges, respectively, both from Biotage (Uppsala, Sweden).

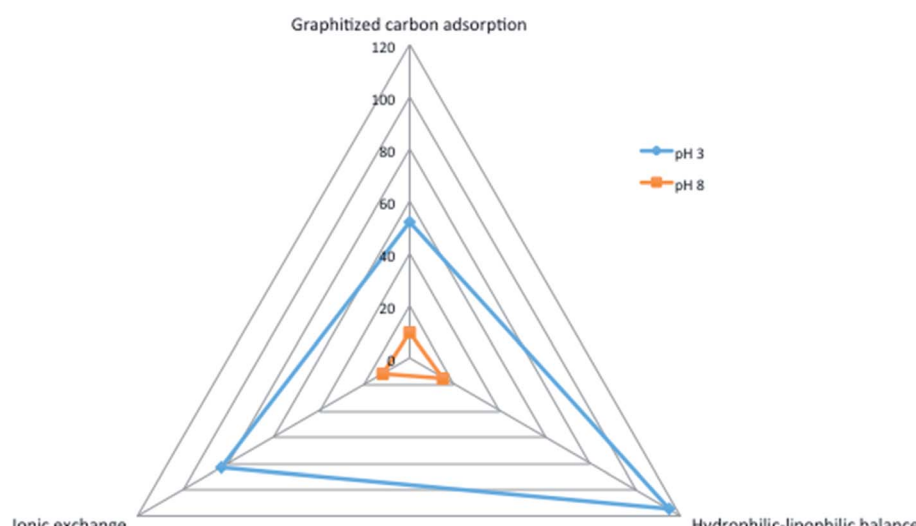


Fig. 1 Values of recovery obtained in the different SPE treatments.



All cartridges were conditioned with 6 mL of methanol and equilibrated with 6 mL of water adjusted at the same pH of the sample. Loading and elution conditions were optimized for each cartridge independently:

For HLB, 100 mL spiked seawater was loaded at pH 3 or pH 8. A washing step after loading was performed with 6 mL of water at pH 3 or pH 8, and elution was carried out with 20 mL of methanol.

For ENVI CARB, 100 mL spiked seawater at pH 3 and pH 8 was loaded into the cartridge. A washing step with 20 mL of water at pH 3 or 8 was added. Elution was performed with 25 mL of a mixture of methanol/acetonitrile/acetone (1 : 1 : 1).

For SAX and WCX, 25 mL of spiked seawater was loaded into the cartridges conditioned at pH 8 for anion exchange and at pH 3 for cation exchange. Elution was carried out with 25 mL of methanol.

All extracts from the elution step were concentrated and reconstituted as explained in Section 2.3 for the subsequent analysis.

Recovery values for different SPE treatments are presented in Fig. 1. As can be seen, the best recovery rates were obtained for the HLB stationary phase when the spiked samples were conditioned at pH 3. The graphitized carbon showed as well acceptable values with the samples conditioned at acidic pH but repeatability was higher with HLB. On the other hand, the

elution of DA graphitized carbon required higher volumes of solvent than the HLB. Regarding the cation/anion exchange strategy, the cation exchange allowed better recovery of DA than the anion one, but lower than that with HLB and samples at acidic pH. Considering the results under these conditions HLB cartridges were selected for further optimisation.

The HLB phase was tested by loading different volumes of samples: 100 mL, 250 mL, 500 mL and 1.0 L. A sample volume of 500 mL was selected as the maximum volume without the critical loss of the compounds and the clogging of the cartridge, reaching a value of $69.24 \pm 1.14\%$ and $48.09 \pm 17\%$ at concentration levels of 20 ng L^{-1} and 50 ng L^{-1} , respectively. Although the recoveries were a bit lower when loading higher volumes of samples, a significant improvement in the mLOQ was obtained with 500 mL of seawater instead of 100 mL because of the high pre-concentration of the sample; thus 500 mL was selected as the optimal volume. For quantification purposes, recovery correction was applied for each sample.

For the chromatographic separation, two different types of chromatography techniques were compared: reversed-phase liquid chromatography (RPLC) using a C18 Synergy (50 mm \times 2 mm, 5 μm , 80 \AA) analytical column, and hydrophilic interaction liquid chromatography (HILIC) using a HILIC LUNA[®] (150 mm \times 2 mm, 3 μm , 200 \AA) analytical column, both from

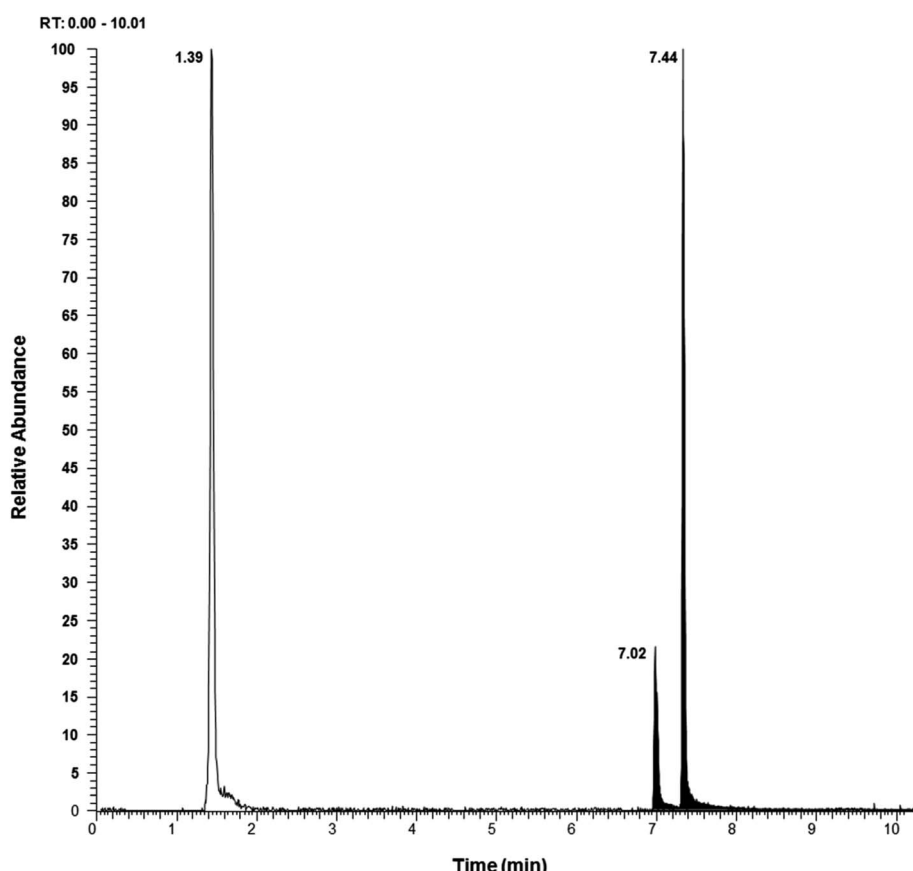


Fig. 2 Extracted ion chromatograms of DA from the two different chromatography techniques: white color corresponds to the reverse-phase column (C₁₈) while the black color corresponds to the hydrophilic interaction column (HILIC) which separates DA (tr 7.44) from its isomer epi-DA (tr 7.02).



Phenomenex. For HILIC, the mobile phase was composed of (A) acetonitrile and (B) water with FA and ammonium formate (see the optimised elution gradient in Section 2.4). RPLC was performed using a mobile phase composed of (A) 90% acetonitrile 0.1% FA and (B) 10% water 0.1% FA with an isocratic gradient. The working flow rate was established to be 0.3 mL min⁻¹ and the total analysis run was for 10 min.

The extracted ion chromatograms of DA using RPLC and HILIC are shown in Fig. 2. As can be seen, a good peak shape was obtained in both cases, but using RPLC the retention time was 1.40 min. Such a short retention time is favourable in terms of rapid screening, but this retention time is too close to the dead time. While RPLC chromatographic approaches have been used for the analysis of DA in shellfish and plankton in combination with some other lipophilic toxins,^{16,17} poor retention of DA may cause its coelution with multiple matrix components that may elute at the beginning of the chromatogram in complex extracts, potentially interfering with its detection and causing matrix effect. Because of this, HILIC has been selected as the optimal column since it increases DA retention and allows the separation of DA from its isomer *epi*-

DA with a retention time of 7.40 min. This type of chromatography gives the possibility of combining the analysis of DA with other hydrophilic marine toxins, allowing a high throughput analysis of polar compounds in seawater. In mussel analysis, HILIC columns have already been employed in the determination of DA along with PSP toxins with a suitable resolution.^{6,18}

Two mass spectrometer analysers were compared for the determination of DA: a triple quadrupole working in the acquisition mode of Selected Reaction Monitoring (SRM), and an Orbitrap operating in PRM mode. A standard solution of DA in MeOH was directly infused into the mass spectrometers by soft ionization with an electrospray ionization (ESI) source. Optimal conditions were obtained for the ionization of DA and its fragmentation in the two instruments. Fragmentation patterns are shown in Table 1.

Sensitivity, resolution and selectivity were compared for both analysers, and Orbitrap was selected for this study. In terms of sensitivity, there is no significant difference between both mass spectrometer analysers, but working in PRM mode provides extra selectivity than working in SRM, thanks to accurate mass measurements. Also, the PRM mode can be combined with

Table 1 Fragmentation pattern of DA at the optimized collision energies for two different analysers, Orbitrap and triple quadrupole. NCE* corresponds to the normalized collision energy stepped at the value of 10. The mass tolerance error in ppm is shown in parentheses

| | Chemical formula | Precursor ion | <i>m/z</i> | Collision energy 1 | Product ion 1 | <i>m/z</i> | Collision energy 2 | Product ion 2 | <i>m/z</i> |
|----------|--------------------|---------------|-----------------------|--------------------|------------------------|---------------------|--------------------|------------------------|-----------------------|
| Orbitrap | $C_{15}H_{21}NO_6$ | $[M + H]^+$ | 312.1442 (1.5 ppm) | 30% NCE* | $[C_{14}H_{20}O_4N]^+$ | 266.1382 (1 ppm) | 30% NCE* | $[C_{14}H_{18}O_3N]^+$ | 248.1275 (1.5 ppm) |
| | | | | | | | | | |
| QqQ | | | 312 | 16.6 V | | 266 | 17.6 V | | 248 |

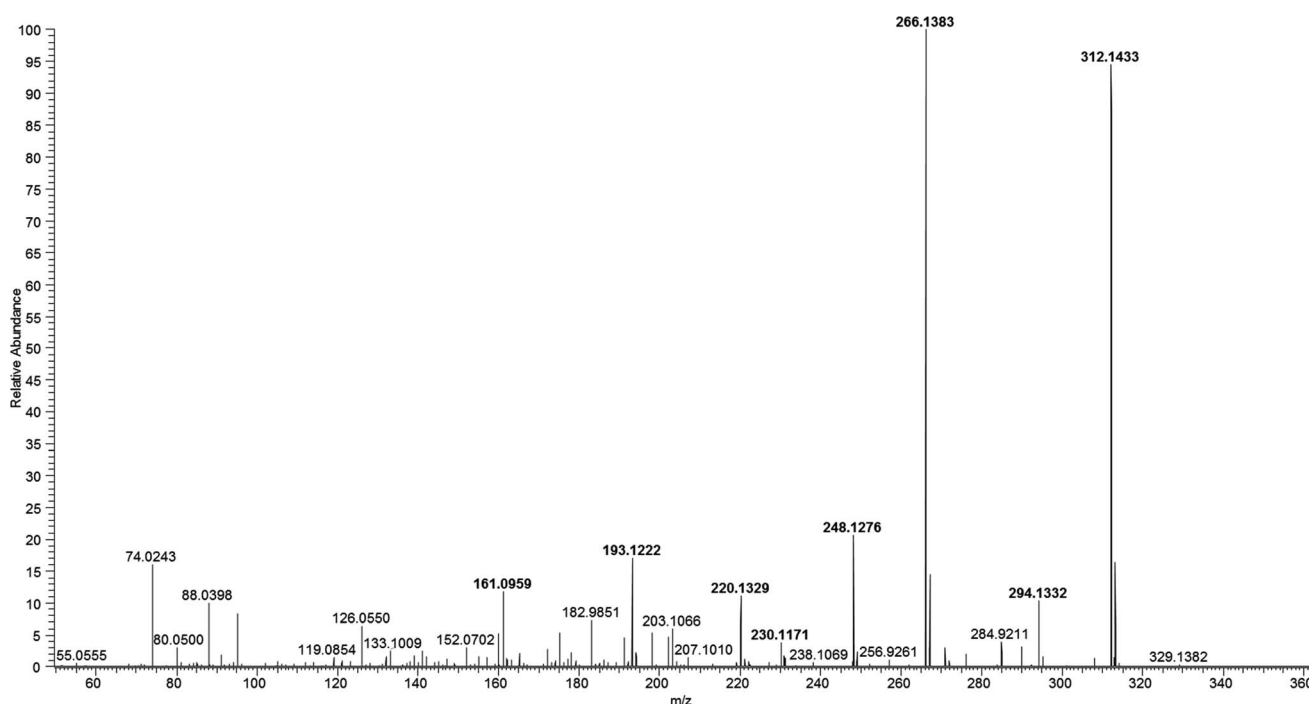


Fig. 3 High resolution mass spectrum of DA obtained with Q Exactive.

parallel full-MS acquisition, which allows the possibility of performing retrospective analysis for the detection of suspected compounds and even non-target analysis at high resolution for the identification of other pollutants. The high-resolution mass spectrum of DA obtained at 30% of NCE is shown in Fig. 3.

In Table 2, the quality parameters of this method are summarized. Good performance of the method has been achieved. The most notable quality is the high sensitivity with the mLOD reaching 0.25 ng L^{-1} for DA. To our knowledge, these are the lowest values reported in the literature. Other sensitive methods based on LC-MS presented mLOD values ranging from 750 ng L^{-1} ,⁸ 30 ng L^{-1} (ref. 7) to the lowest value of 0.3 ng L^{-1} .¹⁹

3.2 Levels of DA in coastal waters

In Table 3, the DA concentrations detected in 34 seawater samples from Ebro's Delta are summarised. DA was present in 65% of the samples at the ng L^{-1} level. The concentration of DA ranged from 0.90 to 69.6 ng L^{-1} with a mean value of 17.3 ng L^{-1} and a median of 5.59 ng L^{-1} . The presence of DA was prevalent in Alfacs and Fangar bays during the seasons analysed while in the lagoons only La Tancada showed a positive result with a concentration of 8.78 ng L^{-1} . The La Tancada lagoon was

similar to the bays in terms of salinity. *Pseudo-nitzschia* spp. are marine and preferably grow in a saline environment.²⁰ Moreover, the production of DA can be favoured at high salinity.²¹ Then, this influence from seawater could explain why DA is present in this lagoon and not in the rest.

The highest concentrations were found in Alfacs bay in the summer season, with a maximum value of 69.6 ng L^{-1} , followed by 67.2 ng L^{-1} , which was detected at the same sampling site during wintertime. This last value was unexpected, considering that during the coldest months the upwelling phenomenon takes place, driving the nutrients from the surface to deeper waters, and the temperatures are lower, having a negative influence on the phytoplankton growth. But these bay areas constantly receive agricultural discharges of freshwater with a high content of nutrients that can disrupt the annual variation of nutrient availability. In other studies, DA has also been detected in shellfish during winter months along the Catalan coast. Giménez-Papiol *et al.* reported concentrations of DA in February and April exceeding the EFSA suggested regulation levels ($4.5 \text{ mg DA per kg shellfish}$) and even the maximum permitted in the current legislation ($20 \text{ mg DA per kg shellfish}$).¹³ These values are in the same range as other studies that determined DA in estuarine areas in the Mediterranean Sea.

Table 2 Analytical parameters for the method validation

| Instrumental parameters | | Intra-day precision | Inter-day precision | |
|----------------------------------|--|---|--|-------------------|
| iLOD (pg on column) | Linearity range (ng L^{-1}), R^2 | $10 \mu\text{g L}^{-1}$ (RSD%, $n = 6$) | $10 \mu\text{g L}^{-1}$ (RSD%, $n = 3$) | |
| 1 | 50–50 000 (0.999) | 5.53 | 2.91 | |
| Method parameters | | | | |
| mLOD/mLOQ (ng L^{-1}) | Linearity range (ng L^{-1}) and linearity (R^2) | Recovery (%), $n = 3$ | | Matrix effect (%) |
| 0.25/0.75 | 2–50 (0.995) | 69.24 ± 1.14 | 48.09 ± 7 | 73 |

Table 3 Concentration of DA in ng L^{-1} in Ebro's Delta bays and lagoons

| | Coordinates | Sampling campaign | | |
|----------------|--------------------------|-------------------|------------|------------|
| | | 1 (Oct-15) | 2 (Feb-16) | 3 (Jun-16) |
| Alfacs | 40.610, 0.724 (beach) | 11.6 | 67.2 | 14.6 |
| | 40.601, 0.610 (shore) | 18.4 | 14.3 | 69.6 |
| | 40.584, 0.579 (open sea) | 15.0 | 21.6 | 41.7 |
| | 40.605, 0.609 (open sea) | No sample | 8.08 | 24.7 |
| Fangar | 40.800, 0.700 (beach) | 2.20 | 8.11 | 5.1 |
| | 40.798, 0.713 (shore) | 4.16 | 2.29 | 11.2 |
| | 40.815, 0.743 (open sea) | 6.09 | 0.9 | 10.1 |
| | 40.817, 0.751 (open sea) | No sample | <LOQ | 13.9 |
| La Tancada | 40.645, 0.742 (lagoon) | 8.78 | <LOQ | <LOD |
| Illa de Buda | 40.703, 0.841 (lagoon) | <LOD | <LOD | <LOD |
| L'Encanyissada | 40.657, 0.673 (lagoon) | <LOD | <LOD | <LOD |
| Canal Vell | 40.745, 0.788 (lagoon) | <LOD | <LOD | <LOD |



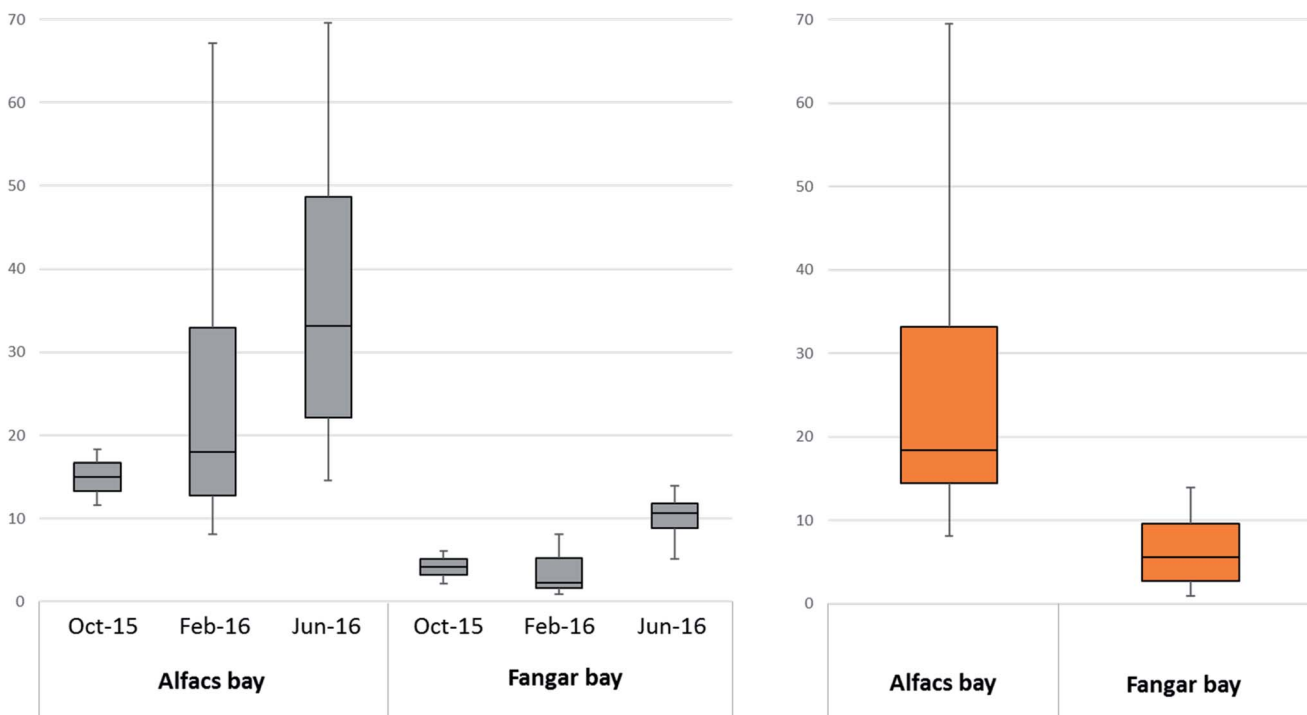


Fig. 4 Box plot graphs of the DA concentration: grey color for the two bays during the three sample campaigns; October 2015, February and June 2016; orange for the total concentration in three campaigns in the two different bays.

For example, Barbaro E. *et al.* registered concentrations ranging from 1.5 to 16.2 ng L^{-1} in the Venice lagoon that is strongly influenced by the seawater from the Adriatic Sea.¹⁹ More focused on the studied area of Ebro's Delta, Busch *et al.* reported concentrations of DA between 3.2 and 18.8 ng L^{-1} in Fangar bay but not in Alfacs.¹²

Comparing the results of the two bays, Alfacs had, in general, higher levels of DA than Fangar. The differences between the total concentration of DA along with the three sampling campaigns in the two bays are displayed in Fig. 4. The location of the bays could be the factor that explains this difference since both have a semi-closed shape but water residence time and stratification differ significantly in both cases: Fangar bay is in the northern part of the delta and influenced by the littoral current coming from the north to the south (Northern current²²), whereas Alfacs is a bay located in the southern part where marine recirculation of water is slower because of the physical barrier that the delta imposes,²³ and freshwater inputs have longer residence times. Water stratification,¹⁴ the occurrence of inorganic components (*i.e.* silicates and Cu^{2+})²⁴ and organic nutrients²⁵ are known to be correlated with the abundance of *Pseudo-nitzschia* spp. (or other algae responsible for their emission, as hypothesized by Busch *et al.*), and all these parameters in the Ebro Delta embayments are substantially affected by the dynamics of irrigation channels. Then, growth of algal phytoplankton could be more favoured in Alfacs, and consequently, the release of DA.

Regarding seasonal variability, a slight tendency can be observed from the lowest DA concentrations in autumn months to the highest in summer months. The differences in the DA

concentrations in each bay during different months of sampling are shown in Fig. 4. This tendency can be observed when considering the median values of all the concentrations detected in each bay across different months. More in detail, this tendency seems to be clearer for the samples taken in the open sea than for those samples from a beach and shore, where the concentration of DA did not seem to follow a clear temporary pattern. This tendency can be as well observed in the box-and-whisker plot for the tree sampling campaign presented in Fig. S2.† However, the *P*-value was 0.1196 that can be considered insignificant. Therefore, the variation among campaign medians is not significantly greater than expected by chance. The Kruskal-Wallis test was applied because the distribution of the results did not pass the normality test for all sampling campaigns.

As commented previously, these semi-closed bays are constantly influenced by variable freshwater discharges from agricultural activities, leading to no regular values of salinity, nutrients and turbidity in the nearshore.

4. Conclusions

A highly sensitive analytical method for the detection and quantification of DA in seawater has been developed and applied to the analysis of water samples from the marine bays of the Ebro Delta.

For the method development, several stationary phases were tested and Oasis HLB cartridges were selected. Mean recovery values ranging from 48 to 69% were obtained when treating 500 mL samples acidified with FA. A low matrix effect was

observed, indicating the elimination of interference and salt from seawater. Anyway, this effect of ion suppression was assessed by quantification with a matrix-matched calibration curve. High sensitivity has been achieved, thanks to the 2000-fold concentration of the samples, allowing a mLOD and mLOQ of 0.25 and 0.75 ng L⁻¹, respectively.

The high polarity of DA allows the option of using both RPLC and HILIC with suitable resolution and peak shapes. However, HILIC has been selected in this study to determine DA because of its higher retention factor and for a possible combination in the joint analysis of other hydrophilic compounds present in seawater. Moreover, the separation of DA and its isomer *epi*-DA was possible.

The Orbitrap and triple-quadrupole mass spectrometers presented good sensitivity and selectivity. However, the Orbitrap-MS was selected for the method, not only for its highest resolution but also for its capability to acquire comprehensive sample screenings in the full-MS mode at high resolution and the possibility of conducting retrospective analyses and carrying out non-target screening of other MBTs and contaminants.

Final validation of the method showed excellent performance in terms of sensitivity, linearity, repeatability, selectivity and recovery. Application of the method to real samples was successful.

DA was present in 65% of the samples analysed with a mean value of 17.3 ng L⁻¹. The concentration of DA ranged from 0.90 to 69.6 ng L⁻¹ and was higher during the summer campaign. Highest levels were found in Alfacs bay in every season. The occurrence of DA was not observed in the lagoons except in La Tancada where DA was detected in autumn and winter, at concentrations of <mLOQ and 8.78 ng L⁻¹.

This study confirms the presence of DA in Ebro's Delta bays during all seasons. Although this marine toxin is present at trace level concentrations, it is essential to monitor and control DA due to its toxic properties and capability to accumulate in feeder filters which are necessary for the mariculture activities of the area.

Conflicts of interest

There are no conflicts to declare.

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