

Monitoring Domoic Acid in Marine Phytoplankton by Disposable Immunosensors

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Abstract

Advances in sensor technology have allowed the significant progress in the monitoring of noxious compounds in the sea, providing real-time detection so as to prevent risks associated with the diffusion and dispersion of toxic substances in the environment. An important element in the overall picture is the harmful algal blooms which pose serious threats to marine ecosystems through the production of toxins that accumulate in filter-feeders and ultimately impact both human health and fisheries. Domoic acid is a neurotoxic amino acid produced by marine planktonic diatoms of the genus *Pseudo-nitzschia*. Here we monitored domoic acid production by natural *Pseudo-nitzschia* populations in phytoplankton samples collected along the Middle Tyrrhenian coast, over the course of one year, using selective immunosensors based on screen-printed electrodes, using differential pulse voltammetry as the electrochemical technique, to yield quantitative outputs. In this work, disposable devices have been applied for monitoring the production of domoic acid on algal extracts and the results have been validated by conventional high pressure liquid chromatography-ultraviolet detection methods. The data obtained revealed the presence of domoic acid in Italian phytoplankton, especially in coastal impacted areas, highlighting the potential risk of toxin entering into marine food webs and the environment. Immunosensors based on screen-printed electrodes prove to be effective tools for annual monitoring of domoic acid in seawater samples, thus providing a reliable early warning system relative to health and economic impact of algal toxins.

Keywords

Domoic Acid Monitoring, Immunosensors, Screen Printed Electrodes, Toxic *Pseudo-nitzschia*

1. Introduction

Harmful Algal Blooms (HABs) are an important environmental and public health problem all over the world and their economic impact can produce losses of great magnitude in the affected areas [1] [2]. HABs have been increasingly frequent, in recent years, endangering marine coastal life and food resources through massive growth of microalgae [3] [4]. Some HAB species produce toxins that may accumulate beyond the defined safety level in filter-feeders ultimately causing acute or chronic syndromes in humans. Marine biotoxins can also directly bring about animal mortalities, causing extensive seafood mortalities and seafood market closures involving the contaminated products [5] [6].

The marine biotoxin domoic acid (DA) is an analogue of the excitatory amino acid glutamate and acts as a neurotoxin in animals. DA is primarily produced by planktonic diatoms belonging to the cosmopolitan genus *Pseudo-nitzschia*. During *Pseudo-nitzschia* blooms, DA can contaminate shellfish and other filter feeders and can be transferred by ingestion to higher trophic levels that include marine mammals, seabirds and humans, causing the neurologic syndrome known as Amnesic Shellfish Poisoning (ASP) [7] [8] [9].

Blooms of *Pseudo-nitzschia* are recurrent phenomena in Italian coastal waters, with densities reaching up to 10^6 cell/L for prolonged periods of time and exceeding the regulatory level of 10^5 cell/L [10] [11] [12]. Ultrastructural examination of natural phytoplankton samples highlighted mass densities of at least six potentially toxic species in the Middle Tyrrhenian Sea (Mediterranean Sea) [12] and toxicity had been proven in cultures of *Pseudo-nitzschia galaxiae* and *P. multistriata* strains isolated from the Gulf of Naples [13] [14]. However, potentially toxic *Pseudo-nitzschia* species appear more widespread than the toxicity events in the whole Mediterranean basin, which indicates there is a high risk that toxicity may suddenly arise in some areas as a consequence of the increased exploitation of marine resources.

Following the first documented ASP event in Canada [15], the European Commission Directive 2002/226/EC implemented a maximum permitted level (MPL) of 20 mg of domoic acid/kg in whole shellfish intended for human consumption [16]. High performance liquid chromatography with ultraviolet detection (HPLC-UV) is the first chemical analytical method employed for DA detection and is still the most commonly used for monitoring shellfish [17]. The HPLC-UV method has been validated and standardised through the AOAC International Official Methods Program (AOAC method 991.26 and European Committee for Standardization (CEN) method 14176) [18]. Several other chemical methods have been developed to improve analytical sensitivity [19] and include liquid chromatography-mass spectrometry using reaction monitoring (MRM) [20]. Although these techniques provide good accuracy and reproducibility, several drawbacks remain: they require time-consuming sample preparation and the Electrospray Ionization (ESI) interface of the mass spectrometry turns out to be susceptible to salt effects with a decrease in instrumental sensitivity and dynamic range [21]. Thus, procedures based on HPLC combined with ultraviolet diode array detec-

tion (UV-DAD), ESI or MS appear suitable for laboratory assays and confirmatory investigations. However, due to the increasing exploitation of marine coastal resources high throughput screening and/or on-site testing would be more desirable but the standard laboratory instrumentation is often inadequate for such operation as it is expensive, time consuming and not portable. In addition, a method based on enzyme-linked immunosorbent assay (ELISA) has been validated (AOAC method 2006.02) and is officially allowed to be used in the EU for screening purposes [16]. The assay is primarily intended for using in routine monitoring of DA levels in bivalve molluscs to comply with the regulatory MPL, but is also applicable for DA quantification in other marine matrices like algal samples, seawater and body fluids of marine mammals.

Recently ELISA assays have been coupled with screen-printed electrodes (SPEs) to assemble disposable immunosensors for simple and fast measurement of several types of analytes (pesticides, toxins, heavy metals and hormones) found in the environment and food [22]-[29]. These systems combine the selectivity and sensitivity of immunological assays with a short analytical time and can be inexpensively produced in a “ready to use” form for field analysis by means of a portable electrochemical detector. However, few immunosensors are commercially available at present and they have yet to be established as research or routine tools, due to the lack of validated protocols for a wide range of sample matrices.

In this study, we have directly applied a detection method based on a disposable immunosensor coupled to a highly sensitive electrochemical technique, previously developed for DA determination in mussels [22], to marine phytoplankton samples. The developed immunosensor required assay procedure optimization for application to the phytoplankton matrices. The method basically involves the use of screen-printed electrodes (SPEs) with an “indirect competitive ELISA format” and the differential pulse voltammetry (DPV) for toxin detection. To our knowledge, this is the first study to determine DA in natural microalgal samples collected from Italian coastal waters over an annual cycle. Results indicate the potential of this SPEs-based method for monitoring DA at critical sites and during particular times of the year in order to provide an early warning of toxin presence in seawater phytoplankton and thus to prevent poisoning of humans and other animals.

2. Materials and Methods

2.1. Chemical and Reagents

DACS-1C certified calibration solution (100 µg DA/mL) was obtained from the Canadian National Research Council (<http://www.nrc-cnrc.gc.ca/>, Halifax, Canada). Affinity purified anti-goat IgG (H + L, from mouse), alkaline phosphatase conjugate (Ab_{II}-AP), polyvinyl alcohol (PVA) and all other reagents were from Sigma (www.sigma-aldrich.com, St Louis, MO, USA). 1-naphthyl phosphate (1-NPP) was obtained from Fluka Chemie (www.sigma-aldrich.com, Sigma-Aldrich, Milan, Italy). Bovine serum albumin conjugated with DA (BSA-DA) and

the polyclonal antibodies against DA (PAb_I from sheep) were kindly provided by Toxicology and Food Safety Research (AgResearch Limited, www.agresearch.co.nz/, New Zealand). Domoic acid (DA), used to prepare the standards and the spiked samples, was from Biomol (www.enzolifesciences.com/, Plymouth Meeting, USA). Single use syringe filters were purchased from Sartorius AG (www.sartorius.it, Bagno a Ripoli, Italy).

2.2. Apparatus

All electrochemical measurements were performed using a computer-controlled system, AUTOLAB model PGSTAT 12 with GPES software (ECO-CHEMIE, The Netherlands). The screen-printed electrodes (SPEs) were produced with a 245 DEK (Weymouth, UK) screen printing machine. Graphite based ink (Electrodag 423 SS) from Acheson (Milan, Italy) was used to print the working and counter electrode. Silver ink (Electrodag 477 SS) was used to print the reference electrode. The substrate was a flexible polyester film (Autostat HT5) obtained from Autotype Italia (Milan, Italy). The electrodes were home produced in foils of 48. The diameter of the working electrode was 0.3 cm resulting in a geometric area of 0.07 cm².

The HPLC system consisted of a modular Chromoquest spectral system form Thermoquest (San Jose, CA, USA), equipped with two P4000 pumps, a Shimadzu UV-VIS spectrometer, model (SPD-10AV) as detector. An SN 4000 controller operated the HPLC system working under software HPLC CL VP 4.3 software. The chromatographic separation was performed using a reverse phase C18 RESTEK Pinnacle II™ C18, (250 × 4.6 mm, D 3 μm) stainless steel column.

The Ultrasonic Cell Disruptor system was a SONIFER B12 (G. Heinemann, Germany).

2.3. Procedure for Immunosensor Production

The immunosensor for DA determination was realized as described by Micheli *et al.* [22] based on screen-printed electrodes (SPEs). Immunoassays were performed on the carbon-working surface of an SPE, which was modified in order to obtain a device to react with specificity and selectivity towards analyte.

The working electrode was coated with 7 μl of 0.1 M of 50 mM carbonate buffer pH 9.6 (CB), containing BSA-DA conjugate (30 μg/mL) and incubated for 1 h at room temperature. After washing with 160 μl of 15 mM phosphate saline buffer with Tween 20 (5%) PBS-T (2 min) and then 160 μl of PBS (2 times, 2 min), the electrode was treated with blocking reagent (1% PVA in CB) for 30 min at room temperature.

Competition reactions involved the addition a PBS-M solution with a fixed concentration of PAb_I (1:500 v/v) and with various concentrations of DA standard for 30 min at room temperature. After the washing step, the Ab_{II}-AP (1:1000 v/v, 6 μL) was incubated on the working electrode for 15 min. After another washing step, the enzymatic reaction was performed by addition of the substrate solution (80 μl) to each electrode.

Differential pulse voltammetry (DPV) was used to measure the analytical signal (potential range 0 - 600 mV, scan rate 300 mV/s, pulse amplitude 70 mV, pulse width 50 ms). 80 μL of solution containing the enzymatic substrate (1 mg mL^{-1} 1-NPP in diethanolamine buffer, DEA, with 1 mM MgCl_2) was applied on the sensor surface. After 2 min of incubation, the DPV measurement was started and the resulting current recorded.

2.4. HPLC Analysis

For immunosensor validation, separation and chromatographic analyses were performed according to the recommended instructions given in MUS-1 Certified Reference Material, Mussel Tissue Reference Material for Domoic Acid, NRC-CNRC [30]. The mobile phase was aqueous acetonitrile with 0.1% trifluoroacetic acid and a flow rate of 0.7 $\text{mL}\cdot\text{min}^{-1}$ (loop 20 μL) for 15 min. Detection was accomplished by monitoring absorbance at 242 nm with a 10 nm bandwidth [19] [30]. Quantification of DA was obtained by comparing the areas of peaks from extracts with those from a dilution series of the DACS-1 calibration solution. The calibration curve for DA was linear over 3 orders of magnitude and reproducibility of this external calibration procedure was 3% RSD for 6 repeated analysis of a single extract.

2.5. Sampling and Sample Preparations

Samples were collected fortnightly, when possible, at 6 stations, 500 m offshore, along the Latium coast, in a 300 km stretch (Middle Tyrrhenian Sea), from January 2001 to December 2002 (Foce Marta, VTA; Ladispoli, RMB; Fiumicino, RMC; Rio Martino, LTD; Monte d'Argento LTE; Zannone Island, LTF, **Figure 1**) in the framework of a research collaboration between the University of Rome "Tor Vergata" and the Regional Agency for Environmental Protection of Latium (ARPA Lazio) on coastal environment assessment and algal surveillance activities [Programma di Monitoraggio per il Controllo dell'Ambiente Marino Costiero (L. 979/82), Programma di Sorveglianza Algale (D.P.R. 470/82) from 1998]. Closing plankton net (mouth diameter of 0.21 m and 20 μm mesh size) was towed obliquely for 300 m intervals to obtain concentrated phytoplankton samples. After gentle mixing to homogenize the suspension, each sample was subdivided into two aliquots. A 5 mL aliquot was immediately preserved with 2.5% glutaraldehyde for electron microscopy to confirm species identification [12]. A second aliquot (50 to 200 mL suspension) was concentrated, within 48 h from collection, onto a 5.0 μm membrane filter. The filters were then transferred to a graduated tube and stored at -20°C for DA determination according to Bates *et al.* [31].

For DA measurements, a fixed volume (5.2 mL) of $\text{CH}_3\text{OH}:\text{H}_2\text{O}$ (3:1 v/v) solution was added to membrane filters. Filters, after mixing (1 min), were sonicated for 2 min at 28 μs amplitude and the suspension then centrifuged at 3000 rpm (10 min) to remove the particulate. Then, the supernatant was filtered through a 0.22 μm filter and analyzed, without dilution, by immunosensor and

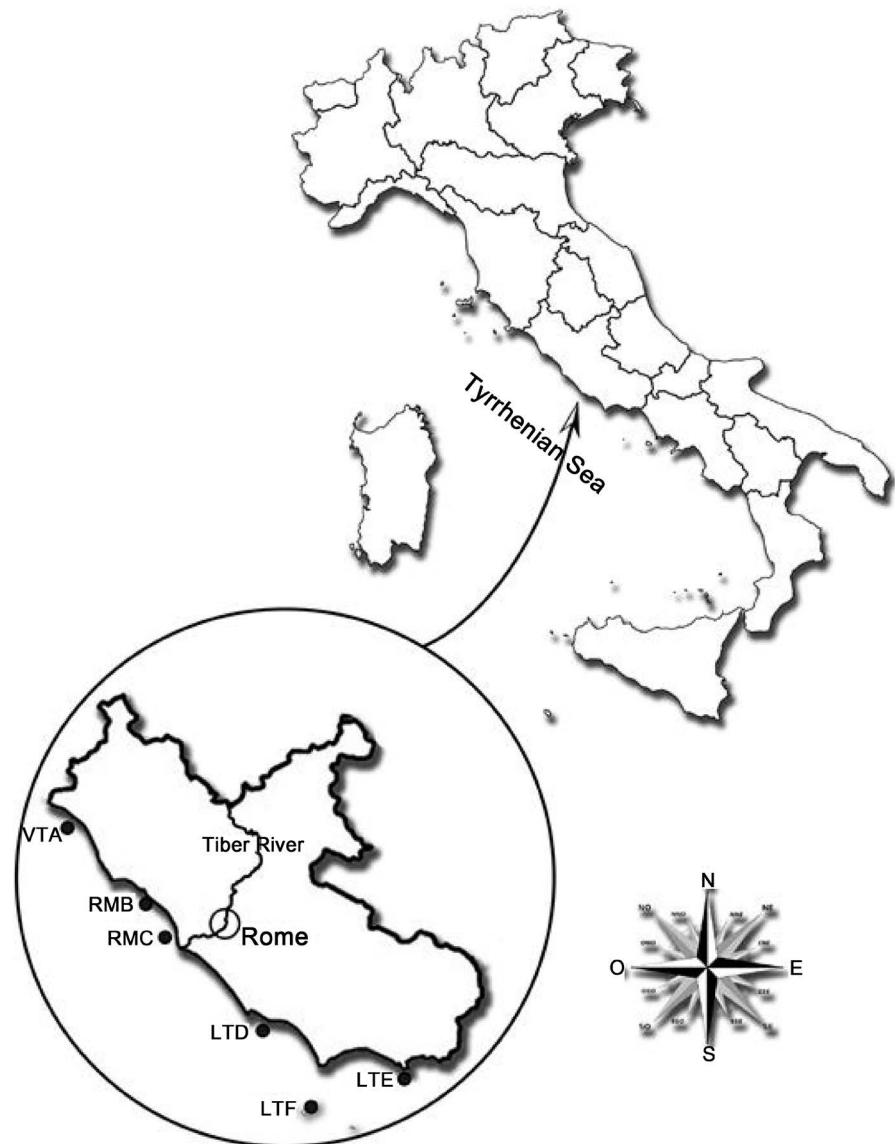


Figure 1. Study area along the Middle Tyrrhenian coast and sampling sites in the inset.

by HPLC methods following Micheli *et al.* [22] and the procedure reported in [19] [30].

2.6. Determination of Recovery

DA recovery from phytoplankton was determined by adding DACS-1 followed by 1 mL of sonicated phytoplankton samples with no *Pseudo-nitzschia* present (after microscopy measurements) and domoic acid (determined by HPLC measurements) to a 13 mm × 100 mm disposable screw-cap glass test tube at 3 DA concentrations (0.5, 20, 60 ng·mL⁻¹), each in triplicate. The spiked samples were concentrated on membrane filters and stored at -20°C according to the procedure described above.

The analysis of each spiked sample was repeated five times in the same day to obtain repetitively values and five times on three different days to obtain repeat-

ability values. Confirmation of the immunosensor results was obtained by analyzing the same extracts (sample or spiked phytoplankton) using a previously validated HPLC method [19] [30].

2.7. Calibration Plots and Analysis of Samples

Standard curves were obtained using DA standard solutions prepared in 15 mM phosphate saline buffer (PBS), pH 7.4, with 10% CH₃OH (PBS-M) in the range 0 - 500 ng mL⁻¹ for immunosensor and HPLC analysis, respectively.

The data obtained for each curve (measured current *versus* competitor concentration) for the immunosensors were fitted using a “non-linear 4 parameter logistic calibration plots” [32] and Sigma Plot software (SPSS). The four parameter logistic function (Equation (1)) is:

$$f(x) = \left\{ \frac{[1-a]}{[1+(x/c)^b]} \right\} + d \quad (1)$$

the parameters “*a*” and “*d*” are the asymptotic maximum and minimum values, respectively; “*c*” is the value at the inflection point (IC₅₀) and *b* is the slope. The detection limit (LOD) was defined as the decrease of the maximum signal equal to three times the value of the standard deviations (*I*₀ - 3SDs), measured in the absence of DA (*I*₀, no competition point). Before analysing samples, the matrix effect was evaluated using phytoplankton samples with no *Pseudo-nitzschia* present (after microscopy observations) and domoic acid (HPLC analysis), called “blank samples” from now on in the text. The matrix effect on immunosensor measurements was evaluated by comparison of the two calibration curves realised with the same known amount of DA standard: the first was obtained with DA standard prepared in PBS-M (assay buffer); the second in blank samples spiked with the same amount of DA standard used for the first curve.

3. Results and Discussion

A total of 105 phytoplankton samples were collected and aliquots immediately observed by light microscopy for a phytoplankton survey. Subsamples for DA analysis were filtered and stored without any washing or other treatment (as described in the experimental section).

Before analyzing samples collected on filters (particulate DA), the effect of this matrix on the immunosensor analysis was evaluated using two spiked blank samples with the same concentration of DA standard utilised for calibration curves in buffer. The standard curve for DA diluted in PBS-M (●, **Figure 2**) was constructed and compared with the calibration curves obtained in spiked blank samples (see material and methods) (△ and ×, **Figure 2**). The linear regression of the calibration curves obtained in buffer (PBS-M; $Y = 167.80 - 90.58X$, $r^2 = 0.972$) and in both algal blank samples ($Y_1 = 162.25 - 86.20X$, $r^2 = 0.979$; $Y_2 = 161.11 - 85.38X$, $r^2 = 0.979$) showed comparable slopes, indicating that there was no significant matrix effect by the various components present in phytoplankton samples on electrochemical measurements. The slope of the linear regression of the three calibration curves were similar, but not equal indicating that the

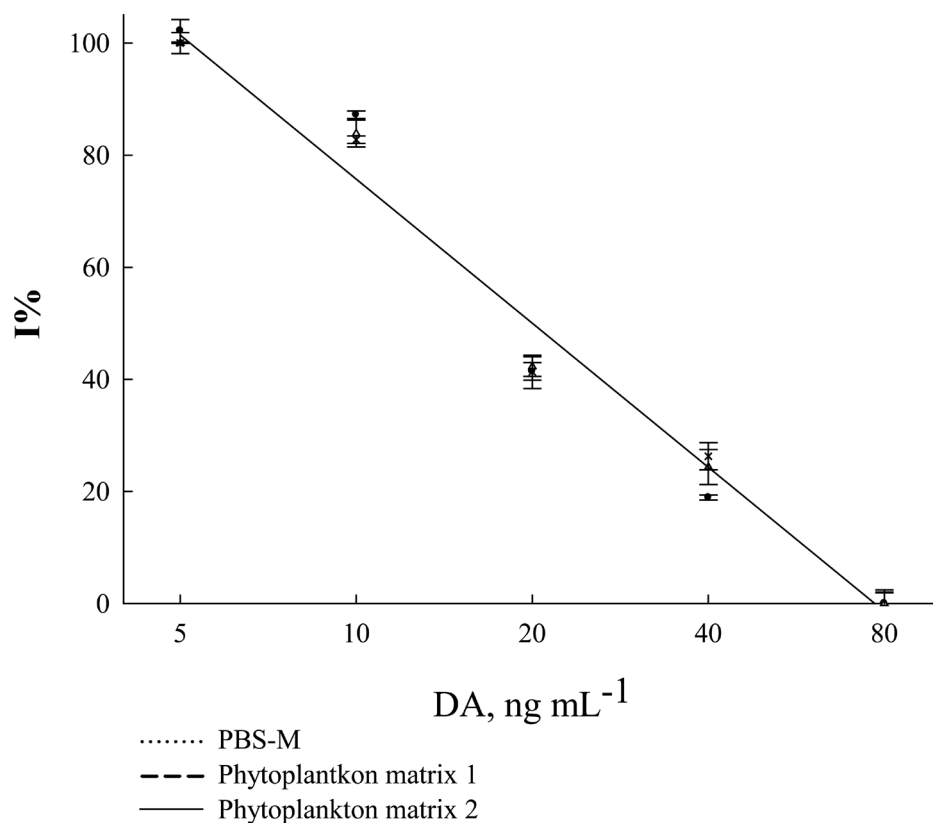


Figure 2. Study of matrix effect: comparison of calibration curves for DA obtained in buffer (saline phosphate buffer-methanol; PBS-M) (●), and in two blank algal matrixes of the matrix (△ and x).

sensitivity of the analysis (determined by the slope) was not significantly affected by the sample matrix. This effect is very little, as can be observed by the linear regression lines that are superimposed for the two matrices, and parallel but not coincident with that in buffer, probably due to the different composition (ionic strength and matrix components) of the matrix from the algal buffer.

In all cases, the linear range and detection limit (LOD) were, respectively, between 5 and 80 ng·mL⁻¹ and around 1 ng·mL⁻¹. Precision assays were carried out at three levels (5, 20, 60 ng·mL⁻¹) and the results were satisfactory. The RSD% values (n = 5) obtained ranged from 2% to 6% for repeatability (intra-day precision) and from 2% to 4% for intermediate (inter-day) precision. The intra-electrode reproducibility, expressed as %RSD, was of 6.6%, for a concentration of 10 ng·mL⁻¹ with n = 30.

Recovery studies were carried out using validation samples collected on the same membrane filters as the ones used for phytoplankton analysis, in order to evaluate the accuracy of the method. Samples were spiked (artificially contaminated phytoplankton) with DA concentrations equal to 5, 20, 60 ng·mL⁻¹ before the extraction. Due to the lack of toxin limit for phytoplankton samples, we decided to validate results using the three concentration levels comprised in the immunosensor linear range. Confirmation of the immunosensor results for spiked samples was obtained analyzing the same extracts by the HPLC method

Table 1. Study of the recovery and relationship with HPLC results.

Standard	Immunosensor				HPLC			
	DA, $\mu\text{g/mL}$	DA observed, $\mu\text{g/mL}$	Recovery %	RSD%*	DA observed, $\mu\text{g/mL}$	Recovery %	RSD%*	RE%**
5.0		4.9	98	8	4.9	98	2	0
2.5		2.0	80	9	2.5	100	3	-20
0.5		0.4	80	8	0.6	120	2	-30

*RSD% (relative standard deviation) = (standard deviation/media of concentration) \times 100; **RE% (relative error%) = [(measuredvalue – truevalue)/truevalue] \times 100.

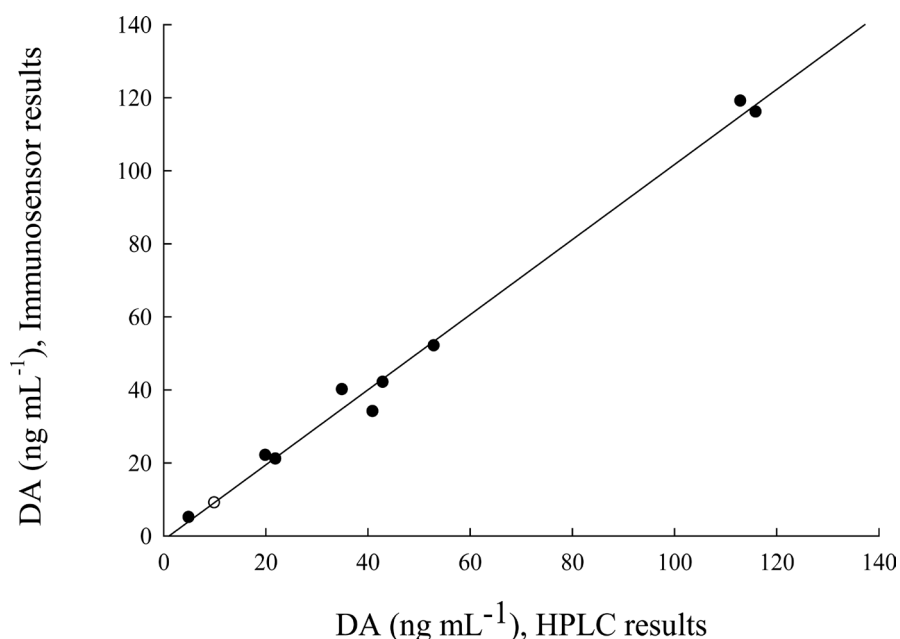


Figure 3. Comparison of the results obtained with immunosensor and HPLC in order to evaluate the accuracy of the proposed electrochemical immunosensor versus chromatographic analyses [18] using spiked phytoplankton samples (\bullet correlation point).

following the AOAC protocol [17]. The recovery of DA ranged between 100% - 80%, with good agreement between the proposed and the conventional methods. **Table 1** reports the accuracy of immunosensor *versus* HPLC analyses for artificially contaminated phytoplankton and evaluated resulting around 20%.

After the evaluation of the proposed electrochemical results in algal matrix, phytoplankton samples were analyzed by the immunosensor method and the data validated by HPLC analyses (**Figure 3**: $Y = 1.03X - 1.08$, for $n=10$), obtaining a scatterplot satisfactory agreement ($R = 0.993$). 26 samples were found to be positive using the immunosensors and 31 using HPLC (**Figure 4**, **Table 2**).

This is the first study to document the presence of DA and its concentration in natural phytoplankton samples from the Italian seawater. Although this toxin has been found in shellfish from the Adriatic Sea, on both Italian [33] and Croatian [34] coasts, as well as in cultured strains of the major representative *Pseudo-nitzschia* species from Italian waters, no previous research aimed at investigating

Table 2. Comparison of the results, in terms of DA concentration found, obtained for the phytoplankton samples analysed by the immunosensor and HPLC.

Immunosensor	HPLC	RE%
DA found (ng/mL)	DA found (ng/mL)	
9	10	-10
34	41	-17
119	113	5
21	22	-5
42	43	-2
52	53	-2
22	20	-10
116	116	0
5	5	0
40	35	14

the temporal history (occurrence and seasonal variations) of DA directly in the potential producing organisms. In any case, there have not been any reported incidences of ASP in Italian waters. This highlights the fact that the presence or absence of DA contamination in local food webs may not reflect either the magnitude of the local population of toxic *Pseudo-nitzschia* or their local production of DA, as has also been hypothesized for some stretches of the Californian coast [35] [36], where most of the studies on DA in phytoplankton were carried out.

The results presented here document the particulate DA (pg DA L^{-1}), as evaluated using electrochemical immunosensors in phytoplankton samples from five coastal stations, known to be impacted, and in one control site. We also carried out parallel electron microscopic analyses of samples and assessed the *Pseudo-nitzschia* diversity therein. Results revealed the presence of at least eight different *Pseudo-nitzschia* species, 6 of which are known DA producers in culture [12].

DA concentration varied between the monitored stations, and in addition, intra-annual variability was observed at each single station. This highlighted certain periods and conditions that may influence both the potential impact over the year as well as variations along the sampled stretch of coast. In particular, the VTA and LTE stations showed the highest number of positive samples (seven out of 19 and 15, respectively), while samples from the control site, Zannone Island (LTF), revealed no presence of DA whereas at LTD only 2 winter records of DA, below 7 pg L^{-1} , were detected (Figure 4). There was a distinct north-south differentiation between sampling sites in relation to frequency of DA detected, with the stations located to the north of the Tiber River mouth (VTA, RMB, RMC) showing higher numbers of positive results. The absolute maximum DA concentration in the extracts ($7830 \text{ pg DA L}^{-1}$) was registered at RMC in the late June sampling. It is very interesting to note that the maximal concentrations measured at the Italian stations studied were substantially lower (on average

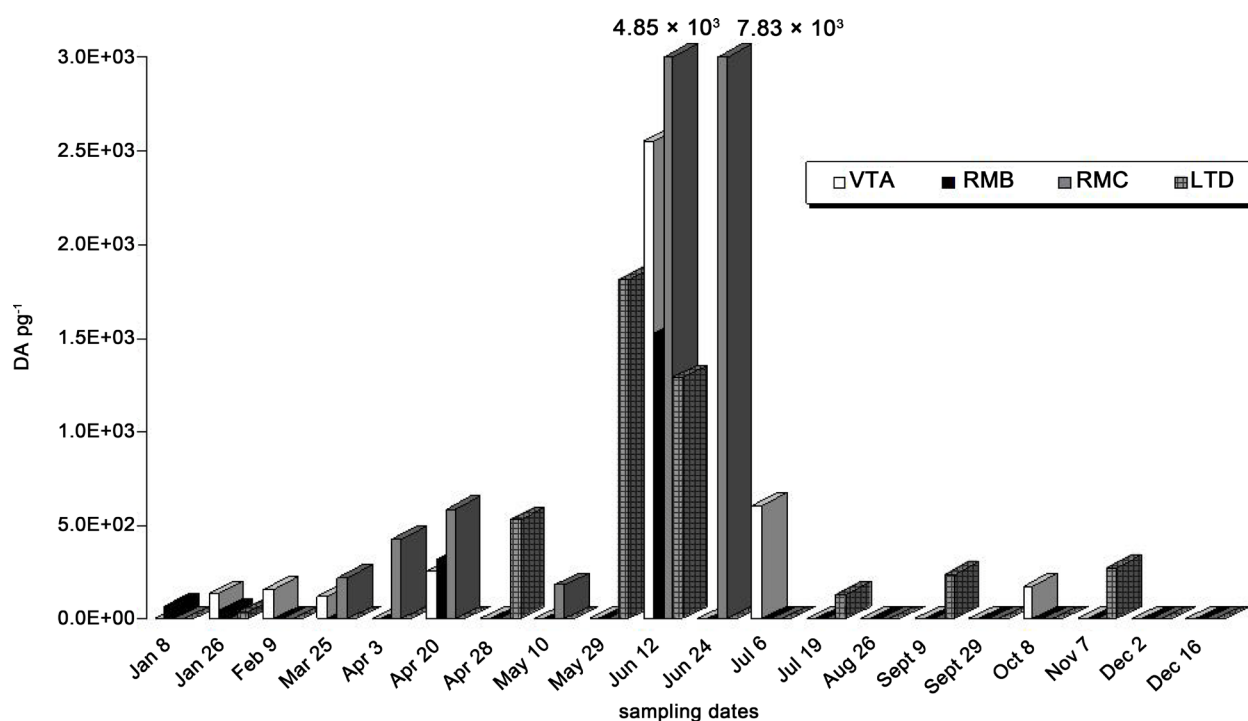


Figure 4. DA concentration in the samples, data (only two records) for station LTD are not shown.

500-fold) than values recorded during surveys conducted in Californian surface waters, from 1991 to 2007 [35] [36] and more recently in the Eastern English Channel (Normandy, France [37]). However, the DA levels recorded in our study were comparable to concentrations determined in natural plankton assemblages from the Chesapeake Bay [36] [38], where the relatively low values measured, despite a massive presence of toxic *Pseudo-nitzschia* demonstrated also in that area, was considered to be the result of both genetic and environmental factors (e.g. light and salinity). The highest relative levels of DA at the stations VTA, RMB and LTE were also measured in June, but earlier than the maximum record at RMC. Despite this homogeneity in the occurrence of the relative DA peak values, temporal distribution of DA production showed a degree of variability between northern and southern stations. In the north, particulate DA had a winter-spring occurrence, whereas a more diffuse presence throughout the year was detected at south of the mouth of Tiber. In addition, the concentration of domoic acid appeared to be influenced by river runoff and by the prevailing, northward, coastal current [39] with consequent higher availability of nutrients in the sites affected by the Tiber plume. River discharge and nutrient loading have been considered to influence *Pseudo-nitzschia* population development all over the world, but a direct association between DA fluxes and nutrients in nature definitely deserves further investigation [36] [38].

4. Conclusion

The data reported in this paper are first to result from the application of relatively simple, practical immunosensors for the monitoring domoic acid in microalgal natural populations of the Mediterranean basin. These sensors showed

an operating range and detection limits that made them highly suitable for controls in the field. Thus, they represent an important alternative tool to assist monitoring of potentially harmful DA presence in the marine environment, although further development could contribute to reducing the gap between the HPLC and immunosensor results obtained in this study. In any case, the electrochemical immunosensors were previously developed for DA detection in mussels [22] and were here optimised and applied to detect DA directly in phytoplankton samples, with no matrix effect. Commercially available ELISA kits offer highly sensitive toxin determination in algal extracts and water [40] but our method has proved to be more rapid (1 vs. 2.5 h), simpler and easy to perform, with good accuracy and reproducibility. Thus, this method has a high potential to be an effective screening analytical technique to “sense” DA directly in algal producers and to trace toxin transfer and magnification through the food chain in real time [41].

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