

*OSTREOPSIS FATTORUSSOI* SP. NOV. (DINOPHYCEAE), A NEW BENTHIC TOXIC  
*OSTREOPSIS* SPECIES FROM THE EASTERN MEDITERRANEAN SEA<sup>1</sup>

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The new benthic toxic dinoflagellate, *Ostreopsis fattorussoi* sp. nov., is described from the Eastern Mediterranean Sea, Lebanon and Cyprus coasts, and is supported by morphological and molecular data. The plate formula, Po, 3', 7'', 6c, 7s, 5''', 2''''', is typical for the *Ostreopsis* genus. It differs from all other *Ostreopsis* species in that (i) the curved suture between plates 1' and 3' makes them approximately hexagonal, (ii) the 1' plate lies in the left half of the epitheca and is obliquely orientated leading to a characteristic shape of plate 6''. The round thecal pores are bigger than the other two Mediterranean species (*O. cf. ovata* and *O. cf. siamensis*). *O. fattorussoi* is among the smallest species of the genus (DV:  $60.07 \pm 5.63 \mu\text{m}$ , AP:  $25.66 \pm 2.97 \mu\text{m}$ , W:  $39.81 \pm 5.05 \mu\text{m}$ ) along with *O. ovata*. Phylogenetic analyses based on the LSU and internal

transcribed spacer rDNA shows that *O. fattorussoi* belongs to the Atlantic/Mediterranean *Ostreopsis* spp. clade separated from the other *Ostreopsis* species. *Ostreopsis fattorussoi* produces OVTX-a and structural isomers OVTX-d and -e, *O. cf. ovata* is the only other species of this genus known to produce these toxins. The Lebanese *O. fattorussoi* did not produce the new palytoxin-like compounds (ovatoin-i, ovatoin-j<sub>1</sub>, ovatoin-j<sub>2</sub>, and ovatoin-k) that were previously found in *O. fattorussoi* from Cyprus. The toxin content was in the range of 0.28–0.94 pg · cell<sup>-1</sup>. On the Lebanon coast, *O. fattorussoi* was recorded throughout the year 2015 (temperature range 18°C–31.5°C), with peaks in June and August.

**Key index words:** benthic dinoflagellates; harmful algae; Mediterranean Sea; nutrients; *Ostreopsis*; ovatoin; palytoxins; phylogeny; taxonomy

**Abbreviations:** AP, anterioposterior diameter; DV, dorsoventral diameter; ITS, internal transcribed

<sup>1</sup>Received 12 May 2016. Accepted 2 August 2016.

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Editorial Responsibility: R. Wetherbee (Associate Editor)

spacer; L:D, Light:Dark; LC-HR MS<sup>n</sup>, Liquid Chromatography-High Resolution Multiple Stage Mass Spectrometry; ML, Maximum Likelihood; MP, Maximum Parsimony; NJ, Neighbor Joining; OVTX, ova-toxin; PLTX, palytoxins; RP, resolving power; UV, Ultraviolet; W, transdiameter

The genus *Ostreopsis* belongs to the family of Ostreopsidaceae (Gonyaucales, Dinophyceae), which includes two genera of benthic dinoflagellates (i.e., *Ostreopsis* and *Coolia*). The type species, *Ostreopsis siamensis* Schmidt, was first described in the Gulf of Siam (Thailand) in 1900 (Schmidt 1901). In the following years, several other species have been described by other authors: *O. lenticularis* Fukuyo, *O. ovata* Fukuyo (Fukuyo 1981), *O. heptagona* Norris, Bomber & Balech (Norris et al. 1985), *O. mascarensis* Quod (Quod 1994), *O. labens* Faust & Morton (Faust and Morton 1995), *O. marina* Faust, *O. belizeana* Faust and *O. caribbeana* Faust (Faust 1999, Hoppenrath et al. 2014). Of late, however, the validity of some *Ostreopsis* species has been questioned. Although the taxonomy of *Ostreopsis* is based on morphological characters (thecal plate pattern, shape and size), high morphological variability is reported within the same species and the original descriptive characters of some species are often considered inadequate to discriminate among most *Ostreopsis* species in field samples (Penna et al. 2005, Parsons et al. 2012, David et al. 2013). In addition to the confusion on the morphological features, no genetic data were provided in these original descriptions and the submission of any further new species description accompanied by a sequence and comparison with other sequences in GenBank has been recommended (Parsons et al. 2012). Lately, several phylogenetic studies based on LSU and internal transcribed spacer (ITS) analyses have been carried out on *Ostreopsis* species, supplying an increasing number of molecular clades (and subclades) that may represent cryptic species (Penna et al. 2010, 2014, Sato et al. 2011, Tawong et al. 2014).

*Ostreopsis* species were reported since a long time in tropical ciguatera endemic areas (Carlson and Tindall 1985, Bomber and Aikman 1989) and were often wrongly implicated in incidences of ciguateric syndrome (e.g., Tosteson 1995). Indeed, some *Ostreopsis* species are toxic, but their toxins (mostly belonging to the palytoxin group) are not those that cause ciguatera. Among the nine species of the genus *Ostreopsis*, toxicity has been demonstrated in *O. siamensis* (also for *O. cf. siamensis*), *O. mascarensis*, *O. lenticularis* and *O. cf. ovata* (Nakajima et al. 1981, Yasumoto et al. 1987, Holmes et al. 1988, Mercado et al. 1994, Meunier et al. 1997, Lenoir et al. 2004, Ciminiello et al. 2006, Scalco et al. 2012, Uchida et al. 2013, García-Altarets et al. 2015, Brissard et al. 2015). Moreover, *O. heptagona* was determined to be toxic, as methanol extracts of this

species isolated from Knight Key (Florida) were weakly toxic to mice (Babinchak, according to Norris et al. 1985).

In the past decade, *Ostreopsis* blooms have also become common in temperate areas during the summer-autumn period (Chang et al. 2000, Rhodes et al. 2000, Pearce et al. 2001, Vila et al. 2001, Taniyama et al. 2003, Turki 2005, Aligizaki and Nikolaidis 2006, Mangialajo et al. 2008, Shears and Ross 2009, Totti et al. 2010, Illoul et al. 2012, Ismael and Halim 2012, Pfannkuchen et al. 2012, Selina et al. 2014). In these areas, *Ostreopsis* blooms are often associated with noxious effects on the health of humans (Tichadou et al. 2010, Del Favero et al. 2012) and benthic marine organisms (Accoroni et al. 2011, Pagliara and Caroppo 2012, Gorbi et al. 2013, Carella et al. 2015).

In the Mediterranean Sea, two genotypes corresponding to the morphotypes *O. cf. ovata* and *O. cf. siamensis* have been recorded to date (Penna et al. 2010, 2012). Almost all knowledge about the ecology of *Ostreopsis* species in the Mediterranean Sea mainly refers to *O. cf. ovata* due to its dominance over the other species in this area (Battocchi et al. 2010, Perini et al. 2011). Several environmental parameters have been recognized to strongly influence bloom dynamics, such as hydrodynamics, water temperature and nutrients (Vila et al. 2001, Shears and Ross 2009, Totti et al. 2010, Mabrouk et al. 2011, Mangialajo et al. 2011, Cohu et al. 2013).

The genotype *O. cf. siamensis* has been detected along the Catalan coast, in the eastern Atlantic coast of Morocco, Portugal, northern Spain and southern Italy (Vila et al. 2001, Amorim et al. 2010, Benouna et al. 2010, Laza-Martinez et al. 2011, Ciminiello et al. 2013, David et al. 2013) and its morphotype has also been reported along the Tunisian and Lebanese coasts (Turki 2005, Turki et al. 2006, Mabrouk et al. 2011, 2012, Abboud-Abi Saab et al. 2013). In addition to these two species, recently Penna et al. (2012) found a new genotype probably corresponding to a new species of *Ostreopsis*, in both the Atlantic Ocean (Canary Islands) and Mediterranean Sea (Greece and Cyprus Island). Strains of this new genotype collected in Cyprus have been reported to produce OVTX-a, -d, -e, and isobaric palytoxin, so far found only in *O. cf. ovata*, adding three new palytoxin-like compounds to those already identified, i.e., OVTX-i, OVTX-j<sub>1</sub>, OVTX-j<sub>2</sub>, and OVTX-k (Tartaglione et al. 2016). This genotype will be described here as a new species.

Descriptions of new species are often based on culture material rather than field samples (e.g., Litaker et al. 2009, Tillmann et al. 2009, Fraga et al. 2011, Percopo et al. 2013, Fraga and Rodríguez 2014). This approach, however, can have drawbacks when cells in culture strongly change their morphology producing aberrant cells and/or modifying the shape and the size of the thecal plates and the entire cell, as occurs in *Ostreopsis* (Aligizaki and

Nikolaidis 2006, Laza-Martinez et al. 2011, Nascimento et al. 2012, Scalco et al. 2012, David et al. 2013). Moreover, under certain culture conditions, in addition to the typical vegetative morphology, different stages of life cycle (characterized by different size, shape, and plate tabulation) may appear (Ali-gizaki and Nikolaidis 2006, Bravo et al. 2012, Accoroni et al. 2014). On the other hand, the observation of culture material is necessary to obtain genetic material for gene sequencing and may be necessary to describe the whole range of morphological variability of a species and its different life stages in order to avoid diverse life stages of the same species being described as different species (Coats 2002). It is, therefore, crucial to use both culture and natural samples in the description of a new species.

In this study, we describe *Ostreopsis fattorussoi*, a new toxic benthic dinoflagellate found in the Lebanon and Cyprus coasts (Eastern Mediterranean Sea), on the basis of molecular and ultrastructural features of both natural and culture samples. We also investigated the ecology of this new species along the Lebanon coast and analyzed its toxin content and profile.

#### MATERIALS AND METHODS

**Sampling and sample treatment.** Macroalgal samples were collected at three sites from Cyprus and Lebanon in 2013 and 2015, respectively: site 1, Vassiliko Bay, southern coast of Cyprus (34°43' 19.61" N, 33°18' 6.67" E), site 2, Batroun, Lebanon (34°15' 5.40" N, 35°39' 24.78" E), and site 3, Byblos, Lebanon (34°06' 51.84" N, 35°38' 53.76" E; Fig. 1). All sites were located in shallow and rocky shores.

Macroalgae (mainly *Halopteris scoparia* (Linnaeus) Sauvageau in Cyprus and *Corallina elongata* Ellis & Solander in Lebanon) were sampled and then processed to detach epiphytic cells (see below). Samples for microscope observations were fixed with 0.8% neutralized formaldehyde (Thronsen 1978) and stored at 4°C in the dark until the analyses.

At site 2 (Batroun, Lebanon), an ecological study about the temporal trend of *Ostreopsis* abundances and the relationships with some environmental parameters was carried out.

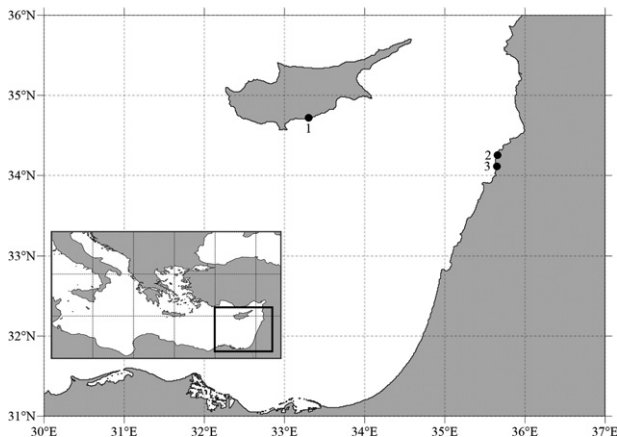


FIG. 1. Map of study area, showing the sites where *Ostreopsis fattorussoi* was found. Site 1: Vassiliko Bay (Cyprus). Site 2: Batroun (Lebanon). Site 3: Byblos (Lebanon).

This site is characterized by a rocky bottom and is affected by the discharge of phosphate into the sea from a chemical factory.

The sampling was carried out monthly from January to April 2015, then every 3–7 d until October 2015. The temperature was recorded using an alcohol thermometer (precision  $\pm 0.1^\circ\text{C}$ ). Salinity was measured using an induction salinometer (Beckman RS7-C) once back in laboratory. Surface water samples for nutrient analyses were collected through plastic bottles. Samples were immediately stored at  $-20^\circ\text{C}$ . Orthophosphates (P- $\text{PO}_4$ ) were analyzed according to Murphy and Riley (1962), nitrites (N- $\text{NO}_2$ ) and nitrates (N- $\text{NO}_3$ ) to Strickland and Parsons (1968).

Macrophytes and seawater samples were collected in triplicate following the protocol of the ENPI-CBCMED project M3-HABs (<http://m3-habs.net>). Water samples were collected in plastic bottles (250 mL) at 20 cm depth (30 cm above the macroalgae), before the collection of the benthic substrata, in order to avoid resuspension. Portions of macroalgal thalli (~15 g) were collected using 250 mL plastic bottles. Seaweed samples in the storage water were shaken vigorously to dislodge the epiphytic cells, then the thalli were re-rinsed with filtered seawater to completely remove *Ostreopsis* cells. Finally, the samples were fixed by adding Lugol's solution at 1% V/V and stored at 4°C in the dark until the analyses.

**Light microscope analysis.** Light microscopy observations for the new species description were carried out using an inverted microscope (Zeiss Axiovert 135, Carl Zeiss, Oberkochen, Germany) equipped with phase contrast, differential interference contrast and epifluorescence with a UV lamp. Field samples were analyzed after staining with the DNA-specific dye SYBR Green to visualize the nucleus and the cellulose specific dye Calcofluor-White M2R to elucidate the thecal morphology.

Microphotographs were taken with a Canon EOS 6D (Canon Inc., Tokyo, Japan) digital camera. In the event that the field depth was not enough for the whole subject, multiple images taken at different focus distances were merged using image manipulation software.

*Ostreopsis* cells from field samples were measured at 400 $\times$  magnification using a micrometric ocular. In samples of both Cyprus and Lebanon, 141 cells were measured along the dorsoventral diameter (DV) and the transdiameter (W); for a subset of cells (101) the anterioposterior diameter (AP) was also measured, using a needle to turn them over.

Abundances of *Ostreopsis* cells at site 2 were estimated both in benthic and in planktonic subsamples (1 and 50 mL, respectively) after homogenization. Counting was carried out in Utermöhl chambers (Edler and Elbrachter 2010) for seawater samples or Sedgewick-Rafter chambers (Guillard 1978) for benthic samples, through an inverted light microscope (Wild M40, Wild Heerbrugg AG, Heerbrugg, Switzerland) and a light microscope (Motic BA 400, MoticEurope S.L.U., Barcelona, Spain) respectively, both equipped with phase contrast, at 200 $\times$  magnification. Counting was performed on random transects, or on the half chamber, in order to count a representative cell number. Cell abundances were expressed as cells  $\cdot$  g $^{-1}$  fw and cells  $\cdot$  L $^{-1}$  for water column.

Estimates of the net growth rates ( $\mu \cdot \text{d}^{-1}$ ) of the natural populations of *Ostreopsis* throughout the year were calculated according to:

$$\mu = (\ln N_2 - \ln N_1) / (t_2 - t_1)$$

where  $N_2$  and  $N_1$  are *Ostreopsis* abundances on each benthic substrata at respective sampling day,  $t_1$ , or  $t_2$ .

The abundances of *Ostreopsis* in benthic samples were tested for significant correlations (Pearson) with all recorded

environmental parameters and with abundance in the water column. Growth rates were tested for significant correlations (Pearson) with all environmental parameters. The statistical analyses were conducted using Statistica (Statsoft) software.

**Scanning electron microscope analysis.** Some preserved subsamples (1–5 mL) were dehydrated by immersion in ethanol at increasing gradations (10%, 30%, 50%, 70%, 80%, 90%, 95%, and 100%). After 1 d in absolute ethanol, the dehydrated samples were filtered on a Nucleopore polycarbonate filter, and treated in a Critical Point Dryer (Polaron CPD 7501, Quorum Technologies, Newhaven, UK). Filters were placed on stubs and sputtered with gold-palladium in a Sputter Coater (Polaron SC 7640) for observation under the scanning electron microscopy (FE-SEM; Zeiss Supra 40).

**Strain isolation.** *Ostreopsis* cell isolates were obtained from microphytobenthos epiphytic on seaweeds collected on July 2013 and June 2014 at site 1, Vasiliko Bay (Cyprus), and site 2, Batroun (Lebanon), respectively, using the capillary pipette method (Hoshaw and Rosowski 1973). After an initial growth in microplates, cells were cultured at  $23^{\circ}\text{C} \pm 1^{\circ}\text{C}$  under a 14:10 L:D photoperiod and an irradiance of  $100 \mu\text{mol photons} \cdot \text{m}^{-2} \cdot \text{s}^{-1}$ , in modified f/10 medium Si free (Guillard 1975) using filtered and autoclaved natural seawater (salinity 35). One (C1005) and five (L1000, L1007, L1008, L1020, and L1022) strains of *Ostreopsis* sp. were isolated from site 1 and 2, respectively (Table 1).

**DNA extraction, PCR amplification, sequence alignment.** Genomic DNA was extracted and purified from 50 mL monoclonal cultures of *Ostreopsis* sp. in logarithmic growth phase using a DNeasy Plant Kit (Qiagen, Valencia, CA, USA) according to the manufacturer's instructions. Briefly, cultures were collected by centrifugation at  $3,000g$  for 20 min. The supernatant was removed and the pellets were transferred to a 1.5-mL tube, centrifuged at  $13,000g$  for 10 min. The pellets were immediately processed or stored at  $-80^{\circ}\text{C}$  until DNA extraction.

The PCR amplification of ribosomal 5.8S gene and non-coding ITS regions, and partial nuclear LSU (D1/D2 domains) were described in Penna et al. (2010). Amplified PCR fragments were purified (Penna et al. 2014) and sent to Eurofins Genomics (Ebersberg, Germany). All nucleotide sequences of ITS-5.8S rDNA and LSU deposited in ENA-EMBL are listed in Table 1.

**Phylogenetic analyses.** The ITS-5.8S and LSU sequences were aligned using MAFFT software. Short aligned sequences and ambiguously aligned positions were excluded from the alignment manually or using Gblocks (<http://molevol.cmima.csic.es/castresana/Gblocks.html>) with default settings.

The jModelTest v.2.1.7 (Darriba et al. 2012) was used to determine the evolutionary model that best fitted data according to Akaike Information Criterion. For both ITS-5.8S and LSU gene rDNA alignment, the most appropriate evolutionary model was found to be HKY+I+G with a gamma distributed rate of variation among sites equal to 1.56 and 1.47, respectively.

Neighbor-Joining (NJ) and Maximum Parsimony (MP) analyses were performed using MEGA6 (Tamura et al. 2013). The MP analyses were performed using the Tree-Bisection-Redrafting algorithm with search level 1 in which the initial trees were obtained by the random addition of sequences (10 replicates). All positions containing gaps and missing data were eliminated. The robustness of NJ and MP trees was tested by bootstrapping using 1,000 pseudo-replicates. Maximum likelihood (ML) analyses were run with Phyml v.3.0 (Guindon et al. 2010). Bootstrap values were calculated with 1,000 pseudo-replicates.

Bayesian analyses (BI) were performed using MrBayes v. 3.2.3 (Ronquist and Huelsenbeck 2003) with the following

settings: four Markov chains were run for 2,000,000 generations with a sampling frequency of 100 generations. Log-likelihood values for sampled trees were stabilized after almost 200,000 generations. The last 18,000 trees were used to estimate Bayesian posterior probabilities, whereas the first 2001 were discarded as burn-in. Results from two-independent runs were used to construct a majority-rule consensus tree containing the posterior probabilities.

The sequences of *Coolia monotis* VGO783 (FN256433) and VGO786 (AM902737) were used as outgroups for the *Ostreopsis* ITS-5.8S and LSU gene phylogenetic analyses, respectively.

**Toxin analysis. Reagents for chemical analyses.** All organic solvents and water (HPLC grade) and glacial acetic acid (Laboratory grade) were by Sigma Aldrich (Steinheim, Germany). A palytoxin standard (100  $\mu\text{g}$ ; lot LAM7122) from Wako Chemicals GmbH (Neuss, Germany) was dissolved in methanol/water (1:1, v/v) and used for quantitative analyses. It should be noted that this standard was not certified and may have contained some contaminants other than the palytoxin and that the quali-quantitative composition of this standard can vary between lots. The standard used here contained 83% of palytoxin itself, 5% of 42-hydroxypalytoxin, and 12% of contaminant(s). A crude extract of Ligurian *O. cf. ovata* (CBA-29, Giussani et al. 2015) containing a pool of OVTXs was used as a reference sample for identification of OVTXs in algal extracts.

**Toxin extraction:** Cell pellets of five *Ostreopsis* strains collected along Lebanon coasts, namely L1002 ( $8.0 \times 10^6$  cells), L1007 ( $1.0 \times 10^7$  cells), L1008 ( $3.0 \times 10^6$  cells), L1020 ( $4.8 \times 10^6$  cells), L1022 ( $1.0 \times 10^6$  cells) were extracted once by adding 1–10 mL of methanol/water (1:1, v/v) to achieve a concentration of around  $1.0 \times 10^6 \text{ cells} \cdot \text{mL}^{-1}$  of extracting solvent. The mixtures were sonicated for 10 min in pulse mode, while cooling in ice bath, and centrifuged at  $3,000g$  for 1 min; the obtained supernatants were then decanted and analyzed by LC-HRMS (5  $\mu\text{L}$  injected).

**Liquid chromatography-high resolution multiple stage mass spectrometry:** MS experiments (positive ions) were carried out on a Dionex Ultimate 3000 quaternary system coupled to a hybrid linear ion trap LTQ Orbitrap XL<sup>TM</sup> Fourier Transform MS (FTMS) equipped with an ESI ION MAX<sup>TM</sup> source (Thermo-Fisher, San José, CA, USA). A Poroshell 120 EC-C18, 2.7  $\mu\text{m}$ ,  $2.1 \times 100 \text{ mm}$  column (Agilent Technologies, Santa Clara, CA, USA) maintained at room temperature was used. It was eluted at  $0.2 \text{ mL} \cdot \text{min}^{-1}$  with water (eluent A) and 95% acetonitrile/water (eluent B), both containing 30 mM acetic acid. A slow gradient elution was used: 28%–29% B over 5 min; 29%–30% B over 10 min; 30%–100% B over 1 min, and held for 5 min (Ciminiello et al. 2015).

HR full scan MS experiments (positive ions) were acquired in the range  $m/z$  800–1400 at a RP of 60,000 (FWHM at  $m/z$  400). The following source settings were used: a spray voltage of 4.8 kV, a capillary temperature of  $290^{\circ}\text{C}$ , a capillary voltage of 17 V, a sheath gas and an auxiliary gas flow of 32 and 4 (arbitrary units). The tube lens voltage was set at 145 V. HR collision induced dissociation MS<sup>2</sup> experiments were acquired at a RP = 60,000 using a collision energy = 35%, isolation width = 3.0 Da, activation Q = 0.250, and activation time = 30 ms. The most intense peak of the  $[\text{M}+\text{H}+\text{Ca}]^{3+}$  ion cluster of OVTX-a ( $m/z$  896.1) and OVTX-d/-e ( $m/z$  901.4) were used as precursors in HRMS<sup>2</sup> experiments. Calculation of elemental formulae was performed using the mono-isotopic peak of each ion cluster using Xcalibur software v. 2.0.7 with a mass tolerance constrain of 5 ppm. The isotopic pattern of each ion cluster was taken into account in assigning molecular formulae. Extracted ion chromatograms of the detected OVTXs were obtained by selecting the  $[\text{M}+\text{H}+\text{Ca}]^{3+}$  ion clusters, using a mass tolerance of 5 ppm and employed

TABLE 1. List of *Ostreopsis* spp. isolates, sampling locations, isolator, ITS - 5.8S, and LSU gene sequence accession numbers from GenBank and EMBL.

Species	Strain ID	Geographical origin and collecting period	Isolator	Accession no. ITS-5.8S	Accession no. LSU
<i>O. cf. ovata</i>	CBA166	Trieste, Italy, Adriatic Sea, Mediterranean, 2009	Penna A.	JX065557	
<i>O. cf. ovata</i>	CBA1823	Taormina, Italy, Ionian Sea, Mediterranean, 2010	Battocchi C.	JX065555	
<i>O. cf. ovata</i>	CBA1597	Marina di Pisa, Italy, Tyrrhenian Sea, Mediterranean, 2010	Casabianca S.	JX065554	
<i>O. cf. ovata</i>	CBA1502	Alghero, Italy, Tyrrhenian Sea, Mediterranean, 2010	Capellacci S.	JX065553	JX065562
<i>O. cf. ovata</i>	CBA1553	Villefranche, Ligurian Sea, Mediterranean, France, 2010	Battocchi C.	JX065556	JX065565
<i>O. cf. ovata</i>	CBA N	Tyrrhenian, Mediterranean, Italy, 2007	Capellacci S.	FM244631	FM946085
<i>O. cf. ovata</i>	VGO 820	Catalan, Mediterranean, 2005	Fraga S.	FM244634	FM994892
<i>O. cf. ovata</i>	VGO 822	Catalan, Mediterranean, 2005	Fraga S.		FM994893
<i>O. cf. ovata</i>	VGO 884	Catalan, Mediterranean, 2005	Fraga S.		FM994931
<i>O. cf. ovata</i>	VGO960	Llavaneres, Spain, Catalan Sea, Mediterranean, 2008	Fraga S.		JX065567
<i>O. cf. ovata</i>	VGOOS20BR	Rio de Janeiro, Brazil, W Atlantic Ocean, 2000	Fraga S.		FM997919
<i>O. cf. ovata</i>	VGO1001	Famara, Canary Island, Spain, E Atlantic Ocean, 2008	Rodriguez F.	JX065551	
<i>O. cf. ovata</i>	VGO1056	Belize, Caribbean Sea, N Atlantic Ocean, 2009	Holland C.	JX065586	JX065588
<i>O. cf. ovata</i>	KC34	Aegean, Mediterrean, 2004	Aligizaki K.	FM242104	FM946092
<i>O. cf. ovata</i>	KC71	Aegean, Mediterrean, 2004	Aligizaki K.	FM244735	FM946099
<i>O. cf. ovata</i>	OS18BR	West Atlantic, Brazil, 2000	Fraga S.	FM244670	FM997918
<i>O. cf. ovata</i>	CBA4	East Pacific, Indonesia, 2007	Penna A.		FM997921
<i>O. cf. ovata</i>	CBA9	East Pacific, Indonesia, 2007	Penna A.	FM244726	
<i>O. cf. ovata</i>	OvPD06	Indian, Malaysia, 1997	Pin L.C.	AF218455	
<i>O. cf. ovata</i>	OvSA04	Indian, Malaysia, 1997	Pin L.C.	AF218461	
<i>O. cf. ovata</i>	PR-03	Indian, Malaysia, 1997	Pin L.C.	AF076218	
<i>O. cf. ovata</i>	OvSA06	Indian, Malaysia, 1997	Pin L.C.	AF218463	
<i>O. cf. ovata</i>	CAWD174	Rarotonga, Cook Island, 2009	Strickland R. Sellwood A.	AB674904	
<i>O. cf. ovata</i>	KabO13	Okinawa,Ishigaki, Kabira, Japan 2010	Suda S.		AB605817
<i>O. cf. ovata</i>	IshiOst50	Okinawa,Ishigaki, Shiraho,Japan, 2009	Shah M.R.		AB605824
<i>O. cf. ovata</i>	QB04	Quang Binh, Vietnam, South China Sea, Pacific, 2009	Nguyen N.L.		JX065571
<i>O. cf. ovata</i>	QB06	Quang Binh, Vietnam, South China Sea, Pacific, 2009	Nguyen N.L.		KC900891
<i>O. cf. ovata</i>	QB03	Quang Binh, Vietnam, South China Sea, Pacific, 2009	Nguyen N.L.		KC900890
<i>O. cf. ovata</i>	LCH001	Thua Thien-Hue, Vietnam South China Sea, Pacific, 2009	Nguyen N.L.		JX065569
<i>O. cf. siamensis</i>	IO-9601	Sines, Portugal, E Atlantic, 2008	Veloso V.	JX065587	
<i>O. cf. siamensis</i>	OS-2V	Alboran, Mediterranean, 1999	Fraga S.	AJ491313	
<i>O. cf. siamensis</i>	OS-5V	Alboran, Mediterranean, 1999	Fraga S.		FN256430
<i>O. cf. siamensis</i>	CSIC-D7	Catalan, Mediterranean, 2000	Garcés E.	AJ491334	FN256431
<i>O. cf. siamensis</i>	CNR-B4	Tyrrhenian Sea, Mediterranean, 2000	Giacobbe M.G.	AJ301643	
<i>O. cf. siamensis</i>	CAWD96	Kerikeri, New Zealand, 1999	North Health Services	AB674915	
<i>O. cf. siamensis</i>	Dn17EHU	Biscay, Spain, 2007–2009	Laza A.		HQ414225
<i>O. cf. siamensis</i>	Dn 20EHU	Biscay, Spain, 2007–2009	Laza A.		HQ414224
<i>O. cf. siamensis</i>	Dn18EHU	Biscay, Spain, 2007–2009	Laza A.		HQ414222
<i>O. cf. siamensis</i>	Dn171EHU	Saint Jean de Luz, France, 2010	David H.	JX987690	
<i>O. cf. labens</i>	VGO897	Indian, Malaysia, 1997	Mohammad N.	FM244728	
<i>O. cf. lenticularis</i>	NT011	Ninh Thuan, Vietnam, South China Sea, 2009	Nguyen N.L.	JX065584	
<i>O. cf. lenticularis</i>	NT013	Ninh Thuan, Vietnam, South China Sea, 2009	Nguyen N.L.		JX065570
<i>O. cf. lenticularis</i>	OLPR01	Indian, Malaysia, 1997	Pin L.C.	AF218465	AF244941
<i>Ostreopsis</i> sp.	TB34OS	Khao Lak, Phang-Nga, Thailand, 2015	Tawong W.	AB841214	
<i>Ostreopsis</i> sp.	TB35OS	Khao Lak, Phang-Nga, Thailand, 2015	Tawong W.	AB841215	
<i>Ostreopsis</i> sp.	TB33OS	Khao Lak, Phang-Nga,Thailand, 2015	Tawong W.	AB841213	
<i>Ostreopsis</i> sp.	KC84	Cyprus, Aegean Sea, Mediterranean, 2008	Aligizaki K.	JX065549	JX065558
<i>Ostreopsis</i> sp.	KC86	Crete, Greece, Aegean Sea, Mediterranean, 2009	Aligizaki K.	JX065550	JX065559
<i>Ostreopsis</i> sp.	CBA0203	Honolulu, Hawaii, N Pacific Ocean, USA, 2010	Capellacci S.	JX065552	JX065561
<i>Ostreopsis</i> sp.	VGO881	Canary Island, East Atlantic, 2005	Fraga S.	FM244637	FM994895
<i>Ostreopsis</i> sp.	Dn83EHU	Crete, Greece, 2010	David H.	JX987673	
<i>Ostreopsis</i> sp.	Dn110EHU	Puerto Rico, North America 2011	David H.	JX987680	
<i>Ostreopsis</i> sp.	TF29OS	Koh Wai, Trat, Thailand, 2011	Tomohiro N.	AB841255	
<i>Ostreopsis</i> sp.	TF25OS	Koh Wai, Trat, Thailand, 2011	Tomohiro N.	AB841254	
<i>O. fattorusoi</i>	CBA C1005	Cyprus, Aegean Sea, Mediterranean, 2013	Capellacci S.	LT220222	
<i>O. fattorusoi</i>	CBA C1012	Cyprus, Aegean Sea, Mediterranean, 2013	Capellacci S.	LN875554	
<i>O. fattorusoi</i>	CBA C1019	Cyprus, Aegean Sea, Mediterranean, 2013	Capellacci S.	LN875552	LT555465
<i>O. fattorusoi</i>	CBA C1020	Cyprus, Aegean Sea, Mediterranean, 2013	Capellacci S.	LN875556	LT555466

(continued)

TABLE 1. (continued)

Species	Strain ID	Geographical origin and collecting period	Isolator	Accession no. ITS-5.8S	Accession no. LSU
<i>O. fattorussoi</i>	CBA C1035	Cyprus, Aegean Sea, Mediterranean, 2013	Capellacci S.	LN875553	
<i>O. fattorussoi</i>	CBA C1036	Cyprus, Aegean Sea, Mediterranean, 2013	Capellacci S.	LN875557	
<i>O. fattorussoi</i>	CBA L1000	Batroun, Lebanon, Mediterranean, 2014	Capellacci S.	LT220223	
<i>O. fattorussoi</i>	CBA L1020	Batroun, Lebanon, Mediterranean, 2014	Capellacci S.	LT220224	LT555467
<i>O. fattorussoi</i>	CBA L1007	Batroun, Lebanon, Mediterranean, 2014	Capellacci S.		LT555468
<i>O. fattorussoi</i>	CBA L1008	Batroun, Lebanon, Mediterranean, 2014	Capellacci S.		LT555469
<i>O. fattorussoi</i>	CBA L1022	Batroun, Lebanon, Mediterranean, 2014	Capellacci S.		LT555470
<i>Ostreopsis</i> sp.	UrGt12	Okinawa, Urasoe, off camp Kinser, Japan, 2009	Suda S.		AB605813
<i>Ostreopsis</i> sp.	UrGm6	Okinawa, Urasoe, off camp Kinser, Japan, 2009	Suda S.		AB605815
<i>Ostreopsis</i> sp.	IkeOst2	Ikejima, Uruma, Okinawa, Japan 2008	Suda S.		AB605814
<i>Ostreopsis</i> sp.	OA21-C10	Okinawajima Island, Japan	Nakashima A.		AB605828

in quantitative analyses. Due to availability of the only palytoxin standard, quantitative determination of OVTX-a, -d, and -e in the extracts was carried out by using a calibration curve (triplicate injection) of palytoxin standard at seven concentrations (1,000, 500, 250, 125, 62.5, 31.2, and 15.6 ng · mL<sup>-1</sup>). OVTX molar responses were assumed to be similar to that of PLTX. Calibration curve equation was  $y = 2,455.6x - 19,184$  and its linearity was expressed by  $R^2 = 0.9976$ . The limit of detection for palytoxin in pure solvent was 13 ng · mL<sup>-1</sup> after correction for the 83% purity of the standard.

## RESULTS

***Ostreopsis fattorussoi* Accoroni, Romagnoli et Totti sp. nov.**

**Diagnosis.** Cells are ovate in shape and ventrally pointed with a DV of 42.5–72.5 µm, AP of 20–32.5 µm and W 26.3–50 µm. The thecal plate formula is Po, 3', 7'', 6c, 7s, 5''', 2'''''. Thecal plates are smooth with evenly distributed same-sized round pores. The apex is strongly eccentric, located on the left dorsal side of the epitheca. The apical pore plate Po is 10–12.5 µm long and slightly curved. Plate 1' is heptagonal, oblique, and is almost entirely in the left half of the epitheca. Plate 3' is hexagonal, almost entirely in the left half of the epitheca. The pentagonal 6'' has the 6''/5''' suture length almost twice as long as the 6''/7''' suture length. The narrow sulcal groove runs obliquely from the left side of the ventral area into the hypotheca. Cells are photosynthetic. The nucleus has a slightly elongated (subspherical) shape positioned obliquely and occupies the dorsal part of the cell. The species is toxic producing ovatoxins.

**Holotype.** SEM stub n. 3/16-UNIVPM deposited at the Botanical Museum of the Università Politecnica delle Marche, Ancona Italy, in the Herbarium Anconitanum (ANC). Figure 5A represents the holotype.

**Isotype.** Preserved sample, deposited at the Università Politecnica delle Marche, Ancona Italy, in the Herbarium Anconitanum (ANC).

Preserved DNA of clonal strain CBA-L1020, bar-coded in ENA-EMBL (EMBL ID: LT220224), held

at the Dipartimento di Scienze Biomolecolari, Università di Urbino, Italy.

**Type Locality.** Batroun, Lebanon (34°15' 5.40" N, 35°39' 24.78" E, Fig. 1).

**Habitat.** Benthic, often epiphytic on seaweeds or living on other substrata; it was shown that it could also be resuspended into the water column.

**Etymology.** The specific epithet honors our dear colleague Prof. Ernesto Fattorusso from the University of Napoli Federico II, who significantly contributed to the study of algal biotoxin structures and elucidation of novel organic metabolites produced by marine algae.

**Distribution:** Eastern Mediterranean (i.e., Cyprus, Lebanon coasts). The same genotype is known also in Crete (David et al. 2013, Penna et al. 2014), Canary Islands, Spain (Penna et al. 2010) and Puerto Rico, USA (David et al. 2013).

**Morphology.** Cells are ovate and ventrally slender with average DV  $60.1 \pm 5.6$  (42.5–72.5) µm, AP  $25.7 \pm 3$  (20–32.5) µm, and W  $39.8 \pm 5.1$  (26.3–50) µm. The DV:AP ratio is  $2.35 \pm 0.22$ , while DV:W is  $1.52 \pm 0.14$ .

The thecal plate formula is Po, 3', 7'', 6c, 7s, 5''', 2'''''. (Figs. 2 and 3). Thecal plates are smooth with evenly distributed round pores visible using LM with Calcofluor-White M2R staining (Fig. 4A). Small perforations are located inside the pores (Fig. 5E). They are scattered on the epi- and hypothecal plates and lined up along the border of the pre- and postcingular plates close to the cingular groove (Fig. 5, A and B) and the borders of the two cingular lists (Fig. 5D). Only one size class pore is visible ranging from 0.26 to 0.53 µm ( $0.38 \pm 0.08$  µm) with 11–15 pores per 100 µm<sup>2</sup>. Cells produce extracellular mucilage and a complex network of filaments is extruded through the thecal pores (Fig. 6D).

Both epitheca and hypotheca are equal in size. The apex is strongly eccentric, located on the left dorsal side of the epitheca. The apical pore plate Po is  $11.67 \pm 1.44$  µm long, slight curved and contacts the three apical plates. Plate 1' is heptagonal

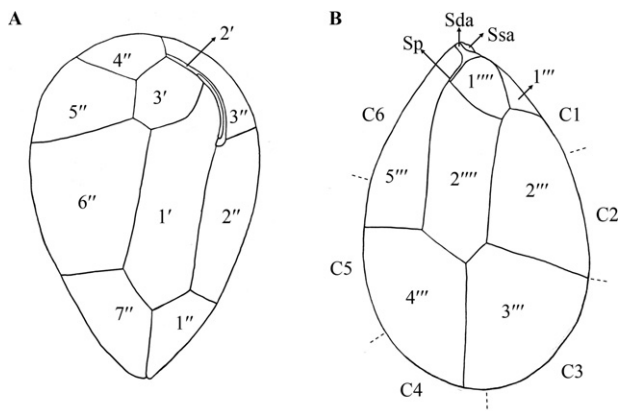


FIG. 2. *Ostreopsis fattorusoi*. Schematic drawings with plate tabulation of (A) epithea, (B) hypotheca and cingulum. Sda, right-anterior sulcal; Ssa, left anterior sulcal plate; Sp, posterior sulcal plate.

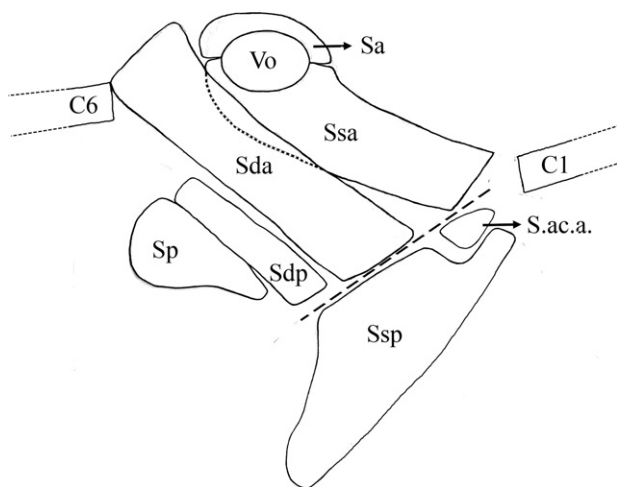


FIG. 3. Interpretation drawing of the sulcus of *Ostreopsis fattorusoi*. Sa, anterior sulcal plate; Ssa, left anterior sulcal plate; Sda, right-anterior sulcal plate; Sdp, right-posterior sulcal plate; Sp, posterior sulcal plate; Ssp, left posterior sulcal plate; S.ac.a., accessory anterior sulcal plate; dotted line indicates the position of the bottom of the sulcus (Sa, Ssa, Sda, Sdp, and Sp are in the upper side of the sulcus groove and Ssp and S.ac.a. are in the inferior side of the sulcus); Vo, ventral opening.

and almost entirely lies in the left half of the epithea and is not parallel to the dorsoventral axis, but its dorsal part is shifted on the left and dorsally pointed. Plate 1' touches plates 2', 3', 1'', 2'', 6'', and 7''. Plate 2' is narrow and almost twice the size of Po, separating the 3' and 3'' plates. Plate 3' is hexagonal, almost entirely located in the left half of the epithea, and it is dorsally displaced.

Among the seven irregularly quadrangular (with the only exception of the plate 6'') precingular plates 1'', and 4'' are the smallest. Because of the left and dorsal displacement of 3', plate 5'' is transversally elongated and 3'' appears as narrow and

dorsally elongated as plate 2'' (Figs. 4, A and C; 5A). The characteristic shape of the pentagonal 6'' is due to the oblique orientation of the 1' and the 6''/5'' suture length is almost twice as long as 6''/7'' suture length, resulting in an oblique 6''/1' suture; moreover, its length:width ratio is  $1.06 \pm 0.11$  (0.94–1.2) and the 3'/6'' suture is dorsally shifted. The first and the last precingular plates (i.e., 1'' and 7'', respectively) are extended and elongated, tapering toward the sulcal area. Plate 2'' is narrow and the 4'' has a similar size of 3'.

The cingulum is descending and displaced one time its width and consists of six plates, almost of the same length.

The narrow sulcal groove runs obliquely from the left side of the ventral area into the hypotheca. The ventral side of the 1'''' plate forms a "wing" that covers almost all the sulcal area of the hypotheca (Fig. 4, D–F) leading well identifiable only 5 plates in intact cells (Fig. 6A): the anterior sulcal plate (Sa), the left anterior sulcal plate (Ssa), the right-anterior sulcal plate (Sda), the right-posterior sulcal plate (Sdp), and the posterior sulcal plate (Sp). In broken cells, a further two plates are identifiable on the inner left side of the sulcus: the left posterior sulcal plate (Ssp) and the accessory anterior sulcal plate (S.ac.a.) (Fig. 6, B and C).

In the upper part of the ventral area, a conspicuous tube-like structure, the ventral tube (Vt), ends with the ventral opening (Vo, Figs. 4D and 6, A and B). The upper half of the Vt is surrounded by the small Sa and the lower half by the elongated Ssa. The Sa is small and touches the ventral part of the 1'' and 7'' plates. The Ssa plate is wide and extends from the lower side of the Sa and Vt to the first cingular plate (C1) and the S.ac.a. plate. The left (i.e., the anterior) margin of the Ssa plate touches the 1'' plate, while the right (i.e., the posterior) margin is in contact with the anterior, sigmoid edge of the Sda plate.

The Sda plate is the most elongated of all the sulcal plates. Its anterior part is curved toward the epithea and contacts, from right to left, the sixth cingular (C6), the 7'' and the Sa plates. It surrounds the right (i.e., the posterior) margin of plate Ssa along its whole side. The left portion of the Sda plate is hidden by the 1'''' and contacts the Ssp plate in the bottom of the sulcal groove. At its posterior part, Sda contacts the ventral end of the 5'' and the narrow Sdp.

In the end of the sulcal groove, the right-anterior part of the irregularly triangular Sp touches the ventral end of the 5'' plate, and its right-posterior part adjoins to a small portion of the ventral end of the 2'''' plate (Fig. 4E). The upper part of the Sp connects with the Sdp and in its posterior part to the Ssp.

The other side of the sulcal groove (and therefore completely covered by the 1'''' plate) consists of the almost quadrangular S.ac.a. plate for the small

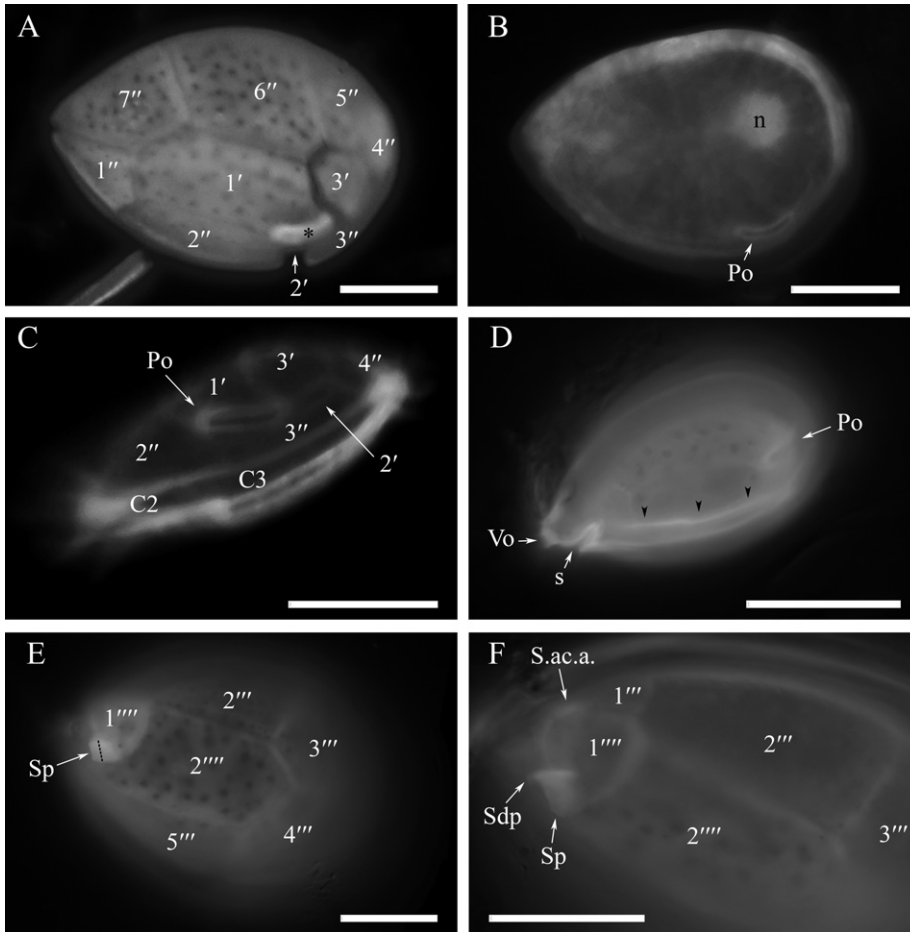


FIG. 4. *Ostreopsis fattorussoi*, light microscopy micrographs. Epifluorescence observations after Calcofluor-White of cells from natural samples. (A) Epitheca (\* = apical pore plate Po). (B) Apical view of a cell stained with SYBR Green highlighting the nucleus (apical pore plate Po is visible). (C) Left lateral view showing the contact of plate 2' with 4''. (D) Merging of several photograms taken at different foci of the left lateral side of a cell showing the depth of the sulcus (s) and the ventral opening (Vo); arrowheads indicate the cingulum border in the epitheca. (E, F) Hypotheca; (E) focus on the posterior sulcal plate (Sp): although the Sp is almost entirely covered by the plate 1'''' (dotted line indicates the position of the 1'''' border on the Sp plate), the focus allows observing the entire plate and its small contact with plate 2''''; (F) focus on the plates Sdp and S.ac.a. under the 1'''''. Scale bars = 20  $\mu$ m.

upper part (Fig. 6C) and the elongated Ssp makes up rest of the groove.

Among the five postingular plates, the triangular 1''' plate is the smallest of the series. The 2''' plate is wide and quadrangular. The quadrangular 3''' and 4''' plates occupy most of the dorsal part of the hypotheca, while the 5''' plate is oblong and quadrangular, although the ventral side is very short. There are two antapical plates which are both pentagonal. The 1'''' plate is much smaller than the 2'''' plate which occupies the center of the hypotheca (Figs. 4, E and F; 5B).

There are numerous elongated chloroplasts located within the cell periphery. The nucleus, has a slightly elongated (often subspherical) shape ( $13.6 \pm 2.0 \mu\text{m}$  long,  $8.8 \pm 1.7 \mu\text{m}$  wide) positioned obliquely in the dorsal part (from the right to the center) of the cell (Fig. 4B).

**Phylogenetic analyses of *Ostreopsis* ITS-5.8S and LSU ribosomal genes.** The final alignments of *Ostreopsis* spp. ribosomal gene sequences, as ITS-5.8S and LSU with *Coolia monotis* VGO783 and VGO786 as outgroups, respectively, were as follows: ITS-5.8S was 419 bp in length (A = 27%, T = 34.8%, C = 18.2%, G = 20%) with 116 polymorphic sites, of which 93 parsimony informative, and a transition/

transversion ratio of 1.7; LSU was 701 bp in length (A = 28.5%, T = 32.9%, C = 16.9%, G = 21.7%) with 343 polymorphic sites, of which 284 parsimony sites, and a transition/transversion ratio of 1.4.

Based on single ITS-5.8S and LSU rDNA sequences, only minor differences between the NJ, MP, ML and Bayesian inference analyses were found; therefore, only the ML phylogenetic trees are presented (Figs. 7 and 8). The ITS-5.8S rDNA phylogeny, based on 46 isolates of *Ostreopsis* spp., identified three major clades within the genus *Ostreopsis*: the first comprising *Ostreopsis* sp. CBA0203 from Hawaii, and strains identified as *O. cf. lenticularis*, *O. cf. labens* and *Ostreopsis* sp. 6 (Tawong et al. 2014) from Pacific Ocean; the second clade comprised the *O. cf. siamensis*, and *Ostreopsis* sp. (now identified as *O. fattorussoi*) from eastern Mediterranean Sea, Greece (KC84, KC86, Penna et al. 2014 and Dn83EHU, David et al. 2013), Lebanon and Cyprus (this study), with *Ostreopsis* sp. VGO881 (Canary Islands) and *Ostreopsis* sp. Dn110EHU (Porto Rico) from Atlantic Ocean. Finally, the third grouping included *O. cf. ovata* complex from Atlantic, Mediterranean, and Pacific. All these clusters were supported by high bootstrap and posterior probability values.



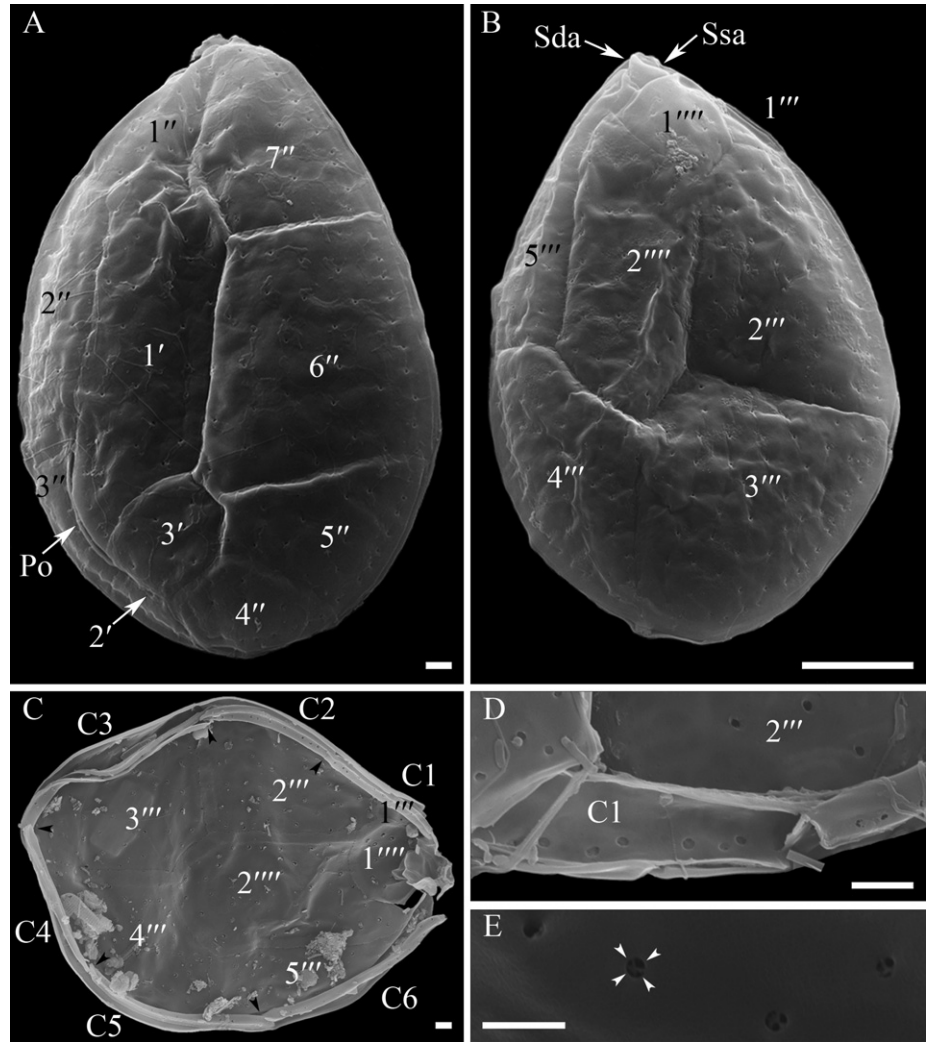


FIG. 5. *Ostreopsis fattorussoi*, SEM micrographs. (A) Epitheca. (B) Hypotheca. (C) Internal view of the hypotheca showing the cingular plates with distinctive borders (arrowheads) between two adjacent plates. (D, E) Magnification on the internal side of the hypotheca showing the thecal pores; (D) pores lined up along both the borders of the two cingular lists; (E) small perforations inside thecal pores. Po, apical pore plate; Sda, right-anterior sulcal; Ssa, left anterior sulcal plate. Scale bars = 2  $\mu\text{m}$  (A, C, D, and E); 10  $\mu\text{m}$  (B).

The LSU rDNA phylogeny that was obtained from 40 isolates of *Ostreopsis* spp. showed some differences in tree topology compared with ITS-5.8S rDNA phylogenetic analysis. The first splitting clade from outgroup *Coolia* included two sub-clades of *O. cf. lenticularis* and *Ostreopsis* sp. along with *Ostreopsis* sp. 5 and *Ostreopsis* sp. CBA0203; all these isolates derived from Indian and Pacific Ocean. The second clade grouped the *Ostreopsis* sp. (now identified as *O. fattorussoi*) from eastern Mediterranean Sea (Lebanon, Greece and Cyprus) and Atlantic VGO881 isolate. The third clade comprised all *O. cf. ovata* isolates collected from many sites worldwide and *O. cf. siamensis*. All these lineages were strongly supported by high bootstrap and posterior probability values.

The difference of base pairs between *O. fattorussoi* and other *Ostreopsis* species ranged from 200 to 231 bp, and from 99 to 82 bp based on LSU and ITS-5.8S rDNA analysis of net nucleotide average difference, respectively.

*Temporal trend and relationships with environmental parameters.* *Ostreopsis* cells were detected at the

Batroun site throughout the year with two blooming periods, the first observed between the end of May and the beginning of July and the second in August (Fig. 9B). Maximum abundances were detected on 2 July 2015 ( $28 \times 10^3 \text{ cells} \cdot \text{g}^{-1} \text{ fw}$ ) and on 22 June 2015 ( $840 \text{ cells} \cdot \text{L}^{-1}$ ) on benthic substrata and in the water column, respectively.

The correlation between *Ostreopsis* abundances on benthic substrata and those in the water column was positive and significant ( $n = 9$ ;  $r = 0.79$ ;  $P < 0.05$ ).

Surface temperature ranged from 18°C to 31.5°C with maximum temperatures of 29.5°C to 31.5°C occurring from the end of July to early October and minimum values of around 18°C in the first months of the year. The maximum values of *Ostreopsis* abundances were recorded when water temperature ranged from 27°C to 29.7°C (Fig. 9B). Throughout the year, the salinity ranged from 38.14 to 39.43. Correlation analyses among *Ostreopsis* abundances, growth rate, temperature, and salinity revealed no significant relationships.

No obvious trends were determined for of  $\text{NO}_2$ ,  $\text{NO}_3$  and  $\text{PO}_4$  (Fig. 9A). Only  $\text{NO}_3$  showed high

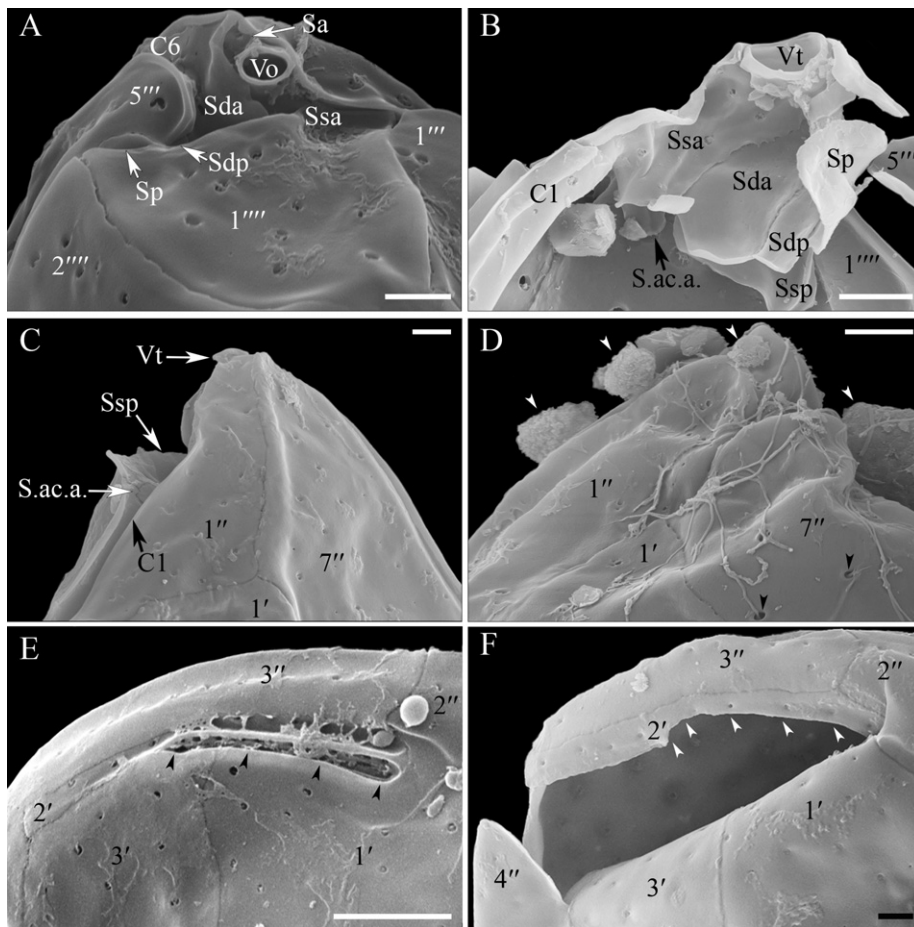


FIG. 6. *Ostreopsis fattorussoi*, SEM micrographs. (A) Ventral area in an intact cell. (B) Internal view of the hypotheca and the sulcus in a broken cell showing all the sulcal plates (with the exception of Sa): Ssa, Sda, Sdp, and Sp are in the upper side of the sulcus groove and partially cover Ssp and S.ac.a. which are in the inferior side of the sulcus; Sp is unnaturally bended leading show the space between the 1''' and 5''' plates. (C) Left lateral view of the ventral area partially showing the inferior side of the sulcus. (D) Apical ventral view of a cell showing a complex network of filaments released by the thecal pores (black arrowheads) and extracellular mucilage (white arrowheads). (E, F) Magnification on the apical dorsal part showing the apical pore plate Po; (E) intact Po (black arrowheads) and the plate 2'; (F) suture between the plate 2' and the Po which is missing (white arrowheads). Scale bars = 2  $\mu\text{m}$  (A, B, C, D, and F); 5  $\mu\text{m}$  (E).

values ( $>1.3 \mu\text{mol} \cdot \text{L}^{-1}$ ) in January and February compared with the rest of the year, when values ranged from 0.156 to  $0.626 \mu\text{mol} \cdot \text{L}^{-1}$ .  $\text{NO}_2$  concentrations were markedly lower than those of nitrates, ranging from 0.015 to  $0.250 \mu\text{mol} \cdot \text{L}^{-1}$ . No significant relationships were found between *Ostreopsis* abundances (or growth rate) and both  $\text{NO}_2$  and  $\text{NO}_3$ .  $\text{PO}_4$  concentrations ranged from 0.015 to  $0.433 \mu\text{mol} \cdot \text{L}^{-1}$  and rarely exceeded  $0.150 \mu\text{mol} \cdot \text{L}^{-1}$  throughout the year. The two maximum values of  $\text{PO}_4^{3-}$ , i.e., 0.433 and  $0.175 \mu\text{mol} \cdot \text{L}^{-1}$ , were recorded in February and June, respectively. Although no correlation between *Ostreopsis* abundances and  $\text{PO}_4$  concentrations was detected, a positive and significant correlation was found between *Ostreopsis* growth rates and  $\text{PO}_4$  concentrations ( $n = 9$ ;  $r = 0.87$ ;  $P < 0.01$ ).

**Toxicity.** Crude extracts of five strains of *O. fattorussoi* collected along the Lebanese coast (i.e., L1002, L1007, L1008, L1020, and L1022) were analyzed by LC-HRMS<sup>n</sup> ( $n = 1, 2$ ) to characterize their toxin profiles and measure their toxin content. The analyses were carried out with a palytoxin standard and a Ligurian *O. cf. ovata* extract containing OVTX-a, -d and -e and isobaric palytoxin (Giussani et al. 2015). The chromatographic conditions

allowed the separation of all of the analogs contained in the extracts.

Toxin extracts from the five Lebanese strain of *O. fattorussoi* revealed three strains to be toxic; L1007, L1020, and L1022 all contained OVTX-a, -d, and -e. Their identity was ascertained based on comparison of their retention times and associated full MS spectra (Fig. 10) with those of OVTX-a, -d, -e, contained in the reference extract. Further confirmation for the identity of OVTX-a and of the structural isomers OVTX-d and -e was provided by their LC-HRMS<sup>2</sup> spectra.

Extracts L1002 and L1008 did not contain any palytoxin congener. Considering the extraction volume, the presence of toxins in pellet ( $1 \times 10^6$  cells), could be excluded at levels  $\leq 15 \text{ fg} \cdot \text{cell}^{-1}$ .

The results of the quantitative analysis are shown in Table 2.

## DISCUSSION

In the Mediterranean Sea, only two *Ostreopsis* species have been recorded until now, *O. cf. ovata* and *O. cf. siamensis* (Vila et al. 2001, Penna et al. 2005, 2010, Battocchi et al. 2010, Totti et al. 2010, Mangialajo et al. 2011, Perini et al. 2011, Mabrouk et al.

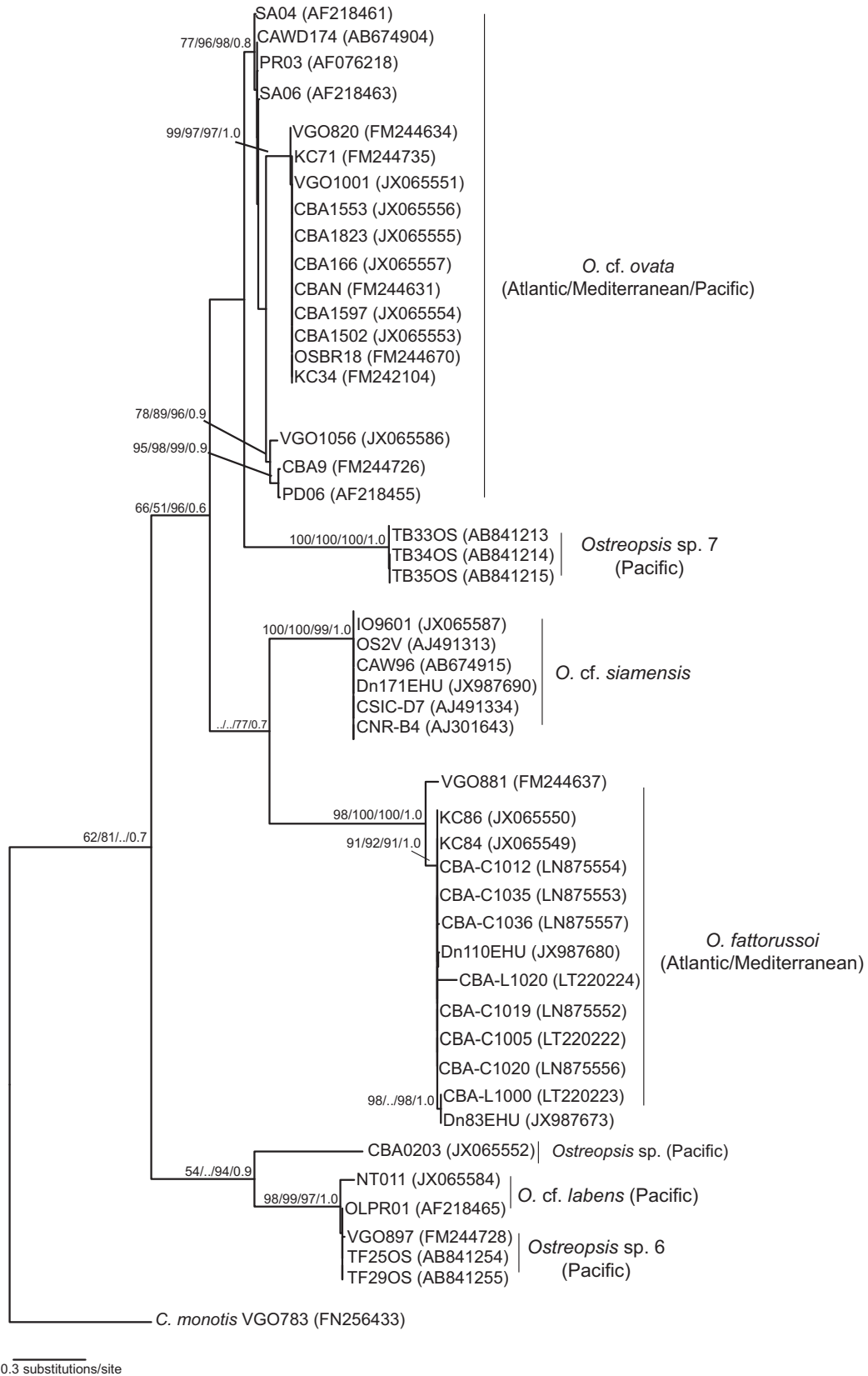


FIG. 7. Maximum likelihood phylogenetic tree of the genus *Ostreopsis* inferred from ITS-5.8S ribosomal gene sequences. The tree is rooted with *Coolia monotis* VGO783 as outgroup. Numbers on the major nodes represent from left to right NJ (1,000 pseudo-replicates), MP (1,000 pseudo-replicates), ML (1,000 pseudo-replicates) bootstrap and Bayesian posterior probability values. Only bootstrap values >50% are shown. Geographical origins of *Ostreopsis* isolates are indicated.

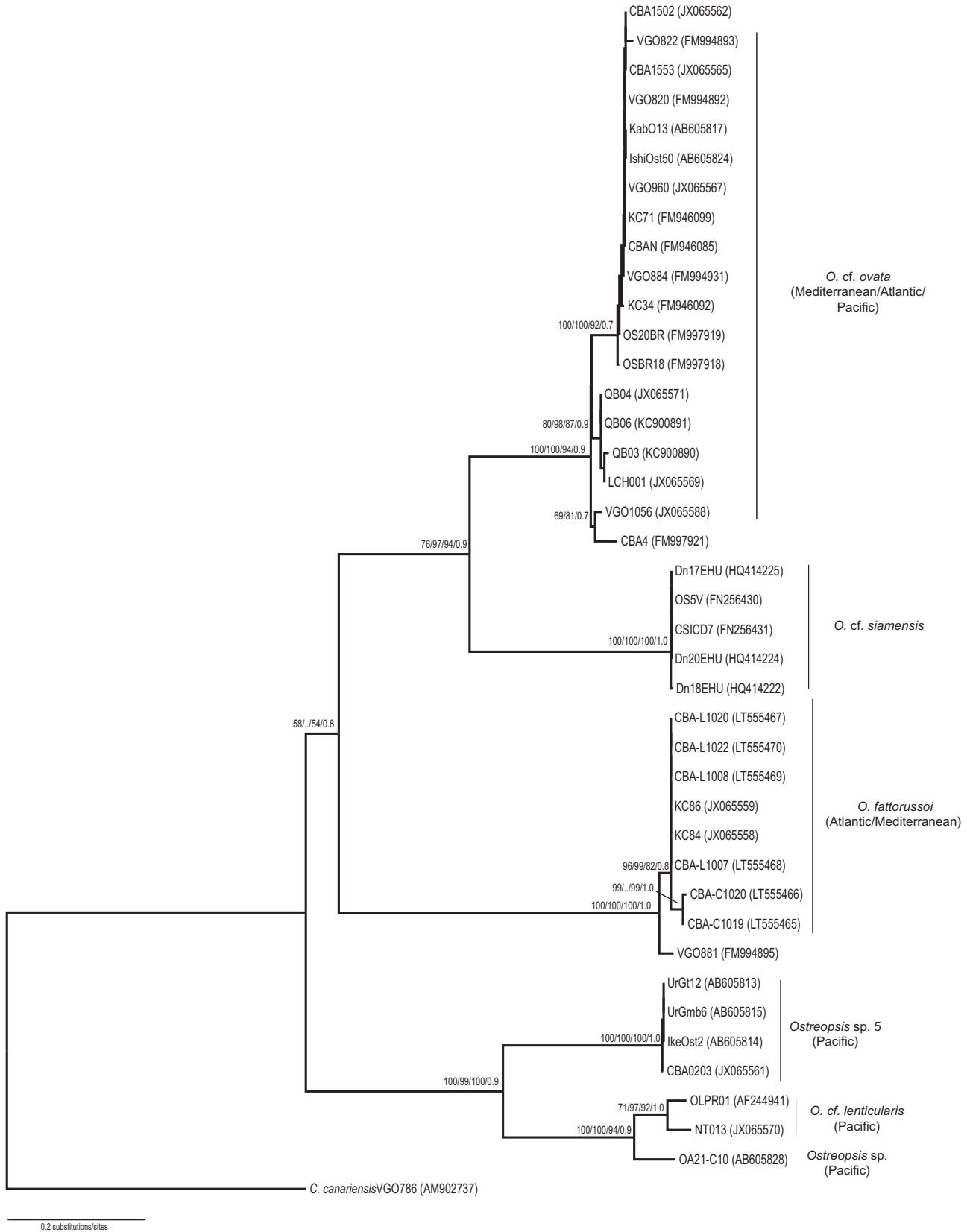


FIG. 8. Maximum likelihood phylogenetic tree of the genus *Ostreopsis* inferred from LSU ribosomal gene sequences. The tree is rooted with *Coolia monotis* VGO786 as outgroup. Numbers on the major nodes represent from left to right NJ (1,000 pseudo-replicates), MP (1,000 pseudo-replicates), ML (1,000 pseudo-replicates) bootstrap and Bayesian posterior probability values. Only bootstrap values >50% are shown. Geographical origins of *Ostreopsis* isolates are indicated.

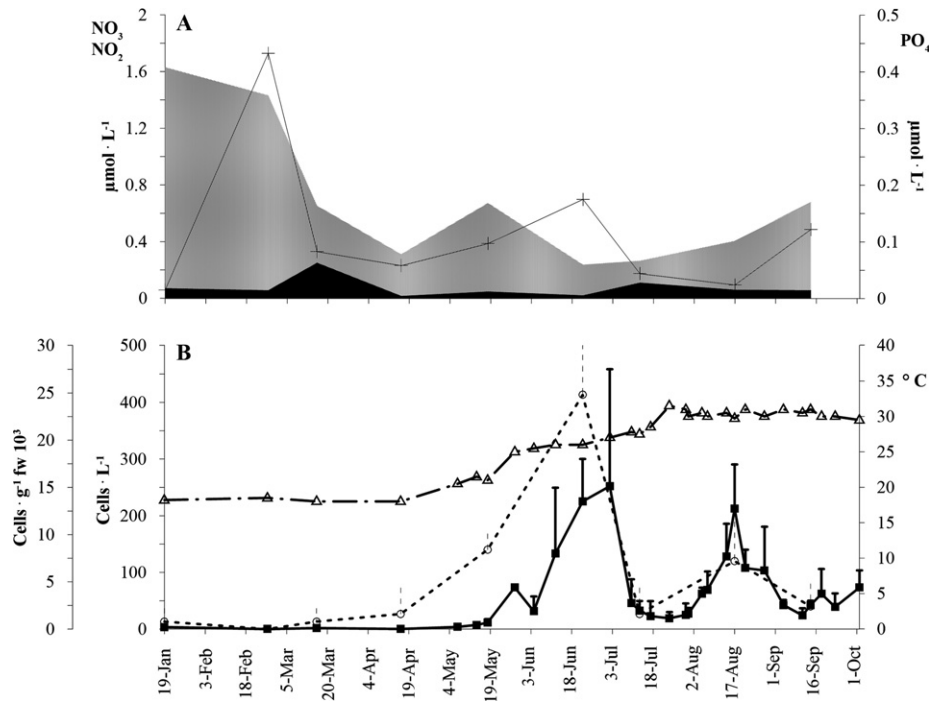


FIG. 9. Trend in nutrient concentration, temperature and *Ostreopsis fattorussoi* abundances both on benthic macroalgae and in water column in Batroun (Lebanon) during 2015. (A) Nutrient concentration ( $\mu\text{mol} \cdot \text{L}^{-1}$ ): inorganic nitrogen with the detail of  $\text{NO}_3$  and  $\text{NO}_2$  (gray and black, respectively, left y-axis), and  $\text{PO}_4$  (line, right y-axis). (B) *O. fattorussoi* abundances on seaweeds ( $\blacksquare$ , 1st left y-axis) and in the water column ( $\circ$ , 2nd left y-axis) expressed in  $\text{cells} \cdot \text{g}^{-1} \text{fw}$  and  $\text{cells} \cdot \text{mL}^{-1}$ , respectively. Temperature values expressed in  $^{\circ}\text{C}$  ( $\blacktriangle$ , right y-axis). Error bars indicate standard deviations.

2012, Abboud-Abi Saab et al. 2013, Accoroni and Totti 2016). Recently, new molecular phylogenetic analyses based on ribosomal DNA identified another clade (*Ostreopsis* sp.) distinct from *O. cf. ovata* and *O. cf. siamensis*, in the eastern Atlantic and Mediterranean coasts, particularly from islands of Cyprus and Crete (Penna et al. 2012, Tartaglione et al. 2016). In this study, we name and describe this new genotype as a new species of *Ostreopsis*, named *O. fattorussoi*, by analyzing samples from coasts of Lebanon and Cyprus (Mediterranean Sea).

In this study, the description of the thecal tabulation and the morphometrical analyses of this new species were performed on field samples only, as cultured specimens of *O. fattorussoi* frequently demonstrated modifications in shape, size and thecal pattern, as already reported in other *Ostreopsis* species (Norris et al. 1985, Laza-Martinez et al. 2011, Nascimento et al. 2012, Scalco et al. 2012, David et al. 2013). For example, a cultured Indonesian strain of *O. cf. ovata* (CBA-6) has been observed either to show (Parsons et al. 2012) or not (Penna et al. 2010) the contact between the  $1'$  and  $5''$  plates, which is a defining plate pattern character of *O. heptagona*. The numerous strains of *O. fattorussoi* were used to characterize the toxic profile and to perform the molecular sequencing of this new species.

The plate tabulation of *Ostreopsis* has been frequently reassessed over the years from its first

descriptions (Table 3). The first interpretations of the thecal plates were given by Schmidt (1901) and later by Fukuyo (1981), who included a partial plate formula of  $\text{Po}$ ,  $3'$ ,  $7''$ ,  $4'''$  (+ accessory small plates), 1 antapical plate and  $\text{Po}$ ,  $3'$ ,  $7''$ ,  $5'''$ ,  $1''''$ , respectively. This plate pattern interpretation based on Kofoidian plate tabulation was adopted by Norris et al. (1985) and Faust et al. (1996) who, however, labeled the large hypothetical plate at the antapex (i.e.,  $1''''$  in Fukuyo (1981)) as a posterior intercalary plate (1p). Over the years, several other authors followed this plate pattern interpretation, although often with a little variation (Penna et al. 2005, Sato et al. 2011, Kang et al. 2013, Hoppenrath et al. 2014).

On the contrary, Besada et al. (1982) did not follow the plate pattern suggested by Fukuyo (1981) but reinterpreted the apical plate pattern, providing for the first time a complete plate formula ( $\text{Po}$ ,  $4'$ ,  $6''$ ,  $6c$ ,  $8s$ ,  $5'''$ ,  $2''''$ ) which was closer to that of gonyaulacoid dinoflagellates. In this view, the epithelial tabulation was strongly rearranged as the plate considered as  $1''$  (precingular) in Fukuyo (1981) and Faust et al. (1996) was considered homologous to the first apical plate of the Gonyaulacales and therefore was named  $1'$ . As a consequence, the plate tabulation included four apical and six precingular plates. This interpretation has been re-adopted by Escalera et al. (2014) for *Ostreopsis cf. ovata* and was applied to *Gambierdiscus*

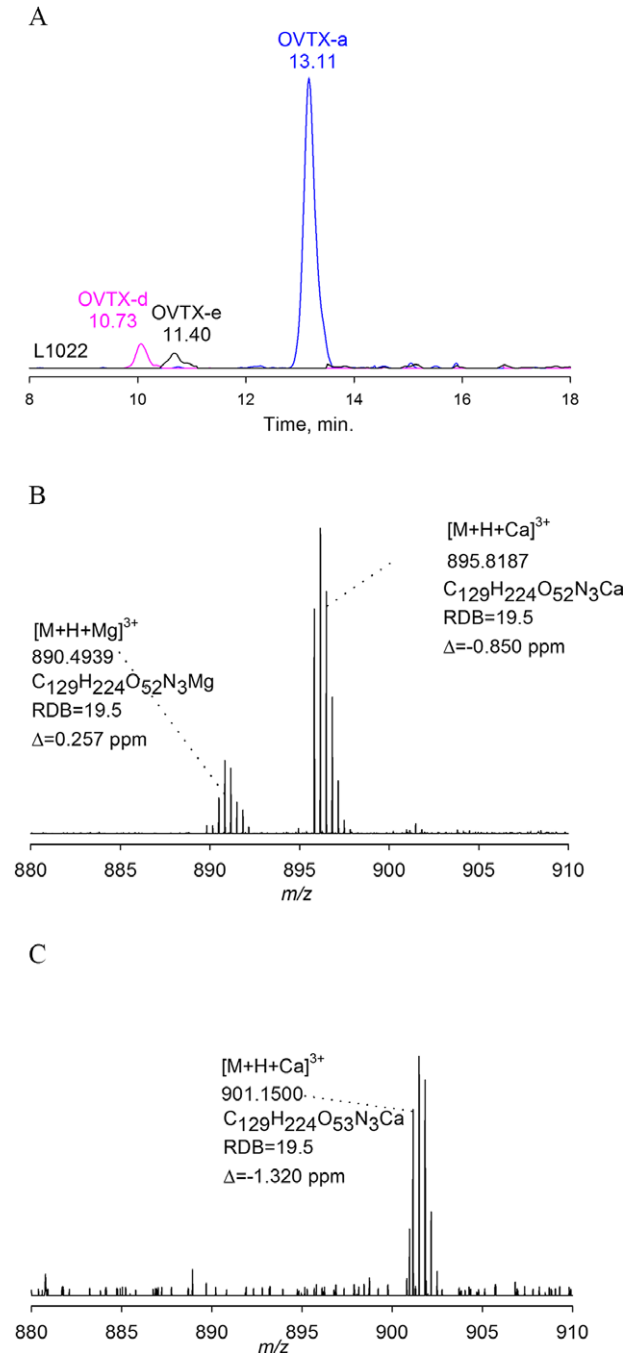


FIG. 10. LC-HRMS analysis of *Ostreopsis fattorussoi* L1022 strain, including (A) extracted ion chromatogram (XIC) of  $[M+H+Ca]^{3+}$  ions of ovatoxin-a (OVTX-a) and of the structural isomers ovatoxin-d and -e (OVTX-d, -e). Enlargements of the full HRMS spectra of (B) OVTX-a and (C) OVTX-d/-e.

and *Coolia* as well as by other authors (Fraga et al. 2011, Mohammad-Noor et al. 2013, Fraga and Rodríguez 2014), arguing that, although this interpretation does not follow the strict relationships among the plates, it takes into account their homology, according to the tabulation of Gonyaulacales (Balech 1980, Fensome et al. 1993).

In this paper, we adopted the interpretation of the thecal plates used by Hoppenrath et al. (2014), who mainly followed the original description of Fukuyo (1981), Norris et al. (1985) and Faust et al. (1996) regarding the epitheca, and Besada et al. (1982) regarding the hypotheca, recognizing that there is no an intercalary plate in *Ostreopsis*. In this regard, we observed that the large hypothecal plate at the antapex contacts the Sp plate (and therefore the sulcus), although the contact area is very narrow and not always visible, confirming that is not an intercalary but an antapical plate (i.e., the  $2''''$ ), as suggested by Hoppenrath et al. (2014). The uncertainty in the attribution of the large hypothetical plate (i.e., 1p vs.  $1''''$ ) may be explained by the

TABLE 2. LC-HRMS analyses of *Ostreopsis fattorussoi* strains from Lebanon.

	Toxin content (pg · cell <sup>-1</sup> )	Relative intensities		
		OVTX-a	OVTX-d	OVTX-e
L1002	nd	—	—	—
L1007	0.28	81.6%	11.7%	6.7%
L1008	nd	—	—	—
L1020	0.47	86.7%	8.8%	4.5%
L1022	0.94	87.8%	7.7%	4.5%

TABLE 3. Different interpretations of plate tabulation in *Ostreopsis* of different authors. n.r., not reported.

	Besada et al. (1982), Escalera et al. (2014)	Norris et al. (1985), Faust et al. (1996)	Hoppenrath et al. (2014)
Epitheca	Po (pp)	Po	Po
	4'	1'	1'
	2'	2'	2'
	3'	3'	3'
	1''	1''	1''
	1''	2''	2''
	2''	3''	3''
	3''	4''	4''
	4''	5''	5''
	5''	6''	6''
	6''	7''	7''
	Hypotheca	1'''	1'''
2'''		2'''	2'''
3'''		3'''	3'''
4'''		4'''	4'''
5'''		5'''	5'''
1''''		1''''	1''''
2''''		1p	2''''
Cingulum	n.r.	n.r.	1c
	n.r.	n.r.	2c
	n.r.	n.r.	3c
	n.r.	n.r.	4c
	n.r.	n.r.	5c
	n.r.	n.r.	6c
	n.r.	n.r.	Sa
Sulcus	n.r.	n.r.	Sda
	n.r.	n.r.	Ssa
	n.r.	n.r.	S.ac.a
	n.r.	n.r.	Sdp
	n.r.	n.r.	Ssp
	ps	n.r.	Sp

contact point of this plate with the sulcus that may not be visible in some species (Hoppenrath et al. 2014), as also within the same species (Sato et al. 2011, Escalera et al. 2014).

*Ostreopsis fattorussoi* could quickly be distinguished from the other two Mediterranean species in LM by its round thecal pores which are easily visible and appear larger than those in both *O. cf. ovata* and *O. cf. siamensis* (Table 4). Moreover, differently from *O. cf. ovata*, which shows pores of two size classes (Penna et al. 2005, Kang et al. 2013), pores of only one size class were visible in *O. fattorussoi*. Another way to distinguish *O. fattorussoi* from *O. cf. ovata* is the Po, which is longer in the former (Table 4). Moreover, the shape of the 6" plate is different: the 6"/5" suture length is almost twice as long as 6"/7" suture length, and the length:width ratio of the plate 6" is  $1.06 \pm 0.11$  (0.95–1.2), while in *O. cf. siamensis* and *O. cf. ovata* is  $1.5 \pm 0.2$  (1.1–2.4) and  $1.6 \pm 0.2$  (1.3–2.1), respectively (David et al. 2013).

Moreover, *O. fattorussoi* is readily distinguishable from the other *Ostreopsis* species because of some peculiar characteristics of its plate tabulation (Fig. 11). (i) in *O. fattorussoi* the 1' plate lies in the left half of the epitheca and is obliquely orientated giving a characteristic shape to the 6" plate and the oblique 6"/1' suture. In all other *Ostreopsis* species, the 1' plate closely occupies the center of the epitheca and is not oblique. (ii) *O. fattorussoi* is readily identifiable by the curved suture between 1' and 3' which makes plates 1' and 3' approximately hexagonal, while in the other *Ostreopsis* species they are pentagonal (with the exception of *Ostreopsis heptagona* which have a quadrangular 3' and a pentagonal 1' plate).

In *O. fattorussoi*, the 2' plate is narrow and almost twice the size of Po, separating the 3' and 3" plates. This characteristic could be useful to distinguish *O. fattorussoi* from many of the other *Ostreopsis* species, as only in *O. heptagona* and *O. labens* does plate 2' seem to divide plate 3' from plate 3" (Fig. 11). This characteristic may also be present in other *Ostreopsis* species but just not reported in their morphological descriptions: for example, in the original

description of *O. ovata*, Fukuyo (1981) indicated that this plate does not touch the plate 4", while this contact was indicated later by Besada et al. (1982). Similarly, *O. cf. ovata* has been described with (Selina and Orlova 2010, Kang et al. 2013) or without this contact (Escalera et al. 2014).

The hypotheca of *O. fattorussoi* does not show differences with that of *O. cf. ovata* (see Selina and Orlova 2010, Hoppenrath et al. 2014), although it does differ from the original drawings of *O. ovata* by Fukuyo (1981) (e.g., the latter has smaller 1'''' plate, a longer 2''''/4'''' suture, a longer and narrower 2'''' plate and a smaller 3'''').

The correct identification of *Ostreopsis* species in field samples based only on morphometric characters is often highly problematic. As the species recorded in Mediterranean Sea until now (*O. cf. ovata* and *O. cf. siamensis*) are very similar in shape and size, the DV/AP ratio was proposed as a characteristic for a quick distinction between the two species. Originally a DV/AP ratio of <2 for *O. cf. ovata* and >4 for *O. cf. siamensis* was proposed (Penna et al. 2005, Aligizaki and Nikolaidis 2006, Selina and Orlova 2010). The situation is now slightly more complex giving that *O. cf. ovata* from the northern Adriatic Sea was shown to have a DV/AP ratio slightly higher than 2, ~2.3–2.4 (Monti et al. 2007, Guerrini et al. 2010, Accoroni et al. 2012b). *Ostreopsis fattorussoi* has a DV/AP ratio of  $2.35 \pm 0.22$   $\mu\text{m}$  so this character is of no use to discriminate between *O. fattorussoi* and *O. cf. ovata*.

Considering cell size, *O. fattorussoi* seems to be on average slightly bigger (DV:  $60.1 \pm 5.6$   $\mu\text{m}$ , AP:  $25.7 \pm 3$   $\mu\text{m}$ , W:  $39.8 \pm 5.1$   $\mu\text{m}$ ) than *O. cf. ovata*, but these dimensions still fall within the upper range of the morphological variability reported for the latter species in natural samples (e.g., Accoroni et al. 2012b, Kang et al. 2013, Carnicer et al. 2015).

*Ostreopsis fattorussoi* and *O. ovata* are the smallest species of the genus and misidentification in field samples, in case both were present, is likely. In this regard, David et al. (2013) who stated their difficulties in distinguishing on a morphological base *Ostreopsis* spp. in field samples, most likely showed

TABLE 4. Comparison of morphometric parameters of *Ostreopsis siamensis*, *O. ovata*, *O. cf. siamensis*, *O. cf. ovata*, and *O. fattorussoi* based on measurements of specimens from literature values and this study.

Species	Apical pore ( $\mu\text{m}$ )	Pore size in thecal plates ( $\mu\text{m}$ )	Reference
<i>O. fattorussoi</i> sp.nov.	10–12.5 ( $11.7 \pm 1.4$ )	0.26–0.53 ( $0.38 \pm 0.08$ )	This study
<i>O. siamensis</i> Schmidt	27	0.1–0.5	Faust et al. 1996,
		0.08–0.38	Chang et al. 2000,
<i>O. cf. siamensis</i>	7.4–9.7	0.11–0.56 ( $0.30 \pm 0.07$ )	Penna et al. 2005,
	$10.9 \pm 1.24$	0.23–0.29	Aligizaki and Nikolaidis 2006,
	11–13	0.14–0.32	Selina and Orlova 2010,
<i>O. ovata</i> Fukuyo	8	0.07	Faust et al. 1996,
	6.9–9.6	0.16–0.55 ( $0.34 \pm 0.10$ )	Penna et al. 2005,
	$10.9 \pm 0.77$	0.07–0.32	Aligizaki and Nikolaidis 2006,
<i>O. cf. ovata</i>	6.6–9.0	0.25	Monti et al. 2007,
	6.3–8.3	0.12–0.25	Selina and Orlova 2010,
	4.8–6.8	0.06–0.26	Kang et al. 2013

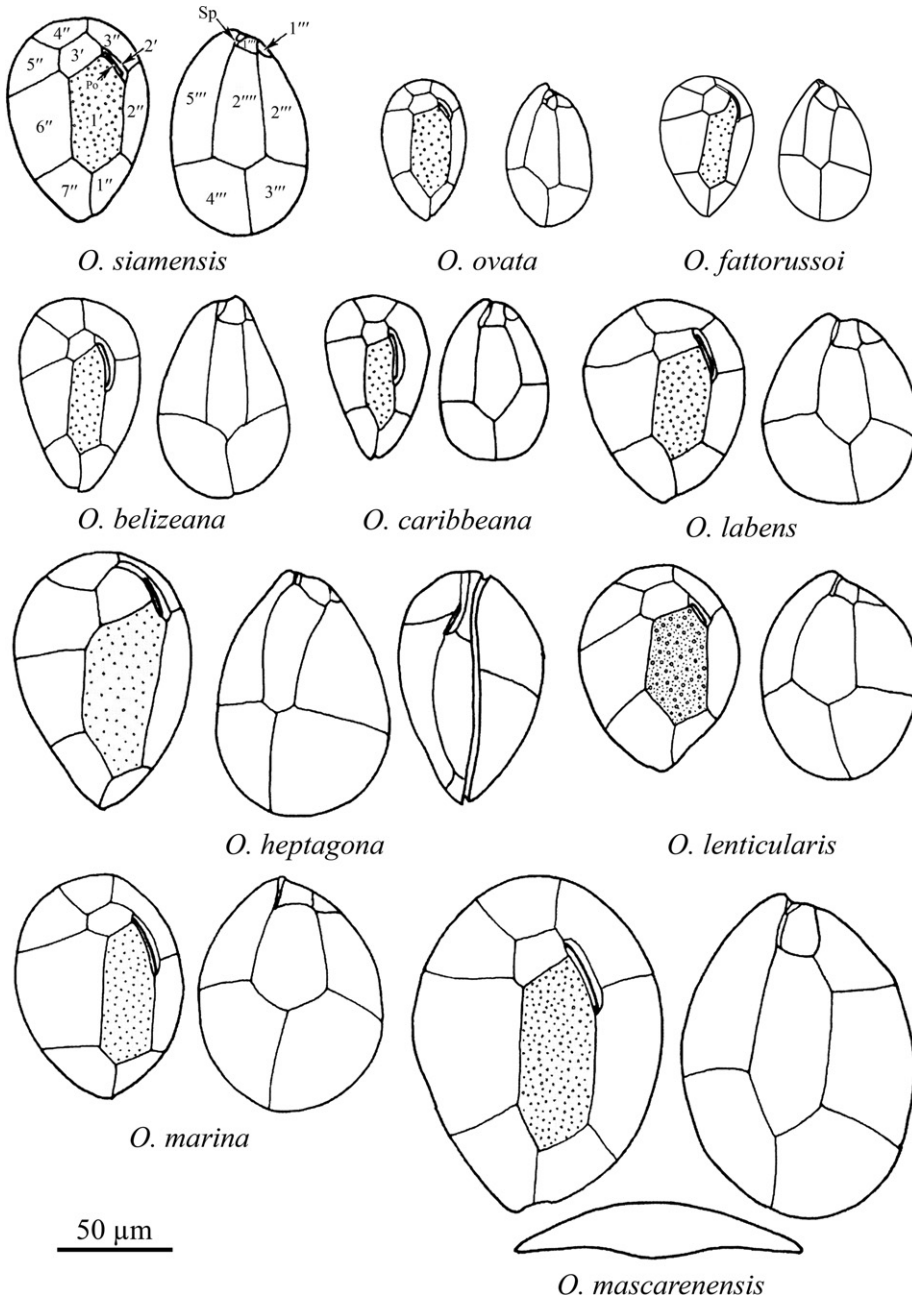


FIG. 11. Drawings of the plate patterns of the ten described *Ostreopsis* spp. modified from Hoppenrath et al. (2014). All species to scale.

*O. fattorussoi* and certainly not *O. cf. ovata* in their Figure 6A, given both the oblique plate 1' and 1'/6'' suture and the wide thecal pores and Po. If true then the distribution of *O. fattorussoi* is likely to include the Atlantic section of the Iberian Peninsula.

The ITS-5.8S and D1/D2 LSU ribosomal genes were able to consistently delineate divergences at the species level among *Ostreopsis* species (Sato et al. 2011, David et al. 2013, Penna et al. 2014, Tawong et al. 2014). In particular, the rDNA phylogeny showed clearly that strains belonging to *O. fattorussoi* were included in a distinct clade; in fact, both ITS-5.8S and LSU rDNA tree topologies were very

similar. The phylogenetic inference obtained from rDNA sequences was robust demonstrating that the clade grouping *O. fattorussoi* strains was supported by high values of bootstraps and Bayesian inferences. Specifically, the ITS phylogeny indicated the existence of *O. fattorussoi*, which included strains from eastern Mediterranean and Atlantic, as sister clade of *O. cf. siamensis*. The LSU phylogeny also indicated the presence of *O. fattorussoi* as a distinct species. This clade represented the second grouping that segregated from the first one comprising *Ostreopsis* sp., *Ostreopsis* sp. 5, and *O. cf. lenticularis*. Furthermore, collectively the rDNA phylogenies indicated the existence of at least other six different



species (Sato et al. 2011, Penna et al. 2014, Tawong et al. 2014).

This is the first report of *O. fattorussoi* in Lebanese coastal waters. The same species (reported as *Ostreopsis* sp.) has previously been reported in Cyprus coastal waters by Tartaglione et al. (2016). Besides Cyprus and Lebanon, molecular analyses revealed that the same genotype was found in Crete, Canary Islands (Spain) and Puerto Rico (USA) (Penna et al. 2010, 2014, David et al. 2013).

In the coastal waters of Lebanon, records of *Ostreopsis* species date back from 1979 (Abboud-Abi Saab 1989), and until now, the only reported species was *O. siamensis*, which has been recorded between July and September in the majority of the rocky coasts (Abboud-Abi Saab et al. 2013). In this study, *Ostreopsis* cells were detected throughout the year at the Lebanese coastal site, differently from the strongly seasonal trends observed in other Mediterranean areas (Accoroni et al. 2016). Molecular quantitative analysis by qPCR carried out in Lebanese samples indicated that *O. fattorussoi* was the dominant (often exclusive) *Ostreopsis* species, while the presence of *O. cf. ovata* was almost negligible (Casabianca S. and Penna A. pers. comm.). Abundances of *O. fattorussoi* were two orders of magnitude lower than those of *O. cf. ovata* recorded in other Mediterranean areas during their summer-late summer blooms (e.g., Mangialajo et al. 2011, Accoroni et al. 2015). However, it should be noted that in our study, the sampling was carried out mainly on *Corallina elongata*, a red alga with calcified (and therefore heavy) thallus that strongly influences the values of *Ostreopsis* abundances when they are expressed as cells · g<sup>-1</sup> fw. *Ostreopsis* maximum abundances were detected during the warmest months with two blooming periods in June–July and August. *Ostreopsis* blooms are generally summer events in temperate areas and maximum abundances are normally associated with the highest recorded temperatures (e.g., Aligizaki and Nikolaidis 2006, Mangialajo et al. 2008). However, some exceptions have been detected in the northern Adriatic Sea (Monti et al. 2007, Accoroni et al. 2012a) and the Sea of Japan (Selina et al. 2014), where blooms develop and reach maximum abundances at temperature values below the summer maximum. Similarly, *O. fattorussoi* showed the highest abundances when temperature values were between 27°C and 29.7°C, i.e., lower than the summer maximum (31.5°C). This would suggest that other environmental factors besides temperature may affect the development of *Ostreopsis* abundances. Several studies have provided increasing evidence of a link between harmful algal events and the nutrient enrichment of coastal waters (Glibert and Burkholder 2006, Heisler et al. 2008, Glibert et al. 2010). Although a correlation between *O. fattorussoi* abundances and nutrient concentrations was not observed, a significant correlation was highlighted between *O. fattorussoi* growth rates and

phosphate concentrations. The link between an influx of phosphorus-rich waters and higher net growth rate of *O. cf. ovata* has already been recognized in the northern Adriatic Sea (Accoroni et al. 2015).

*Ostreopsis* species have been shown to produce different toxins, mostly belonging to the palytoxin group. Mediterranean *O. cf. ovata* strains produce palytoxin analogs, such as isobaric palytoxin, OVTX-a, b, c, d, e, f, g, and h (Ciminiello et al. 2010, 2012b, Scalco et al. 2012, Brissard et al. 2015, García-Altres et al. 2015). On the contrary, the Mediterranean *O. cf. siamensis* strain was shown relatively non-toxic, producing only sub-fg levels of palytoxin (Ciminiello et al. 2013).

Blooms of *Ostreopsis* are well known in the Mediterranean Sea due to their effect on human health, mainly following inhalation of seawater droplets containing *Ostreopsis* cells or fragments and/or aerosolized toxins (Gallitelli et al. 2005, Kermarec et al. 2008, Tichadou et al. 2010, Del Favero et al. 2012, Casabianca et al. 2013, Ciminiello et al. 2014). The typical symptoms of *Ostreopsis* intoxication through aerosol and/or direct contact exposure are broncho-constriction, dyspnoea, fever, conjunctivitis, and skin irritations. Moreover, in the Mediterranean Sea, *Ostreopsis* toxins were found to contaminate seafood (Aligizaki et al. 2008, Ciminiello et al. 2015, Pelin et al. 2016). Several laboratory studies have shown that *Ostreopsis* exerts toxicity also on several marine organisms, both invertebrates and vertebrates (Faimali et al. 2012, Gorbi et al. 2012, 2013, Pagliara and Caroppo 2012, Carella et al. 2015). So far, no reports of human intoxications have been reported from Lebanon coast (Abboud-Abi Saab, pers. comm.).

Recently, Tartaglione et al. (2016) described the toxin profile of *Ostreopsis* sp. (named now as *O. fattorussoi*) isolated from Cyprus waters, showing some strains to produce isobaric palytoxin and OVTX-a, d and e, previously found only in *O. cf. ovata*, while some others produced only new palytoxin-like compounds named OVTX-i, OVTX-j<sub>1</sub>, OVTX-j<sub>2</sub>, and OVTX-k. In this study, we showed that two of the five analyzed Lebanese strains of *O. fattorussoi* were not toxic, and three produced OVTX-a and the structural isomers OVTX-d and -e. The confirmation for the identity of these toxins was provided by their perfectly superimposable LC-HRMS<sup>2</sup> spectra over those of previously characterized OVTX-a, -d, and -e (Ciminiello et al. 2010, 2012a, Dell'Aversano et al. 2015). The toxin profile of these strains also matched those of other *O. fattorussoi* strains from Cyprus (Tartaglione et al. 2016) and those found in ~40% of the Mediterranean *O. cf. ovata* strains analyzed so far (Dell'Aversano et al. 2015). However, Lebanese *O. fattorussoi* strains analyzed here did not produce any new palytoxin-like compounds such as those found in Cypriot *O. fattorussoi* (OVTX-i, OVTX-j<sub>1</sub>, OVTX-j<sub>2</sub>, and OVTX-k, Tartaglione et al. 2016).

So far, the toxin content of *O. fattorussoi* strains has occurred in the range of 0.06–2.8 pg · cell<sup>-1</sup> (Cyprus strains, Tartaglione et al. 2016 and Lebanese strains, this study) which is significantly lower than that of the Mediterranean *O. cf. ovata* strains maintained in the same cultured conditions (up to 44.0 pg · cell<sup>-1</sup>, Tartaglione et al. 2016). This has also been further supported by eco-toxicological tests on *Artemia salina* nauplii showing that *O. fattorussoi* from Cyprus had very low toxicity compared to *O. cf. ovata* (Tartaglione et al. 2016).

In conclusion, despite of the difficulties often reported concerning the identification of *Ostreopsis* species based on morphological characters alone (Penna et al. 2005, Parsons et al. 2012, David et al. 2013), *O. fattorussoi* does show some morphological features that lead unequivocally to its identification. These include:

- 1 the curved suture between plate 1' and 3' which makes plates them approximately hexagonal.
- 2 the 1' plate that lies in the left half of the epitheca and is obliquely orientated
- 3 the characteristic shape of plate 6'': its length: width ratio of  $1.06 \pm 0.11$  (0.95–1.2) and the 6''/5'' suture length is almost twice as long as that of 6''/7'.

Moreover, the phylogenetic analyses show that *O. fattorussoi* belongs to the Atlantic/Mediterranean *Ostreopsis* spp. clade distinct from the other *Ostreopsis* species.

This benthic dinoflagellate has been detected along the Lebanon coast throughout the year 2015 (with temperatures ranging from 18°C to 31.5°C), with bloom occurring in June and August and a significant correlation was highlighted between *Ostreopsis* growth rates and phosphate concentrations.

*O. fattorussoi* is a toxic species producing OVTX-a and structural isomers OVTX-d and -e, so far found only in *O. cf. ovata*, and three exclusive palytoxin-like compounds (OVTX-i, OVTX-j<sub>1</sub>, OVTX-j<sub>2</sub>, and OVTX-k, Tartaglione et al. 2016). All the data collected on this new species about its toxicity so far, however, would suggest a lower risk to human health and marine fauna to that of *O. cf. ovata*.

We gratefully acknowledge Demetris Kletou for the help in collecting Cyprus samples, Antonia Mazzeo for her assistance with the chemical analysis of toxins and Silvia Casabianca for molecular information about Lebanese *Ostreopsis* occurrence. A special thanks to Neil T.W. Ellwood for the English revision. This publication has been produced with the financial assistance of the European Union under the ENPI CBC MED Programme (M3-HABs project). The LC-HRMS study was granted by Programme STAR Linea 1 2013 (VALTOX, Napoli\_call2013\_08) financially supported by UniNA and Compagnia di San Paolo.

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