

Pentaplacodinium saltonense gen. et sp. nov. (Dinophyceae) and its relationship to the cyst-defined genus Operculodinium and yessotoxin-producing Protoceratium reticulatum

Article

Accepted Version

Creative Commons: Attribution-Noncommercial-No Derivative Works 4.0

Mertens, K. N., Carbonell-Moore, M. C., Pospelova, V., Head, M. J., Highfield, A., Schroeder, D., Gu, H., Andree, K. B., Fernandez, M., Yamaguchi, A., Takano, Y., Matsuoka, K., Nézan, E., Bilien, G., Okolodkov, Y., Koike, K., Hoppenrath, M., Pfaff, M., Pitcher, G., Al-Muftah, A., Rochon, A., Lim, P. T., Leaw, C. P., Lim, Z. F. and Ellegaard, M. (2018) Pentaplacodinium saltonense gen. et sp. nov. (Dinophyceae) and its relationship to the cyst-defined genus Operculodinium and yessotoxin-producing Protoceratium reticulatum. Harmful algae, 71. pp. 57-77. ISSN 1878-1470 doi: https://doi.org/10.1016/j.hal.2017.12.003 Available at http://centaur.reading.ac.uk/75005/

It is advisable to refer to the publisher's version if you intend to cite from the work. See <u>Guidance on citing</u>.

To link to this article DOI: http://dx.doi.org/10.1016/j.hal.2017.12.003



Publisher: Elsevier

All outputs in CentAUR are protected by Intellectual Property Rights law, including copyright law. Copyright and IPR is retained by the creators or other copyright holders. Terms and conditions for use of this material are defined in the End User Agreement.

www.reading.ac.uk/centaur

CentAUR

Central Archive at the University of Reading

Reading's research outputs online

- The potentially toxic Pentaplacodinium saltonense gen. et sp. nov. (Dinophyceae), and its 1 relationship to the cyst-defined genus *Operculodinium psilatum* and the yessotoxin-producing 2 Protoceratium reticulatum 3 4 Kenneth Neil Mertens[∗] 5 Research Unit for Palaeontology, Ghent University, Krijgslaan 281 s8, 9000 Ghent, Belgium 6 Now at: Ifremer, LER BO, Station de Biologie Marine, Place de la Croix, BP40537, F-29185 7 8 Concarneau Cedex, France, Tel: (0033)2 98 10 42 82 9 M. Consuelo Carbonell-Moore* 10 Oregon State University, Department of Botany and Plant Pathology, College of Agricultural 11 Sciences, 2082 Cordley Hall, Corvallis, OR 97331-2902, U.S.A. 12 13 Vera Pospelova 14 School of Earth and Ocean Sciences, University of Victoria, OEASB A405, 15 16 P.O. Box 1700 STN CSC, Victoria, BC, V8W 2Y2, Canada 17 Martin J. Head 18 Department of Earth Sciences, Brock University, 1812 Sir Isaac Brock Way500 Glenridge 19 Avenue, St. Catharines, Ontario, L2S 3A1, Canada 20 21
- 22 Andrea Highfield
- The Marine Biological Association of the United Kingdom, Citadel Hill, Plymouth PL1 2PB, 23
- 24 United Kingdom

Declan Schroeder 26 The Marine Biological Association of the United Kingdom, Citadel Hill, Plymouth PL1 2PB, 27 United Kingdom 28 School of Biological Sciences, University of Reading, Reading RG6 6AJ, United Kingdom 29 30 Haifeng Gu 31 Third Institute of Oceanography, SOA, Xiamen 361005, China 32 33 Karl B. Andree 34 Margarita Fernandez 35 IRTA, Crta. Poble Nou, Km 5.5, 43540-Sant Carles de la Rápita, Spain 36 37 38 Aika Yamaguchi Kobe University Research Center for Inland Seas, Kobe 657-8501, Japan 39 40 41 Yoshihito Takano Kazumi Matsuoka 42 Institute for East China Sea Research (ECSER), Nagasaki University, 1551-7, Taira-machi, 43 Nagasaki, 851-2213, Japan 44 45 Elisabeth Nézan 46 Ifremer, LER BO, Station de Biologie Marine, Place de la Croix, BP40537, F-29185 47 Concarneau Cedex, France 48 49 Gwenael Bilien 50

Ifremer, LER BO, Station de Biologie Marine, Place de la Croix, BP40537, F-29185 51 Concarneau Cedex, France 52 53 Yuri Okolodkov 54 Universidad Veracruzana, Instituto de Ciencias Marinas y Pesquerías, Calle Hidalgo núm. 55 617, Colonia Río Jamapa, Boca del Río, 94290 Veracruz, México 56 57 Kazuhiko Koike 58 Graduate School of Biosphere Science, Hiroshima University, Kagamiyama 1-4-4, Higashi-59 Hiroshima, Hiroshima 739-8528, Japan 60 61 Mona Hoppenrath 62 63 Senckenberg am Meer, Deutsches Zentrum für Marine Biodiversitätsforschung (DZMB), Südstrand 44, D-26382 Wilhelmshaven, Germany 64 65 Maya Pfaff 66 Marine Biology Research Center, Ma-RE Institute, Zoology Department, University of Cape 67 68 Town, Rondebosch 7701, South Africa 69 **Grant Pitcher** 70 71 Marine and Coastal Management, Private Bag X2, Rogge Bay 8012, Cape Town, South Africa 72 73 Abdulrahman Al-Muftah Department of Biological and Environmental Sciences, Qatar University, Doha, Qatar 75

76	
77	Rochon, Andr <u>é</u> e
78	Institut des sciences de la mer de Rimouski (ISMER), Université du Québec à Rimouski,
79	310 allée des Ursulines, Rimouski, QC, Canada G5L 3A1
80	
81	Po Teen Lim, Chui Pin Leaw, Zhen Fei Lim
82	Institute of Ocean and Earth Sciences, University of Malaya, 16310 Bachok, Kelantan,
83	Malaysia
84	
85	Marianne Ellegaard
86	Department of Plant and Environmental Sciences, University of Copenhagen,
87	Thorvaldsensvej 40, DK-1871, Frederiksberg, Denmark
88	
89	(*both authors contributed equally)
90	
91	

Strains of a dinoffageliate from the Salton Sea, previously identified as <i>Protoceratium</i>
reticulatum and yessotoxin producing, have been reexamined morphologically and genetically
and Pentaplacodinium saltonense n. gen. et sp. wasis erected to accommodate this species.
Pentaplacodinium saltonense differs from Protoceratium reticulatum (Claparède et
Lachmann 1859) Bütschli 1885 in the number of precingular plates (5five vs. six6), cingular
displacement (2two widths vs. 4one), and distinct cyst morphology. Incubation experiments
(excystment and encystment) show that the resting cyst of Pentaplacodinium saltonense is
morphologically most identical similar to the cyst-defined species Operculodinium
israelianum (Rossignol 1962) Wall 1967 and Operculodinium. psilatum Wall 1967.
Comparative eCollections of comparative material from around the globe (including
Protoceratium reticulatum -and the genus Ceratocorys) and single cell PCR wasere used to
clarify molecular phylogenies. Variable regions in the LSU (3three new sequences), SSU (12
new sequences) and intergenic ITS 1-2 (14 new sequences) were sequenceobtaineded These
show that Pentaplacodinium saltonense and Protoceratium reticulatum form two distinct
clades. Pentaplacodinium saltonense formeds a monophyletic clade with several unidentified
strains from Malaysia. LSU and SSU rDNA sequences of three species of Ceratocorys (C.
armata, C. gourreti, C. horrida) from the Mediterranean and several other unidentified strains
from Malaysia form a well-supported sister clade. The unique phylogenetic position of an
unidentified strain from Hawaii is also documented thatand requires further examination. In
addition, based on the V9 SSU topology (bootstrap values >80%), specimens from Elands
Bay (South Africa), originally described as Gonyaulax grindleyi by Reinecke (1967), cluster
with Protoceratium reticulatum. So far, tThe known range of Pentaplacodinium saltonense is
tropical to subtropical waters, and its cyst is recorded as a fossil in lateupper Cenozoic

117	sediments. Protoceratium reticulatum and Pentaplacodinium saltonense seem to inhabit
118	different niches: motile stages of these species dinoflagellates have not been found in the
119	same plankton sample.
120	
121	Keywords
122	yessotoxins, Protoceratium, Pentaplacodinium, Protoceratium, precingular plates, Salton Sea,
123	Ceratocorys, Operculodinium, Cribroperidinioideae
124	
125	
126	
127	

1. Introduction

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

128

The dinoflagellate genus *Protoceratium* was erected by Bergh (1881, p. 242) with *Protoceratium aceros* as the type species (fig. 36), which was recovered from Strib, Denmark. Bütschli (1885, p. 1007, plate 52, fig. 2) considered *Peridinium reticulatum* as described earlier by Claparède and Lachmann (1858) from Bergen Fjord, Norway, as a senior synonym, and he proposed the combination *Protoceratium reticulatum*. He also considered *Clathrocysta* reticulata as described by Stein (1883) a junior synonym. The plate formula for P. reticulatum, 4', 0a, 6", 6"', 1p, 1"", was first provided by Wołoszyńska (1929) through the study of Baltic Sea specimens. Reinecke (1967) erected the name Gonyaulax grindleyi for specimens from Elands Bay in Cape Town, South Africa, with the tabulation 3', 1a, 6", 6"', 1p, 1"". Based on a detailed study of the theca of *Protoceratium reticulatum* from the North Sea, Stosch (1969) considered G. grindleyi to be a junior synonym of P. reticulatum, although he considered it assignable to the genus Gonyaulax. Dodge (1989) agreed with the tabulation of Reinecke (19679), but retained the genus *Protoceratium* because he considered it different from his emendation of the genus Gonyaulax, by having only one intercalary plate. Hansen et al. (1997) restudied specimens close to the type locality of *P. aceros*, and based on the plate analysis concluded that P. reticulatum, P. aceros and G. grindleyi were conspecific, and agreed with the tabulation of Wołoszyńska (1929). Paez-Reyes and Head (2013) reviewed the morphological variability reported for *P. reticulatum* and concurred with Dodge (1989) in maintaining *Protoceratium* as a distinct genus from *Gonyaulax*. Since the early 1900's, sSeven other *Protoceratium* species have been described since the early 1900s, and the latest review of these taxa was having been performed by Schiller (1937 p. 322–326). Kofoid (1907) described P. areolatum from the tropical Pacific and

emended the genus for the first time. Meunier (1910) described a very similar species from

the Kara Sea that he named *Protoceratium splendens*, which is possibly a junior synonym, as 153 154 suggested by Gómez (2012). Later, Kofoid in Kofoid and Michener (1911) emended Protoceratium once more to include several new species from the eastern tropical Pacific that 155 were described without illustration (P. cancellorum, P. globosum, P. pellucidum, P. pepo, P. 156 promissum), and he suggested a tabulation formula for the genus: 2', 0a, 6" (?c), 6"', 0p, 3"". 157 Schiller (1937) transferred Clathrocysta aculeata as described by Stein (1883) to 158 159 Protoceratium aculeatum, presumably based on the fact that Bütschli (1885) had considered the genus Clathrocysta described by Stein 1883 as a junior synonym of Protoceratium. 160 Schiller (1937) transferred *Peridinium spinulosum* as described by Murray and Whitting 161 162 (1899) to the genus *Protoceratium*. Later, Balech (1988) rediscovered this species in the South—Wwest Atlantic and suggested yet another another variation on the tabulation for 163 Protoceratium, 3', 0a, 6", 6"', 2"", based on his observations of Protoceratium spinulosum. 164 165 Protoceratium reticulatum (Claparède et Lachmann) Bütschli 1885 is a very common dinoflagellate found in cold and warm waters, as well as in oceanic and neritic environments 166 (e.g., as Operculodinium centrocarpum in Zonneveld et al., 2013). -Its resting cyst distribution 167 today reveals a strong link with the North Atlantic Current, an association traceable through 168 the upper Cenozoic fossil record (Hennissen et al., 2017 and references therein). 169 170 Protoceratium reticulatum is considered potentially toxic because of its production of yessotoxins (e.g. Paz et al., 2008; Sala-Pérez et al., 2016). It has been successfully isolated 171 and cultured from many parts of the world, and grown into cultures. Cysts of *P. reticulatum* 172 were first observed in cultures established from motile cells from the inner Oslofjord-, 173 Norway) by Braarud (1945). This cyst was related by Wall and Dale (1966, 1967, 1968) to the 174 cyst-defined species described from the Miocene of Australia, Operculodinium centrocarpum 175 (Deflandre et Cookson 1955) Wall 1967. That assignation was challenged by Head and 176 Wrenn (1992) and Head (1996a) on the grounds that Operculodinium centrocarpum was 177

larger and more robust than the cysts recorded by Wall and Dale (1966) from modern sediments. However, aA restudy of the holotype of Operculodinium centrocarpum refuted this assignation confirmed this, and the name "cyst of Protoceratium reticulatum" was recommended (Matsuoka et al., 1997). Wall and Dale (1968) proposed that P. reticulatum was also related to the cyst-defined *Operculodinium psilatum* Wall 1967 and furthermore possibly to Operculodinium israelianum (Rossignol 1962) Wall 1967. The cyst-defined Pyxidinopsis psilata (Wall et Dale in Wall et al., 1973) Head 1994 was subsequently also linked to Protoceratium reticulatum (Dale, 1996, as Tectatodinium psilatum) although this connection was later questioned later (Mertens et al., 2011). Because of uncertainty regarding the links between the cysts produced by P. reticulatum (see Head, 2006), and cysts-cystdefined species named based on from the fossil records, Head (1996a, 1996b) and subsequent authors used the the term "Operculodinium centrocarpum sensu Wall and Dale, 1966" was used to describe the cysts that had first been observed by Braarud (1945) and Wall and Dale (1966). With the removal of *Pyxidinopsis psilata* as a potential cyst of *Protoceratium* reticulatum, Paez-Reyes and Head (2013) argued on the basis of non-overlapping geographic distribution that the "cyst of Protoceratium reticulatum" was now unambiguous and should replace the term "Operculodinium centrocarpum sensu Wall and Dale, 1966". That approach is followed here. Recent studies of variation in the process length of cysts of *Protoceratium* reticulatum have been related to variations in sea surface salinity and other parameters (e.g., Mertens et al. 2011; Jansson et al., 2014), and the cyst wall appears to be composed of cellulose glucan (Bogus et al. 2014). Resting cyst production through sexual reproduction has recently been demonstrated by Salgado et al. (2017). Protoceratium reticulatum was assigned questionably to the subfamily Cribroperidinioideae by Fensome et al. (1993b) based on the presence of six precingular

plates, L-type ventral organization and possible dextral torsion, which at the time had not

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

193

194

195

196

197

198

199

200

201

been documented. This assignation was confirmed by Paez-Reyes and Head (2013). However, tThe description of the very similar cyst-defined Operculodinium bahamense, with neutral torsion and modified L-type ventral organization, which would be placed allowing placement in the subfamily Leptodinioideae, either challenges challenges such the present subfamilial classification of the Gonyaulacaceae, or implies that *Operculodinium* is polyphyletic, with both outcomes being possible -(Paez-Reves and Head, 2013). Furthermore, molecular phylogenetics show that *Protoceratium reticulatum* is closely related to the family Ceratocoryaceae but not to the other extant cribroperidinean, Lingulodinium polyedra (Saldarriaga et al., 2004). It should also be noted that morphological variation and sequencing of cysts has suggested pseudocryptic speciation in *P. reticulatum* (Mertens et al., 2012a). Howard et al. (2009) investigated the phylogenetic relationships of yessotoxin-producing dinoflagellates, including several strains of *P. reticulatum* from different localities. Using Large Sub Unit (LSU) and Internal Transcribed Spacer (ITS) ribosomal DNA (rDNA) sequencing, they showed that the *P. reticulatum* strains formed a monophyletic clade in both phylogenies. However, oOne particular strain (CCMP404) isolated from the Salton Sea (California) in 1966 showed significant genetic differences from the other strains in both phylogenies. Despite these genetic differences, Howard et al. (2009) considered all the strains to belong to the species *P. reticulatum*. The Salton Sea is the largest saline lake in California with a surface area of 980 km² (Reifel et al., 2002). It has a mean depth of 8 m and a maximum depth of 15 m (Ferrari and Weghorst, 1997). Although originally composed of relatively freshwater, it has become saline due to a lack of outflow and high evaporation rates. During 1997–1999, the salinity was between 41 and 45 g l⁻¹ (Watts et al., 2001), while the temperature varied between about 12

and 40°C seasonally (Watts et al., 2001; Holdren and Montaño, 2002). Oxygen at times was

supersaturated due to phytoplankton photosynthesis, but was also often severely depleted,

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

226

occasionally even in surface waters (Watts et al., 2001; Holdren and Montaño, 2002). Reifel et al. (2002) reported *P. reticulatum* from the Salton Sea without illustration or description.

In the present study, through reevaluation of the CCMP404 strain originated from the Salton Sea and observations of recently collected plankton samples from the Salton Sea, it iswe demonstrated that specimens living in the Salton Sea that have had been previously been identified as *P. reticulatum*, have a dissimilar different tabulation to that of *P. reticulatum*. To resolve this issue and accommodate these organisms, we have erected a new genus and new species, Pentaplacodinium saltonense n. gen_ et sp_ is erected. From the Salton Sea plankton samples, we describe the morphology of the thecate stage is described, showing significant differences with *P. reticulatum*. Similarly, through incubation of cysts from Salton Sea surface sediments, we describe the corresponding cyst is described. Phylogenetic relationships are explored, including those with several unpublished sequences of *P. reticulatum*, Ceratocorys armata (Schütt 1895) Kofoid 1910, Ceratocorys gourretii Paulsen 1931, Ceratocorys horrida Stein 1883, and several unidentified strains. In addition, both the autecology and fossil record of Pentaplacodinium saltonense are examined.

2. Material and Methods

The cyst—theca relationship of *P. saltonense* was established through a germination experiment of a sample from the Salton Sea (CA, USA). To identify differences and similarities between *P. reticulatum* and *P. saltonense*, we compared the morphology of thecate stages of strains present in culture collections and other cells used for sequencing, arewere compared (Table 1, Suppl. Table 1). In addition, we studied the phylogenies arewere constructed of using LSU, ITS and SSU rDNA based sequences of *P. saltonense* and *P. reticulatum* from several of the same cells or cultured strains, as well as three *Ceratocorys*

species isolated from the Mediterranean, and several unidentified strains from Hawaii and Malaysia (Table 1, Suppl. Table 1).

2.1. Morphological study imaging of cells in from plankton samples and strains present in culture collections with microscopy

Plankton samples were obtained from the Salton Sea (California, U.S.A.; 33.50 °N, 115.91 °W) on 24 Oct. 2013 using a plankton net with a 20 µm mesh size. These samples were fixed with ethanol (50% final concentration) and stored cold. Several strains from previously sequenced strains from culture collections established from several other locations were also studied using transmitted light or scanning electron microscopy (Figure 1, Table 1).

For scanning electron microscopy (SEM) of thecate stages by M.C.C.-M., samples were prepared either by filtering a plankton sample or culture, or isolating a single cell under a LeicaTM Inverted Light-light Microscopemicroscope (Germany). When the sSamples were was filtered, by placing an aliquot of ~300 μL aliquot was placed on a MilliporeTM 0.25 mm diameter—5—μm pore—polycarbonate filter at the bottom of a MilliporeTM column. Approximately 7 mL of distilled water were added to remove the fixative (ethanol, lugol or formaldehyde) and seawater. A gentle manual vacuum with a 60 cc syringe was used to speed filtration. Individually isolated ceCells were removed using a glass micropipette under a Leica Inverted inverted Light mMicroscope (Germany) with magnification 10x5x. Individual cells were washed six times with distilled water in double depression microscope slides). After the cells were clean, they were placed on the same kind of filter as for the filtered samples. All filters were air-dried, then adhered affixed to 25 mm diameter aluminium stubs with adhesive tabs (7/16" diameter). The mounted filters were then coated with a mixture of gold-palladium in a Cressington Sputter Coater (U.S.A.) for 60 s.

Observations were performed with a FEI Quanta 3D Dual Beam SEM (Clackamas, Oregon, U.S.A.), at 5 kV. Tilts up to 52° were applied. Digital images were saved in Tiff format (2048 x 1768 pixels). Adobe-_PhotoshopTM software was used to remove the background while maintaining the integrity of the original image.

For scanning electron microscopy (SEM)_of culture CCMP 3243 by K.N.M., the culture was filtered and washed with distilled water and dehydrated in a graded ethanol series (30 to 100% in six steps). The filters were encased in metallic baskets, critical-point dried with CO₂ (CPD Bal-Tec 030), glued onto stubs, sputter coated with platinum/palladium for 90 s (JEOL JFC-2300 HR) and examined in a JEOL 6330F scanning electron microscope (JEOL, Tokyo, Japan) at the University of Copenhagen.

Measurements of thecate thecae stages of the newly described species were done conducted by M.C.C.-M. under SEM. For each motile cell, the length was measured along the center of the longitudinal axis, the width was measured along the middle of the cingulum, perpendicular to the longitudinal axis from one lateral margin to the other. All motile cell measurements in the species descriptions cite the minimum, average (in parentheses) and maximum values (in μm), in that order. The standard deviation (SD) is also provided where appropriate.

Label<u>l</u>ing of tabulation follows a modified Kofoid system that recognizes homologs (e.g., Fensome et al. 1993<u>b</u>). The sulcal plate labelling <u>is</u>-accord<u>ing tos with</u> Balech (1980).

2.2. Germination experiment of cysts of P. saltonense

Sediment samples were collected from the Salton Sea-(CA, U.S.A.) aton the same timeday of as plankton sample collectionon, during the same field campaign_-on 24 October- 2013, using a Petite Ponar Grab at shallow water depths (<0.5 m). All samples were stored in plastic

bags in a refrigerator at 4°C. *In_-situ* sea surface salinities and sea surface temperatures were measured during sampling (Table 1).

About 0.5–1.0 cm³ of wet sediment was immersed in filtered seawater and, after one minute of ultrasonication using an ultrasonic bath, the sediment was rinsed through a 20 μm nylon-mesh sieve using filtered seawater. From this residue, the cyst fraction was separated using the heavy-liquid sodium polytungstate (SPT) at a density of 1.3 g cm⁻¹ (Bolch, 1997). Single cysts were then transferred to Orange Scientific 0.5 mL microwells subjected to an irradiance of 100 μmol photons m⁻² s⁻¹ and 24-hour light, and filled with f/2 medium at room temperature and a salinity of 35 psu. Cysts were regularly checked for germination, and observations were performed under a Leitz DM IL inverted light microscope. Encysted and excysted cysts, as well as motile cells, were photographed and measured using a Leica DM5000B light microscope with 100x oil immersion objectives.

2.3. Morphological study of cysts extracted from surface sediments with using light microscopy and SEM

Surface sediment samples were collected from several sites in the Salton Sea sites were obtained forto study of cysts of *Pentaplacodinium saltonense* (Table 1). Palynological techniques were used for processing (e.g., Pospelova et al., 2010; Mertens et al., 2012b). Material was rinsed twice with distilled water to remove salts. The samples were oven-dried at 40°C and then treated with room-temperature 10% hydrochloric acid (HCl) at room temperature to remove calcium carbonate particles. To dissolve siliceous-silicate particles, samples were treated with 48–50% room-temperature hydrofluoric acid (HF) at room temperature for two days, and then treated for 10 min with room-temperature HCl (10%) to remove fluorosilicates. The residue was rinsed twice with distilled water, ultrasonicated for

~30 sec and finally collected on a 15 µm mesh. Aliquots of residue were mounted on microscope slides using glycerine jelly.

All measurements and light photomicrographs were obtained by K.N.M., and V.P., respectively using an Olympus BX51 with a Nikon digital sight DS-1L 1 module, and a Nikon Eclipse 80i transmitting light microscope with a DS-L2 module, all with 100x oil immersion objectives.

For each cyst, the lengths of the three longest visible processes with the corresponding widths at their base were measured within the focal plane. Process length was measured from the middle of the process base to the process tip. The <u>average</u> distance <u>between processes</u> was determined <u>by measuring the distance</u> between a process <u>near the center of the cyst</u> on the upper surface <u>of the cyst near the centreer</u> and the five processes nearest to it, as measured between the middle of the process bases as seen from the surface of the cyst. The central body wall thickness was measured at two to three positions around the cross section of each cyst. The central body maximum and minimum diameters were also measured unless specimens were overly compressed or broken. Fragments representing less than half of a cyst, and cysts with mostly broken processes, were not measured. All cyst measurements in the species descriptions cite the minimum, average (in parentheses) and maximum values (in µm), in that order. The standard deviation (SD) is also provided where appropriate.

For SEM observation of cysts at Geotop (the Université du Québec à Montréal, Canada), single specimens were picked under an inverted microscope with a micropipette, sputter coated with platinum/palladium for 60 s and observed using a scanning electron microscope (Hitachi S-3400N SEM).

2.4. Single-cell polymerase chain reaction (PCR) amplification and sequencing of culture of Salton Sea culture

376

377

354	Isolated cells were washed three times in serial drops of $0.22\ \mu m$ filtered and sterilized
355	seawater by micropipette. Each cell was transferred to a 200 μm PCR tube containing 10 μL
356	of Quick Extract FFPE DNA Extraction Solution (Epicentre, Madison, WI, USA) and
357	incubated for 1h at 56°C, then for 2 min at 90°C. The resulting extract was used as a DNA
358	template for the initial PCR amplification. Sequences of SSU and partial LSU rDNA were
359	determined from single cells of <i>P. saltonense</i> . The PCR was performed with EconoTaq 2X
360	Master Mix (Lucigen, Middleton, WI, USA) following the manufacture's protocols. The
361	external primers (SR1 and LSU R2) were used for the initial PCR. The first PCR product was
362	used as a DNA template for the second PCR. The following combinations of primer pairs
363	were used separately for the second PCR: SR1 and SR12, 25F1 and LSU R2. Using the
364	second PCR products as the template DNA, the third PCR were was performed by the
365	following combinations of primer pairs: SR1b and SR3, SR1b and SR5TAK, SR4 and
366	SR7TAK, SR6 and SR9p, SR8p and SR12, 25F1 and 25R1, D3A and LSU R2. The dDetails
367	of the primers are described in Takano and Horiguchi (2004) and Yamaguchi et al. (2016).
368	The PCR protocols and sequencing are described in Yamaguchi et al. (2016).
369	
370	2.5. Sequencing of single cells of Protoceratium reticulatum from Elands Bay (South Africa),
371	originally described as Gonyaulax grindleyi by Reinecke (1967)
372	
373	Isolated cells were washed three times in serial drops of 0.22 μm filtered and sterilized
374	distilled water and then transferred to a 0.2 mL PCR tube. Cells were subjected to three
375	rounds of heating to 95°C for 5 minutes and cooling on ice for 5 minutes to induce cellular

lysis. $5\,\mu\text{L}$ of the cell lysate was then used as a template for PCR using primers to amplify a

168 bp region of the SSU, encompassing the V9 region, V9 For (5'-

378	GTACACCGCCCGTC-3') V9 Rev (5'- TGATCCTTCTGCAGGTTCACCTAC-3')
379	(Lane-et al., 1991; Medlin et al., 1988). PCR reactions were carried out in 25 μ L volumes
380	containing 5 μ L DNA template, 10 pmol each primer, 1 x buffer, 1 mM MgCl ₂ , 0.0025 mM
381	dNTPs, 0.5 Unit Gotaq polymerase (Promega). PCR reactions proceeded with an initial
382	denaturation at 95°C for 5 minutes, followed by 35 cycles of denaturation at 95°C for 30
383	seconds, annealing at 54°C for 20 seconds and extension at 72°C for 20 seconds and a final
384	extension step of 72°C for 5 minutes. PCR products were sequenced directly in both
385	directions using the respective primers (Source Bioscience).
386	
387	2.6. Sequencing of unidentified cultured strains
388	
389	For strains from Hawaii and Malaysia, single cells were isolated from plankton samples
390	(Suppl. Table 1) and washed three times with sterilized bi-distillate water and were used as
391	the template to amplify about 1,430 bp of the LSU rRNA gene (D1-D6 domains), using the
392	primers D1R (forward, 5' -ACCCGCTGAATTTAAGCATA-3') (Scholin et al., 1994), 28-
393	1483R (reverse, 5' -GCTACTACCACCAAGATCTGC-3') (Daugbjerg et al., 2000), 1740
394	bp of the SSU rRNA gene, using the primers SR1(forward, $5'$ -
395	TACCTGGTTGATCCTGCCAG-3 $^{\prime}$) and SR12b (reverse, 5 $^{\prime}$ -
396	CGGAAACCTTGTTACGACTTCTCC-3') (Takano & Horiguchi, 2006), and 600 bp of the
397	total ITS1-5.8S-ITS2, using the primers ITSA (forward, 5' -CCTCGTAAC
398	AAGGHTCCGTAGGT-3'), ITSB (reverse, 5'-CAGATGCTTAARTTCAGCRGG)
399	(Adachi et al., 1996). A 50 μL PCR cocktail containing 0.2 μM forward and reverse primer,
400	PCR buffer, 50 µM dNTP, 1U of Taq DNA polymerase (Takara, Dalian, China) was
401	subjected to 35 cycles using a Mastercycler PCR (Eppendorf, Hamburg, Germany). The PCR

reaction procedure was 4 min at 94 °C, followed by 25 cycles of 1 min at 94 °C, 2 min at 45 °C, 3 min at 72 °C, and final extension of 7 min at 72 °C. PCR products were sequenced directly in both directions using the ABI Big-Dye dye-terminator technique (Applied Biosystems, Foster City, CA, USA), according to the manufacturer's recommendations.

DNA extracts from strains collected in Spain (processed at IRTA) were prepared according to the protocol described in Andree et al. (2011). The extracted DNA was used in the amplification of ITS-1, 5.8S, ITS-2 sequences utilizing primers described in Andree et al. (2011), and a partial LSU sequence was amplified utilizing the primers described previously in Hansen et al. (2000). The amplification reactions were carried out in 25μL volume containing: 10 mM Tris-HC1 pH 8.3 (at 25 °C), 50 mM KC1, 2 mM MgC1₂, 0.001 % w/v gelatin, 400 pM dNTP's, 1 μM of each primer, and 1 U Taq polymerase. Amplifications were performed using the following parameters: 94 °C for 5 min followed by 35 cycles of 95 °C for 30 s, 50 °C for 45 s, 72 °C for 1 min, and a final extension of 72 °C for 5 min. The PCR products were purified using Qiagen spin columns (Qiagen PCR Purification Kit) and sent for bi-directional sequencing by a commercial company (Sistemas Genomicos, Valencia, Spain) utilizing the same primers as those used in the original amplification. The resulting nucleic acid sequence data was manually proofed using BioEdit (Hall et al., 1999) to confirm the consensus sequence.

The strain 091223-38_M16 from Helgoland (North Sea) was sequenced by M.H.. The Epicentre MasterPure complete DNA & RNA Purification Kit was used for the DNA extraction. We use puReTaq ready-to-go PCR beats arewere used; annealing temperature was 50°C; 33 cycles; primers: ITS1 (forward) 5' GGTGAACCTGAGGAAGGAT 3'; ITS4 (reverse) 5' TCCTCCGCTTATTGATATGC 3'. The PCR product of the right correct size was gel isolated (QIAquick Gel Extraction Kit). Sequencing was done by Macrogen with the ITS1 primer.

427	Strains and single cells from Japan, sequenced by Yoshihito Takano and Kazuhiko
428	Koike, were sequenced using methods mentioned in Mertens et al. (2012a).
429	DNA was extracted from cultures (strain references K1474, 1476, 1477, 1478, 1479
430	and 0976) acquired from the NCMA (National Centre for Marine Algae) using the DNeasy
431	DNA extraction kit (Qiagen) according to manufacturers' instructions. The 760 bp region of
432	the LSU rRNA gene was amplified using 2 µL DNA in PCR reactions spanning the D1-D2
433	variable region D1R (forward, 5'-ACCCGCTGAATTTAAGCATA-3'), D2C (reverse, 5'-
434	GCTTGGTCCGTGTTTCAAGA-3') (Scholin et al., 1994) a 168 bp region of the SSU rRNA
435	gene (V9) V9 For (5'-GTACACACCGCCCGTC-3') V9 Rev (5'-
436	TGATCCTTCTGCAGGTTCACCTAC-3') (Lane et al., 1991; Medlin et al., 1988) and a 710
437	bp intergenic region ITS1, 5.8S, ITS2, EITS2 For (5'-GTAGGTGAACCTGCVGAAGA-3')
438	EITS2 Rev (5'-TGGGGATCCTGTTTAGTTTC-3') (Guillou et al. 2002). PCR for V9 is
439	detailed in section 2.5. For LSU and ITS, PCR reactions were carried out in 50 μL volumes
440	containing 2 μ L DNA, 20 pmol each primer, 1 x buffer, 1.5 mM MgCl ₂ , 0.0025 mM dNTPs,
441	1 Unit Gotaq polymerase (Promega). PCR reactions proceeded with an initial denaturation at
442	95°C for 5 minutes, followed by 35 cycles of denaturation at 95°C for 30 seconds, annealing
443	at 60°C for 45 seconds and extension at 72°C for 1 min and a final extension step of 72°C for
444	5 minutes. PCR products were sequenced directly in both directions using the respective
445	primers (Source Bioscience) and sequences were manually verified using Chromas
446	(Technelysium Pty Ltd) prior to phylogenetic analysis.
447	Novel sequences were deposited in Genbank under accession numbers
448	XXXXXMG646283-MG646333.
449	
450	2.7. Sequence alignments and phylogenetic analyses

Multiple sequence alignments were constructed for sequences generated for the variable regions V9 (SSU), D1-D2 (LSU) and partial ITS1, 5.8S, ITS2 (intergenic region), respectively, in BioEdit 7.0 (Hall 1999) using ClustalW along with other available sequences from Genbank. Alignments were trimmed accordingly based on the lengths of the sequences acquired and to allow for a sufficient number of sequences to be included in the phylogeny. Phylogenetic analysis based on neighbour-joining and maximum likelihood was undertaken using MEGA 6 (Tamura et al., 2013) using the default parameters. Bootstrap values were retrieved from 1000 replicates and are indicated on the nodes of the trees.

3. Results

3.1. Study of plankton samples, culture strains, germination experiments, and surface sediments

Investigation of plankton samples from the Salton Sea revealed the presence of a species that is superficially similar to *P. reticulatum* and is here assigned to *Pentaplacodinium saltonense* gen. et sp. nov.n. sp. Three process-bearing cysts (Plate 1) were isolated from surface sediments of the Salton Sea, California, USA (Table 1) and identical morphologies emerged from these cysts (Plate 2). These cells started dividing after germination, and one strain was maintained. The cells were identical in morphology to specimens observed in plankton samples from the Salton Sea (Plate 3), as well as to specimens from several culture strains (Plate 4, Suppl. Table 1), as described below.

3.2. Systematics

- 477 Division DINOFLAGELLATA (Bütschli 1885) Fensome et al. 1993b
- 478 Class DINOPHYCEAE Pascher 1914
- 479 Subclass PERIDINIPHYCIDAE Fensome et al. 1993b
- 480 Order GONYAULACALES Taylor 1980
- 481 Suborder Gonyaulacineae autonym
- 482 Family uncertain
- 483 Genus *Pentaplacodinium* Mertens, Carbonell-Moore, Pospelova et Head gen. n. (Plate 3)
- 484 *Type*: Plate 3A, the holotype of *Pentaplacodinium saltonense* gen. et sp. nov.
- 485 Diagnosis: A gonyaulacoid gonyaulacinean genus with roundish to slightly polyhedral thecae
- 486 with bearing heavily reticulated plates without appendices appendages. The tabulation is Po,
- 487 Pt, <u>4' or 2'+*2', 5", 6C6c, 6S6s, *65"</u>", 1p, 1"", cover plate is oval.
- 488 Etymology: The name is derived from the Greek words penta meaning five, plax plate, and
- 489 *dino* whirling; with reference to the five precingular plates that characterize this dinoflagellate
- 490 genus.
- 491 *Pentaplacodinium saltonense* Mertens, Carbonell-Moore, Pospelova et Head gen. et sp. n.
- 492 (Plates 3, 4, Figs 2A, 3, 4A)
- 493 *Synonymy:*
- 494 1970 Protoceratium reticulatum (Claparède et Lachmann); Steidinger and Williams, p. 62,
- 495 plate 38, fig. 140a-c.
- 496 1991 Protoceratium reticulatum (Claparède et Lachmann); Al-Muftah, pp. 180–181, figs.
- 497 246–247.
- 498 ? 2002 Protoceratium reticulatum (Claparède et Lachmann); Reifel et al., p. 275.
- 499 2005 Gonyaulax grindleyi Reinecke; Faust et al., p. 110, figs. 2–4.
- ? 2007 Gonyaulax grindleyi Reinecke; Tiffany et al., p. 582.

```
701 ? 2009 ""-Protoceratium globosum" Kofoid <u>etand</u> Michener; Morquecho et al., p. 18, 20, figs.
```

- 502 13–17.
- 503 Diagnosis: Theca roundish to somewhat polyhedral with tabulation Po, Pt, 2'+*2', 35"+*2",
- 504 6C6c, 6S6s, *65", 1p, 1"", with 3" interpreted as *(3"+4"). The theca has an L-type ventral
- organization and dextral torsion. The plates are heavily reticulated with one pore inside each
- reticulation, although two or more pores might be found in reticulations next to a suture. The
- ends of the descending cingulum are displaced by ~2.0 widths. The cysts have an
- 508 <u>approximately roundish spherical</u> central body with a thin pedium and <u>thicker spongy-fibrous</u>
- luxuria. Processes Process distribution apparently have a nointrantabular distribution, are
- Processes fibrous and distally tapering, and have acuminate to minutely expanded distal ends.
- The archeopyle corresponds to the $\frac{\text{third}}{(3''+4'')}$ precingular plate and has a smooth margin
- with rounded angles. The operculum is free.
- 513 *Etymology*: The specific epithet refers to the type locality for this species.
- 514 Type locality: The Salton Sea, California, U.S.A. (station 1 at 33°30.192<u>"</u> N, 115°54.869<u>"</u> W).
- 515 Gene sequence: The 28S and 18S gene sequence of the cell isolated from culture 2E3,
- established from a cyst extracted from surface sediment from station 2 in the Salton Sea
- 517 (Table 1). —GenBank Accession No. XXXXXX MG646301 (18S) and XXXXXX
- 518 MG646323 (28S). Several other strains are considered to belong to the same species (Suppl.
- 519 Table 1).
- 520 Holotype: Hustrated on Plate 3A. The specimen illustrated is on an SEM stub (designated
- 521 CEDiT2017H62) will be deposited curated at the Senckenberg Research Institute and Natural
- 522 History Museum, Centre of Excellence for Dinophyte Taxonomy, Germany.
- 523 *Description: Motile cells observed in the Salton Sea plankton samples* (Plate 3, except D).
- 524 The cell-shape varies from Thecae have a roundish to somewhat polyhedral shape (Plate 3A,
- 525 C). The thecae have and a typical sexiform gonyaulacoid tabulation (sensu Fensome et al.,

1993b, tText-Fig. 64B) with an L-type ventral organization (sensu Fensome et al., 1993b, Text-Fig. 82A, C) and dextral torsion (sensu Fensome et al., 1993b, Text-Fig. 83C). The epitheca was is often somewhat shorter in length than the hypotheca. The plates are reticulated with one pore inside each reticulation, although two or more pores could may occurbe found in reticulations next to a suture. All pores each contain ~3 minute pores (small arrowhead in Plate 3B). The reticulations are faintly expressed on the sulcus and cingulum (Plate 3C). The cell content is brownish-red owing to the presence of chloroplasts (Plate 2A). Several red bodies are present (Plate 2A–C).

The apical pore complex consists of a cover plate surrounded by a pore plate (Plate 7F,

526

527

528

529

530

531

532

533

534

535

536

537

538

539

540

541

542

543

544

545

546

547

548

549

550

H). The oval cover plate, which is often absent (Plate 3B), is relatively broad and is surrounded by the pore plate. The pore plate is perforated by 5–7 large pores. A low apical collar may encompass the pore plate and is formed by the raised edges of the first and second apical plates, and the fourth apical homolog as well (Plate 3F, H). The first and second apical plates (1' and 2') and the fourth apical homolog (*4') are elongated. The first apical plate (1') is rectangular, while whereas the second apical plate (2') and the fourth apical homolog (*4') are six-sided and irregularly shaped (Plate 3B). The third apical homolog (*3') is small and contacts 2' and *4', but in the specimens that were observed it never contacted the apical pore plates (Plate 3B). There is a large ventral pore located posteriorly between 1' and *4' (Plates 3A, B, 4A–C). The precingular series consists of five large plates, where 2" is the largest, *(3"+4") forms the keystone plate, and *6" is the smallest. Plates 1", *(3"+4"), and *5" are five-sided, 2" is four-sided, while *6" is six-sided (the suture with the anterior right sulcal is very small (Fig. 3) (Plates 3B, 4A—C). External views of the theca can could suggest that there would be no contact between the anterior sulcal plate and 1' (e.g., Plate 3A, B). However, pProperly oriented external views and internal views, however, show however a narrow contact between both plates (Plate 3D). This contact between the anterior sulcal plate

and 1', in combination with the contact between *6" and 1' therefore results in an insert configuration (sensu Fensome et al. 1993b, Text-Fig. 62A). The cingulum is left-handed (descending), lined with narrow lists, and comprises six cingular plates. The ends of the cingulum do not overhang, and are displaced by ~2.0 widths (Plates 3A, 4B).

The sulcus is narrow anteriorly and slightly widens posteriorly. It consists of six plates (Plate 3D, Fig. 3) — the first postcingular plate 1" is treated as a sulcal and labeled the anterior left sulcal plate (Ssa). The anterior sulcal plate (Sa) is relatively large and anteriorly intruded between plates 1" and *6" and barely contacts 1' (Plate 3D). The anterior left sulcal plate (Ssa) is similar in size to the anterior right sulcal plate (Sda). Immediately below these two plates, lay the small posterior right sulcal (Sdp) and a much larger plate, the left posterior sulcal. Finally, there occurs the large posterior sulcal (Sp) is found, which presents lines of pores around its sutures with the adjacent non-sulcal plates (Plate 3D, Fig. 3).

The hypotheca is asymmetrical as a consequence of dextral torsion (Plates 3A). There are five homolog postcingular plates. Plate *2" is irregularly shaped and the smallest in the series. All other postcingular plates are large, though *6" is relatively smaller; in addition, they are trapezoidal and four-sided (Plates 3E, 4E). The posterior intercalary plate (1p) bears a conspicuous flange on its right margin (Plates 3A, 4C). The plate overlap is typical for gonyaulacoids, with 3" (in our case *(3"+4")) forming the keystone plate (the plate that overlaps all adjacent plates) oin the epitheca, and *4" forming the keystone plate oin the hypotheca (Fig. 4, Plate 3E).

Cysts from the Salton Sea surface sediments (Plates 1, 5). The central body is approximately spherical roundish. The wall is thick, consisting of a thin, solid pedium that has a smooth inner surface, and a thicker spongy-fibrous luxuria that appears loosely granular in surface view.

Processes are numerous and are solid and fibrous along their entire length, often loosely

fibrous at the base. Process bases are expanded, and larger processes may be concave in lateral profile for at least half of their length. Some closely adjacent processes are joined at the base. Most processes usually have a minute distal expansion, observed under SEM as a concave platform ~1.0 µm or less in diameter with strongly irregular margins that may be approximately perpendicular to the shaft. Alongside these, some processes on most specimens taper to distal points, and such processes occasionally predominate on eyst specimens individual specimens. Processes are mostly of even height, but shorter and thinner processes may be interspersed. The process length/central body ratio is about 0.06. Processes are not evenly spaced, and their parallel alignment and bands devoid of processes observed in many specimens suggest intratabular distribution. However, tThere is however no clear evidence of tabulation except for the archeopyle and often parallel alignment along the cingulum cingular margins. The archeopyle is formed from the loss of plate *(3"+4"), is reduced, and has slightly rounded angles and straight margins. The usual archeopyle that is moderately wide and reflects the precingular thecal plate *(3"+4"), whereas the operculum is released as a single piece, and has well defined to moderately rounded angles and straight margins, as illustrated on Plate 1G-H. -A reentrant angle along the anterior margin of the archeopyle, signaling the fusion of plates 3" and 4", was not seen in the thecal or cyst tabulation of P. saltonense although this might not in fact be expected (see Below, 1987, p. 36, fig. 18a; translated in Fensome et al., 1993a, p. 844). An unusually wide archeopyle that seems to reflect two adjacent precingular thecal plates, *(3"+4") and 2", where the operculum is again released as a single piece, is illustrated on Plate 1C-E. If this interpretation is correct, then the component representing 2" in the archeopyle/operculum is reduced in size, because on the theca the second precingular plate is actually similar or larger in size than the *(3''+4'')plate.

576

577

578

579

580

581

582

583

584

585

586

587

588

589

590

591

592

593

594

595

596

597

598

```
Dimensions: The holotype cell illustrated in Plate 3A, the holotype figure, is 44 µm in length,
601
      41μm in width and 38 μm in depth. Germinated motile cells: length, 48.1 (53.7) 63.4 μm
602
      (SD=6.0, n=5); width, 38.5 (42.4) 47.5 µm (SD= 3.2, n=5). Cells observed in plankton from
603
604
      St. 2 in the Salton Sea: length, 37.8 (46.1) 59.8 \mu m (SD = 5.5, n=28); width, 31.0 (39.5) 48.5
      \mum (SD=4.2, n=28).
605
      Two single cysts germinated to give the identifiable thecae: maximum central body diameter,
606
      52.3 (53.5) 54.7 µm (SD=1.7, n=2); minimum central body diameter, 51.1 (52.2) 53.3 µm
607
      (SD=1.6, n=2); average length of three randomly chosen processes per cyst, 2.4 (3.0) 3.6 µm
608
      (SD=0.4, n=6); process width at base 1.4 (2.2) 2.7 (SD=0.6, n=6) and wall thickness 1.3 (1.7)
609
      2.1 (SD=0.3, n=6). Palynologically treated cysts from surface sediments of the Salton Sea:
610
      maximum central body diameter, 48.6 (56.3) 70.9 µm (SD=5.3, n=23); minimum central body
611
      diameter, 45.7 (52.1) 61.4 µm (SD=3.8, n=22); average length of three processes per cyst, 1.0
612
613
      (3.1) 5.7 μm (SD=1.2, n=66); process width at base 1.0 (2.2) 3.9 (SD=0.6, n=66) and wall
      thickness 0.9 (1.6) 2.4 (SD=0.4, n=66).
614
      Comments: Pentaplacodinium saltonense n. gen et sp. is defined primarily from the characters
615
616
      of the motile stage, these distinguishing it from species of the genus Protoceratium. The
      morphology of several thecae observed from off Yucatan (Gulf of Mexico), the Indian River
617
      Lagoon (Florida, USA), and off Qatar (Persian Gulf) (Table 1; Plate 2) and from cultures
618
      established from cells from Biscayne Bay (Florida, USA) (CCMP1720, CCPM1721), the
619
      Indian River Lagoon (Florida, USA) (CCMP3241, CCMP3243) and the Salton Sea
620
      (California, USA) (CCMP404) (Suppl. Table 13, Plate 1) agree with the description of P.
621
      saltonense given above. Cysts formed from cultures established from a strain from the Indian
622
      River Lagoon (Florida, USA) (CCMP3243) have the same morphologies (Plate 6). The
623
      observed cysts correspond most closely to the fossil based taxon-species Operculodinium
624
      israelianum (Rossignol 1962) Wall 1967 described from the Pleistocene of Israel, and
625
```

Operculodinium psilatum Wall 1967 described from the postglacial (Holocene) of the
Caribbean, p. 111–112, Plate 6, figs. 6–8. However, Operculodinium israelianum has longer
processes (6–10 µm; Rossignol, 1964), and O. psilatum has a psilate surface interrupted by
minute and sparsely distributed processes, and a pronounced cingulum (Wall, 1967). Both
have archeopyles that are less wide than for the cyst of <i>P. saltonense</i> .

3.3. Phylogenetic position of P. saltonense and other studied strains

The SSU rDNA sequences for all *P. reticulatum* strains analysed were identical, forming a distinct clade separated from the *P. saltonense* sequences which were identical to the unidentified Malaysian sequences (Fig. 5). *P. reticulatum* and *P. saltonense* sequences shared 92% nucleotide identity for the V9 region analysed.

For the LSU rDNA V4 analysis (Figure 6), *P. reticulatum* sequences were identical apart from a couple of sporadic nucleotide substitutions which were <u>called_identified_as</u> ambiguous bases by the sequencing software. The unidentified strain from Hawaii had 12 nucleotide substitutions across the 570_bp multiple sequence alignment compared to *P. reticulatum*. *P. saltonense* sequences shared more similarity with the unidentified GgSm strains from Malaysia (96%) compared to that of *P. reticulatum* (94%).

The ITS (intergenic region between ITS1 and 2) was the only marker to resolve intraspecific diversity within the *P. reticulatum* species, with strain E12 (Baffin Bay, Arctic) sharing 98% nucleotide similarity with strain VG0757 isolated from Spain. The phylogeny separates *P. reticulatum* into two large subclades: subclade 1A that regroups several strains from warmer waters, and subclade 1B that regroups several strains from colder waters (Fig 7).

The three phylogenies (Figures 5—7) show that strains identified as *Protoceratium reticulatum* form a monophyletic group (Clade 1), as well as strains identified as *P*.

saltonense that form a clade with the unidentified GgSm strains from Malaysia (Clade 2), as well as the *Ceratocorys* species that form a clade with PrTT strains from Malaysia (Clade 3) supported by high bootstrap values (>70). The unidentified strain from Hawaii does not group with the *Protoceratium reticulatum* or *Pentaplacodinium saltonense* clades. The topology of the trees are is not consistent between the three phylogenies (i.e. the relatedness between clades), however but the three clades identified are consistently formed. The trees furthermore highlight the unexplored diversity within this group of dinoflagellates, and further incubation and plankton studies from these locations should reveal whether the unidentified strains are new species or not.

In addition, the phylogenies show that V9 SSU sequences from cells from Elands Bay (South Africa), (bootstrap values >80%), that have been previously identified as *G. grindleyi* by Reinecke (1967), clusters with *Protoceratium reticulatum* (Fig.-ure_5).

The three studied species of *Ceratocorys* (*C. armata, C. gourreti, C. horrida*) share high nucleotide similarity for the SSU (100%) and LSU sequences (>99% identity) (Figure 6).

4. Discussion

4.1. Comparison of the cell theca of P. saltonense

Pentaplacodinium saltonense differs from Protoceratium reticulatum because it bears five precingular plates, whereas P. reticulatum has six-precingular plates. Furthermore, P. saltonense has a larger cingulum-cingular displacement (2 widths vs. 1 width respectively) and an oval cover plate, as opposed to a sigmoidal cover plate in P. reticulatum. In addition, the theca of P. saltonense is mostly roundish, whereas in P. reticulatum it is always polyhedral. Both species have an insert configuration, but in P. saltonense the contact

between Sa and 1' is very narrow whist whereas in P. reticulatum this contact is wide —_this
causes an evident conspicuous separation between 1" and 6" in <i>P. reticulatum</i> , when in <i>P.</i>
saltonense there is an apparentalmost a small point of contact between those two plates (Plate
4D3D). Gonyaulax grindleyi Reinecke 1967 is here considered shown to be a synonym of P.
reticulatum, as it has been already suggested by von Stosch (1969) and Hansen et al. (1997),
and is now confirmed by the LSU rDNA phylogeny in this study (see below).

Several other *Protoceratium* species have been described (e.g. Schiller, 1937, p. 322–326). *Protoceratium splendens* Meunier 1910 from the Kara Sea has six precingular plates; it is possibly a junior synonym of *Protoceratium reticulatum*, as suggested by Gómez (2012). *Protoceratium aculeatum* (Stein 1883) Schiller 1937 bears antapical spines and an apical horn. *Protoceratium areolatum* Kofoid 1907 and *Protoceratium spinulosum* (Murray and Whitting 1899) Schiller 1937 have fewer reticulations in both the epitheca and hypotheca than *P. saltonense*. Of the five species described by Kofoid and Michener (1911):-), *Protoceratium cancellorum*, *Protoceratium pellucidissimum*, *Protoceratium pepo*, *Protoceratium globosum* and *Protoceratium promissum*, none have has illustrations and it is therefore it is impossible to compare them to *P. saltonense*.

Pentaplacodinium saltonense differs from Ceratocorys anacantha Carbonell-Moore 1996 because it is not as polyhedral. In addition, in contrast to the insert epithecal configuration of *P. saltonense*, *C. anacantha* has an episert type I epithecal configuration, meaning that 1' does not contact the anterior sulcal plate and that 1" and 6" are in contact (Paez-Reyes and Head, 2013).

4.2. Comparison of the cyst of P. saltonense

The cyst of *Pentaplacodinium saltonense* corresponds toompares with *Operculodinium psilatum* because its cysts display an alignment of processes along the cingulum, it bears short processes (1.0–5.7 μm-vs. 2 μm (Wall, 1967) or 0.0–2.9 μm (Head, 1996)), its body diameter is of similar size (45.7–70.9 μm vs. 50–60 μm-(, Wall, 1967)-; or 62–79 μm-(, Head, 1996b)) and its wall thickness (0.9–2.4 vs. 1.4–2.2 μm-(; Head, 1996b)) is similar. *Operculodinium psilatum* differs, however, in having processes that in general are shorter (2 μm, Wall, 1967; 0.0–2.9 μm, Head, 1996b) and sparsely distributed. The cingulum and sulcus are also more conspicuously expressed in *Operculodinium psilatum* (Wall, 1967; Head et al., 1996b), and *O. psilatum* lacks the wide archeopyle of *P. saltonense*.

700

701

702

703

704

705

706

707

708

709

710

711

712

713

714

715

716

717

718

719

720

721

722

723

724

The cyst of P. saltonense is also similar to Operculodinium israelianum (Rossignol 1962) Wall 1967 as described by Rossignol (1964, as *Baltisphaeridium israelianum*); however, although the processes of the latter species are longer (6–10 µm) than of the specimens from Salton Seafor P. saltonense (1.0-5.7 µm). At this time, we do notilt is not presently known whether variation in process length is related to variations in ecology, similar toas demonstrated for the cysts of Lingulodinium polyedra (=Lingulodinium machaerophorum) (Mertens et al., 2009), cysts of Protoceratium reticulatum (Mertens et al., 2011) and cysts of *Pyrodinium bahamense* (=*Polysphaeridium zoharyi*) (Mertens et al., 2015). In addition, tThe process distribution appears to be is-intratabular for the cysts of P. saltonenseand often shows parallel alignments, which is not readily visible and this is likely to be the case also for-on- O. israelianum (e.g., O. cf. israelianum of Head, 1997, fig. 17.2), although the study of topotype material will be needed for confirmation. The rounded angles of the archeopyle in O. israelianum, O. psilatum and the cysts of Pentaplacodinium saltonense, and the shared presence of a spongy-fibrous to fibroreticulate luxuria, accentuate the overall similarities between these cysts, although the relatively wider archeopyle in P. saltonense cysts distinguishes them from these other species.

We t<u>Therefore, it is not considered</u> it not to be a morphological variant of

Operculodinium israelianum. There is reasonable doubt that Operculodinium israelianum is

related to another species: Wall and Dale (1968) (and subsequently Dale, 1983) have

suggested that O. israelianum can be related to Protoceratium reticulatum. Operculodinium

israelianum is very similar toresembles the Miocene Operculodinium centrocarpum

(Deflandre et Cookson 1955) Wall 1967, which is also has a spongy-fibrous luxuria, although

it is somewhat larger (54–80 µm) (; Deflandre and Cookson, 1955) and has longer processes.

Head (1996b) noted an intergradation in size and process length between O. israelianum and
O. centrocarpum in Pleistocene assemblages of eastern England. It is of interest to remark

that specimens from the Ludham borehole intergrade between O. israelianum and O.

centrocarpum Deflandre et Cookson 1955 (Head, 1996). A restudy of both holotypestopotype

material as well as their needed to confirm the range of variability within each species would

be needed to resolve this issue.

The cysts of *P. saltonense* differs from the cystthose of *P. reticulatum*, due to the in having a thick spongy-fibrous luxuria wall-(vs. thin, fibrous luxuria), less developed distal ends of the processes, larger central its-body size diameter (48.6–70.9 μm vs. 33–48 μm; Rochon et al., 1999), and generally shorter process length (1.0–5.7 μm vs. typically 7–14 μm; Rochon et al., 1999)(48.6 (56.3) 70.9 μm (SD=5.3, n=23)) for *P. saltonense*, as opposed to 33–48 μm for *P. reticulatum* (Rochon et al., 1999) and its process length (1.0 (3.1) 5.7 μm (SD=1.2, n=66)) for *P. saltonense*, as opposed to typical 7–14 μm for *P. reticulatum* (Rochon et al., 1999); although the cysts of *P. reticulatum* shows very wide variation vary widely sizes, with formation of some being completely bald cysts (e.g., Mertens et al., 2012a; Jansson et al., 2014).

Several Numerous other *Operculodinium* species have been described and <u>a detailed</u> comparison is given by we refer to Marret and Kim (2009) for their detailed comparison, as; none of these closely resembles the cysts of *P. saltonense*.

4.3. Phylogenetics, evolution and position and relationships of Protoceratium,

Pentaplacodinium, and Ceratocorys

There are sSeveral morphological characteristics of the theca that are important to in understanding the evolution of <u>Protoceratium</u>, <u>Pentaplacodinium</u>, and <u>Ceratocorys</u> the three genera in question (Plate 7). The shape of the cover plate of <u>Ceratocorys</u> is more similar to that of the cover plate of <u>Pentaplacodinium</u>, but less similar to the sigmoidal cover plate of <u>Protoceratium</u>.

Pentaplacodinium and Ceratocorys can be considered closer to Gonyaulax than

Protoceratium, because the anterior intercalary is always well-separated from the apical pore plates, whilst whereas in Protoceratium reticulatum it is closer and has even been suggested to contact the apical pore plates (Hansen et al. 1997). However, It should however-be noted, however, that Protoceratium-reticulatum and Gonyaulax have six precingular plates, whereas whilst Ceratocorys and Pentaplacodinium have five precingulars. So it is not surprising that in the molecular phylogenies, Pentaplacodinium has an intermediate position between Ceratocorys and Protoceratium (Figs. 5–7); the relation to other gonyaulacoids at this time is unclear and further molecular studies of related genera are required, particularly to understand how to resolve the position of Protoceratium at family level. Another issue regards a conflict in the dual nomenclature: the cyst of P. reticulatum and P. saltonense both are considered to belong to the cyst-defined genus Operculodinium, whereas whilst the thecate stages belong to

two different genera; further cyst—theca experiments within this group of related species should help to understand how the <u>genus generic</u> concepts can be <u>made conformrationalized</u>.

In addition, the ITS marker was able to separate two large subclades within *P. reticulatum*: strains that are predominantly associated with warmer waters (Sub-clade 1A), and other strains largely associated with colder waters (Sub-clade 1B) (Fig. 7). Do these subclades reflect pseudocryptic speciation in *Protoceratium reticulatum* as previously suggested by Mertens et al. (2012a)?

Other morphological characteristics of the theca are conserved in *Protoceratium*, *Pentaplacodinium*, and *Ceratocorys* and other gonyaulacoids. For instance, there is no difference in overlap pattern between *Protoceratium*, *Pentaplacodinium*, *Ceratocorys*, *Gonyaulax* and *Lingulodinium* (Fig. 4).

4.4. Biogeography and ecology of P. saltonense

According to the plankton observations, *P. saltonense* can be found in tropical to subtropical regions. We have not observed *P. saltonense* and *P. reticulatum* have not been observed in the same samples, which suggests that both these species possibly inhabit different niches, where *P. saltonense* has a preference for higher temperatures and salinities, and *P. reticulatum* has a preference for somewhat lower temperatures and salinities. This difference would need to be established quantified through culture experiments.

4.5. Toxicity

Strains identified as *Pentaplacodinium saltonense* (CCMP404, CCMP1720 and CCMP1721), have been identified as yessotoxin producers using fluorescence HPLC (Paz et al., 2004). A

later toxin analysis by LC-MS of the same strains was negative (Paz et al., 2007), and the authors considered that these strains had lost their toxicity after a number of years in culture. The produced toxins produced by these strains of *Pentaplacodinium saltonense* are similar to toxins produced by strains we identified as *Protoceratium reticulatum*, all of which are yessotoxin producers, such as strains from Chile (Alvarez et al., 2011), Jervis Inlet, British Columbia, Canada (Cassis, 2005), German Bight, the North Sea (Röder et al., 2011, 2012), Okkirai Bay, Japan (Koike et al., 2006) and Spain (Paz et al., 2007, 2013). The presence of *P. saltonense* in the Salton Sea has been considered a potential causative agent of mortality events in the Salton Sea indicates that it is potentially toxic (Reifel et al., 2002, whom identified it as *Protoceratium reticulatum*). However, there have not however not been no reports of toxic events knowingly involving *P. saltonense*.

Several other studies <u>have</u> investigated the toxicity of strains <u>that</u> they designate as *Protoceratium reticulatum*, but <u>for which we could not verify</u> the identifications <u>could not be</u> <u>verified</u> (e.g., Satake et al., 1999; Ciminiello et al., 2003; Samdal et al., 2004; Finch et al., 2005; Eiki et al., 2005; Mitrovic et al., 2005; Guerrini et al., 2007; Suzuki et al., 2007).

5. Conclusions

Pentaplacodinium saltonense gen. et sp. nov. is described from the Salton Sea (CA, USA). The distinct cover plate (similar to *Ceratocorys*, but sigmoidal in *Protoceratium*), five precingular plates (also fiveas in *Ceratocorys*, but six in *Protoceratium*), the very narrow contact between 1' and Sa (wide contact in *Protoceratium*, no contact in *Ceratocorys*), a more roundish rounded eell-thecal shape, the displacement of the cingulum by two widths (vs. one width in *Protoceratium*), as well as the clear separation and distances seen in the three phylogenies, justifies the creation of a new genus. The chorate cysts of *P. saltonense* bear

short processes often with parallel alignments. These cysts,—correspond to the cyst-defined species genus Operculodinium Wall 1967, and are most similar to O. israelianum and O. psilatum-with short processes and often parallel alignments. The geographic distribution of mMotile stages of Pentaplacodinium saltonense wasare confirmed infrom four widely dispersed locations and its distribution is therefore considered, suggesting a subtropical to tropical distribution for this species. , and does not overlap with that of Protoceratium reticulatum and Pentaplacodinium saltonense are not known to inhabit the same environments. Similar to As with the -yessotoxin-producing Protoceratium reticulatum, Pentaplacodinium saltonense is potentially a yessotoxin producer, as shown by previous studies.

Note added: While this paper has been going through the process of final acceptance to

Harmful Algae, another study was accepted (Salgado et al., accepted) that addresses similar

Acknowledgements

scientific questions.

V.P. and M.J.H. <u>each</u> acknowledge support from a Discovery Grant of the Natural Sciences and Engineering Research Council of Canada (NSERC). K.B.A. and M.F. <u>acknowledge were</u> supported from by the Instituto Nacional de Investigación y Tecnología Agraría y Alimentaria of the Spanish Government (project RTA2005-00109-00-00). We thank CINVESTAV, Merida is thanked for supplying information about the Yucatan samples. Beatriz Paz is acknowledged for interesting discussions on toxicity. We would like to thank Anke Kremp, Maija Hutunnen, Jacob Larsen, Paul Hargraves and Ximena Vivanco <u>are thanked forkindly providing provided</u> plankton samples. Nancy Lewis and Margaret Beaton <u>are thanked</u>

forgenerously sharing shared information on strains from Nova Scotia, Canada. We are 847 grateful Thanks to to Captain Brown, crew of the MSV Strickland, Ms. Sarah Thornton, and 848 EOS313-2010 (University of Victoria) students are all thanked for their participation in 849 sediment sample collection. Paul Hargraves is acknowledged forkindly providing provided 850 SEM images of *P. saltonense* from cultures established from the Indian River Lagoon. 851 852 853 References 854 Adachi, M., Sako, Y., Ishida, Y., 1996. Analysis of Alexandrium (Dinophyceae) species using 855 sequences of the 5.8S ribosomal DNA and internal transcribed spacer regions. J. Phycol. 32, 856 424–432. 857 858 859 Al-Muftah, A.R., 1991. Dinoflagellates of Qatari Waters waters (PhD thesis), vol. 2. University College of North Wales, Bangor University, pp. 261. 860 861 Alvarez, G., Uribe, E., Díaz, R., Braun, M., Mariño, C., Blanco, J., 2011. Bloom of the 862 Yessotoxin producing dinoflagellate *Protoceratium reticulatum* (Dinophyceae) in Northern 863 northern Chile. J. Sea Res. 64, 427–434. 864 865 Andree, K.B., Quijano-Scheggia, S.; Fernández, M., Elandaloussi, L.M., Garcés, E., Camp, 866 J., Diogene, J.—(, 2011.) Quantitative PCR Coupled with Melt-melt Curve curve 867 Analysis analysis for Detection detection of Selected Selected Pseudonitzschia spp. 868 (Bacillariophyceae) from the Northwestern northwestern Mediterranean Sea. App. and Env. 869 870 Micro, 77(5) 1651–1659.

872	Balech, E., 1980. On thecal morphology of dinoflagellates with special emphases emphasis on
873	circular and sulcal plates. Anales del Centro de Ciencias del Mar y Limnología, UNAM 7(1),
874	57–68.
875	
876	Balech, E., 1988. Los dinoflagellados del Atlantico sudoccidental. Publicaciones Especiales
877	Instituto Español de Oceanografia 1, 1–310.
878	
879	Below, R., 1987. Evolution und Systematik von Dinoflagellaten-Zysten aus der Ordnung
880	Peridiniales. I. Allgemeine Grundlagen und Subfamilie Rhaetogonyaulacoideae (Familie
881	Peridiniaceae). Palaeontographica, Abteilung B 205, 1–164.
882	
883	Bergh, R.S., 1881. Der Organismus der Cilioflagellaten. Eine phylogenetische Studie.
884	Morphologisches Jahrbuch 7(2), 177–288.
885	
886	Bogus, K. Mertens, K.N., Lauwaert, J., Harding, I.C., Vrielinck, H., Zonneveld, K.A.F.,
887	Versteegh, G.J.M., 2014. Differences in the chemical composition of organic-walled
888	dinoflagellate resting cysts from phototrophic and heterotrophic dinoflagellates. J. Phycol. 50,
889	254–266.
890	
891	Bolch, C.J.S., 1997. The use of polytungstate for the separation and concentration of living
892	dinoflagellate cysts from marine sediments. Phycologia 37, 472–478.
893	
894	Braarud, T., 1945. Morphological observations on marine dinoflagellate cultures (Porella
895	perforata, Goniaulax tamarensis, Protoceratium reticulatum). Avh. Utgit. Nor. Vidensk.
896	Akad. Oslo Mat.–Naturvidensk. Kl. 11, 1–18.

including the erection of three new genera of unarmoured dinoflagellates. Phycologia 39, 922 302-317. 923 924 Deflandre, G., Cookson, I.C., 1955. Fossil microplankton from Australian Late Mesozoic and 925 Tertiary sediments. Australian Journal of Marine and Freshwater Research 6, 242–313. 926 927 Dodge, J.D., 1989. Some revisions of the family Gonyaulaceae (Dinophyceae) based on a 928 929 scanning electron microscope study. Bot. Mar. 32, 275–298. 930 Eiki, K., Satake, M., Koike, K., Ogata, T., Mitsuya, T., Oshima, Y., 2005. Confirmation of 931 yessotoxin production by the dinoflagellate *Protoceratium reticulatum* in Mutsu Bay. Fish. 932 Sci. 71, 633–638. 933 934 Faust, M.A., Litaker, R.W., Vandersea, M.W., Kibler, S.R., Tester, P.A. 2005. Dinoflagellate 935 936 diversity and abundance in two Belizean coral-reef mangrove lagoons: a test of Margalef's 937 mandala. Atoll Research Bulletin 534, 105-132. 938 Fensome, R.A., Gocht, H., Stover, L.E., Williams, G.L., 1993a. The Eisenack Catalog of 939 940 Fossil Dinoflagellates. New Series. Volume 2. p.829–1461; E. Schweizerbart'sche Verlagsbuchhandlung, Stuttgart, Germany. 941 942 Fensome, R., Taylor, F., Norris, G., Sarjeant, W., Wharton, D., Williams, G., 1993b. A 943 classification of fossil and living dinoflagellates. Micropaleontology Special Publication 7, 1– 944 945 245.

- 947 Ferrari, R.L., Weghorst, P., 1997. Salton Sea 1995 hydrographic GPS survey. U.S. Bureau of
- 948 Reclamation, Water Resources Services. Denver, Colorado, 23 pp.

- 950 Finch, S.C., Wilkins, A.L., Hawkes, A.D., Jensen, D.J., MacKenzie, A.L., Beuzenberg, V.,
- 951 Quilliam, M.A., Olseng, C.D., Samdal, S.A., Aasen, J., Selwood, A.I., Cooney, J.M., Sandvik,
- 952 M., Miles, C.O., 2005. Isolation and identification of (44-R,S)-44,55-dihydroxyyessotoxin
- 953 from *Protoceratium reticulatum*, and its occurrence in extracts of shellfish from New
- 254 Zealand, Norway and Canada. Toxicon 46, 160–170.

955

- 956 Gómez, F., 2012. A checklist and classification of living dinoflagellates (Dinoflagellata,
- 957 Alveolata). CICIMAR Océanides 27(1), 65–140.

958

- Guerrini, F., Ciminiello, P., Dell'Aversano, C., Tartaglione, L., Fattorusso, E., Boni, L.,
- Pistocchi, R., 2007. Influence of temperature, salinity and nutrient limitation on yessotoxin
- production and release by the dinoflagellate *Protoceratium reticulatum* in batch-cultures.
- 962 Harmful Algae 6, 707–717.

963

- 964 Guillou, L., Nezan, E., Cueff-Gauchard, V., Barbier, G., 2002. Genetic diversity and
- molecular detection of three toxic dinoflagellate genera (Alexandrium, Dinophysis and
- 966 Karenia) from French coasts. Protist. 153, 223-238

967

- 968 Hall, T.A., 1999. BioEdit: a user-friendly biological sequence alignment editor and analysis
- 969 program for Windows 95/98/NT. Nucl. Acids. Symp. Ser. 41+, 95—98.

971	Hansen, G., Moestrup, Ø. and Roberts, K.R., 1997. Light and Electron electron Microscopical
972	microscopical Observations observations on Protoceratium reticulatum (Dinophyceae). Arch.
973	Protistenk. 147, 381–391.
974	
975	Hansen, G., Daugbjerg, N., Henriksen, P., 2000. Comparative study of <i>Gymnodinium</i>
976	mikimotoi and Gymnodinium aureolum, comb. nov. (= Gyrodinium aureolum) based on
977	morphology, pigment composition, and molecular data. J. Phycol., 36, 394–410.
978	
979	Head, M.J., Wrenn, J.H. (eds.), 1992. A forum on Neogene–Quaternary dinoflagellate cysts:
980	The edited transcript of a round table discussion held at the Second Workshop on Neogene
981	dinoflagellates. In: Head, M.J., Wrenn, J.H. (eds.), Neogene and Quaternary Dinoflagellate
982	Cysts and Acritarchs. American Association of Stratigraphic Palynologists Foundation,
983	Dallas, Texas, pp. 1–31.
984	
985	Head, M.J., 1996a. Modern dinoflagellate cysts and their biological affinities. In: Jansonius,
986	J., McGregor, D.C. (eds.), Palynology: principles and applications. American Association of
987	Stratigraphic Palynologists Foundation, Dallas, Texas, vol. 3, pp. 1197–1248.
988	
989	Head, M.J., 1996b. Late Cenozoic dinoflagellates from the Royal Society borehole at
990	Ludham, Norfolk, eastern England. J. Paleontol. 70, 543–570.
991	
992	Head, M.J., 1997. Thermophilic dinoflagellate assemblages from the mid Pliocene of eastern
993	England. J. Paleontol. 71, 165–193.
994	

995	Hennissen, J.A.I., Head, M.J., De Schepper, S., Groeneveld, J., 2017. Dinoflagellate cyst
996	paleoecology during the Pliocene–Pleistocene climatic transition in the North Atlantic.
997	Palaeogeogr., Palaeoclimatol., Palaeoecol. 470, 81–108.
998	
999	Holdren, G.C., Montaño, A., 2002. Chemical and physical characteristics of the Salton Sea,
1000	California. Hydrobiol. 473, 1–21.
1001	
1002	Howard, M.D.A., Smith, G.J., Kudela, R.M., 2009. Phylogenetic relationship of yessotoxin-
1003	producing dinoflagellates based on the Large Subunit and Internal Transcribed Spacer
1004	Ribosomal DNA Domains domains. Appl. Environ. Microbiol. 75, 54–63.
1005	
1006	Jansson, IM., Mertens, K.N., Head, M.J. with contributions from de Vernal, A., Londeix, L.,
1007	Marret, F., Matthiessen, J., Sangiorgi, F., 2014. Statistically assessing the correlation between
1008	salinity and morphology in cysts produced by the dinoflagellate Protoceratium reticulatum
1009	from surface sediments of the North Atlantic Ocean, Mediterranean-Marmara-Black Sea
1010	region, and Baltic-Kattegat-Skagerrak estuarine system. Palaeogeogr. Palaeoclimatol.
1011	Palaeoecol. 399, 202–213.
1012	
1013	Kofoid, C.A., 1907. Reports on the scientific results of the expedition to the eastern tropical
1014	Pacific, in charge of Alexander Agassiz, by the U.S. Fish Commission steamer ",Albatross",
1015	", from October, 1904, to March, 1905, Lieut Commander L.M. Garrett, U.S.N.,
1016	commanding. IX. New species of dinoflagellates. Bull. Mus. Comp. Zool. 50(6), 163–207.
1017	
1018	Kofoid, C.A., Michener, J.R., 1911. Reports on the Scientific Results of the Expedition to the
1019	Eastern Tropical Pacific, in Charge of Alexander Agassiz, by the U.S. Fish Commission

```
Steamer "-, Albatross ALBATROSS", from October 1904, to March, 1906, Lieut. L.M.
1020
        Garrett, U.S.N., Commanding. XXII. New genera and species of
1021
       Dinoflagellates dinoflagellates. Bull. Mus. Comp. Zool. 54(7), 267–302.
1022
1023
       Koike, K., Horie, Y., Suzuki, T., Kobiyama, A., Kurihara, K., Takagi, K., Kaga, S.N.,
1024
       Oshima, Y., 2006. Protoceratium reticulatum in northern Japan: environmental factors
1025
1026
       associated with seasonal occurrence and related contamination of yessotoxin in scallops. J.
1027
       Plankton Res. 28, 103–112.
1028
       Lane, D. J., 1991. 16S/23S sequencing. In: Nucleic Acid Technologies in Bacterial
1029
       Systematic. Stackebrandt E, Goodfellow M (Eds) pp. 115–175, Wiley, NY.
1030
1031
1032
       Marret, F., Kim, S.-Y., 2009. Operculodinium aguinawense sp. nov., a dinoflagellate cyst
       from the late Pleistocene and recent sediments of the east Equatorial Atlantic ocean.
1033
1034
       Palynology 33, 125–139.
1035
       Matsuoka, K., McMinn, A., Wrenn, J.H., 1997. Restudy of the Holotype of Operculodinium
1036
       centrocarpum (Deflandre & Cookson) Wall (Dinophyceae) from the Miocene of Australia,
1037
       and the Taxonomy of Related related Species Palynology 21, 19–33.
1038
1039
       Medlin, L. K, Elwood, H. J., Stickel, S., Sogin, M. L., 1988. The characterization of
1040
1041
       enzymatically amplified eukaryotic 16S-like rRNA-coding regions. Gene. 71, 491–499.
1042
1043
       Mertens, K.N., Ribeiro, S., Bouimetarhan, I., Caner, H., Combourieu Nebout, N., Dale, B., de
        Vernal, A., Ellegaard, M., Filipova, M., Godhe, A., Goubert, E., Grøsfjeld, K., Holzwarth, U.,
1044
```

- Kotthoff, U., Leroy, S.A.G., Londeix, L., Marret, F., Matsuoka, K., Mudie, P.J., Naudts, L.,
- 1046 Peña-Manjarrez, J.L., Persson, A., Popescu, S.-M., Pospelova, V., Sangiorgi, F., van der
- Meer, M., Vink, A., Zonneveld, K.A.F., Vercauteren, D., Vlassenbroeck, J., Louwye, S.,
- 2009. Process length variation in cysts of a dinoflagellate, *Lingulodinium machaerophorum*,
- in surface sediments: Investigating its potential as salinity proxy. Mar. Micropaleontol. 70,
- 1050 54–69.

- Mertens, K.N., Dale, B., Ellegaard, M., Jansson, I.-M., Godhe, A., Kremp, A., Louwye, S.,
- 2011. Process length variation in cysts of the dinoflagellate *Protoceratium reticulatum*, from
- surface sediments of the Baltic-Kattegat-Skagerrak estuarine system: a regional salinity
- 1055 proxy. Boreas 40, 242–255.

1056

- Mertens, K.N., Bringué, M., Van Nieuwenhove, N., Takano, Y., Pospelova, V., Rochon, A.,
- de Vernal, A., Radi, T., Dale, B., Patterson, R.T., Weckström, K., Andrén, E., Louwye, S.,
- Matsuoka, K., 2012a. Process length variation of the cyst of the dinoflagellate *Protoceratium*
- 1060 reticulatum in the North Pacific and Baltic-Skagerrak region: calibration as an annual density
- proxy and first evidence of pseudo-cryptic speciation. J. Quaternary Sci. 27, 734–744.

1062

- Mertens, K.N., Price, A., Pospelova, V., 2012b. Determining the absolute abundance of
- dinoflagellate cysts in recent marine sediments II: further tests of the *Lycopodium* marker-
- grain method. Rev. Palaeobot. Palynol. 184, 74–81.

- Mertens, K.N., Wolny, J., Carbonell-Moore, C., Bogus, K., Ellegaard, M., Limoges, A., de
- Vernal, A., Gurdebeke, P., Omura, T., Mohd. A Al-Muftah, A., Matsuoka, K., 2015.
- Taxonomic re-examination of the toxic armoured dinoflagellate *Pyrodinium bahamense* Plate

- 1906: can morphology or LSU sequencing separate *P. bahamense* var. *compressum* from var.
- 1071 bahamense? Harmful Algae 41, 1–24.

- Meunier A., 1910. Microplancton des mers de Barents & de Kara. In: Duc d''Orléans
- 1074 Campagne Arctique de 1907. Bulens, Bruxelles, 355 pp.

1075

- 1076 Mitrovic, S.M., Hamilton, B., McKenzie, L., Furey, A., James, K.J., 2005. Persistence of
- 1077 yessotoxin under light and dark conditions. Mar. Environ. Res. 60, 397–401.

1078

- Morquecho, L., Góngora-González, D.T. & Okolodkov, Y.B. 2009. Cyst-theca relationships
- 1080 of Gonyaulacales and Peridiniales (Dinophyceae) from Bahía Concepción, Gulf of California.
- 1081 Acta Botanica Mexicana 88, 9–29.

1082

- Murray, G., Whitting, F.G., 1899. New Peridiniaceae from the Atlantic. Trans. Linn. Soc.
- 1084 Lond., Bot. 5, 321–342.

1085

- 1086 Paez-Reyes, M., Head, M.J., 2013. The Cenozoic Gonyaulacacean gonyaulacacean
- 1087 Dinoflagellate dinoflagellate Genera genera Operculodinium Wall, 1967 and Protoceratium
- Bergh, 1881 and Their their Phylogenetic Phylogenetic Relationships relationships. J. Paleo.
- 1089 87(5), 786–803.

1090

- Paz, B., Riobó, P., Fernández, M.L., Fraga, S., Franco, J.M., 2004. Production and release of
- 1092 yessotoxins by the dinoflagellates *Protoceratium reticulatum* and *Lingulodinium polyedrum*
- in culture. Toxicon 44, 251–258.

```
Paz, B., Riobó, P., Ramilo, I., Franco, J.M., 2007. Yessotoxins profile in strains of
1095
        Protoceratium reticulatum from Spain and USA. Toxicon 50, 1–17.
1096
1097
1098
       Paz, B., Daranas, A.H., Norte, M., Riobó, P., Franco, J.M., Fernández, J.J., 2008.
1099
        Yessotoxins, a Group group of Marine marine Polyether polyether Toxinstoxins: an
        Overviewoverview. Mar. Drugs 6, 73–102.
1100
1101
       Paz, B., Blanco, J., Franco, J.M., 2013. Yessotoxins production during the culture of
1102
       Protoceratium reticulatum strains isolated from Galician Rias Baixas (NW Spain). Harmful
1103
1104
       algae 21–22, 13–19.
1105
        Pospelova, V., Esenkulova S., Johannessen S.C., O'Brien O'Brien M. C. &, Macdonald
1106
1107
        R.W., 2010. Organic-walled dinoflagellate cyst production, composition and flux from 1996
        to 1998 in the central Strait of Georgia (BC, Canada): a sediment trap study. Mar.
1108
       Micropaleontol. 75:, 17–37.
1109
1110
       Reifel, K.M., McCoy, M.P., Rocke, T.E., Tiffany, M.A., Hurlbert, S.H., Faulkner, D.J., 2002.
1111
1112
       Possible importance of algal toxins in the Salton Sea, California. Hydrobiologia 473, 275–
       292.
1113
1114
1115
       Reinecke, P., 1967. Gonyaulax grindleyi sp. nov.: a dinoflagellate causing a red tide at Elands
       Bay, Cape Province in December 1966. S. Afr. J. Bot. 33, 157–160.
1116
1117
        Rochon, A., de Vernal, A., Turon, J.-L., Matthiessen, J., Head, M.J., 1999. Distribution of
1118
       Recent-recent dinoflagellate cysts in surface sediments from the North Atlantic Ocean and
1119
```

adjacent areas in relation to sea-surface parameters. American Association of Stratigraphic

Palynologists, Contributions Series, no. 35, 146 p.

1122

1123 Röder, K., Fritz, N., Gerdts, G., Luckas, B., 2011. Accumulation and Depuration depuration of Yessotoxin-yessotoxin in Two-two Bivalvesbivalves. J. Shellfish Res. 30, 167–175.

1125

1127

1128

Röder, K., Hantzsche, F.M., Gebühr, C., Miene, C., Helbig, T., Krock, B., Hoppenrath, M.,

Luckas, B., Gerdts, G., 2012. Effects of salinity, temperature and nutrients on growth, cellular

characteristics and yessotoxin production of *Protoceratium reticulatum*. Harmful Algae 15,

1129 59–70.

1130

1131

1132

Rossignol, M., 1964. Hystrichosphères du Quaternaire en Méditerranée orientale, dans les

sédiments Pléistocènes et les boues marines actuelles. Revue de micropaléontologie 7, 83–99.

1133

1135

Sala-Pérez, M., Alpermann, T.J., Krock, B., Tillmann, U., 2016. Growth and bioactive

secondary metabolites of arctic *Protoceratium reticulatum* (Dinophyceae). Harmful Algae 55,

1136 85–96.

1137

Saldarriaga, J.F., Taylor, F.J.R., Cavalier-Smith, T., Menden-Deuer, S., Keeling, P.J., 2004.

Molecular data and the evolutionary history of dinoflagellates. Eur. J. Protistol. 40, 85–111.

1140

Salgado, P., Figueroa, R.I., Ramilo, I., Bravo, I., 2017. The life history of the toxic marine

dinoflagellate *Protoceratium reticulatum* (Gonyaulacales) in culture. Harmful Algae 68, 67–

1143 81.

1144

1145	Salgado, P., Fraga, S., Rodríguez, F., Riobó, P., accepted. Ceratocorys mariaovidiorum sp.
1146	nov. (Gonyaulacales), a new dinoflagellate species previously reported as Protoceratium
1147	reticulatum. Journal of Phycology, accepted.
1148	
1149	Samdal, I.A., Naustvoll, L.J., Olseng, C.D., Briggs, L.R., Miles, C.O., 2004. Use of ELISA to
1150	identify <i>Protoceratium reticulatum</i> as a source of yessotoxin in Norway. Toxicon 44, 75–82.
1151	
1152	Satake, M., Ichimura, T., Sekiguchi, K., Yoshimatsu, S., Oshima, Y. 1999. Confirmation of
1153	Yessotoxin and 45,46,47-Trinoyessotoxin production by <i>Protoceratium reticulatum</i> collected
1154	in Japan. Natural toxins 7, 147–150.
1155	
1156	Schiller, J., 1937. Dinoflagellatae (Peridineae) in monographischer Behandlung. In: Dr. L.
1157	Rabenhorst's Kryptogamen-Flora von Deutschland, Österreich und der Schweiz. Bd. 10(3).
1158	Teil 2(3), pp. 321–480.
1159	
1160	Scholin, C.A., Herzog, M., Sogin, M., Anderson, D.M. 1994. Identification of group- and
1161	strain-specific genetic markers for globally distributed Alexandrium (Dinophyceae). II.
1162	Sequence analysis of a fragment of the LSU rRNA gene. J. Phycol. 30, 999–1011.
1163	
1164	Steidinger K.A. & J. Williams (1970) Dinoflagellates. Memoirs of the Hourglass Cruises 2,
1165	1–251. Published by Marine Research Laboratory, Florida Department of Natural Resources,
1166	St. Petersburg, Florida.
1167	

- Stein, F. von, 1883. Der Organismus der Infusionsthiere. 3. Abt. Der Organismus der
- Arthrodelen Flagellaten nach eigenen Forschungen in systematischer Reihenfolge bearbeitet.
- 2. Hälfte. Einleitung und Erklärung der Abbildungen, W. Engelmann, Leipzig, pp. 1–30.

- Stosch, H.A. von, 1969. Dinoflagellaten aus der Nordsee I. Über *Cachonina niei* Loeblich
- 1173 (1968), Gonyaulax grindleyi Reinecke (1967) und eine Methode zur Darstellung von
- 1174 Peridineenpanzern. Helgoländ. Wiss. Meer. 19, 558–568.

1175

- Suzuki, T., Horie, Y., Koike, K., Satake, M., Oshima, Y., Iwataki, M., Yoshimatsu, S., 2007.
- 1177 Yessotoxin analogues in several strains of *Protoceratium reticulatum* in Japan determined by
- 1178 liquid chromatography–hybrid triple quadrupole/linear ion trap mass spectrometry. Journal of
- 1179 Chromatography A 1142, 172–177.

1180

- Takano, Y., Horiguchi, T., 2004. Surface ultrastructure and molecular phylogenetics of four
- unarmoured heterotrophic dinoflagellates, including the type species of the genus *Gyrodinium*
- 1183 (Dinophyceae). Phycol. Res. 52, 107–116.

1184

- 1185 Takano, Y., Horiguchi, T. 2006. Acquiring scanning electron microscopical, light
- microscopical and multiple gene sequence data from a single dinoflagellate cell. J. Phycol. 42,
- 1187 251–256.

1188

- Tamura, K., Stecher, G., Peterson, D., Filipski, A., Kumar, S. 2013 MEGA6: Molecular
- 1190 Evolutionary Genetics Analysis Version 6.0. Mol Biol Evol. 30: 2725—2729

```
Tiffany, M.A., González, M.R., Swan, B.K., Reifel, K.M., Watts, J.M., Hurlbert, S.H. 2007.
1192
        Phytoplankton dynamics in the Salton Sea, California, 1997-1999. Lake and Reservoir
1193
        Management 23, 582–605.
1194
1195
        Wall, D., 1967. Fossil microplankton in deep-sea cores from the Caribbean Sea.
1196
        Palaeontology 10, 95-123.
1197
1198
1199
        Wall, D., Dale, B., 1966. "Living fossils" in western Atlantic plankton. Nature 211 (5053),
        1025-1026.
1200
1201
        Wall, D., Dale, B., 1968. Modern dinoflagellate cysts and evolution of the Peridiniales.
1202
1203
       Micropaleontology 14, 265–304.
1204
        Watts, J.M., Swan, B.K., Tiffany, M.A., Hurlbert, S.H., 2001. Thermal, mixing and oxygen
1205
1206
       regimes of the Salton Sea, California, 1997–1999. Hydrobiol. 162, 159–176.
1207
        Wołoszyńska, J., 1929. Dinoflagellatae Polskiego Baltyku i blot nad Piasnica. Archivum
1208
1209
       Hydrobiologji i Rybactwa 3, 153–278 [In Polish].
1210
        Yamaguchi, A., Yoshimatsu, S., Hoppenrath, M., Wakeman, K.C., Kawai, H. 2016.
1211
       Molecular phylogeny of the benthic dinoflagellate genus Amphidiniopsis and its relationships
1212
        with the family Protoperidiniaceae. Protist. 167, 568—583.
1213
1214
1215
        Zonneveld, K.A.F., Marret, F., Versteegh, G.J.M., Bogus, K., Bonnet, S., Bouimetarhan, I.,
       Crouch, E., de Vernal, A., Elshanawany, R., Edwards, L., Esper, O., Forke, S., Grøsfjeld, K.,
1216
```

- Henry, M., Holzwarth, U., Kielt, J.-F., Kim, S.-Y., Ladouceur, S., Ledu, D., Chen, L.,
- 1218 Limoges, A., Londeix, L., Lu, S.-H., Mahmoud, M.S., Marino, G., Matsouka [sic], K.,
- Matthiessen, J., Mildenhal [sic], D.C., Mudie, P., Neil, H.L., Pospelova, V., Qi, Y., Radi, T.,
- 1220 Richerol, T., Rochon, A., Sangiorgi, F., Solignac, S., Turon, J.L., Verleye, T., Wang, Y.,
- Wang, Z., Young, M., 2013. Atlas of modern dinoflagellate cyst distribution based on 2405
- datapoints. Rev. Palaeobot. Palynol. 191, 1–197.

Figure captions 1224 1225 Figure 1. Sites of studied plankton samples and cultured strains containing thecate stages of 1226 1227 Pentaplacodinium saltonense (in red) and Protoceratium reticulatum (in blue). The locations are listed in Table 1. 1228 1229 1230 Figure 2. Line drawings of extant members of the subfamily *Cribroperidinioideae* in dorsal view to show the dextral torsion typical of these gonyaulacoids. A. Pentaplacodinium 1231 saltonense. B. Protoceratium reticulatum. C. Lingulodinium polyedra. Labeling of tabulation 1232 1233 follows a modified Kofoid system that recognizes homologs. 1234 **Figure 3.** Line drawing of sulcal area of *Pentaplacodinium saltonense*. FP: flagellar pore; Sa: 1235 1236 anterior sulcal plate; Sda: right anterior sulcal plate; Sdp: right posterior sulcal plate; Ssa: anterior left sulcal plate; Ssp: posterior left sulcal plate; Sp: posterior sulcal plate; c: cingular 1237 1238 plates. 1239 Figure 4. Line drawings of epithecal overlapping plate patterns of gonyaulacoids discussed in 1240 this paper. Arrows indicate direction of overlap. A. Pentaplacodinium saltonense. B. 1241 Protoceratium reticulatum. C. Lingulodinium polyedra. D. Ceratocorys horrida. E. 1242 Gonyaulax spinifera. 1243 1244 **Figure 5**. Neighbour-joining tree of *P. reticulatum*, *P. saltonense* and related strains 1245 sequenced in this study and sequences from Genbank based on an 80 bp alignment of the V9 1246 1247 region of the SSU gene. Bootstrap values were retrieved from 1000 replicates and those >70% are indicated at the nodes for neighbour-joining and maximum likelihood respectively. 1248

Strain names are indicated followed by their geographic origin and accession number (Genbank).

Figure 6. Neighbour-joining tree of *P. reticulatum, P. saltonense* and related strains sequenced in this study and sequences from Genbank based on a 571 bp alignment of the V4 region of the LSU gene. Bootstrap values were retrieved from 1000 replicates and those >70% are indicated at the nodes for neighbour-joining and maximum likelihood respectively. Strain names are indicated followed by their geographic origin and accession number (Genbank).

Figure 7. Neighbour-joining tree of *P. reticulatum*, *P. saltonense* and related strains sequenced in this study and sequences from Genbank based on a 356 bp alignment of the ITS 1–2 region. Bootstrap values were retrieved from 1000 replicates and those >70% are indicated at the nodes for neighbour-joining and maximum likelihood respectively. Strain names are indicated followed by their geographic origin and accession number (Genbank).

Plate Captions

Plate 1. Light microscope images of *Pentaplacodinium saltonense* based on cyst-theca experiment from the Salton Sea. A. Living cyst from the Salton Sea St. 1. B–F. Germinated cyst from St. 2 (culture 2E3 used for single-cell PCR). B. Cross section, showing attached operculum. C. Focus on elongated simple operculum reflecting plates 2"+*(3"+4"). D–E. Focus on archeopyle, after removal of operculum. F. Cross section showing processes. G–I. Germinated cyst from St. 2 (culture 1A7 used for single-cell PCR). G. Focus on precingular

archeopyle reflecting plate *(3"+4"), showing attached operculum. H. Focus on operculum. I.

Cross section, showing opened operculum. Scale bars = $20 \mu m$.

1275

1274

1276 Plate 2. Light microscope images of cyst-theca experiment from the Salton Sea. A–I. Images

of living cells of *Pentaplacodinium saltonense* germinated from cyst depicted in Plate 1, Figs.

B-F (culture 2E3). A. Globular cell. B. Angular cell. C. Fusiform cell. D. Epitheca. E.

Hypotheca. F. Ventral view showing configuration of apical plates. G–I. Sulcal plates. Scale

1280 bars = $20 \mu m$.

1281

1282

1283

1284

1285

1286

1287

1288

1289

1290

1278

1279

Plate 3. Scanning electron microscope images of *Pentaplacodinium saltonense*, all different

cells from the Salton Sea, except D. A. Holotype. Ventral view. Arrowhead points to ventral

pore between plates 1' and *4'. Arrow shows flange on plate 1p. B. Apical view, missing the

cover plate. Small arrowhead points to small pores inside the thecal pores. Large arrowhead

points to ventral pore between plates 1' and *4'. Small arrowhead points to the three minute

pores inside most pores. C. Dorsal view, showing dextral torsion. Note the cell roundness. D.

Sulcal plates of a cell from culture SSCAP K-1479 (Indian River Lagoon, Florida).

Arrowhead shows the narrow point of contact between the Sa and 1' plates. E. Antapical

view. Scale bars A–C, $E = 10 \mu m$; $D = 5 \mu m$.

1291

1292

1293

1294

1295

1296

1297

Plate 4. Scanning electron microscope images of *Pentaplacodinium saltonense* from the

Indian River Lagoon. A. Apical view of a cell from culture SSCAP K-1479. Arrowhead

points to ventral pore between plates 1' and *4'. B. Same specimen as in A. Ventral view.

Arrowhead points to ventral pore between plates 1' and *4'. C. A different cell from a

plankton sample courtesy of Paul Hargraves. Ventral view. Arrowhead points to ventral pore

between plates 1' and *4'. Arrow shows flange on plate 1p. D. Ventral view of a cell from a

culture established by Paul Hargraves. E. Antapical view of a cell from the same culture as in D. E. Apical view of a cell from the same culture as in D. D–F: SEMs by Paul Hargraves.

Scale bars = $10 \mu m$.

Plate 5. Scanning electron microscope images of cysts of *Pentaplacodinium saltonense* extracted from Salton Sea sediment (St. 2) using palynological methods. A–C. Views showing shape of archeopyle, reflecting plate *(3"+4"). D. Specimen that is torn along the cingulum. E. Specimen showing alignment of processes along the cingulum. F. Specimen with relatively large openings in cyst wall. G. Specimen with distinct intratabular processes. H. Specimen with relatively coarsely reticulated wall surface. I. Internal view of smooth cyst wall. Scale bars = 10 µm.

Plate 6. Scanning electron microscope images of cysts of *Pentaplacodinium saltonense* formed in culture of strain 3243 (Indian River Lagoon). A. Specimen showing preformed archeopyle and margins of principal archeopyle suture with reduced ornament. B. Specimen with attached thecal plate. C. Specimen with partly developed processes. D. Specimen with processes clearly reflecting tabulation. E. Specimen with preformed archeopyle. F. Specimen showing reflection of the sulcus. G–H. Specimen with well-developped wall texture. I. Wall texture of specimen with 'spider-web' microreticulation. Scale bars = $10 \mu m$, except H, I, scale bars = $1 \mu m$.

Plate 7. Scanning electron microscope images of *Protoceratium reticulatum* cells and of the apical pore plates of the gonyaulacoids discussed in this study. A. *Protoceratium reticulatum*. Cell from Greenland, ventral view. B. Same cell in apical view. C. *Protoceratium reticulatum*. Cell from Elands Bay, South Africa. Dorsal view, note the dextral torsion. D. Apical pore

plates of a different cell of *Protoceratium reticulatum* (Greenland). E. Apical pore plates of a cell of *Ceratocorys horrida* (Central equatorial Pacific). F. Apical pore plates of a cell of *Pentaplacodinium saltonense* from culture SSCAP K-1479 (Indian River Lagoon). G. Apical pore plates of a cell of *Ceratocorys gourretii*. H. Apical pore plates of another cell of *Pentaplacodinium saltonense* from culture SSCAP K-1479 (Indian River Lagoon). Scale bars A-C = 10 μm; D-H = 5 μm.

Figure 1 Click here to download high resolution image

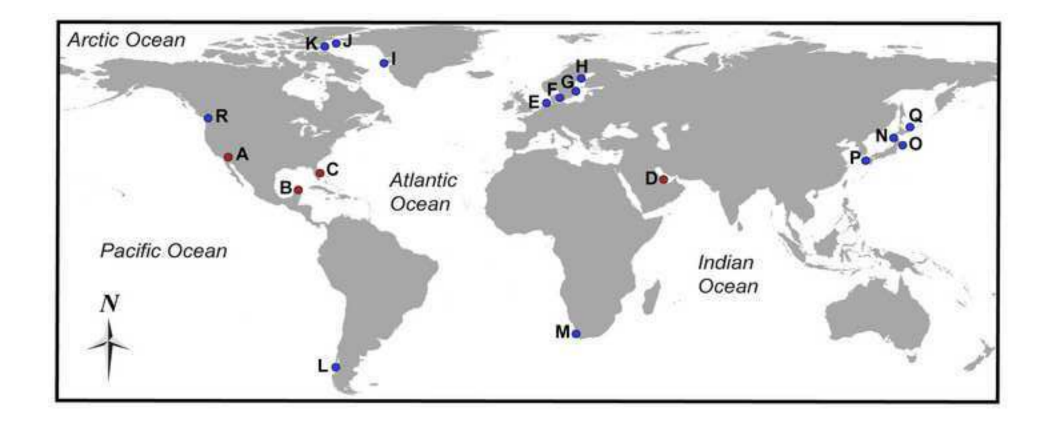


Figure 2 Click here to download high resolution image

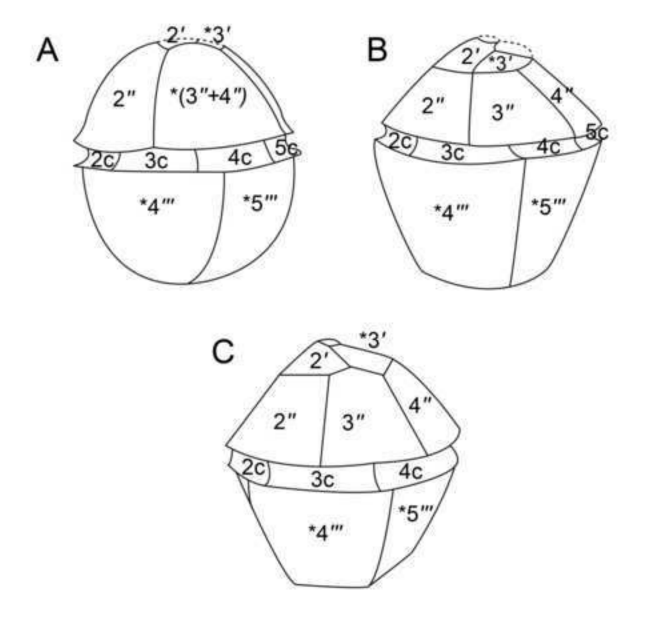


Figure 3
Click here to download high resolution image

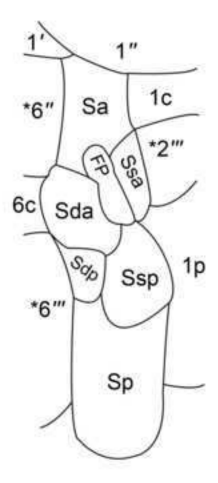


Figure 4 Click here to download high resolution image

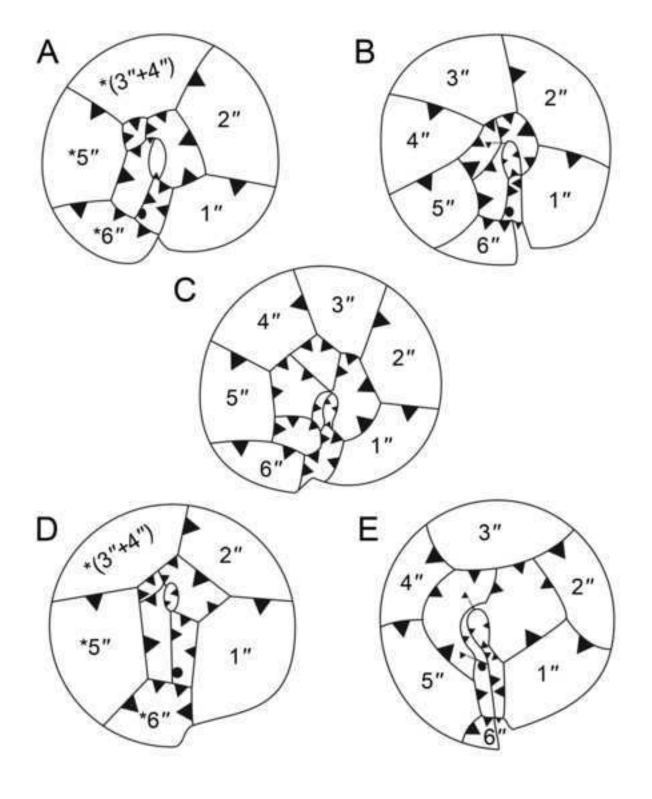


Figure 5
Click here to download high resolution image

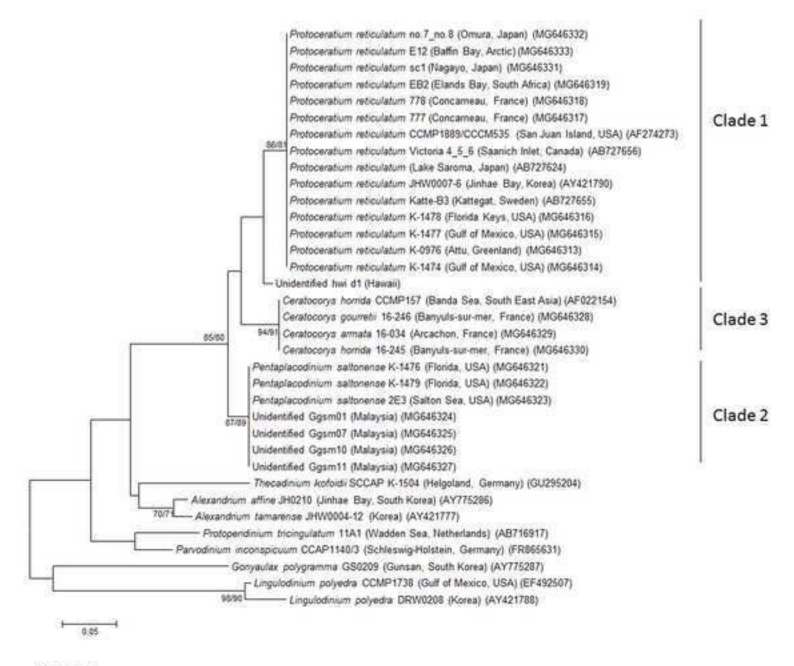


Figure 5

Figure 6
Click here to download high resolution image

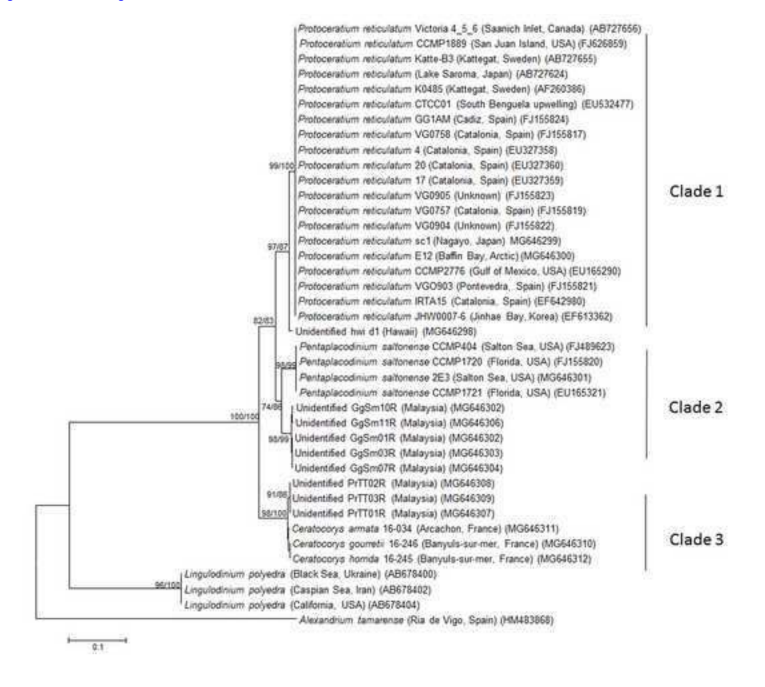


Figure 6

Figure 7
Click here to download high resolution image

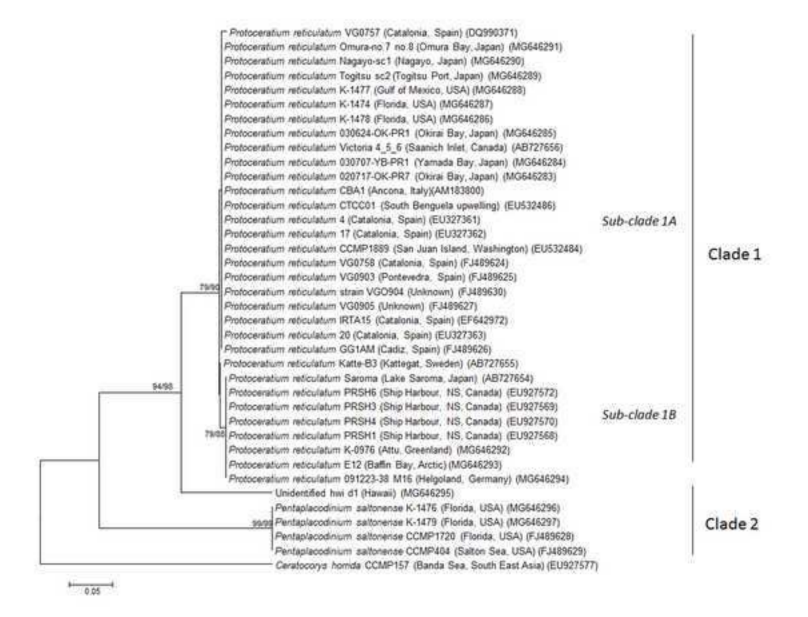


Figure 7

Plate 1 Click here to download high resolution image

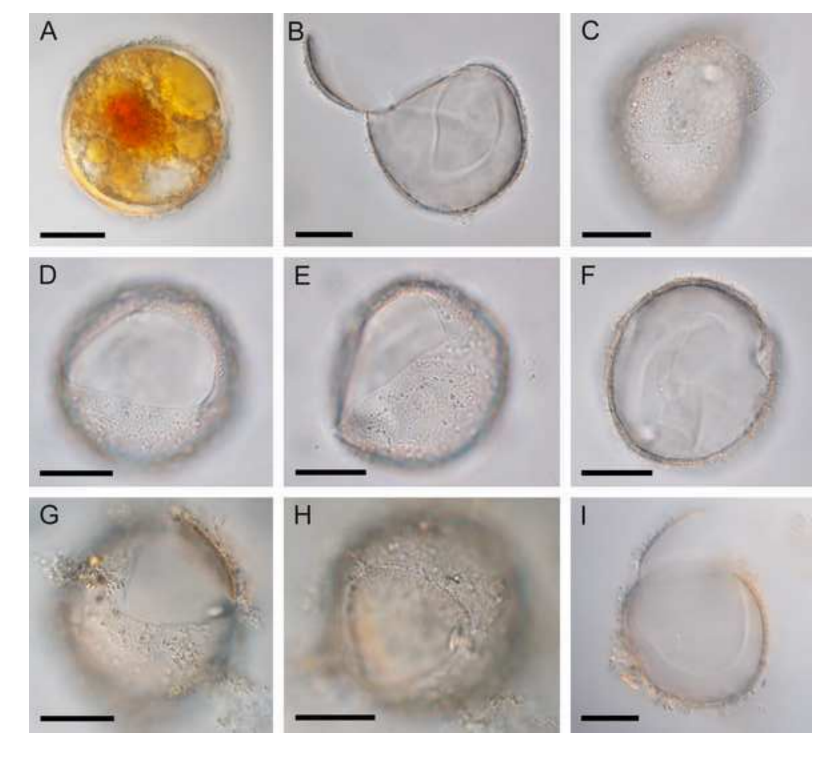


Plate 2 Click here to download high resolution image

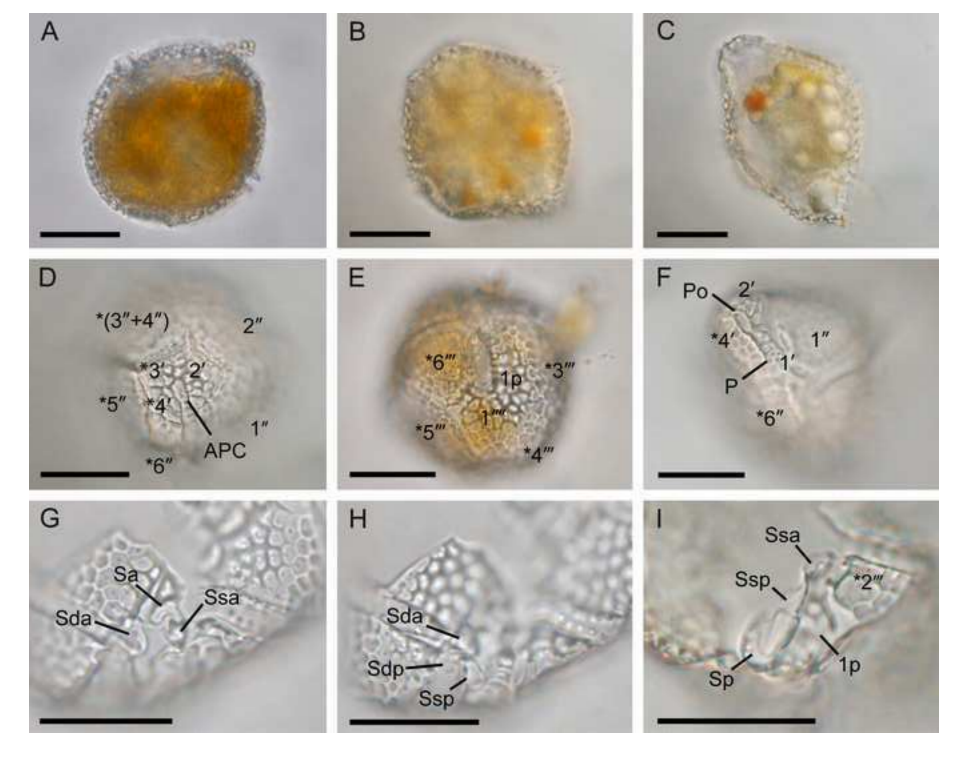


Plate 3 Click here to download high resolution image

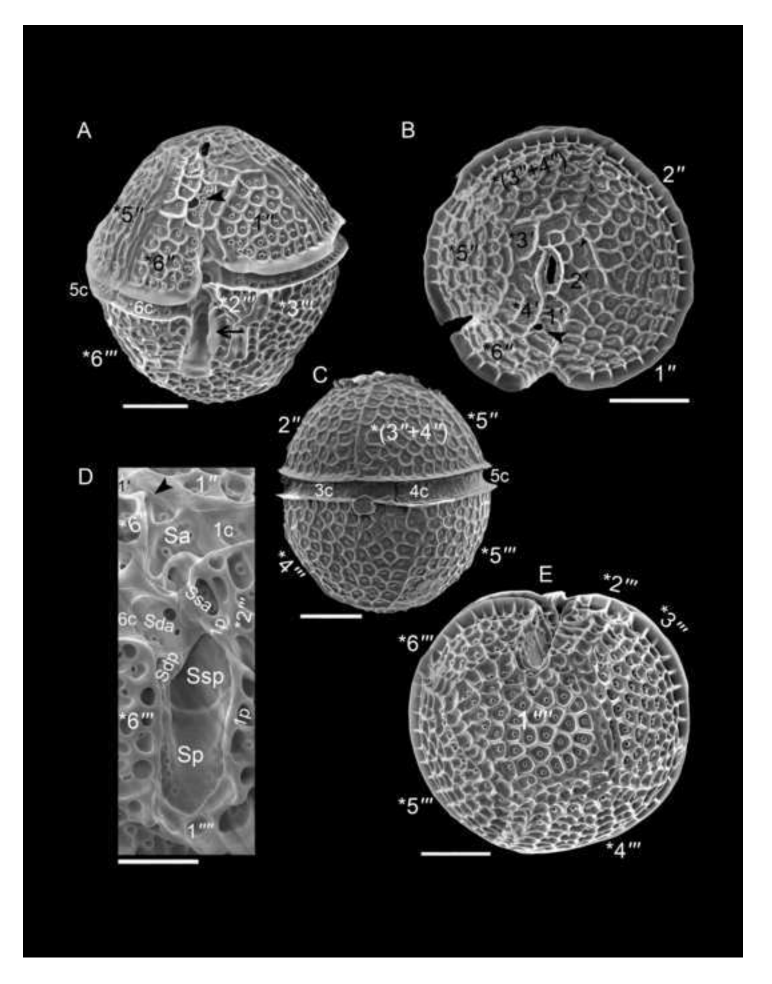


Plate 4 Click here to download high resolution image

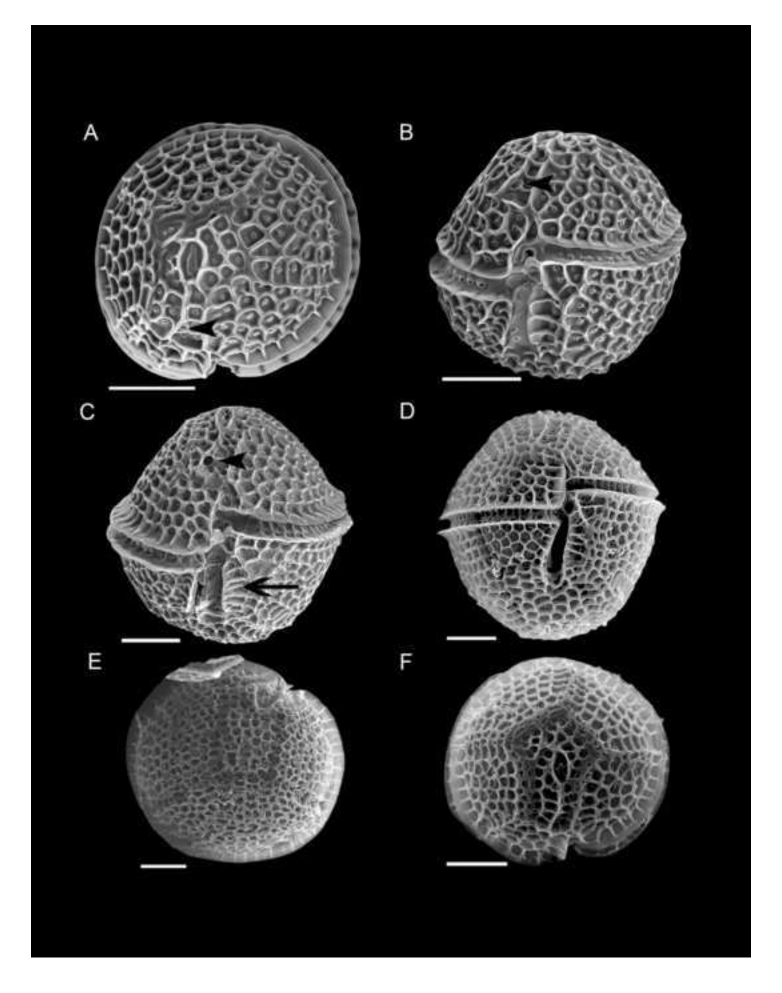


Plate 5 Click here to download high resolution image

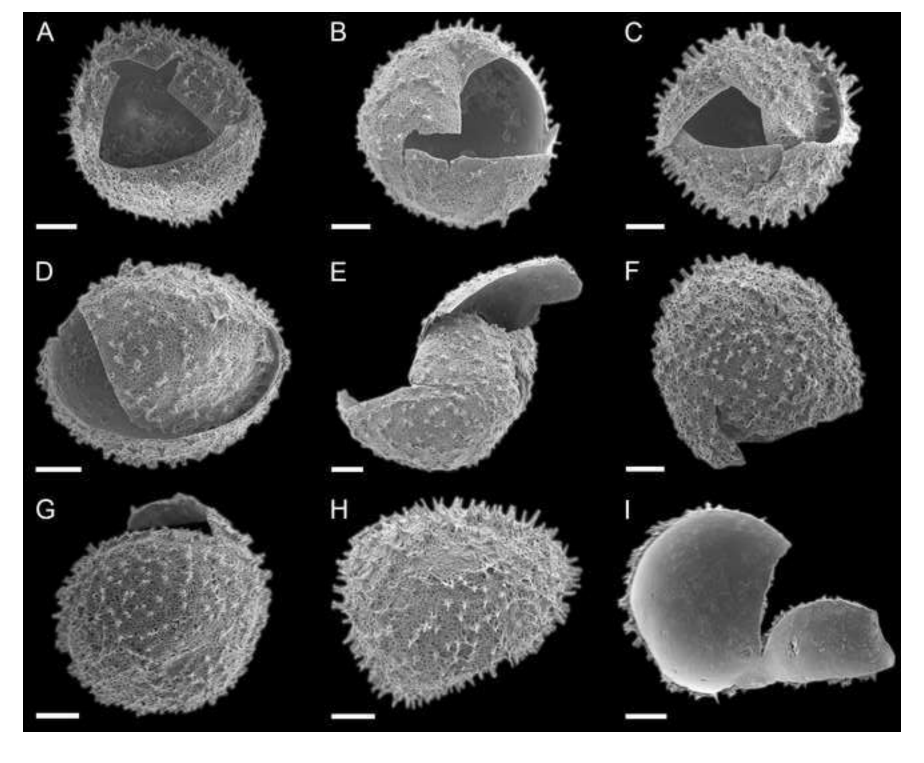


Plate 6 Click here to download high resolution image

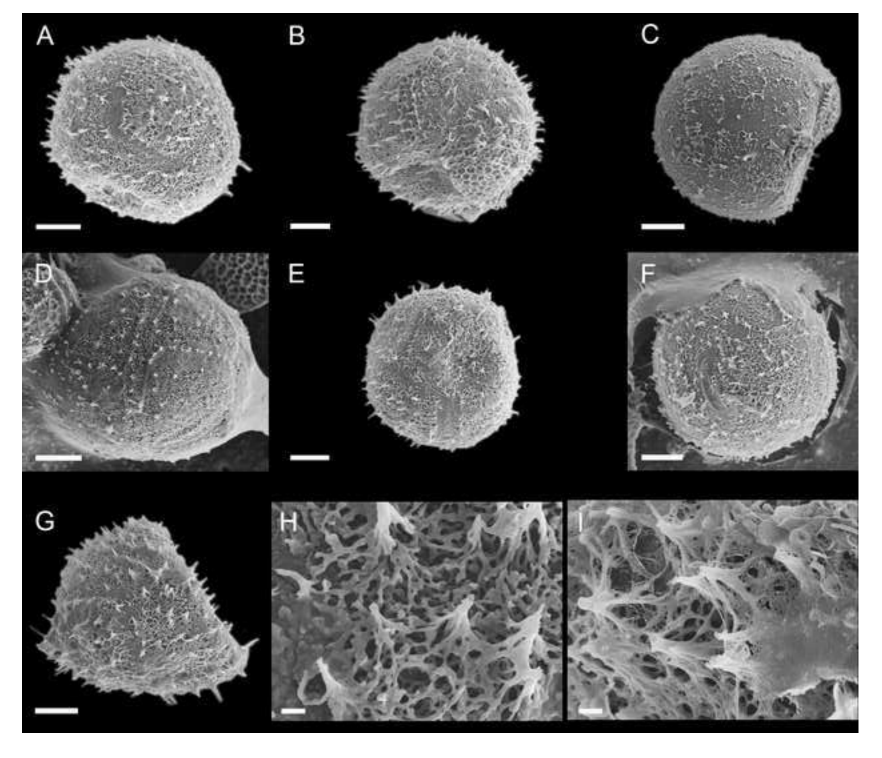


Plate 7 Click here to download high resolution image

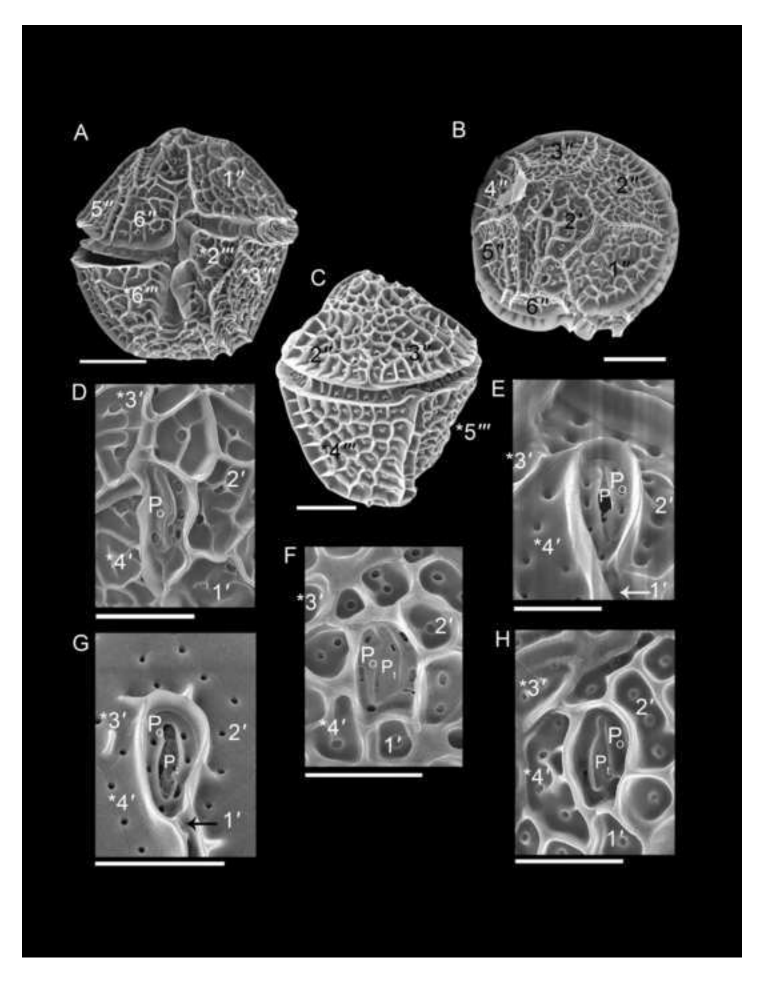


Table 1. Site location of plankton samples investigated, location mark on Figure 1, sampling date, latitude, longitude, sea surface salinity (psu), sea surface temperature ($^{\circ}$ C), sampling device, used fixative, and name of persons who did the sampling.

Sampling site	Location mark on Figure 1	Sampling date	Latitude (°)	Longitude (°)	SSS (psu)	SST (°C)	Sampling device	Fixative used	Species present	Sampled by
Salton Sea, St. 1, California, USA	A	24-Oct-09	33,50	-115,91	>40.2 (62)	23,1	Plankton net \geq 20 μm	Ethanol 100%	PS	KM, VP, MH, MCCM
Salton Sea, St. 2, California, USA	A	24-Oct-09	33,50	-115,91	>40.2 (64)	25,5	Plankton net > 20 μm	Ethanol 100%	PS	KM, VP, MH, MCCM
Salton Sea, St. 3, California, USA	A	24-Oct-09	33,50	-115,92	>40.2 (65)	25,5	Plankton net > 20 μm	Ethanol 100%	PS	KM, VP, MH, MCCM
Salton Sea, St. 4, California, USA	A	24-Oct-09	33,50	-115,91	>40.2 (56)	30,3	Plankton net $> 20 \ \mu m$	Ethanol 100%	PS	KM, VP, MH, MCCM
Salton Sea, St. 5, California, USA	A	24-Oct-09	33,50	-115,91	>40.2 (65)	~30	Plankton net > 20 μm	Ethanol 100%	PS	KM, VP, MH, MCCM
Off Yucatan, St. 4, Gulf of Mexico, Mexico	В	19-May-09	21,39	-88,08	31,0	24,1	Plankton tow >20 μm	Formaldehyde	PS	YO
Off Yucatan, Gulf of Mexico, Mexico	В	19-May-09	21,40	-88,84	31,0	24,1	Plankton tow >20 μm	Formaldehyde	PS	YO
Indian River Lagoon, St. TR, Florida, USA	С	28-May-08	27,50	-80,34	37,4	29,5	Plankton tow >20 μm	Formalin 2%	PS	PH
Off Qatar, Persian Gulf	D	Sept. 1991	25,29	51,54	38-43	20-35	Plankton net	Formalin 5%	PS	AA
North Sea, Helgoland, Germany	E	3-Jun-02	54,19	7,90	32,0	12,5	Plankton net >20 µm	Lugol	PR	MHO
Kattegat, St. Central, Denmark	F	NA	56,92	11,28	NA	NA	Plankton net	Lugol & Formalin	PR	JL
Kattegat, St. 431, Denmark	F	22-Jun-05	55,77	12,75	NA	NA	Plankton net	Lugol	PR	JL
Kattegat, St. 925, Denmark	F	17-Aug-00	56,08	11,02	NA	NA	Plankton net	Lugol	PR	JL
Baltic Sea, St. F64, Finland	G	20-Aug-10	60,18	19,13	5,7	17,6	Plankton net	Lugol	PR	AK, MHU
Gulf of Bothnia, Baltic Sea, St. US5B, Finland	Н	20-Aug-10	62,58	19,98	5,1	16,0	Plankton net	Lugol	PR	AK, MHU
Western Greenland, St. 516, Denmark	I	30-Jul-12	69,20	-54,10	33,2	6,9	Water Bottle-CTD	Formaldehyde	PR	UT
Baffin Bay, St. 2008-029-0039/9/0039A, Canada	J	6-Sep-08	76,57	-73,96	31,0	2,2	Plankton net >20 µm	Formaldehyde	PR	AR
Baffin Bay, St. 2008-029-027A, Canada	J	5-Sep-08	77,29	-74,34	30,8	2,2	Plankton net >20 µm	Formaldehyde	PR	AR
Baffin Bay, St. 2008-029-0043A, Canada	K	7-Sep-08	75,58	-78,63	31,0	2,0	Plankton net >20 µm	Formaldehyde	PR	AR
Baffin Bay, St. 2008-029-0035A, Canada	J	6-Sep-08	76,33	-71,43	31,0	4,5	Plankton net >20 μm	Formaldehyde	PR	AR
Off Puerto Aguirre, Chile	L	21-May-06	-44,99	-73,53	28,8	10,0	Plankton net > 26 µm	Formaldehyde	PR	XV
Elands Bay, South Africa	M	16-Mar-13	-32,31	18,32	NA	NA	Plankton net	Formaldehyde	PR	MP
Off Cape Town, South Africa	M	NA	-33,89	18,42	NA	NA	Plankton net	Lugol + formalin Glutaraldehyde	PR	JL
Mutsu Bay, Aomori, Japan	N	5-Apr-10	40,92	141,12	32,7	5,9	Plankton net >20 μm	1% Glutaraldehyde	PR	KK
Okkirai Bay, Iwate, Japan	О	16-Jul-03	39,08	141,85	32,6	15,8	Plankton net >20 μm	5% Glutaraldehyde	PR	KK
Okkirai Bay, Iwate, Japan	0	28-Aug-03	39,08	141,85	32,3	20,0	Plankton net >20 μm	5% Glutaraldehyde	PR	KK
Okkirai Bay, Iwate, Japan	0	23-Jun-04	39,08	141,85	32,7	16,0	Plankton net >20 μm	5% Glutaraldehyde	PR	KK
Okkirai Bay, Iwate, Japan	0	12-Aug-04	39,08	141,85	33,5	23,0	Plankton net >20 μm	5%	PR	KK
Omura Bay, Nagayo-ura, Japan	P	9-May-11	32,85	129,87	32,7	20,2	Plankton net >20 μm	Formalin	PR	KMA
Omura Bay, Inoura, Japan	P	12-May-11	33,05	129,74	25,6	18,5	Plankton net >20 μm	Formalin	PR	KMA
Omura Bay, Togitsu Port, Japan	P	23-May-11	32,85	129,87	32,6	19,5	Plankton net >20 μm	Formalin	PR	KMA
Kagoshima Bay, St. 1, Japan	P	20-Jun-11	31,55	130,57	23,8	22,9	Plankton net >20 μm	Formalin	PR	KMA
Saroma Lake (Lagoon), St. 1, Japan	Q	22-Jul-11	44,12	143,82	32,2	17,1	Plankton net	Ethanol	PR	KM

Saanich Inlet, St. S2, BC, Canada Saanich Inlet, St. S5, BC, Canada	R R	13-Jul-10 13-Jul-10	48,55 48,71	-123,53 -123,46	27,8 27,8	14,8 13,6	Plankton net Plankton net	Formaldehyde Formaldehyde	PR PR	VP VP
Saanich Inlet, St. S3, BC, Canada	R	13-Jul-10	48,59	-123,48	27,8	14,8	Plankton net	Formaldehyde	PR	VP
Saanich Inlet, St. S4.5, BC, Canada	R	14-Jul-10	48,67	-123,49	28,0	13,7	Plankton net	Formaldehyde	PR	VP
Saanich Inlet, St. S11, BC, Canada	R	14-Jul-10	48,73	-123,54	28,0	13,7	Plankton net	Formaldehyde	PR	VP
Saanich Inlet, St. S4, BC, Canada	R	13-Jul-10	48,63	-123,49	27,4	15,1	Plankton net	Formaldehyde	PR	VP
Saanich Inlet, St. Pat UVic 262, BC, Canada	R	10-Aug-11	48,65	-123,44	29,0	17,8	Plankton net	Formaldehyde	PR	VP
Saanich Inlet, St. Pat UVic 264, BC, Canada	R	21-Aug-11	48,65	-123,44	31.0	17,9	Plankton net	Formaldehyde	PR	VP

Abbreviations: NA = Not available, PS = Pentaplacodinium saltonense, PR = Protoceratium reticulatum, AK = Anke Kremp, AR = André Rochon, AA = Abdulrahman Al-Muftah, MCCM = Consuelo Carbonell-Moore, JL = Jacob Larsen, KK = Kazuhiko Koike, KM = Kenneth Neil Mertens, KMA = Kazumi Matsuoka, MH= Martin J. Head, MHO = Mona Hoppenrath, MHU=Maija Huttunen, MP = Maya Pfaff, PH = Paul Hargreaves, UT = Urban Tillmann, VP = Vera Pospelova, XV = Ximena Vivanco, YO = Yuri B. Okolodkow

Supplementary Material

 $Suppl.\ Table\ 1.\ Culture\ strains\ and\ cells\ or\ cyst\ picked\ for\ SEM\ and/or\ phylogenetic\ analysis.$

CODI (Strain ID)	Identified here by its morphology as	Geographic Origin	Latitude (°N)	Longitude (°E)	Isolation date	Isolated by	LSU (28S) Genbank	ITS Genbank	SSU (18S) Genbank	Sequenced in this study b
CCMP404 = ALO011	P. saltonense	Salton Sea (California)	33.375	-116.0	1966	Dodson, A.	EU532476	FJ489629	FJ489629***	
Polton Coo 2E2 *	D	Salton San (California)	33°	1150 54 970	10/11/2012	Montono V	MC(4(201	Not	MG(4(222	Att- W
Salton Sea 2E3 *	P. saltonense	Salton Sea (California)	30.122'	-115° 54.879'	19/11/2013	Mertens, K.	MG646301	sequenced	MG646323	Aika Yamaguchi
CCMP1720 = K-1475 =	P. saltonense	Biocoura Boy (Florido)	25.8	-80.3333	15/02/1005	Hamamariaa D	FJ155820	FJ489628	FJ489628***	
ALO013	P. sattonense	Biscayne Bay (Florida)	25.8	-80.3333	15/02/1995	Hargraves, P.	FJ155820		FJ489028***	
CCMP1721 = K-1480 =	D - I	Discours Desc (Florida)	25.8	-80.3333	01/06/1004	II D	EU165321	Not	N-4	
ALO012	P. saltonense	Biscayne Bay (Florida)	25.8	-80.3333	01/06/1994	Hargraves, P.	EU105321	sequenced	Not sequenced	
CCMP3241 = K-1479	P. saltonense	Indian River Lagoon (Florida)	27.83	-80.45	11/06/2008	Hargraves, P.	Not sequenced	3.500.4030	MG646322	Andrea Highfield / Decla
CCMP3241 = K-1479	r. sationense	indian River Lagoon (Fiorida)	21.83	-60.43	11/00/2008	naigiaves, r.	Not sequenced	MG646297	MG040322	Schroeder
CCMD2242 IV 1476	D - I	Indian Discoul account (Florida)	27.92	90.45	05/06/2000	II D	N-4	3.500.4030.0	3400040331	Andrea Highfield / Decla
CCMP3243 = K-1476	P. saltonense	Indian River Lagoon (Florida)	27.83	-80.45	05/06/2008	Hargraves, P.	Not sequenced	MG646296	MG646321	Schroeder
										Andrea Highfield / Decla
CCMP3031 = K-1474	P. reticulatum	Off coast Florida, Gulf of Mexico	25.0167	-81.4003	NA	NA	Not sequenced	MG646287	MG646314	Schroeder
Victoria-no.4_5_6 *	P. reticulatum	Brentwood Bay, Saanich Inlet	48.57	123.47	Oct. 2011	Mertens, K.	AB727656	AB727656	AB727656	
CCMP1889 = K-0634 =		Friday Harbor, San Juan Island,								
CCCM535	NA	Washington USA	48.544	-123.01	1983	Taylor, F.J.R.	EU532475	EU532484	FJ626858***	
										Andrea Highfield / Decla
CCMP2776 = K-1477	P. reticulatum	Gulf of Mexico, U.S.A.	25,0167	-81,4003	NA	Sinigalliano, C.	EU165290	MG646288	MG646315	Schroeder
										Andrea Highfield / Decla
CCMP3113 = K-1478	P. reticulatum	Marquesa Keys, Florida Keys	-24.58	-82.1	NA	NA	Not sequenced	MG646286	MG646316	Schroeder
/GO757	NA	Ebro Delta (Catalonia, Spain)	NA	NA	NA	Fernandez-Tejedor, M.	FJ155819	DQ990371	DQ990371***	
/GO758	P. reticulatum**	Alfacs Bay, Ebro Delta (Catalonia Spain)	NA	NA	NA	Fernandez-Tejedor, M.	FJ155817	FJ489624	FJ489624***	
VGO903 = ALO014	P. reticulatum**	Ría de Ponteverde (Bueu)	NA	NA NA	NA NA	Fernandez-Tejedor, M.	FJ155821	FJ489625	FJ489625***	
VGO903 = ALO014 VGO904	NA	NA	NA	NA NA	NA NA	Fernandez-Tejedor, M.	FJ155822	FJ489630	FJ489630***	
VGO904 VGO905	NA NA	NA NA	NA NA	NA NA	NA NA	Fernandez-Tejedor, M.	FJ155823	FJ489627	FJ489627***	
IRTA015	P. reticulatum	Ebro Delta (Catalonia Spain)	NA NA	NA NA	NA NA	Fernandez-Tejedor, M.	EF642980	EF642972	EF642972***	
20			NA NA	NA NA				EU327363		
	NA NA	Catalan Coast	NA NA	NA NA	NA NA	Fernandez-Tejedor, M.	Not sequenced FJ155824		EU327363***	
GG1AM		La Atunara (Cádiz, Spain)		NA NA	NA	Fernandez-Tejedor, M.		FJ489626	FJ489626***	
17 1	NA	Catalan Coast	NA	· ·	NA	Fernandez-Tejedor, M.	Not sequenced	EU327362	EU327362***	
ŀ	NA	Catalan Coast	NA	NA	NA	Fernandez-Tejedor, M.	Not sequenced	EU327361	EU327361***	
Katte-B3 *	D C L ((() ())	Water of Com Pale and	57.5	11.8	N 2011	Masters V	A D727.655	A D727655	A D 727 65 5	
	P. reticulatum (cyst-based)	Kattegat, Swedish coast	37.3	11.8	Nov. 2011	Mertens, K.	AB727655	AB727655	AB727655	
091223-38_M16_Protocer3- TS1	P. reticulatum	Helgoland, Germany	54.19	7.9	avr-03	Hoppenrath, M.	Not sequenced	MG646294	Not sequenced	Karin Röder
	1. Tellellitation	Trongomma, Germanny	5.,17	,,,,	u.i. 03	торрешин, т.	riot sequenced		rior sequenced	111111111111111111111111111111111111111
Lake Saroma *	P. reticulatum (cyst-based)	Lake Saroma, Japan	44.12	143.87	Aug. 2011	Mertens, K.	AB727654	AB727654	AB727654	
	-				-					Andrea Highfield / Decla
K-0976	P. reticulatum	Attu, Greenland	67.924068	-53.649824	21.08.2005	Moestrup, Ø.	Not sequenced	MG646292	MG646313	Schroeder
Arctic E12 = K1-1-1 - K-1-3										
F	P. reticulatum	Station 323, Northern Baffin Bay	74.12	79.45	02-mai-11	Mertens, K.	MG646300	MG646293	MG646333	Yoshihito Takano
ζ-0485	P. reticulatum	Southern Kattegat, Bouy St.	56.20	12.04	03-avr-89	Hansen, G.	AF260386	Not sequenced	Not sequenced	
PRSH6 (NRC Halifax)	P. reticulatum	Ship harbour, NS, Canada	NA	NA	NA	Ferrell, J. F.	Not sequenced	EU927572	EU927572***	
PRSH3 (NRC Halifax)	P. reticulatum	Ship harbour, NS, Canada	NA	NA NA	NA NA	Ferrell, J. F.	Not sequenced	EU927569	EU927569***	
PRSH4 (NRC Halifax)	P. reticulatum	Ship harbour, NS, Canada Ship harbour, NS, Canada	NA NA	NA NA	NA NA	Ferrell, J. F.	Not sequenced	EU927570	EU927570***	
RSH1 (NRC Halifax)	P. reticulatum	Ship harbour, NS, Canada	NA NA	NA NA	NA NA	Ferrell, J. F.	Not sequenced	EU927576 EU927568	EU927568***	
TCC 01	P. reticulatum	Southern Benguela upwelling region	NA NA	NA NA	NA NA	NA	EU532477	EU532486	EU532486***	
10001	1. renemmant	Southern Bengueia apweining region	IVA	11/1	11/1	11/1	1.0332411		EU332400 · · ·	A
	P. reticulatum	Elands Bay, South Africa	- 32°18.618'	18°19.267	16 March 2013	Carbonell-Moore, C.	1	Not sequenced	MG646319	Andrea Highfield / Decla Schroeder

***************************************	NT4	1	NIA	NT A	1	77' 17 37	EE(12262	Not	A37421700	1
JHW0007-6	NA	Jinhae Bay, Korea	NA	NA	juil-00	Kim, KY.	EF613362	sequenced	AY421790	1
CBA-1	NA	Adriatic Sea, Ancona, Italy	NA	NA	NA	Totti C.	Not sequenced	AM183800	AM183800***	1
020717-OK-PR7	P. reticulatum	Okkirai Bay, Iwate, Japan	39.08	141.85	17-Jul-02	Koike, K.	Not sequenced	MG646283	Not sequenced	Kazuhiko Koike
030707-YB-PR1	P. reticulatum	Yamada Bay, Iwate, Japan	39.46	141.97	7-Jul-03	Koike, K.	Not sequenced	MG646284	Not sequenced	Kazuhiko Koike
030624-OK-PR1	P. reticulatum	Okkirai Bay, Iwate, Japan	39.08	141.85	24-Jun-03	Koike, K.	Not sequenced	MG646285	Not sequenced	Kazuhiko Koike
Omura-no.7_no.8 *	P. reticulatum	Omura Bay, Japan	32.85	129.87	2 may 2011	Takano, Y.	Not sequenced	MG646291	MG646332	Yoshihito Takano
Nagayo-sc1 *	P. reticulatum	Nagayo, Japan	32.85	129.87	24 May 2011	Takano, Y.	MG646299	MG646290	MG646331	Yoshihito Takano
Togitsu-sc2 *	P. reticulatum	Togitsu Port, Japan	32.85	129.87	23 May 2011	Takano, Y.	Not sequenced	MG646289	Not sequenced	Yoshihito Takano
777	P. reticulatum	Concarneau large, France	47.83	-3.95	16 July 2008	Nézan, E.	Not sequenced	Not sequenced	MG646317	Gwenael Bilien
778	P. reticulatum	Concarneau large, France	47.83	-3.95	16 July 2008	Nézan, E.	Not sequenced	Not sequenced	MG646318	Gwenael Bilien
							· ·	Not		
16-034	Ceratocorys armata	Bouée 7, Arcachon, France	44.54	-1.26	18 Dec 2015	Nézan, E.	MG646311	sequenced	MG646329	Gwenael Bilien
16-246	Ceratocorys gourretii	Station B70, Banyuls-sur-mer, French Mediterranean	42.48	3.18	26 Sept 2016	Nézan, E.	MG646310	Not sequenced	MG646328	Gwenael Bilien
	· [Station B70, Banyuls-sur-mer, French	1	r	1		1	Not	1	1
16-245	Ceratocorys horrida	Mediterranean	42.48	3.18	26 Sept 2016	Nézan, E.	MG646312	sequenced	MG646330	Gwenael Bilien
CCMP157	Ceratocorys horrida	Banda Sea, South East Asia	-5,00	130,00	22 April 1975	Sweeney, B.	Not sequenced	EU927577	AF022154	1
2001 D IZ 1504	m	71.1.1.0	54.10	7.0	4 2002	77L M	GU205207***	Not	CTIONSON	1
SCCAP K-1504	Thecadinium kofoidii	Helgoland, Germany	54.19	7.9	Aug. 2002	Hoppenrath, M.	GU295207***	sequenced Not	GU295204	
JH0210	Alexandrium affine	Jinhae Bay, South Korea	NA	NA	NA	NA	Not sequenced	sequenced	AY775286	1
JHW0004-12	Alexandrium tamarense	Korea	NA	NA	NA	NA	Not sequenced	Not sequenced Not	AY421777	1
CCMP116	Alexandrium tamarense	Ria de Vigo, Spain	42.23	-8.8	1 June 1984	Yentsch, C.M.	HM483868	sequenced	Not sequenced	1
CCMP1738	Lingulodinium polyedra	Gulf of Mexico, USA	27.8	-97.13	NA	Buskey, E.	Not sequenced	Not sequenced	EF492507	1
	1 2	· ·						Not		1
DRW0208	Lingulodinium polyedra	Korea	NA	NA	Aug. 2001	NA To a second	Not sequenced	sequenced	AY421788	1
BLACK1	Lingulodinium polyedra	Black Sea, Ukraine	49.90	30.29	2011	Takano, Y.	AB678400	AB678399***	AB693195***	1
CASP1	Lingulodinium polyedra	Caspian Sea, Iran	37.51	49.91	2011	Takano, Y.	AB678402	AB678401***	AB693194***	1
SANPEDRO1	Lingulodinium polyedra	San Pedro Harbor, California, USA	33.74	-118.24	2011	Takano, Y.	AB678404	AB678403***	AB693196***	1
GS0209	Gonyaulax polygramma	Gunsan, South Korea	NA	NA	NA	NA	Not sequenced	Not sequenced	AY775287	1
	Protoperidinium				1		T	Not		1
11A1	tricingulatum	Wadden Sea, Netherlands	53,60	6,58	2006	Kawami, H.	Not sequenced	sequenced	AB716917	1
CCAP1140/3	Parvodinium inconspicuum	Kl. Ukleisee, Schleswig-Holstein, Germany	NA	NA	NA	Meyer	Not sequenced	Not sequenced	FR865631	<u> </u>
hwi d1	Unidentified	Hawaii	21,59	-158,10	04-mars-14	Anne de Vernal and Geneviève Vautour	MG646298	MG646295	MG646320	Haifeng Gu
GgSm10R	Unidentified	Malaysia	1,60	110,32	17-janv-13	Bao Juan Kam	MG646305	Not sequenced	MG646326	Haifeng Gu
GgSm11R	Unidentified	Malaysia	1,60	110,32	17-janv-13	Bao Juan Kam	MG646306	Not sequenced	MG646327	Haifeng Gu
GgSm01R	Unidentified	Malaysia	1,60	110,32	10-août-10	Toh Hii Tan	MG646302	Not sequenced	MG646324	Haifeng Gu
GgSm03R	Unidentified	Malaysia	1,60	110,32	22-sept-10	Toh Hii Tan	MG646303	Not sequenced	Not sequenced	Guat Ru Liow
GgSm07R	Unidentified	Malaysia	1,60	110,32	17-janv-13	Toh Hii Tan	MG646304	Not sequenced Not	MG646325	Zhen Fei Lim
PrTT02R	Unidentified	Malaysia	1,92	109,77	28-mars-13	Sing Tung Teng	MG646308	Not sequenced Not	Not sequenced	Haifeng Gu
PrTT03R	Unidentified	Malaysia	1,92	109,77	28-mars-13	Sing Tung Teng	MG646309	Not sequenced	Not sequenced	Haifeng Gu
PrTT01R	Unidentified	Malaysia	1,92	109.77	28-mars-13	Sing Tung Teng	MG646307	Not	Not sequenced	Haifeng Gu

	'	sequenced	

^{* =} single cells or cysts sequenced through single-cell PCR; Accession numbers in bold denote sequences from this study. ** = these cultures showed presence of 5 precingular plates and are considered aberrant. *** = sequence not used in phylogenies. NA = Not acknowledged.