**Nitzschia omanensis** sp. nov., a new diatom species from the marine coast of Oman, characterized by valve morphology and molecular data

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**Abstract:** An unidentified diatom species belonging in the genus *Nitzschia* isolated from samples in the Omani coastal region of the Arabian Sea is the focus of the present study. This species has been successfully cultivated in the laboratory and DNA was harvested and sequenced. Light microscopy (LM) and scanning electron microscopy (SEM) examination of the strain revealed some distinct features, of which the ultrastructure of canal raphe and the absence of central nodule are the most remarkable ones. In terms of morphology, the strain being studied resembles taxa belonging in *Nitzschia* sect. *Lanceolatae*. However, our phylogenetic tree based on a three–gene dataset, comprising concatenated nuclear–encoded small–subunit ribosomal DNAs (SSU) and chloroplast encoded (rbcL and psbC) shows that our strain is sister to *N. filiformis*, which belongs in *Nitzschia* sect. *Obtusae*. This species that we have named *Nitzschia omanensis*, is thus far only known from the harsh coastal waters of Oman, which are characterized by very high daily temperatures and extremely low precipitation.

**Key words:** Arabian Sea, biodiversity, diatoms, new species, *Nitzschia omanensis*, Oman

**INTRODUCTION**

Species belonging in the diatom genus *Nitzschia* Hassall have long received much attention from scientists due to their importance in water quality assessment and biomonitoring (Lange–Bertalot & Simonsen 1978; Krammer & Lange–Bertalot 1997; Trobajo et al. 2004, 2013; Kelly et al. 2015; Lange–Bertalot et al. 2017). In recent decades, the exploration of *Nitzschia* biodiversity has expanded to include biochemical research, especially since evidence of domoic acid production was recorded among some *Nitzschia* species such as *N. navis–varinigica* Lundholm et Moestrup from the Vietnamese coast (Kotaki et al. 2000; Lundholm & Moestrup 2000) and *N. bizertensis* Smida, Lundholm, Hlaili et Mabrouk from a Mediterranean lagoon along the coast of Tunisia (Smida et al. 2014). Although the use of SEM and molecular data for consistent identification are now commonplace (e.g. Lundholm & Moestrup 2000, 2002; Knattrup et al. 2007), species delimitation within *Nitzschia* remains very problematic, both among marine and freshwater taxa (Mann 1978; Krammer & Lange–Bertalot 1997; Witkowski et al. 2004; Trobajo et al. 2013, 2015; Rimet et al. 2014). This can be attributed to the increasing number of recently described taxa, the extreme range and poor documentation of intra–specific variability of both morphological and molecular characters, and the absence of detailed documentation on nomenclatural types (Jahn et al. 2004; Tudesque et al. 2008; Trobajo & Cox 2006; Trobajo et al. 2013).

Species of the genus *Nitzschia* are ubiquitous as they occur in many types of aquatic ecosystems, from freshwater to brackish or marine habitats (Hasle 1964; Krammer & Lange–Bertalot 1997; Witkowski et al. 2000; Spaulding et al. 2010; Lange–Bertalot et al. 2017). Despite their ubiquitous distribution, studies on *Nitzschia* species have mainly covered fresh waters and to a lesser extent oceanic and marine habitat (e.g. Cleve & Grunow 1880; Gandhi 1960; Mann 1978; Kaczmarska et al. 1986; Krammer & Lange–Bertalot 1997; Trobajo et al. 2013; Mann & Trobajo 2014; Witkowski et al. 2015). Research on the identification and distribution of *Nitzschia* species in the Arabian Sea is also very scarce. Simonsen (1974) studied planktonic
Material and Methods

Study sites description. The new species was isolated from three sites in Al Wusta region in Oman. This coastal area from Duqm to Ras Madrakah is a relatively large intertidal zone that is dominated by high cliffs with large salt marshes (‘Sabkha’) behind. It is exposed to strong monsoon winds in the summer. Soils along the beach are white and sandy. The inshore marine topography is characterized by a narrow continental shelf and a steep continental slope (Middle East Scientific Equipment Corporation et al., 2010). The sampling sites are considered non-polluted due to the very low population density and to the local economic activity that is based on artisan fishery and animal husbandry. However, the recent construction of a large port in Duqm for minerals export (Asian Infrastructure Investment Bank 2016) and the imminent construction of a crude oil storage terminal in Ras Markaz (Business Gateways International 2017) could have severe impacts on the marine biodiversity in the future. The characteristics of the sampling sites are summarized in Table 1.

Collection of samples and culture conditions. Seawater samples were collected from different locations between January and February 2015 using sterile 6 l bottles. Salinity, pH and dissolved oxygen were measured in situ with a multi-parameter handheld meter (Horiba, U52, Mfr # 3200164501). Immediately after sampling, the seawater bottles were transported to the laboratory in Muscat and kept at room temperature during the transportation period of around 6 h. Upon return to the laboratory, the samples were partially decanted, and unialgal cultures were prepared by isolation and purification techniques using serial dilution. Cultures were grown in f/2 medium (Guillard 1975) at 24 °C, with an irradiance of ~200 μmol.m⁻².s⁻¹ and 18:6 h light:dark cycle. Afterwards, they were transferred to Szczecin Culture Collection (SZCZ) where they were maintained in f/2 culture medium.

Microscopic examination. The chloroplast morphology of live diatoms during the Indian Ocean Expedition of RV “Meteor” 1964–1965 from different stations including 6 stations in the middle of the Gulf of Oman. Simonsen (op. cit.) identified 32 Nitzschia taxa with two species and one variety described as new to science. Later, Witkowski et al. (2000) studied several species from Qurum Beach on the coast of Oman, and species of the genus Nitzschia were poorly represented. More recently, in a study dedicated to benthic diatoms located northward Kuwait marine waters, Al-Yamani & Saburova (2011) recorded less than 20 Nitzschia species, but their identification was mostly limited to the genus level.

Oman, with a coastline on the Arabian Sea, is one of the most biologically productive regions of the world’s oceans (Longhurst 2007). Such high primary biological productivity has been evidenced by remotely-sensed satellite data on chlorophyll a concentration (Al-Azri et al. 2010; Piontkovski et al. 2011; Sarma et al. 2012). The coastal zone of Oman, with over 3,000 km in length, includes different topographic and geological features such as ‘Sabkha’ (salt-flats), ‘khawrs’ (lagoons), and stretches of sand, mountains, islands, and mangrove forests (Directorate General of Nature Conservation 2010). In general, the main characteristics of the area are high salinity (ca. 36 ppt), arid climate, large daily and seasonal temperature fluctuations, and a vigorous system of monsoon winds (Jupp et al. 1996). The Northeast (NE) monsoon that affects the region from November till April is weak and dry, whereas the Southwest (SW) monsoon that occurs from June to September is strong and moist. The strong upwelling induced by the Southwest monsoon is classified as one of the five most intense upwellings in the world (Currie 1992; Ormond & Banaimoon 1994). It brings up nutrient-rich deep-water masses along the Somalia and Arabian coast in response to the Ekman transport and vorticity balance (Swallow 1984) and results in spectacular phytoplankton blooms (Haake et al. 1993a,b; Tudhope et al. 1996).

In view of the poor representation of Nitzschia species in marine environments in general, and the Arabian Sea in particular, we aim in the present investigation to isolate specimens of Nitzschia from 3 different sites along the coast of Oman, specifically from Al Wusta region and to provide an exact identification based on a detailed morphological description using scanning electron microscopy (SEM) and sequence information of three genes (SSU, psbC and rbcL). In this study, we describe Nitzschia omanensis Barkia I. et Ch. Li as a species new to science.

Table 1. Water characteristics and description of the selected sampling sites.

| Name of site | GPS coordinates         | Salinity (ppt) | pH      | DO (mg.l⁻¹) | General Observations                          |
|--------------|-------------------------|---------------|---------|-------------|-----------------------------------------------|
| Duqm         | 19°30'32.98"N 57°42'14.36"E | 34.9          | 6.83    | 4.56        | Wide extended coastal beach, sandy soil, flat tidal pool |
| Ras Markaz   | 19°11'04.62"N 57°45'47.04"E | 33.8          | 7.36    | 6.19        | Wide extended coastal beach, sandy soil, calm beach |
| Ras Madrakah | 18°58'06.2"N 57°45'01.9"E     | 34.5          | 6.95    | 5.38        | Exposed to strong wind in summer, swell breaking on rocks |
cells was observed using Nikon TS300 inverted microscope (Nikon Corporation, Tokyo, Japan) equipped with an ×100 Plan Apochromatic oil immersion objective (n.a.=1.40) and differential interference contrast (DIC) optics. For LM of the valves, permanent slides were prepared from rinsed and cleaned material. Accordingly, a cell culture suspension was boiled with 30% hydrogen peroxide (H₂O₂) for a few hours at 150°C and then rinsed 5 times with deionized water. The resulting cleaned diatom material was pipetted onto coverslips, dried and mounted on glass slides using Naphrax® mounting medium (Brunel Microscopes Ltd, Wiltshire, U.K.). LM observations of the cleaned material were made with a Zeiss Axiol Imager M2 (Carl Zeiss, Jena, Germany) using a ×100 Plan Apochromatic oil immersion (n.a.=1.40) objective. The DNA sequence consisted of a matrix of 4362 nucleotide sites from the diatom Bolidomonas pacifica (Brunel Microscopes Ltd, Wiltshire, U.K.). 4362 nucleotide sites were partitioned according to different genes, different codon positions (in case of chloroplast markers), and paired and unpaired sites (in case of SSU markers) with a GTR+G+I model. Phylogenetic tree was reconstructed with 1,000 bootstrap replicates using rapid Bootstrap analysis in RAxML v8.1 (Stamatakis 2014). The best-scoring ML tree was chosen as the final tree, rooted with two strains of Bolidomonas pacifica with bootstrap values added to their corresponding nodes.

**Results**

*Nitschia omanensis* Barkia I. et Ch. Li sp. nov.

**Description**

**Light microscopy (Figs. 1–6):** The frustules are rectangular in girdle view with 2 chloroplasts per cell, one towards each apex (Fig. 6). The valves are linear lanceolate with parallel, sometimes slightly constricted margins and acutely rounded apices. The length and width ranges are 31.0–46.0 μm and 4.5–6.5 μm, respectively (n=15). The canal raphe is strongly eccentric. Transapical striae are parallel, and resolvable in LM with a stria density of 32–37 in 10 μm (arrow in Fig. 3). They are composed of small areolae, difficult to resolve in LM. The fibulae are short and delicate, and in general somewhat unequally distributed (8–13 in 10 μm, arrowhead in Fig. 2). The two middle fibulae are not consistently more distant than other fibulae (arrowhead in Fig. 2).

**DNA extraction and PCR amplification.** Genomic DNA was extracted from clone SZCZCH1763 (*N. omanensis*) during the exponential growth phase using the Genomic DNA Nucleospin® Plant II Kit (Macherey–Nagel, Germany) according to the manufacturer’s instructions. Nuclear–encoded small subunit ribosomal RNAs (SSU) and chloroplast–encoded (rbcL and psbC) were amplified by PCR using the primers described in Li et al. (2015). The 25 μl PCR reaction contained 2 μl (20 ng) purified DNA template; 2.5 μl 10× Dream Taq buffer (including 20 mM MgCl₂); 1 μl Ultrapure dNTPs Set (5 mM each); 0.5 μl each primer (10 μM); 0.15 μl Dream Taq DNA polymerase (5 U μl⁻¹); and 0.05 μl of the remaining volumes were added with sterile distilled water. PCR conditions for SSU were as follows: 94 °C for 2 min, 35 cycles of (94 °C for 15 s, 55 °C for 15 s, 72 °C for 1 min and 35 s), and final extension at 72 °C for 7 min. PCR conditions for psbC and rbcL were the same as with SSU but with 53°C for annealing temperature and 1 min and 15 s for extension time. PCR products were purified using Exonuclease I & Polar–BAP (EURx, Gdańsk, Poland). Sequencing reaction was conducted in Oligo.pl DNA Sequencing Laboratory IBB PAS, (Warsaw, Poland) using Big Dye Terminator v. 3.1 sequencing chemistry and run on an ABI 3730 xl sequencer.

**Phylogenetic analyses.** Maximum likelihood (ML) analysis was performed using a combined three–gene (SSU, rbcL and psbC) dataset including 263 diatom taxa (Table S1), which encompass centric diatoms, araphid diatoms and raphid diatoms. The DNA sequence consisted of a matrix of 4362 nucleotide (Supplementary data S1) composed of SSU (1–708 bp), rbcL (1758–3230 bp) and psbC (3231–4362 bp). The secondary structural alignment of SSU sequences was performed by SSU–align (Nawrocki et al. 2015) using covariance models and different interference contrast (DIC) optics. Bootstrap values were added to their corresponding nodes.

**Results**

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**DNA extraction and PCR amplification.** Genomic DNA was extracted from clone SZCZCH1763 in the Szczecin Diatom Culture Collection, University of Szczecin, Faculty of Geosciences, Poland. Strain ID: SZCZCH1763 deposited in the Andrzej Witkowski Diatom Collection.

**Type habitat:** coastal zone of Oman (19°30’32.98”N, 57°42’14.36”E), collected by Ines Barkia, February 2015.

**Etymology:** This new species is named for the country Oman from which it has been sampled.

**Isotype:** Slide no. BM 101963 deposited in the Natural History Museum, Cryptogamic Herbarium, London.

Valves are strongly eccentric (n=20). Raphe canal is slightly elevated above the valve face. The raphe is filiform. External proximal raphe endings (central nodule) are absent with a raphe slit running through (Figs 7, 8, black arrow in Fig. 8). At the apices, the terminal raphe endings are strongly hooked in the same side of the valve and terminate on the mantle (arrowheads in Figs 12, 13). Valve face is flat; proximal valve mantle is shallow with two rows of areolae; distal valve mantle is formed by very narrow, hyaline stripe of silica. Transapical striae are parallel in the middle, becoming slightly radiate towards the apices, 32–37 in 10 μm (Figs 7, 12, 13). Areolae are small and oblong to circular, 31–37.5 in 10 μm. The areolae on the canal raphe are distinctly longer than those of the valve face and are transapically elongate (white arrow in Fig. 8). Each row of transapical striae corresponds to one elongate areola on canal raphe (arrows in Fig. 8).

Internally, the valve mantle is perpendicular to the valve face with two rows of areolae above the raphe slit as enclosed by the raphe canal. Fibulae are irregularly distributed along the valve length (arrowheads in Fig. 9).

In SEM, the spaces between the fibulae are variable with the tendency to become larger towards the valve apices (Fig. 9). Fibulae are narrow and similar in shape, 8–13 in 10 μm (Fig. 11). Due to the flat position of the specimens in SEM, which mostly displays the valve face towards the observer, we were unable to find a specimen
that clearly showed the internal raphe slit. Thus far, in the two slightly tilted specimens showing the raphe in the valve middle, the internal proximal raphe endings were absent (arrows in Figs 10, 11). At apices, the raphe terminates in small helictoglossae. Internally, the transapical striae are composed of small transapically elongate areolae, positioned at the same level as the virgae (Fig. 11).

**Ecology and Distribution**

*Nitzschia omanensis* was sampled from the upper subtidal zone (depth <1m) of three sites in Al Wusta region in February 2015. The sampling involved water samples containing planktonic and tychoplantonic forms. This period of the year corresponds to the beginning of the NE monsoon, which triggers huge growth of phytoplankton. In this particular survey, an accurate abundance of phytoplankton was not measured. However, the phytoplankton populations from the collected water samples were dominated by medium sized chain-forming diatoms (up to 50 µm), including species of *Chaetoceros*, *Bellerochea* and *Thalassiosira*, as well as dinoflagellates such as *Noctiluca scintillans*. Satellite images from the sampling sites at that period could be found in the following link (https://earthobservatory.nasa.gov/IOTD/view.php?id=85718).

*N. omanensis* was collected from water samples with a salinity of 34 ppt, a neutral pH, and DO from 4.56 to 6.19 (Table 1). In view of the limited number of samples, it was difficult to define precisely the ecological amplitude and preferences of the species. Yet, the cultivation of *N. omanensis* in open ponds under ambient conditions (Barkia et al. 2018) revealed that our new taxon had a high tolerance to desiccation as well as a strong resistance to bleaching agents. In fact, when the ponds were emptied for cleaning purposes, *N. omanensis* survived in the joints or cracks of the ‘Plexiglas’ ponds and reappeared in co-culture with other uni-algal cultures. Such resistance to harsh and dry conditions might be attributed to its natural habitat which is classified as an extremely stressful marine environment (Sheppard et al. 1992).

### Table 2. Morphometric data of species that have the closest similarity to *Nitzschia omanensis* sp. nov.

| Species          | Length (µm) | Width (µm) | Striae (in 10 µm) | Fibulae (in 10 µm) | Central nodule | Valve shape                          |
|------------------|-------------|------------|-------------------|-------------------|----------------|--------------------------------------|
| *N. omanensis*   | 31–46       | 4.56.5     | 32–37             | 8–13              | No             | Linear lanceolate, acutely rounded apices |
| *N. filiformis*  | 27.0–150.0  | 4–5        | 29–34             | 7–9               | Yes            | linear to linear-lanceolate, apices acutely rounded |
| *N. sociabilis*  | 22.4–49.5   | 3.5–4.5    | Invisible         | 8–11              | Yes            | lanceolate to rhomboid with acute apices |
| *N. palea*       | 44.0        | 4.3        | Invisible         | 11–13             | Yes            | Lanceolate with subcapitate apices    |
| *N. intermedia*  | 50–101      | 5.2–7.4    | 23–25             | 9–12              | No             | linear with short, wide, subcapitate apices |

### Phylogenic analysis

In the present study, the phylogenetic tree was constructed with 263 diatoms taxa of which 72 were exclusively within the Bacillariaceae family. Eventually, the genera used for the study involved *Nitzschia*, *Pseudonitzschia*, *Fragilariopsis*, *Denticula*, *Psammodictyon*, *Cylindrotheca*, *Tryblionella*, *Hantzschia* and *Bacillaria*. Our phylogenetic analysis showed that the new taxon was positioned in a sister clade to *N. filiformis* with low support value (bv=43, not shown in the branched node, Figs. 14, S1). This low bootstrap value could be attributed to the missing molecular data of the newly-added Bacillariaceae clones, specifically SSU and psbC sequences. Obviously, the phylogenetic position of *N. omanensis* is still unresolved. However, it could be stated that *N. omanensis* belongs in the Bacillariaceae family within a larger, strongly-supported clade (bv = 97%) that is composed of *Nitzschia dravellensis*, *N. paleacea*, *N. acicularis*, *N. aequorea*, *N. sigmaformis*, *N. filiformis*, *N. palea*, *N. capitellata*, *N. lembiformis*, *N. pusilla*, *N. viretlae* and *N. bergii*. The phylogenetic tree (Fig. 14) demonstrated also that *Nitzschia* is paraphyletic genus, whereas *Pseudonitzschia* and *Tryblionella* are monophyletic.
Figs 1–13. LM and SEM micrographs of *Nitzschia omanensis* Barkia I. et Chunlian Li sp. nov.: (1–5) LM micrographs of cleaned valves of *Nitzschia omanensis* strain SZCZCH1763. Specimen illustrated in Fig. 1 is holotype; note the presence of irregularly distributed fibulae (arrowhead, Fig. 2) and fine striae (arrow, Fig. 3); (6) LM micrographs of living specimens; (7–13) SEM micrographs of *Nitzschia omanensis*, (7) SEM external view of the whole specimen, (8) Close up of specimen illustrated in Fig. 7, note the continuous raphe external slit (black arrow) and canal raphe areola which are transapically elongate (white arrow), (9) SEM illustration of the valve interior; note the presence of irregularly distributed fibulae (arrowheads); (10, 11) Close up of the valve internal view; note the continuous raphe slit (black arrows in both images) and rib between canal raphe and valve face (black arrowheads in both images); (12, 13) Close up of the external view of the valve apical part; note the presence of strongly hooked apical raphe endings (arrowheads). Scale bar 10 μm (1–7, 9); 2 μm (8, 10–13).
Fig. 14. Maximum likelihood phylogeny (with bootstrap values at nodes) inferred from three gene dataset, and only the topology with Bacillariaceae + Achnanthes + Craspedostauras was shown. The whole phylogenetic tree was illustrated in supplementary data Fig. S1. Nitzschia omanensis is bolded for clarity. Support values lower than 50% were not included in the tree.

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**Discussion**

Until today, the number of established *Nitzschia* species is estimated between 900 (Mann 1986) and 1314 (Kociolek 2017). However, due to the high degree of intraspecific morphological variability, the older descriptions of *Nitzschia* species have sometimes led to unreliable morphological diagnoses (Rovira et al. 2015) and underestimated the ecology of some species (Kelly et al. 2015).

Difficulties in the identification of taxa belonging in *Nitzschia* arise broadly from the lack of unambiguous characters to group them. The routine identification of *Nitzschia* as bioindicators in ecological assessment is based upon the measurements of size (length, width), fibula, stria and areola linear density. But, in general the identification of *Nitzschia* species relies basically on morphological features such as the presence/absence of the break in the raphe slit, the central nodule, and the distance between the two middle fibulae (Lange–Bertalot & Simonsen 1978; Trobajo et al. 2004). However, this approach can sometimes lead to unsatisfactory results (Mann 1982). In fact, some characters (e.g. size, stria and fibula density) may change drastically due to daily or seasonal variation in environmental conditions or within the cell cycle (Trobajo et al. 2012). For instance, it was demonstrated that length, width, stria and fibula density in *N. frustulum* (Kützing) Grunow were significantly affected by some physico-chemical parameters such as salinity, nutrients and water movement (Trobajo et al. 2004).

Since morphological data on their own are insufficient to provide a rigorous taxonomic resolution, the use of a combined morphological and molecular examination is proposed to offer useful tools in troublesome cases like those of small *Nitzschiae Lanceolatae* or the species complex of *N. palea* or *N. capitellata*. The distinction between two taxa, one possessing a central nodule (*N. capitellata*) and the other lacking a central nodule (*N. palea*) has a high phylogenetic and evolutionary significance (Trobajo et al. 2009).

However, in the present study, the likely position of *Nitzschia omanensis* within the section of *Nitzschiae Lanceolatae* and its phylogenetic relationship to a representative of *Nitzschiae Obtuseae* demonstrates a low significance of sections in the taxonomy of *Nitzschia* as a genus. While the representative of *Nitzschia sect. Lanceolatae* may possess or not central nodule, all known *Nitzschia* sect. *Obtuseae* possess a central nodule (Lange–Bertalot & Simonsen 1978; Mann 1982; Kramer & Lange–Bertalot 1997).

The study of the morphological characteristics showed that *Nitzschia omanensis* is similar to the species of *Nitzschia sect. Lanceolatae*, such as *N. capitellata* Hustedt, and *N. palea* (Kützing) W. Smith (Kramer & Lange–Bertalot 1997), particularly in terms of the strongly eccentric position of the raphe, the narrow mantle and the linear lanceolate valve shape (Trobajo & Cox 2006). However, *Nitzschia omanensis* is distinct from *N. capitellata* mainly by the absence of central raphe endings—the central nodule. Moreover, *Nitzschia omanensis* is easily distinguishable from *N. palea* in terms of the stria structure and canal raphe. In fact, *Nitzschia palea* has very fine striae composed of relatively robust virgae and significantly narrower areolae positioned between virgae, while the virgae in *N. omanensis* are somewhat narrower than areolae (Fig. 11). Besides, the raphe canal in *N. omanensis* is separated from the valve face by somewhat elevated narrow siliceous rib running between the fibulae (arrowhead in Fig. 12), whereas the contact between canal raphe and valve face in *Nitzschia palea* is flat (cf. Fig. 31 in Trobajo & Cox 2006, and SEM Figures of valve interior in Trobajo et al. 2009). *Nitzschia omanensis* was also found to share similar morphological features with larger specimens of *N. filiformis* (sect. *Obtuseae*), especially in terms of the linear to linear–lanceolate valve shape and stria density. While the absence of the central nodule in *N. omanensis* is argued sufficient to distinguish it from *N. filiformis* possessing a central nodule, the SEM microscopy depicts additional different features namely, the fibulae shape. In fact, fibulae are slightly irregular in *N. omanensis* but distinctly irregular in *N. filiformis*.

The comparison of some morphometric data between species having similarity with *Nitzschia omanensis* sp. nov. is illustrated in Table 2.

Our phylogenetic tree showed that *Nitzschia* is a paraphyletic genus. This result corroborates well the studies of Trobajo et al. (2009), Rimet et al. (2011), Rovira et al. (2015) and Witkowski et al. (2015). Species belonging in *Nitzschia* cluster into several clades, each with moderate bootstrap support. The clade into which our strain is comprised contains species belonging in sections *Lanceolatae* (e.g. *N. palea*, *N. paleaeacea*, *N. capitellata*, *N. pusilla*, *N. bergii*, *N. aequorea*, *N. acicularis*), *Obtuseae* (e.g. *N. filiformis*), and *Sigmaota* (N. sigmaformis). Similar results based on 18S rDNA sequences were obtained by Rimet et al. (2011) who reported that the species from the above–mentioned sections were all found in one single clade. However, contrary to the works of Rimet et al. (2011) where some representatives of *Nitzschia* (e.g. *N. palea*, *N. capitellata* and *N. aequorea*) were well–supported towards the terminal branches of the tree (bv = 100%), our new taxon was found in a clade with *N. sigmaformis* Hustedt and *N. filiformis* with low bootstrap support. Similarly, our results showed that *N. omanensis* formed a subclade with *N. filiformis* (sect. *Obtuseae*) with significantly lower bootstrap value than those reported in Barka et al. (2018), bv = 99%. This discrepancy could be explained by the lack of molecular and phylogenetic data describing either unknown or new species. The sister relationship between our species and *N. filiformis* underscores the true diversity of ecological and physiological tolerances among *Nitzschia* species. In fact, *N. omanensis* sp. nov. is a marine species by
occurring in marine tropical waters, while *N. filiformis* is a brackish–water species that occurs in the tidal flats of the moderate climate zone, or in the inland waters with moderately high mineralization (Krammer & Lange–Bertalot 1997; Lange–Bertalot et al. 2017). Information from the phylogenetic tree showed also that *N. omanensis* was genetically distant from the domoic acid–producing diatom *Pseudo–nitzschia galaxiae* (Cerino et al. 2005). Although it could be plausible to argue on the safety of our species, further biochemical analyses remain necessary to endorse the absence of neurotoxins (e.g. domoic acid).

The study of environmental data of the collected samples didn’t show any striking features. In fact, pH was close to neutral and DO values were observed within the normoxia conditions (4.5–6.2 mg l⁻¹). The phytoplankton communities that support the marine ecosystem in the Arabian Sea at the sampling period (i.e. winter monsoon season) are largely understudied. A study conducted in Muscat region (Oman) showed that *Nitzschia* species represented only 13% of the total phytoplankton population in February, but were more abundant in July and October (Al–Hashmi et al. 2012). Similarly, another study done in the northwestern Arabian Gulf showed that *Nitzschia* genus with 14 taxa identified, had the lowest contribution (0.79%) to the total phytoplankton community in winter season (Polikarpov et al. 2016). Thus, it is suggested that an extensive year–round sampling from different geographic locations along the coast of Oman would be useful to determine the distribution and ecology of *Nitzschia omanensis*.

In conclusion, the new species *Nitzschia omanensis* sp. nov. sampled from marine tropical waters of the Arabian Sea was found to present similar morphological features with representatives of *Nitzschia* section Lanceolatae and was distinctive by the absence of central nodule. Nonetheless, the phylogenetic analysis suggests that our strain is sister to *N. filiformis* that belongs in *Nitzschia* sect. Obtusae.

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Supplementary material

the following supplementary material is available for this article:

Table S1. Strains used for phylogenetic tree construction and their accession numbers.

This material is available as part of the online article (http://fottea.czechphycology.cz/contents)