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Citation for published version:
Fahrunnida, Sayekti, P, Nuhamunada, M, Suyono, E & Alam, P 2020, 'Self-assembly of cellular micro-bio machine parts', Journal of Micro-Bio Robotics, pp. 1-11. https://doi.org/10.1007/s12213-020-00125-4

Digital Object Identifier (DOI):
10.1007/s12213-020-00125-4

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published in:
Journal of Micro-Bio Robotics

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Self-assembly of cellular micro-bio machine parts

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Abstract
This paper focusses on intra- and inter-species connections between diatoms; hard bioglass microalgae that adhere through the secretion of sticky extracellular polymeric substances (EPS). We identify entirely new diatom attachment mechanisms, and the associated structures that develop from them. Further, we consider these findings in light of potential strategies for the self-assembled manufacture of micro-bio machine parts, and discuss their possible first-order end uses.

Keywords Diatom technology · Inter-species cellular adhesion · Micro-bio machine · Biomimetic design

1 Introduction

R&D efforts into micro-machine design is on an upward trajectory. This is because micro-machines are understood to have strong exploitation potential within high impact areas, Sitti [1]. One area receiving greater attention involves the integration of biological organisms to micro-machine design. This is because some of the challenges in micro-machine development including; propulsion, energy/power creation, and sense/response mechanisms, are already naturally present in biological microorganisms. As such, microorganisms are seen as potential solutions to the numerous scaling-challenges associated with the development of functional micro-machines and robots.

The mobility of micro-robots for example, is one area showing considerable promise, since many micro-organisms have inherently advanced biological mechanisms that enable self-energised movement. Ghanbari and Bahrami [2] proposed that there are benefits in exploiting the ciliary swimming mechanism of Paramecium sp., and by doing so, circumventing well-known scaling difficulties that relate to micro-robot mobility. The novel cilia-based methodology for enabling swimming in a micro-robot showed higher simulated efficiency in propulsion than an equivalent flagella-based swimming micro-robot. This was reported to be because the length and diameter of cilia are much larger than those of flagella, and as such they have higher propulsive forces at equivalent accelerations. Xu et al. [3] considered helically swimming micro-robots with potential in minimally invasive in vivo medical applications. They concluded that the effectiveness of helical micro-swimming is affected primarily by the angle of the helical twist, as opposed to other geometrical parameters such as the width and thickness of the helical object. Khalil et al. [4] experimentally demonstrated that a sperm driven microtubule could be used as a micro-bio robot with applications in drug delivery and as a micro-actuator. The direction of travel could be controlled using a magnetic force guiding the microtubule, and the authors further proposed that the efficiency of the micro-bio robot might theoretically be increased by reducing the length of the microtubule, as it would result in a larger (tail) beat-amplitude. Hogg [5] proposed that the design of shape-changing micro-bio robots could be guided by their energy dissipation in viscous biological fluids, since surface forces direct both translational and rotational kinematics.

Diatoms have gained recent popularity as cellular structures suitable for use in micro-bio robotics. Diatoms are eukaryotic microalgae that grow strong, hard, bio-glass (silica-rich) exoskeletons known as frustules [6, 7]. The glassy frustules are attractive in micro-bio robotics as they are rigid hollow bodies, ornamented with systematically arranged nanopores, which makes them ideal in a number of micro-bio robot applications. These might include but are not limited to; the development of self-propelled micro-motors [8], microscale
Panda et al. [8] made use of diatoms with geometrical anisotropy to enable movement in a single direction under a low concentration of \( \text{H}_2\text{O}_2 \), which could then be terminated by EDTA addition. Gale et al. [9] functionalised *Cyclotella frustules* with model antibody rabbit immunoglobulin G (IgG) to amplify the intrinsic blue photoluminescence by a factor of six. Delalat et al. [10] genetically engineered the diatom *Thalassiosira pseudonana* to exhibit GB1 protein on the siliceous surface, which could then bind to targeted antibodies on cancer cells, and were able to interact electrostatically with liposome-encapsulated cancer drug molecules.

Diatoms have essentially been considered thus far, as entire micro-bio machines. To date there are no records of micro-bio robots that are composed of conjoined parts. Conjoining micro-parts with specific geometries increases our potential for building larger, structurally and functionally more complex, micro-bio machines. Diatoms may be enablers in this regard, as they are extremely stiff [11], yet have distinct surface flexibility [12] via an extracellular polymeric substance (EPS) layer that attaches the frustule to other materials. EPS is usually produced by pennate diatoms since they possess raphe, in contrast to centric diatoms that tend to be entangled within a biofilm matrix due to the absence of raphe in their cells [13]. Attached diatoms are found in almost all environments, both on land and in water [14], and their biological functions include stabilising the intertidal mudflat [15] and protecting the diatom from salinity induced stresses [14].

They are fundamentally considered an expensive nuisance in their cells [13]. Attached diatoms are found in almost all environments, both on land and in water [14], and their biological functions include stabilising the intertidal mudflat [15] and protecting the diatom from salinity induced stresses [14]. They are fundamentally considered an expensive nuisance in the maritime sector [16] as they are highly effective biofouling organisms. Nevertheless, their properties of biofouling in combination with their excellent mechanical properties have also been identified as beneficial in micro-bio technologies [17] as they have a combination of stiff, compliant and sticky properties. The properties of diatom adhesion varies with respect to species [18, 19] and attachment substrate [20]. Atomic force microscopy (AFM) can be a useful means of characterising the strength of EPS adhesion. Higgins et al. [21] reported pull-off forces of \( 3.58 \pm 1.97 \) nN of EPS from the girdle of the diatom *Craspedostauros australis*, while in the raphe region, the adhesion forces were significantly higher at \( 15.4 \pm 11.2 \) nN, with maximum values recorded at 60nN. These forces are similar to those required to pull polypropylene (PP) from silica glass (14nN), which is strongly correlated to van der Waals interactions [22, 23], though electrostatic interactions have also been quantified between diatom EPS molecules and silica [12]. Though the Young’s modulus of EPS varies from species to species, it can generally be observed as being within range of ca. 0.5 MPa, confirming its bulk properties are similar to those of other soft polymers [24].

The sticky properties of diatoms allow them to connect to one another, by which means they form colonies. Cell-to-cell attachment has been studied extensively for diatoms of the same species (intra-species attachment) including; *Phaeodactylum tricornutum* [25], *Thalassiosira* [26] and *Skeletonema costatum* [27]. Though diatoms also attach between species (inter-species attachment), there are only a handful of studies covering their attachment characteristics, and for only a small number of interacting species. *Amphora* and *Cymbella* are for example, critical contributors to the development of biofilms. *Amphora* acts essentially, as an initial coloniser forming a direct substrate-EPS linkage for stalked *Cymbella*, which in turn traps other centric and non-EPS forming diatoms [13]. Manoylov [28] demonstrated that the stalked structure of *Cymbella* facilitates *Achnanthidium* attachment, allowing both species to occupy shaded environments, and consequently winning space over its local competition, *Cocconeis placenta*. *Amphora* and *Nitzschia* are also known to interact and attach, such that the former selectively chooses the latter species as a host, utilising the motile character of *Nitzschia* to transport itself as an attached diatom to a more favourable environment [29, 30].

In this paper, we consider diatom interactions as a tangible future approach to the construction of micro-bio machines, robots, and their parts. Diatoms are hard-bodied but have flexible attachment polymers (EPS), which potentialises their utility as stiff-compliant-stiff micro-bio machine parts. The Young’s modulus of diatom frustules typically ranges between 7 and 20GPa [31] while its strength in tension and compression has been recorded as high as 560 MPa and 680 MPa, respectively [32]. Importantly, we consider the characteristic geometries that result from both inter- and intra-species diatom attachments (using both pennate and centric diatoms), as these will guide the design functionality of self-assembling multi-cellular-component micro-bio machines.

## 2 Materials and methods

### 2.1 Cell culturing

Five species of microalgae were cultured in species-specific media for optimal growth. *Phaeodactylum* sp. (pennate), *Skeletonema costatum* (centric) and *Nitzschia* sp. (pennate) obtained from Situbondo Marine Aquaculture, East Java were cultured in f/2 medium (1 mL/L NaNO\(_3\), 1 mL/L NaH\(_2\)PO\(_4\), 1 mL/L Na\(_2\)SiO\(_3\), 1 mL/L trace metal solution and filtered natural seawater) pH 7.5, *Navicula* sp. (pennate) obtained from Jepara Marine Aquaculture, Central Java was cultured in Walne medium (1 mL/L nutrient solution, 1 mL/L stock vitamin solution and 1 L filtered natural seawater) pH 7.5 [33], while *Thalassiosira* sp. (centric) obtained from Gondol Marine Aquaculture, Bali was cultured in Na medium (3 mL/L NaNO\(_3\), 1 mL/L Na\(_2\)HPO\(_4\), 1 mL/L Na\(_2\)SiO\(_3\), 1 mL/L stock vitamin solution) pH 8.5 (Gondol Marine Aquaculture, 2018). The
culturing process was performed under 3000 lx lighting at 24 °C in several 500 mL glass bioreactors under a constant stream of air bubbles to ensure diatom cells in the media were adequately agitated. After the diatoms reached their exponential growth phase, the colonies formed by attaching intra- and inter-species diatoms were observed under both light and phase-contrast microscopes.

### 2.2 Intra and inter-species diatom cultures

Intra-species cultures included the following individual diatom species; *Phaeodactylum* sp., *Nitzschia* sp., *Navicula* sp., *Thalassiosira* sp. and *Skeletonema costatum*. Each diatom species was cultured individually in a 500 mL bioreactor with the medium described above for the specific species. Intra-species diatoms reached an exponential phase of growth within 7 days, after which these cultures were used to make inter-species cultures. Inter-species cultures were conceived by mixing 100 mL of diatom culture from a specific individual diatom species with 100 mL diatom culture from different specific individual diatom species. Each inter-species culture consisted of only two specific individual species of diatom since the specific attachments and geometries would be easier to assess in this way. The specific paired diatom couplings are shown in Table 1.

Both inter- and intra-species diatoms were cultured for seven days, after which the diatoms reach their exponential growth phase.

### 2.3 Light and phase-contrast microscopy

Following seven days of growth, diatom cultures were mounted onto microscope slides together with immersion oil on the cover glass to enhance the resolving power of the microscope. The slides were then observed using an Olympus CX22LED light microscope and an Olympus BX40 phase-contrast microscope with a ×100 objective. The images were captured using an Optilab device and camera. Light microscopy was used for early examination of the diatoms, after which the phase contrast microscope was used to differentiate the bioglass and EPS phases of both inter- and intra-species attached diatoms.

### 2.4 Image analysis

Micrograph images from both light and phase-contrast microscopy were probed using image analysis softwares Fiji v.1.52p and ImageRaster. This enabled a clearer view of the connections that arose between diatoms in both inter- and intra-species attachments. Fiji v.1.52p was used to superimpose images captured using a different aperture focus, while ImageRaster was used to correctly scale the images. Since diatoms are not flat, images were taken using different aperture focuses so that important attachment geometries could be captured at different depths in 3-dimensional space. Images were overlaid in Fiji v.1.52p using different colours as it made it easier to differentiate between the different depths of focus, as well as between the frustule and EPS. The scaling was calibrated in each image using the pre-calibrated scale bar from the original microscope image to avoid dimensioning errors.

### 3 Results and discussion

#### 3.1 Biological attachments

Intra-species diatom attachments revealed several interesting connection points and resultant diatom-diatom geometries (Figs. 1, 2 and 4). Inter-species attachments were predominantly found to occur between raphid diatoms at their tips, to the walls of cylindrical and centric diatoms (Figs. 5 and 6).

Intra-species attachments between *Phaeodactylum* sp. revealed four different patterns of connection (Fig. 1). In the first pattern, the cells connected across the entire length of each *Phaeodactylum* sp. cell (both the two tip regions and the central region) (Fig. 1a1-a2), and this pattern of connection was also noted to occur in intra-species connections between *Nitzschia* sp. cells (Fig. 2a). Another intra-species attachment pattern observed between *Nitzschia* sp. was similar to that in Fig. 2a, but differed in that it only connected at the centre of the adjoined diatoms (Fig. 2b). *Phaeodactylum* sp. cells were also noted to attach at only the tip region, creating a V-shape (Fig. 1b1-b2), which was also noted to occur between both *Nitzschia* sp. cells (Fig. 2c1-c2) and *Navicula* sp. cells (Fig. 2d). Intra-species *Phaeodactylum* sp. cells also showed tip-to-tip attachments however, rather than forming V-shapes, they formed a staggered chain-like geometry (Fig. 1c1-c2). Contrarily, another tip-to-tip pattern observed between intra-species *Phaeodactylum* sp. cells formed a cross-like structure of four diatoms or a linear tip-linked structure between two diatoms (Fig. 1d1-d3).

Both *Phaeodactylum* sp. and *Navicula* sp. are classified as raphid diatoms, since they possess raphe in their frustules [34, 35] from which EPS is secreted. Nevertheless, raphid diatoms are reported to secrete EPS through the raphe, as well as through their girdles and through other pores, such as apical pores [35, 36]. The location of secreted EPS is a factor determining how the diatoms will attach to one another, and the consequent geometries that form. For example, we note that raphe secreted EPS connects only at the centre of the valve face [37] (cf. Fig. 2b); girdle secreted EPS connects in only the region around the tip (cf. Fig. 1b and c), and apical pore secreted EPS facilitates specific connections at only the very end of a raphid diatom (cf. Fig. 1d). Figure 3 shows the locations of the girdle, apical pores, raphe and valves in a raphid diatom.

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[34] [35] [36] [37]
Figure 4 shows the results of intra-species connections between cylindrical centric diatoms. *Thalassiosira* sp. cells were noted to join directly at the valve face (Fig. 4a1-a2). This type of attachment occurs between centrally strutted process secreted β-chitin fibrils [36, 38]. Fryxell et al. [26] noted that some *Thalassiosira* sp. formed gelatinous colonies, while certain other species of *Thalassiosira*, such as *T. weissflogii*, cannot connect unless the connections are facilitated by EPS secreting bacteria [39, 40]. There is therefore importance in committing to a focussed species selection during the design process of self-assembling cells. Unlike *Thalassiosira* sp., *Skeletonema costatum* connects through a siliceous structure that extends from tubular processes called fultoportulae [41, 42]. These connections were evident in our intra-species cultures of *S. costatum* and examples are provided in Fig. 4b1-b2.

Interestingly, when we mixed two different araphid species of diatoms in a single culture (*Thalassiosira* sp. and *S. costatum*), we found that they would not connect to each other (Fig. 5a). In turn, attachments between raphid and araphid diatoms occurred for the most part, between the tips of the raphid diatoms and the sides of the centric diatoms (Figs. 5b-e and 6a-b).

Though both cylindrical species of *Thalassiosira* sp. and *S. costatum* were cultured together, they did not attach to one another (Fig. 5a). These cells only formed clusters of cells between their own species. By contrast, raphid-araphid diatoms did self-assemble together to form inter-species geometries. *Phaeodactylum* sp. for example would tend to connect to the side of the cylindrical *Thalassiosira* sp. via its (apical pore) tip (Fig. 5b1-b2). *Navicula* sp. (Fig. 5c1-c8) were noted to connect anywhere to *Thalassiosira* sp. via both its tip region (girdle/apical pore) and its centre (valve). *Nitzschia* sp. was observed to connect to *Navicula* sp. via its tip region (Fig. 5e), indicating the importance of the EPS secretions from the apical pores for attachment, Hoagland et al. [36].

Girdle EPS initiated attachments [35] were also noted in *Navicula* sp./*S. costatum* pairs (Fig. 5d1). Thus far, all observed attachment mechanisms between diatoms have been

| Pair Number | Diatom Species 1            | Diatom Species 2            |
|-------------|-----------------------------|-----------------------------|
| 1           | *Thalassiosira* sp. (centric)| *Skeletonema costatum* (centric) |
| 2           | *Thalassiosira* sp. (centric)| *Phaeodactylum* sp. (pennate) |
| 3           | *Thalassiosira* sp. (centric)| *Navicula* sp. (pennate)    |
| 4           | *Skeletonema costatum* (centric)| *Navicula* sp. (pennate)    |
| 5           | *Nitzschia* sp. (pennate)    | *Navicula* sp. (pennate)    |
diatom-EPS-diatom (i.e. stiff-compliant-stiff structures). One specific pairing (S. costatum with Navicula sp.) exhibited a very different attachment mechanism. In this pairing, Navicula sp. mechanically interlocked itself between the fultoportulae of S. costatum (Fig. 5d2), creating a diatom-diatom (i.e. stiff-stiff) connection, a combined schematic and CAD interpretation for which is provided in Fig. 7.

In nature, both intra- and inter-species diatom attachments may be a way of advancing species success in a competitive environment [43], since colony formation is hypothesised to be critical in this respect [44, 45]. This said, cell-to-cell attachment depends largely on cellular motility, and raphid diatoms from the genera Phaeodactylum, Navicula and Nitzschia are noted herein to have greater characteristics of motility than cylindrical centric Skeletonema and Thalassiosira species.

### 3.2 Conceptual design of micro-bio machine parts

This section aims to propose possible applications of primarily, the inter-species cellular connections observed and discussed in the previous section. Much of the previous research on micro-bio robots consider individual cells as motile biological robotic carries/transporters [2, 4, 8, 10]. The cellular connections elucidated herein are entirely different in structure, properties and purpose of previously proposed conceptual designs of micro-bio robots. Our connected diatoms self-assembled into both stiff-compliant-stiff, and stiff-stiff (interlocked) structures. The stiff-compliant-stiff structures have the advantage of having flexible EPS polymer joints that enable coupled rotations and translations of the adjoined stiff bioglass diatoms. The stiff-stiff (interlocked) diatoms are micro-scale bioglass orthogonally connected structures, presumably with no rotational or translational freedom.

Our first conceptual designs were inspired by raphid-araphid interactions, whereby the raphid diatom tip attaches to the body of a cylindrical araphid diatom. The final structures resemble micro-bio propellers, Fig. 8, with a rotational point at the centroid of the planar end of the cylindrical araphid diatom. Rotation for propulsion could theoretically be borne from magnetic sources as reported by Fu et al. [46] and Todd et al. [47], or through the incorporation of cilia micro-bio technologies as reported by Ghanbari and Bahrami [2]. A micro-bio machine part such as this, could hypothetically, form part of a device that is either powered to turn and thus propel, or, that turns in accordance with local fluid movements to produce power for other micro-devices. From the point of view of propulsion, hydrodynamic motion at the microscale and below, occurs at low Reynolds numbers (laminar), and requires that the ‘swimming object’

### Fig. 2 Intra-species connection patterns between raphid diatoms (here: Nitzschia sp. (a-c) and Navicula sp. (d)). Scale bar = 5 μm in all images

### Fig. 3 Schematic of a raphid diatom showing the locations of the raphe, valves, apical pores and the girdle

A = valve view  
B = girdle view  
C = raphe  
D = apical pore field  
E = girdle band
essentially able to outrun diffusion [48] such that \( l > D/v \) where \( l \) is the swim-length, \( D \) is the diffusion constant \( (10^{-5} \text{ cm}^2/\text{s} \text{ in water}) \) and \( v \) is the swimming velocity. Rotating swimmers such as *E. coli* for example, are able to outrun diffusion reaching speeds of ca. 30 \( \mu \text{m/s} \) by rotating their bodies at ca. 20 Hz [49].

**Fig. 4** Intra-species connections between the cylindrical centric diatoms (a) *Thalassiosira* sp. and (b) *Skeletonema costatum*. Scale bar = 10 \( \mu \text{m} \) in all images.

**Fig. 5** Inter-species attachment between raphid and araphid diatoms: a *Thalassiosira* sp. (upper) and *Skeletonema costatum* sp. (lower); b *Thalassiosira* sp. (centric cell in the middle) and *Phaeodactylum* sp. (pennate cells attaching to centric cell by their tips); c *Thalassiosira* sp. (cylindrical cell) and *Navicula* sp. (pennate cells); d *S. costatum* (cylindrical cells) and *Navicula* sp.; (pennate cells); e *Nitzschia* sp. (upper) and *Navicula* sp. (lower). Scale bar = 5 \( \mu \text{m} \) in all images.
We also noted unique stiff-compliant-stiff arm-like structures that formed through the inter-species connections of Thalassiosira sp. (sides) and Navicula sp. tips, followed by intra-species attachments between Navicula sp. cell-tips, Fig. 9a. We hypothesise these structures show potential as micro-bio robotic arms (Fig. 9b) as the cellular structures themselves are hard (bioglass diatoms), while the connections between them are flexible EPS polymer. As such, the arms have a degree of translational and rotational freedom.

The interlocked connection between S.costatum and Navicula sp., as shown in Fig. 10, provides inspiration for stiff-stiff micro-bio machine applications. In this inter-species connection, half of the body of Navicula sp. is mechanically interlocked between the fultoportulae of S.costatum (cf. Figure 7), both of which in combination create a stiff orthogonally structured assembly. Since these structures are both mechanically interlocked, and are likely to continue to secrete EPS, we can assume that these structures are well attached and thus physically very stable in their hook-
like conformation. A stable stiff-stiff micro-hook has many potential applications. These might include micro-hooks used for climbing on micro-rough or soft surfaces, as a complementary part for micro-robotic arm, and attached in tandem to enable multi-hook devices for stable gripping micro-bio technologies, Fig. 10. The integration of such diatom assemblies into larger or connected systems could be possible using nanowire techniques reported by Agarwal et al. [50]. Using a wall-climbing robot equipped with hooked-claws and claw tips, as inspired by the spiny structures on cockroach legs, Xu et al. [3, 51] reported upon the importance of surface topography, material, and structure on robotic-climbing effectiveness. They moreover noted that the mobility of their robots was constrained despite being subjected to vibration. The implication is therefore, that size is a critical parameter in the design of a climbing robot, with smaller sized robots being lighter with respect to grip strength and less affected by its own inertia. Herein we postulate therefore, that due to their microscopic sizes, diatoms turned into mobile micro-bio robots, might exhibit improved grip and stability than larger robots.
4 Conclusions

This paper was concerned with intra- and inter-species attachments between self-assembling diatoms, and the structures that develop from them. We find several structures from both pennate and centric diatoms that provide inspiration for future micro-bio propellers, translating and rotating arms, and micro-hooks. Our results revealed that cell-to-cell attachments within the same species of raphid diatoms follow four typical, but fundamentally different patterns, while those occurring amongst cylindrical diatoms only attach at their valves in *Thalassiosira* sp., and via the siliceous structures of *Skeletonema costatum*. The connections formed between different species of diatom, by contrast, typically entailed raphid diatoms connected at their tips, to the sides of cylindrical diatoms. Our work shows there is potential in manufacturing micro-bio machine parts by the self-assembly of diatoms. Diatoms have hard bioglass shells and secrete soft, flexible polymeric substances, and as such, provide ideal possibilities for hard bodied micro-machines with flexible joints, or in the case of interlocking diatoms, hard-bodied micro-machine parts with unconventional shapes.

Acknowledgements

We acknowledge Sukirno, Ph.D. for his assistance in teaching Fahrunnida and Puspa Restu Sayekti on the usage of phase contrast microscope in the Laboratory of Parasitology, Faculty of Biology, Universitas Gadjah Mada. We acknowledge Thoriq Teja Samudra, M.Sc. for the assistance in project funding management in Faculty of Biology, Universitas Gadjah Mada.

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References

1. Sitti M (2017) Mobile microrobotics. MIT Press, Cambridge
2. Ghanbari A, Bahrami M (2011) A novel swimming microrobot based on artificial cilia for biomedical applications. J Intell Robot Syst 63:399–416
3. Xu T, Hwang G, Andref N, Negrier S (2016). Influence of geometry on swimming performance of helical swimmers using DoE. J Micro-Bio Robot, 11(1–4): pp 57–66
4. Khalil ISM, Magdanza S, Sanchez S, Schmidt OG, Misra S (2014). Biocompatible, accurate, and fully autonomous: a sperm-driven micro-bio-robot. J Micro-Bio Robot, 9(3–4):79–86
5. Hogg T (2016). Energy dissipation by morphemic micro-robots in viscous fluids. J Micro-Bio Robot, 11(1–4): 85–95
6. Strzepk RF, Harrison P (2004) Photosynthetic architecture differs in coastal and oceanic diatoms. Nature 403:689–692
7. Yang W, Lopez P, Rosengarten G (2011) Diatoms: self assembled silica nanostructures, and templates for bio/chemical sensors and biomimetic membranes. Analyst 136:42–53
8. Pand A, Reddy AS, Venkateswarlu S, Yoon M (2018) Bio-inspired self-propelled diatom micromotor by catalytic decomposition of H2O2 under low fuel concentration. Nanoscale 10:16268–16277
9. Gale DK, Guti T, Jiao J, Chang C, Rorrer GL (2009). Influence of geom- mechancial properties of diatom frustules using Nanoindentation. J Phycol 45:933–949
10. Delalat B, Sheppard VC, Ghaemi SR, Rao S, Prestidge CA, Kroger N, Voelker NH (2015). Targeted drug delivery using genetically engineered diatom biosila. Nat Commun, 6:8791
11. Subash G, Yao S, Bellinger B, Gretz MR (2005) Investigation of mechanical properties of diatom frustules using Nanindentation. J Nanosci Nanotechnol 5:50–56
12. Sanka I, Suyono EA, Alam P (2017) The effects of diatom pore-size on the structures and extensibilities of single mucilage molecules. Carbohydr Res 448:35–42
13. Richard C, Mitbavkar S, Landoi JS (2017). Diagnosis of the diatom community upon biofilm development on stainless steels in natural freshwater. Scanning, 5052646, p 13
14. Steele DJ, Franklin DJ, Underwood GJC (2014) Protection of cells from salinity stress by extracellular polymeric substances in diatom biofilms. J Bioadhesion Biofilm Res 30:987–998
15. Stal LJ, Brouwer JFC (2003) Biofilm formation by benthic diatoms and their influence on the stabilization of intertidal mudflats. Berichte Forschungszentrum Terramare 12:109–111
16. Callow JA, Callow ME (2011) Friendly fouling-resistant marine coatings. Nat Commun 2:210
17. Zusron M, Sanka I, Nuhamunada M, Damayanti Y, Nopitasari S, Kamila AD, Kusuma AB, Ningrum EFC, Rahmawati AR, Zakiyah G (2015). Investigation of toxic substances in marine diatoms on a silicone elastomer. Biofouling 20(6):323–329
18. Passy SI (2017) Diatom ecological guilds display distinct and pre-dictable behavioral characteristics. J Phycol 53:579–586
19. Hoagland KD, Rosowski JR, Gretz MR, Roemer SC (1993) Diatom extracellular polymeric substances: function, fine structure, chemistry, and physiology. J Phycol 29:537–566
20. Borowiezka MA, Chiappino ML, Volcani B (1977) Ultrastructure of a chain-forming diatom *Phaeodactylum tricornutum*. J Phycol 13:162–170
21. stormer EF & Julia NL (2003). Centric Diatoms. In: John D. Wehr & Robert G. Sheath, Freshwater Algae North America: Ecology and Classification. Elsevier, London, pp 559–594
22. Crocker KM, Passow U (1994) Differential aggregation of diatoms. Mar Ecol Prog Ser 71:249–257
23. Mar Ecol Prog Ser 71:249–257
24. Gardes A, Iversen MH, Grossart H-P, Passow U, Ullrich MS (2017) Diatom extracellular polymeric substances: function, structure, and chemistry. J Phycol 53:579–566
25. Borowiezka MA, Chiappino ML, Volcani BE (1977) Ultrastructure of a chain-forming diatom *Phaeodactylum tricornutum*. J Phycol 13:162–170
26. Storem EF &Julia LM (2003). Centric Diatoms. In: John D. Wehr & Robert G. Sheath, Freshwater Algae North America: Ecology and Classification. Elsevier, London, pp 559–594
27. Crocker KM, Passow U (1994) Differential aggregation of diatoms. Mar Ecol Prog Ser 71:249–257
28. Mar Ecol Prog Ser 71:249–257
29. Mar Ecol Prog Ser 71:249–257
30. Roubeix V, Coste M (2017) A case of close interspecific interaction between diatoms: selective attachment on a benthic motile species. Aquat Microb Ecol 80:55
31. Almquist N, Delamo Y, Smith BL, Thompson NH, Bartholdson A, Lal R, Brzezmski M, Hanska PM (2001) Micromechanical and structural properties of a pennate diatom investigated by atomic force microscopy. J Microsc 202:518–532
32. Hamn CE, Merkel K, Springer J, Jurkoj P, Maier C, Prechel K, Smetacek V (2003) Architecture and material properties of diatom shells provide effective mechanical protection. Nature 421:841–843
33. Andersen RA (ed) (2005) Algal Culturing Techniques. Elsevier Academic Press, Cambridge, pp 507–518
34. Wang J, Cao S, Du C, Chen D (2013) Underwater locomotion strategy by a benthic Pennate diatom *Navicula* sp. Protoplasma 250:1203–1212
35. Willis A, Chiovitti A, Dugdale TM, Wetherbee R (2013) Characterization of the extracellular matrix of *Phaeodactylum tricornutum* (Bacillariophyceae): structure, composition, and adhesive characteristics. J Phycol 49:937–949
36. Hoagland KD, Rosowski JR, Gretz MR, Roemer SC (1993) Diatom extracellular polymeric substances: function, fine structure, chemistry, and physiology. J Phycol 29:537–566
37. Borowiezka MA, Chiappino ML, Volcani BE (1977) Ultrastructure of a chain-forming diatom *Phaeodactylum tricornutum*. J Phycol 13:162–170
38. Storem EF & Julia LM (2003). Centric Diatoms. In: John D. Wehr & Robert G. Sheath, Freshwater Algae North America: Ecology and Classification. Elsevier, London, pp 559–594
39. Crocker KM, Passow U (1994) Differential aggregation of diatoms. Mar Ecol Prog Ser 71:249–257
40. Mar Ecol Prog Ser 71:249–257
41. Carradec E, Avers M, Gobet B, Passow U, Ullrich MS (2011). Diatom-associated bacteria are required for aggregation of *Thalassiosira weissflogii*. The ISME Journal, 5:436–445
42. Castillo MA, Meave del Castillo ME, Hernández-Becerril DU (1995) Morphology and distribution of species of the diatom genus *Skeletonema* in a tropical coastal lagoon. Eur J Phycol 30:107–115
43. Round FE, Crawford RM, Mann DG (2007) Diatoms: biology and morphology of the genera. Cambridge University press, Cambridge, pp 507–561
44. Verhaegh W, Aerts E, Kors J (2006) Intelligent algorithms in ambient and biomedical computing. Springer Netherlands, Dordrecht, p 16
45. Passy SI (2017) Diatom ecological guilds display distinct and predict-able behavior along nutrient and disturbance gradients in running waters. J Aqu Botany 86:171–178
46. Riet F, Bouchez A (2012) Life-forms, cell-sizes and ecological guilds of diatoms in European Rivers. Knowl Manag Aquat Ecosyst 448:35–42
47. Todd T, Zhen Z, Tang W, Chen H, Wang G, Chung YJ, Deaton K, Pan Z, Xie J (2014) Iron oxide nanoparticle encapsulated diatoms
for magnetic delivery of small molecules to tumors. Nanoscale 6(4): 2073–2076

48. Purcell E (1977) Life at low Reynolds numbers. Am J Phys 45:3–11

49. Darnton NC, Turner L, Rojevsky S, Berg HC (2006) On torque and tumbling in swimming Escherichia coli. J Bacteriol 189:1756–1764

50. Agarwal R, Ladavac K, Roichman Y, Yu G, Lieber C, Grier D (2005) Manipulation and assembly of nanowires with holographic optical traps. Opt Express 13:8906–8912

51. Xu F, Shen J, Wang B (2015) Analysis, design, and experiments of a Rough Wall climbing robot based on grabbing claws. In: Su H, Wang T, Tokhi MO, Virk GS (eds) Assistive robotics. World Scientific Publishing, Singapore, pp 191–198

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