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Published in:
PLoS Biology

Publication date:
2018

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Citation for published version (APA):

Kamennaya, N. A., Kennaway, G., Fuchs, B. M., & Zubkov, M. V. (2018). “Pomacytosis”—Semi-extracellular phagocytosis of cyanobacteria by the smallest marine algae. *PLoS Biology*, 16(1), 1-13. Article e2003502. <https://doi.org/10.1371/journal.pbio.2003502>

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“Pomacytosis” – semi-extracellular phagocytosis of cyanobacteria by the smallest marine algae

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Author's summary

For microorganisms their global significance is a consequence of their astronomical numbers. This is certainly true for the smallest planktonic algae less than 3 μm in diameter, who are the most numerous eukaryotic organisms of the world ocean. At variance with the perception that algae use only sunlight and dissolved mineral nutrients to grow, these microscopic plants consume large numbers of bacteria. Their acting as mini-predators on bacteria of nearly their own size is hard to imagine. A tiny algal cell is cramped with organelles (e.g. nucleus, mitochondrion, chloroplast). There is simply no space inside this cell to engulf a large bacterium in a usual manner. To find out how the smallest oceanic algae feed we studied them using high resolution electron microscopy. Indeed we found that prey handling by the dominant alga differs from all other types of cell feeding. This 1.3 μm diameter haptophyte alga with two chloroplasts and a symbiont holds the 0.8 μm diameter prey in the open rather than closed cytostome (cell mouth). Among planktonic bacteria the alga apparently selects a ball-shaped *Prochlorococcus* (the abundant cyanobacteria responsible for a major part of global photosynthesis) that tightly fits into the open cytostome like a plug. Instead of full prey digestion the alga leaves the doughnut-shaped, spent prey. We suggest that such unusual feeding of this tiny predatory alga is caused primarily by the space limitation inside its cell.

Abstract

The smallest algae less than 3 μm in diameter are the most abundant eukaryotes of the World Ocean. Their feeding on planktonic bacteria of similar size is globally important but physically enigmatic. Tiny algal cells, tightly packed with the voluminous chloroplasts, nucleus and mitochondrion, appear to have insufficient organelle-free space for prey internalization. Here we present the first direct observations of how the 1.3 μm algae, which are only 1.6 times bigger in diameter than their prey, hold individual *Prochlorococcus* cells in their open hemispheric cytostomes. We explain this semi-extracellular phagocytosis by the cell size limitation of the predatory alga, identified as the *Braarudosphaera* haptophyte with a nitrogen (N_2)-fixing

endosymbiont. Because the observed semi-extracellular phagocytosis differs from all other types of protistan phagocytosis we propose to name it “pomacytosis” (from the Greek πώμα for plug).

Introduction

In conventional phagocytosis the caught prey is internalized, i.e. enclosed by a phagocytic membrane inside the predator cell to form a food vacuole, within which prey is digested and its contents are absorbed through the vacuole membrane [1]. Apart from secure isolation of the prey from the environment full closure of the food vacuole benefits the predator in a number of ways. The fully closed vacuole allows the predator to pump excess water to reduce the vacuole volume, to adjust pH inside the vacuole to facilitate prey digestion by lytic enzymes and to contain lysed prey for efficient nutrient assimilation. Only refractory prey material, e.g. moieties of cell wall, is egested when the closed food vacuole finally fuses back with the plasma membrane [2]. Thus, conventional phagocytosis of internalized prey requires enzymes, microfilament, microtubule and membrane investments and can be limited by the predator size [3].

Phagocytosis of prey of similar size or bigger is difficult but achievable for protists. For example, some dinoflagellates use a feeding tube to inject lytic enzymes into prey and to extract digested prey contents [4, 5]. Other dinoflagellates and several haptophytes form extracellular, yet closed food vacuoles [6-8]. Such extensive extracellular vacuoles can only be completed by large predatory cells, which can produce and stock sufficient amounts of the required investments. Compared to extracellular phagocytosis internalization of similar sized prey requires from the predator fewer investments but sufficient intracellular space free from organelles. In protists the nucleus, mitochondria and chloroplasts (the latter in algae) can vary in size but these organelles cannot be smaller than a certain minimal volume. Owing to the presence of such “non-scalable” organelles [9] the intracellular volume available for investment storage and prey internalization shrinks as a power function of the predator cell size. Consequently, small protists may be unable to internalize (conventionally phagocytose) similar sized prey. To test that we focused on feeding of the

smallest algae (<3 μm in diameter), which cells packed with chloroplasts in addition to the nucleus and mitochondria should have the minimal organelle-free space among free-living protists.

According to our morphometric estimates organelles occupy approximately 70% of a haptophyte alga with a cell volume of $2.8 \pm 0.8 \mu\text{m}^3$ ($n=10$; S1 Fig). Even after taking in account scalable but vital cell components, e.g. endoplasmic reticulum rich in ribosomes and enzymes, both the haptophyte alga as well as the smallest known prasinophyte alga with a cell volume of 1.1-5.7 μm^3 [10] are still capable to internalize a bacterial cell of 0.1-0.3 μm^3 [11]. This is in agreement with the substantial indirect experimental evidence that despite their diminutive size the smallest (1-3 μm diameter) algae are the main predators of bacterioplankton in the open ocean [12, 13]. However, because of insufficient resolution of optical microscopy phagocytosis by these algae could only be inferred [14].

In order to find out how less than 3 μm size algae phagocytose similar sized bacteria we chose to study the smallest oceanic picoeukaryotic algae (PES) separated from other protists and bacteria living in seawater by flow cytometry. Using high resolution electron microscopy to observe fine cellular details of the sorted algae we found that their semi-extracellular bacterial phagocytosis – “pomacytosis” differs from all other types of phagocytosis.

Results

Low concentrations of bacterioplankton and PES (6×10^5 cells ml^{-1} and 4×10^2 cells ml^{-1} , respectively; S2 Fig) in the studied region of the Eastern subtropical North Atlantic Ocean were typical for open ocean waters [11, 13].

The main PES population was well-defined by flow cytometry and selected for sorting (S3 Fig). High-throughput barcoding analysis of flow sorted PES cells (S3 Fig) yielded 10,416 high-quality (≥ 300 nt) 16S rRNA gene reads and identified the dominant taxa: 51% of the amplicons were sequences of cyanobacteria, comprised of *Prochlorococcus* (26%) and UCYN-A (25%), and 38% of the amplicons were chloroplast sequences, the majority of which (58%) belonged to the

Braarudosphaeraceae, a coccolithophore family of the Haptophyta (Fig 1). The remaining chloroplast sequences belonged to 10 other types of small algae, each of which represented only a minor fraction of the PES cells (Fig 1). The negligible number of sequences of SAR11 alphaproteobacteria (Rickettsidae) – the most abundant bacteria in the samples (hence the most probable by-sorted cells) validated the high purity of PES sorting.

Fig 1. The diversity of 16S rRNA gene amplified from the flow sorted PES cells. The 16S rRNA gene-based semi-quantitative analysis identified three major constituents – the *Prochlorococcus* and UCYN-A cyanobacteria and the haptophyte *B. bigelowii*. The remaining reads represented heterotrophic bacteria (deltaproteobacteria - 3%, Actinobacteria - 3%, Pseudomonadales - 1% and Rickettsiales - 1% and others) and 10 other genera of small eukaryotic algae (*Ochromonas* - 9%, *Pelagomonas* - 2%, *Triparma* - 2%, *Imantonia* - 1%, *Chrysochromulina* - 1% and *Rhizochromulina* - 1% and others).

Analyses of nearly full-length ribosomal gene sequences confirmed the phylogenetic affiliation obtained with shorter amplicons. Full-length sequences of the 16S rRNA gene of *Prochlorococcus* and UCYN-A were 99% identical to high light-adapted *Prochlorococcus marinus* sp. MIT9301 and 100% identical to the *Candidatus Atelocyanobacterium thalassa* isolate ALOHA [15], respectively. The 18S rRNA gene sequence was 99% identical to a calcifying *Braarudosphaera bigelowii* isolate TMRscBb7 [16] (S4 Fig) and to a small non-calcifying alga collected from oligotrophic waters of the South East Pacific Ocean [17] (S4 Fig) confirming the chloroplast 16S rRNA gene-based identification.

Scanning and transmission electron microscopy (SEM and TEM, respectively) showed no curved rod-shaped cells of the most abundant SAR11 bacteria (S5A Fig) among flow sorted PES cells. The absence of by-sorted SAR11 bacteria re-affirmed the high sorting purity. The majority (95%) of the imaged PES cells (185 out of 195) were ball-shaped small cells with an estimated diameter of $1.3 \pm 0.22 \mu\text{m}$ ($n=33$, size corrected for 30% linear cells shrinkage during sample dehydration [18]). Some of them bore organic, non-calcified scales (S6 Fig). These morphotypes represent coccolithophore

life cycle stages found in nutrient-poor waters [19-21]. Among the sorted PES cells there were no cells with external mineral investments, i.e. pentagonal-shape liths characteristic of *Braarudosphaera* species found in nutrient-replete waters [22]. A few morphologically different cells (10 out of 195 examined cells) had one or two well-preserved flagella (S7 Fig) that ruled out the artificial loss of external investments by the dominant alga.

Out of 185 cells of the dominant alga, 155 (84%) were associated with smaller coccoid cells $0.81 \pm 0.08 \mu\text{m}$ ($n=10$, size corrected for 30% linear cell shrinkage during cell dehydration) in diameter (Fig 2). An additional intracellular body of the dominant algal cells was observed using TEM (Fig 2E). The $0.47 \pm 0.05 \mu\text{m}$ ($n=4$) diameter body occupied a particular location at the cell periphery next to one of the two chloroplasts. When the body was absent a rupture in the algal cell wall was observed (Fig 2F, thick arrow; S5B Fig), confirming that the body was intracellular but could be lost under mechanical stress caused by sorting PES cells directly on TEM grids. A similar intracellular “spheroid body” in *Braarudosphaera bigelowii* isolate TMRscBb7 was identified as an obligate N_2 -fixing UCYN-A endosymbiont [16] – cyanobiont. Contrary to the cyanobiont, the molecularly identified *Prochlorococcus* associated with PES is a free-living planktonic cyanobacterium, which was numerous in the studied seawater ($1.7 \times 10^5 \text{ cells ml}^{-1}$).

Fig 2. Electron microscopy observations of selective feeding by naked *Braarudosphaera* algae on *Prochlorococcus* cyanobacteria.

(A). The collage depicting that the sorted smallest picoeukaryotic algae were dominated by a single morphological type of cells, molecularly identified as *B. bigelowii* JC142. Note that the majority of the *B. bigelowii* JC142 cells are associated with prey cells, molecularly identified as *Prochlorococcus*. (B-M). Representative micrographs that depict *B. bigelowii* cells with *Prochlorococcus* prey (E-M, groups **a** and **b**) or with lost *Prochlorococcus* (B-D, group **c**). In E-H, less than half of the captured *Prochlorococcus* cell is covered with a cytostome (group **a**) and in I-M more than half of the *Prochlorococcus* cell is embraced (group **b**). Ch – chloroplast; C – cytostome; P – *Prochlorococcus*; S – UCYN-A cyanobiont; D – doughnut-shape deformation of the consumed *Prochlorococcus* cell. Scale

bar = 0.5 μm . The figure combines the SEM (A, C, G, H, I, J, K and M) and TEM (B, D, E, F and L) micrographs.

Synthesising the above evidence we concluded that the UCYN-A amplicon derived from the “intracellular body” and the *Prochlorococcus* amplicons represented the extracellular cocci attached to the PES cells. We interpreted the latter association as phagocytosis of *Prochlorococcus* by the naked haptophyte (further referred to as *B. bigelowii* JC142). We assigned the observed *B. bigelowii* cells to two major groups and one minor group: (a) alga with an associated *Prochlorococcus*, of which less than 50% cell surface is inside the cytostome (49%); (b) alga with an associated *Prochlorococcus*, of which more than 50% cell surface is inside the cytostome (35%); (c) alga with a cytostome but without prey (16%) (Fig 2).

The cytostome is most likely used for shape- (and possibly surface-) selective prey recognition and capture. In support of the notion of selection neither molecular nor microscopic evidence suggested that *B. bigelowii* JC142 fed on SAR11 alphaproteobacteria – the most abundant free-living bacteria in the studied seawater (2.8×10^5 cells ml^{-1} , S2 Fig). The algae preferred to feed on less abundant *Prochlorococcus* (1.7×10^5 cells ml^{-1}), which comprised only 27% of total bacterioplankton in the seawater (6×10^5 cells ml^{-1} , S2 Fig), i.e. the haptophyte selected on average one out of four encountered free-living bacterial cells.

Because of high purity PES sorting the individual *Prochlorococcus* observed by SEM were not by-sorted cells but were in fact cells detached from the haptophytes during sorting (e.g. Fig 2D). Both intact and doughnut-shaped, deformed *Prochlorococcus* cells were observed (S5 Fig). The intact, spherical (14 observed cells) *Prochlorococcus* were probably at the start of pomacytosis, whilst the doughnut-shaped *Prochlorococcus* with a central small spot of depressed surface area (23 cells) were at the end of pomacytosis (S5 Fig). Similarly deformed *Prochlorococcus* cells were observed by SEM and TEM (Fig 2L and 2M), affirming that the deformation was a result of pomacytosis rather than an artefact of SEM sample preparation.

High-power TEM revealed that in the groups (a) and (b) the prey *Prochlorococcus* cell is fitted into a semi-circular cytostomic depression, which according to SEM is in reality hemispherical, and is anchored in the cytostome between the two algal chloroplasts (Fig 2C and 2E). In all 155 specimens observed with the *Prochlorococcus* cell the latter remains at least partially free of the algal cytostome membrane (Fig 2, S8 Fig). To our knowledge, this is the first observation of semi-extracellular phagocytosis of prey by a protist using a partially opened cytostome.

Discussion

The *B. bigelowii* JC142 is the smallest haptophyte that was directly observed to phagocytose free-living bacteria. However, *B. bigelowii* ability to internalize the selected bacterium is evolutionary evidenced – its cyanobiont is of a phagocytic origin. The intracellular UCYN-A symbiont cell in *B. bigelowii* isolate TMRscBb7 is surrounded by a food vacuole membrane [16]. The presence of the UCYN-A cyanobiont further reduces the intracellular space of *B. bigelowii* available for prey internalization. The size of the UCYN-A symbiont of *B. bigelowii* JC142 is at the lowest end of the reported UCYN-A size range [15, 16, 23, 24]. The cyanobiont occupies less than 5% of the estimated volume of the *B. bigelowii* JC142 cell, whilst the *Prochlorococcus* prey measures more than 20% of the algal volume. Perhaps, the choice between conventional phagocytosis and pomacytosis depends on the size ratio between the alga and its prey. In order to overcome its space limitation, 1.3 μm *B. bigelowii* JC142 cell, instead of whole-cell phagocytosis, pomacytoses 0.8 μm *Prochlorococcus*.

Selective feeding of *B. bigelowii* JC142 on *Prochlorococcus* implies that despite the internal supply of fixed nitrogen by the UCYN-A cyanobiont, or perhaps, owing to this supply as well as to metabolic demands of the symbiont, the haptophyte could be limited in other main inorganic nutrients [25], e.g. phosphorus and iron. However, this limitation is unlikely because *B. bigelowii* JC142 was collected in the Eastern subtropical North Atlantic Ocean fertilised by aeolian dust from the Saharan desert. Consequently, the surface waters in the studied area are enriched in phosphate and iron [26] but are poor in nitrogen salts [27] – the environment that facilitates growth of N_2 -fixing

photoautotrophs. Instead of photoautotrophy *B. bigelowii* JC142 cells unconstrained by inorganic nutrients including nitrogen (fixed by its cyanobiont) pomacytose *Prochlorococcus*. Hence, the main nutrient the haptophytes gain from *Prochlorococcus* prey is, perhaps, fixed carbon.

B. bigelowii may require fixed carbon because it has the cyanobiont. The UCYN-A cyanobiont lost its photosystem II complex (PSII) but retained its photosystem I (PSI) [28] to use light energy to fix N₂. In return for the shared fixed nitrogen the *Braarudosphaera* host should share its fixed carbon with the cyanobiont [15, 29]. Furthermore, to minimize inhibition of the cyanobiont N₂-fixation a *B. bigelowii* cell needs to keep its intracellularly dissolved O₂ concentration low. Large host cells, e.g. *Rhizosolenia* and *Rhopalodia* diatoms, do that by spatially segregating their chloroplasts from N₂-fixing cyanobionts within their cells [30]. In the 1.3 μm *B. bigelowii* JC142 cell (Fig 2E) O₂ produced by the adjacent chloroplast could directly inhibit N₂-fixation by the cyanobiont and the haptophyte needs to reduce [29] if not to halt photosynthesis by its own chloroplasts. Consequently both the host and cyanobiont become starved of fixed carbon and require its alternative, external source. In order to acquire that fixed carbon, *B. bigelowii* JC142 selectively pomacytose free-living *Prochlorococcus* cyanobacteria.

Based on our observations (Fig 1 and 2), we suggest to interpret the reported association between the “unknown structure” and UCYN-A-bearing haptophyte (Fig 6 in [29]) as *Prochlorococcus* cell pomacytosed by the haptophyte. Low CO₂ fixation by the haptophyte chloroplasts compared to high CO₂ fixation by the “unknown structure” – *Prochlorococcus* (Fig 6 in [29]) supports our suggestion that the *Braarudosphaera* could acquire fixed carbon from its prey rather than from its own chloroplasts. Perhaps, because a CO₂-fixing *Prochlorococcus* cell also produces O₂, the *B. bigelowii* JC142 cell does not internalize it. Instead, live *Prochlorococcus* is kept segregated from the O₂-sensitive cyanobiont and the haptophyte keeps the cytostome semi-open to allow O₂ dissipation. Hence, pomacytosed *Prochlorococcus* could be viewed as a temporary chloroplast substitute.

Conventional phagocytosis is a relatively quick process that usually takes seconds (e.g. [8]) and one seldom observes a protist predator in the process of internalizing prey. Because the

majority of the *B. bigelowii* JC142 collected during six hour sampling was in a process of feeding (84% held prey), pomacytosis should be a slow process that takes hours. The absence of internalized *Prochlorococcus* cells and nearly 1:1 ratio between pomacytosed *Prochlorococcus* with more than half cell surface exposed (group (a)) and with less than half cell surface exposed (group (b)) suggest that the haptophyte controls exposure of the prey cell to seawater. During slow pomacytosis the predator could gain extra benefit from the prey that fixes CO₂ and takes up nutrients through the cell wall exposed to seawater (Fig 2E-K). Unlike conventionally phagocytosing cells, the pomacyting *B. bigelowii* JC142 detained *Prochlorococcus* in their cytostome without full internalization, perhaps, harvested fixed carbon released by prey and egested the deformed, spent prey without full digestion (S5 and S8 Fig).

Thus, a combination of primarily intracellular space limitation and secondly physiological requirements of the tiny predatory alga leads to semi-extracellular phagocytosis of selected prey.

Materials and Methods

Ethics statement

The study does not involve human participants or tissues. The study does not involve animal research. This is an oceanographic study carried out in the international waters. This research does not require special permission.

Sampling collection and sorting strategies. The study was carried out in the Eastern subtropical North Atlantic Ocean (23° 37' N, 20° 43' W) on board the Royal Research Ship “James Cook” during the research cruise JC142 in November-December 2016. Seawater samples from 25 m (a representative depth of the surface mixed layer) were collected using a rosette of 20-l Niskin bottles mounted on a conductivity-temperature-depth (CTD) profiler. All plastic- and glass-ware for handling seawater were pre-washed with 10% HCl and rinsed with sampled seawater.

Concentrations of total bacterioplankton, *Prochlorococcus* and SAR11, the latter as a population of cells with low nucleic acid content [31], were determined by flow cytometry. Routinely

samples were fixed with 1% (w/v) paraformaldehyde (PFA) final concentration, stained with SYBR Green I DNA dye [11, 32] and analysed with the custom-modified FACSort instrument (Becton Dickinson, Oxford, UK), equipped with the blue diode laser (488 nm, 50 mW, Quantum Analysis, Munster, Germany) using the CellQuest software.

For determining concentrations of PES and *Synechococcus* and for cross referencing microbial populations in the concentrated samples (used for flow sorting) seawater samples were fixed with 2% PFA, stained with $0.1 \mu\text{g ml}^{-1}$ Hoechst 33342 (final concentration) and analysed with the custom-built MoFlo XDP instrument (Beckman-Coulter, High Wycombe, UK) (S2 Fig) using the Summit 5.4 software. The first UV diode laser (355 nm, 100 mW, JDSU, CY355-100, Thailand) and the second blue diode laser (488 nm, 240 mW, Cobolt, Solna, Sweden) were aligned through the first and third pin hole, respectively. Shallow angle light scatter (forward scatter, FSC) of the UV light was detected using the 351 ± 5 nm optical filter and the H957-18 photomultiplier (Hamamatsu, Japan). More sensitive H957-27 photomultipliers (Hamamatsu) were used for detecting particle fluorescence at four wavelengths (457 ± 25 nm, 530 ± 20 nm, 580 ± 15 nm, >643 nm) and the three wavelengths (505-550 nm, 580 ± 15 nm, 670 ± 15 nm) excited by the first and second laser, respectively.

A reference mixture of yellow-green (505/515 nm) $0.5 \mu\text{m}$ beads (Life Technologies, Eugene, USA) and multi-fluorescence $1.0 \mu\text{m}$ beads (Fluoresbrite Microparticles, Polysciences, Warrington, USA) were used as an internal standard for both fluorescence and flow rates. The absolute concentration of beads in the stock solution was determined using syringe pump flow cytometry [33].

For flow sorting microbes were gravity concentrated $\sim 10^3$ folds using sterile $0.2 \mu\text{m}$ pore size Sterivex filter units (Millipore, Watford, UK) attached directly to Niskin bottles. For molecular identification concentrated microbial samples were fixed with Lugol iodine solution [34] and stored at $+4^\circ\text{C}$ before being flow sorted within 48 hours. Samples were discoloured with thiosulfate [34] and stained with Hoechst 33342 prior to sorting. For electron microscopy analyses concentrated

samples were fixed with 2% PFA and stained with Hoechst 33342 prior to sorting. The same dominant distinct population of the smallest picoeukaryotic algae – PES was flow sorted with the MoFlo XDP instrument (S3 Fig) using the Summit 5.4 software. The instrument was optically aligned and its sorting purity and recovery was optimised using blue (350/440 nm) 1.0 µm beads (Life Technologies). Only PES cells gated by both gates (S3B-C and S3E-F Fig) were sorted. Purity of sorted PES cells was validated by the molecular and electron microscopy analyses.

Microscopy. For TEM analyses $1-2 \times 10^3$ target PES cells were flow sorted directly on formvar/carbon covered 200 mesh copper grids (Agar Scientific, Stansted, UK), stained with 2% w/v Gadolinium (aqueous solution), rinsed with pure deionised water and stored in a desiccator for analysis ashore. The grids were examined at 200 keV with the Jeol 2011 LaB6 TEM instrument fitted with a Gatan UltraScan 1000 camera at The Research Technology Platform in Advanced Bioimaging, The University of Warwick, UK.

For SEM analyses 20×10^3 target cells were flow sorted into sterile 1.5 ml microcentrifuge tubes containing aqueous solution of 1% glutaraldehyde (Electron Microscopy Sciences). The tubes were stored at 4°C and brought ashore. The sorted cells were collected onto 0.2 µm pore size 13 mm polycarbonate filters under low vacuum, dehydrated in the ethanol series and critical point dried using 99.9 % hexamethyldisilazane (Sigma-Aldrich). The dehydrated filters were stored in a desiccator at room temperature. Prior to SEM analyses the filters were sputtered with Au/Pd (3:2) to a thickness of 10 nm using the High-Resolution (208hr) Sputter Coater coupled with the MTM20 film thickness controller (Cressington). The filters were examined with the high-resolution SEM UltraPlus instrument (Zeiss Gemini) at 5 keV using the secondary electron detector at the Imaging and Analysis Centre, The Natural History Museum in London, UK.

Cell dimensions were measured on both TEM and SEM micrographs using the ImageJ software [35]. The values obtained from the SEM micrographs were corrected to account for a ~30% cell shrinkage [18]. Average cell volumes were calculated assuming a ball or spheroid shape of algal

cells ($4/3\pi a^2b$), a spherical segment for chloroplasts ($\pi h^2(b-1/3h)$), an ellipsoid for a nucleus ($4/3\pi(a-h)^2b$) and half of this ellipsoid for a mitochondrion (S1 Fig).

Molecular identification. For molecular analyses $20\text{-}50 \times 10^3$ PES cells were flow sorted into sterile 1.5 ml microcentrifuge tubes. An aliquot of 2 μl containing $\sim 2 \times 10^3$ cells was added into 0.2 ml PCR tube containing 30 μl of Q5 High Fidelity Master Mix (New England BioLabs) complemented with primers and nuclease-free water (Ambion). For full-length 16S or 18S rRNA gene amplification, we used 27f/1492r [36] or 63f/1818r [37] primers with annealing temperature of 59°C. The amplicons were added with A-tails (OneTaq DNA polymerase, New England BioLabs), ligated to the pGEM[®] T-Easy vector (Promega) and transformed into the NEB 5-alpha competent *E. coli* cells (New England BioLabs). Plasmids from the positive colonies were sequenced with T7 and SP6 primers to cover the full amplicons length. The 18S rRNA gene sequences were aligned with 18 reference sequences of haptophytes (1400 positions) and phylogenetic relationships for the dataset were calculated with MrBayes software [38].

For a massively parallel sequencing, hyper variable regions V3-V4 (490 bp) were amplified by polymerase chain reaction using S-D-Bact-0341-b-S-17 and S-D-Bact-0785-a-A-21 primers [39]. The forward primer included the PGM barcode adapter (Ion Xpres Barcode Adapters 1-96 Kit, ThermoFisher Scientific) and both primers were tailed with the Ion Torrent sequencing adapters to allow direct downstream multiplexed sequencing. Following amplification, PCR products of ~ 490 bp were gel purified with NucleoSpin Gel and PCR Cleanup kit (Macherey-Nagel) and 1.5 ng of the product were used for template preparation with the Ion Torrent OneTouch System (ThermoFisher Scientific). The templates were sequenced on an Ion Torrent PGM sequencer (ThermoFisher Scientific) using the Hi-Q sequencing chemistry.

After sequencing, the individual sequence reads were first quality trimmed using the Ion Torrent software suite and then further processed using the bioinformatics pipeline of the Silva NGS project [40]. This involved quality controls for sequence length (≥ 300 bp) and the presences of ambiguities ($< 2\%$) and homopolymers ($< 2\%$). The remaining reads were split into individual sample

FASTA files using mothur [41] and aligned against the SSU rRNA seed of the SILVA database release 119. The classification was done by a local BLAST search against the SILVA SSU Ref 115 NR database using BLAST 2.2.22+ with standard settings. The analysis gave (semi-) quantitative information (number of individual reads representing in a taxonomic pool) on the composition of the original PCR amplicon pool [39]. The classification of plastidic SSU rRNA sequence reads was done by nucleotide BLAST search against the nonredundant (nr) database at The National Center for Biotechnology Information (NCBI) (www.ncbi.nlm.nih.gov).

Acknowledgements

We gratefully acknowledge the chief scientist – Bramley Murton, the captain, officers and crew aboard the RRS “James Cook” for their help during the JC142 cruise of the MarineE-tech project of the UK Natural Environment Research Council (NERC). We thank Greta Reintjes, Greta Giljan and Anke Meyerdierks for help with the high throughput sequencing and Ian Hands-Portman, Tomasz Goral and Alex Ball for support with the electron microscopy imaging. We thank Michael Sleight, Michael Cohen and Adrian Martin for their suggestions and critical comments on the earlier drafts of the paper.

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