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Silicified cell walls as a defensive trait in diatoms

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Diatoms contribute nearly half of the marine primary production. These microalgae differ from other phytoplankton groups in having a silicified cell wall, which is the strongest known biological material relative to its density. While it has been suggested that a siliceous wall may have evolved as a mechanical protection against grazing, empirical evidence of its defensive role is limited. Here, we experimentally demonstrate that grazing by adult copepods and nauplii on diatoms is approximately inversely proportional to their silica content, both within and among diatom species. While a sixfold increase in silica content leads to a fourfold decrease in copepod grazing, silicification provides no protection against protozoan grazers that directly engulf their prey. We also found that the wall provides limited protection to cells ingested by copepods, since less than 1% of consumed cells were alive in the faecal pellets. Moreover, silica deposition in diatoms decreases with increasing growth rates, suggesting a possible cost of defence. Overall, our results demonstrate that thickening of silica walls is an effective defence strategy against copepods. This suggests that the plasticity of silicification in diatoms may have evolved as a response to copepod grazing pressure, whose specialized tools to break silicified walls have coevolved with diatoms.

1. Introduction

Higher plants have a wide range of morphological and chemical defensive traits to avoid or reduce consumption by herbivores [1]. Similarly, unicellular organisms of the pelagic realm have evolved a stunning array of defences, including escape behaviours, toxin production and morphological traits, such as colony formation and strengthened cell walls [2]. In many cases, these traits are inducible, meaning that they are activated or intensified in response to the presence of grazers or grazer cues, which strongly supports their defensive role [3-6]. However, often the benefits to the defended organism remain somewhat speculative, and quantitative estimates of the trade-offs are rare [2]. This also applies to the role of silicified cell walls in diatoms, whose protective function together with a number of other adaptive roles have been suggested. For example, the siliceous wall may provide necessary support for the large vacuole [7–9], enhance nutrient uptake [10], facilitate light harvesting [11] and/or protect the cell against UV radiation [12]. Hence, the silica wall may not necessarily only, or mainly, represent an adaptation to reduce herbivory, as is the case in higher plants [13], and the scarcity of experimental evidence for its defensive role in diatoms makes ample room for speculations.

The grazers of diatoms are diverse and include both metazoans (e.g. copepods and euphausiids) and protozoans (e.g. ciliates and heterotrophic dinoflagellates) [7,14,15]. Copepods are key grazers in the ocean since they account for approximately 80% of the mesozooplankton biomass [16]. Calanoid copepods bear powerful mandibles lined with silica-reinforced teeth, well suited to crack diatom shells [17]. Although diatoms that are smaller than the copepod mouth may be directly ingested and survive the passage through the copepod guts [18–20], it has been suggested that the copepods taste and

Table 1. Functional group of phytoplankton, prey species, abbreviated species name, cell size expressed as equivalent spherical diameter (ESD) and longest dimension, and type of grazer (adult copepods, nauplii NIII-IV and dinoflagellates). ESD was measured with a Multisizer 3 Coulter Counter and the longest dimension of pennate diatoms with an optical microscope (Leica, magnification 200×; measured number of cells > 130).

prey group	prey species	abbreviated name	ESD (µm)	longest dimension (μm)	grazer
pennate diatom	Amphiprora paludosa	AP	8	15	copepod, dinoflagellate
centric diatom	Cyclotella cryptica	СС	9	9	copepod, nauplii, dinoflagellate
centric diatom	Cyclotella meneghiniana	CM	8	8	copepod, dinoflagellate
pennate diatom	Navicula incerta	NI	7	13	copepod, dinoflagellate
pennate diatom	Nitzschia laevis	NL	6	14	copepod
centric diatom	Thalassiosira guillardii	TG	6	6	copepod
centric diatom	Thalassiosira weissflogii	TW	13	13	copepod, nauplii
nanoflagellate	Heterosigma akashiwo	NFL	11	11	dinoflagellate
nanoflagellate	Isochrysis galbana	NFL	4	4	dinoflagellate
nanoflagellate	Rhodomonas salina	NFL	6	6	nauplii, dinoflagellate
nanoflagellate	Tetraselmis sp.	NFL	8	8	dinoflagellate

break diatoms before ingestion [21]. However, in some cases, the siliceous wall is able to withstand the crushing forces exerted by the copepod mandibles [21–23]. Planktonic protozoans, such as heterotrophic dinoflagellates and large ciliates, are another major group of grazers in the ocean [14]. Although the silica content in diatoms can affect the palatability and digestibility to protozoan grazers [24], little is known about the potential defensive role of silicified walls against these grazers.

While all diatoms have silicified cell walls, there is a large variation in silica content among and even within species [25–30]. This variation in the silica content may imply different levels of mechanical strength and consequently susceptibility to copepod grazing. Although some studies have suggested that diatoms with a weakly silicified frustule experience higher grazing mortality [23,30], the quantitative relationship between silica content and predation mortality remains unknown. On the other hand, the silica shell is probably of little consequence to heterotrophic dinoflagellates and large ciliates [14]—thus far, very little is known about the potential defensive role of silicified walls against protozoan grazers.

According to optimal defence theory, the advantage of a defence mechanism must come at some cost. Since diatoms have evolved an obligate requirement for dissolved silica, low concentrations of silicon in the environment will limit their growth, which can be regarded as a potential cost of silicification [9,31]. Furthermore, silicon deposition depends on the growth rate of diatoms. Many diatom species do not store sufficient amounts of silicon for the formation of a new valve and must, therefore, harvest most of the required quantities immediately before the cell division [25]. Consequently, fast growth will lead to a reduced period available for silicon uptake, which will result in lighter silicification of the walls, even under non-limiting silicon conditions [32]. Therefore, if increased silicification in response to grazer cues requires reduced growth, this may represent a direct cost of defence. Other costs appear to be less important, and include biochemical costs related to the uptake and deposition of dissolved silica [33], and the costs associated

with the potential losses due to elevated sinking rates of heavy silicified cells [2,34].

Here, we experimentally investigated how different degrees of silicification within and among diatom species affect predation by different types of planktonic grazer: adult copepods, copepod nauplii and heterotrophic dinoflagellates that engulf their prey. Moreover, we examined whether the survival of diatoms during the gut passage of copepods is related to the silica content of the cells. Finally, we assessed the potential cost of diatom silicification in terms of growth reduction.

2. Material and methods

(a) Organisms and culture conditions

The diatom species used as prey were Amphiprora paludosa (CCMP125), Cyclotella cryptica (CCMP331), Cyclotella meneghiniana (CCMP336), Navicula incerta (CCMP542), Nitzschia laevis (CCMP559), Thalassiosira guillardii (CCMP988) and Thalassiosira weissflogii (unknown strain, UNK). The first six species were obtained from the NCMA at Bigelow Laboratory, Maine, while T. weissflogii was obtained from DTU Aqua, Denmark. The species are all non-chain forming, have no spines or other cell structures, are non-toxic and of similar size to minimize other effects than silica content on grazing (table 1). We used different levels of light to acquire different degrees of silicification within a species, as silica content in cells increases under conditions of light limitation [35]. Specifically, each of the seven monocultures was split into two subcultures that were placed under low (15 μ mol photons m⁻² s⁻¹) or high (120 μ mol photons m⁻² s⁻¹) light conditions (14:10 L:D cycle). All subcultures were maintained at a temperature of 15°C in B1 medium with silica [36], and grown until the stationary phase, where the cells contained maximum concentrations of biogenic silica.

We also measured grazing on various species of nanoflagellates (non-silicified prey) in the experiments with copepod nauplii and heterotrophic dinoflagellates to separate size- and silica effects on grazing rates. The nanoflagellate species used as prey were *Heterosigma akashiwo* (UNK), *Isochrysis galbana* (UNK), *Rhodomonas salina* (UNK) and *Tetraselmis* sp. (table 1).

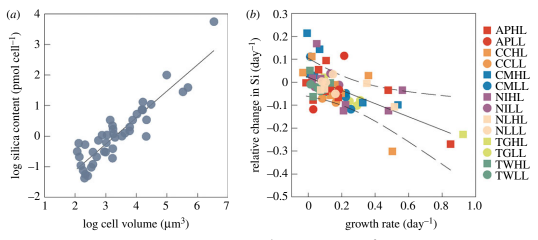


Figure 1. (a) Log-log relationship between biogenic silica content (pmol cell⁻¹) and cell volume (μ m³). The regression line is log(Si) = 0.87 log(V) - 2.89 ($r^2 = 0.84$; p < 0.05). Data are from this study (average silica content and cell volume of diatoms grown under high light conditions) and [25,37–40]. (b) Relationship between relative change in biogenic silica content (SiR; d⁻¹) and growth rate (GR; d⁻¹). The regression line (with 95% confidence limits) is SiR = -0.29GR + 0.02 ($r^2 = 0.6$; p < 0.05). Details about experimental organisms are provided in table 1. HL, high light conditions; LL, low light conditions. (Online version in colour.)

All species were obtained from DTU Aqua (Denmark) and maintained in cultures in B1 medium.

Grazers were adults and nauplii of the copepod *Temora longicornis* (approx. 1 mm cephalothorax length and 0.3 mm body length, respectively) isolated from the Danish coastal waters and maintained in continuous laboratory cultures at $15-18^{\circ}$ C and a salinity of 32 psu in the dark, on a mixed diet of phytoflagellates and diatoms for several generations. To obtain nauplii cohorts of *T. longicornis*, adults were separated from the stock culture with a 200- μ m-mesh sieve and placed in a new tank. After 48 h, the adults were removed and newly produced eggs were transferred to a new tank with food *ad libitum* for 4 days until nauplii stages NIII-IV were obtained. The dinoflagellate *Oxyrrhis marina* (ESD 17 μ m) was obtained from DTU Aqua (Denmark) and maintained in culture on a diet of the cryptophyte *Rhodomonas salina* for many generations.

New diatom batch cultures were established from inoculums of the above subcultures at initial densities of $16~(\pm 6) \times 10^6~\mu\text{m}^3~\text{ml}^{-1}$ and maintained as described above. The growth rates, cellular silica contents, and copepod or dinoflagellate grazing on the diatoms were monitored every second day for several days. Because silica contents in diatoms scale approximately with their volume (figure 1a), we normalized cellular silica content with cell volume and computed the daily relative change in silica content as

$$\Delta Si = \frac{\ln(Si_1) - \ln(Si_0)}{t_1 - t_0},$$
(2.1)

where Si_0 and Si_1 are biogenic silica concentrations (fmol Si_1 μm^{-3}) at times t_0 and t_1 .

(b) Copepod and nauplii grazing experiments

Every second day (days 1, 3, 5, 7 and 9), ingestion rates of *T. long-icornis* adult copepods and nauplii on diatoms were measured. To separate the size and silica effects on grazing, the ingestion rates of nauplii were additionally measured on small non-silicified nanoflagellates for 6 days (days 1, 3 and 5) (table 1). Prey suspensions at concentrations of 3.5×10^6 and $5 \times 10^5 \ \mu m^3 \ ml^{-1}$, respectively, were prepared in B1 + Si medium. For each diatom species and treatment (low and high light), the prey suspension was added to nine 68 ml glass bottles: triplicate initials, triplicate controls (without grazers) and triplicate grazing bottles (with grazers). From each initial bottle, a subsample of 5 ml was removed for cell enumeration and fixed

with acidified Lugol's solution; a subsample of 10 ml was withdrawn for the biogenic silica analysis and treated as described below; and a subsample of 15 ml for measurement of the cell volume. On the experimental days, 15 copepods or 40 nauplii that had starved for 24 h were added to each grazing bottle, and control and grazing bottles were then mounted on a slowly rotating (0.5 rpm) plankton wheel (diameter of 37 cm) for 24 h in darkness and at a temperature of 15°C. At the end of the incubation, subsamples of 5 ml for cell enumeration were fixed with acidified Lugol's solution. The number of copepods or nauplii was counted and their ingestion rates calculated according to [41].

The remaining volume from the grazing bottles with adult copepods was dispensed into culture plates, and approximately 30 faecal pellets from each bottle picked under a dissecting microscope. Faecal pellets were immediately examined under a fluorescence microscope (Olympus, blue filter, magnification 200×), where the number of viable cells per faecal pellet was counted. Afterwards, the remaining contents were fixed with acidified Lugol's solution, and the number of faecal pellets determined. Survival of diatoms during gut passage was calculated as a fraction of ingested diatoms that survived gut passage.

(c) Dinoflagellate grazing experiment

Predation by the dinoflagellate *Oxyrrhis marina* was measured on four diatom species every second day for 10 days (days 1, 3, 5, 7 and 9) and on nanoflagellates for 4 days (days 1 and 3). Diatom and nanoflagellate prey suspensions at concentrations of $1\times 10^6~\mu\text{m}^3~\text{ml}^{-1}$ were prepared in B1 + Si and B1 medium, respectively. The prey suspensions were then dispensed in twelve 68 ml glass bottles: six controls and six grazing bottles with 100 dinoflagellates ml^{-1} . Three bottles from each treatment were immediately harvested for determination of initial prey and grazer concentrations as well as biogenic silica content in diatoms. All the other bottles were mounted on a plankton wheel and harvested at the end of the 24 h incubation. The ingestion rates of dinoflagellates were calculated according to [41,42].

(d) Biogenic silica analysis

The total cellular silica content (biogenic silica) was determined according to [37]. Briefly, 10 ml subsamples were filtered on polycarbonate filters with a pore size of 3 μ m, and washed twice with acidic MilliQ water (pH of MilliQ was adjusted to 5.0 using concentrated hydrochloric acid to minimize dissolution

of silica). Filters were then placed in $50\,\mathrm{ml}$ sterile polyethylene tubes, dried at $65^\circ\mathrm{C}$ for $90\,\mathrm{min}$, and stored at $-20^\circ\mathrm{C}$ until analysis. Dissolution of biogenic silica was later achieved by adding $15\,\mathrm{ml}$ of 0.5% (w/v) sodium carbonate solution to the tubes and heating to $85^\circ\mathrm{C}$ for $90\,\mathrm{min}$. When cool, the pH was adjusted to 5.0-6.0 by adding concentrated sulfuric acid. The reactive silicate concentration was determined according to the molybdate method of [43]) using a SmartChem $200\,\mathrm{wet}$ chemistry analyser (Unity Scientific, MA).

(e) Statistical analysis

R package *lmerTest* [44] was used to perform a linear mixed-effects analysis on the relationship between ingestion rates, silica content and size, with species as random effects. The Akaike information criterion (AIC) was used to select the best model. The model was validated by visual inspection of residual plots.

3. Results

(a) Growth and biogenic silica content

Our data together with the data compiled from the literature show that the silica content is nearly proportional to the cell volume (figure 1a). Hence, we normalized cellular silica content (i.e. pmol cell $^{-1}$) with cell volumes (i.e. pmol μm^{-3}).

Diatoms cultured at low light intensities experienced slower growth compared to those cultured under high light conditions (electronic supplementary material, figure S1). The silica contents of all diatom species decreased during the development of the cultures, and were in most cases higher in light-limited than in light-saturated cells (electronic supplementary material, figure S2). The silica content of cells was not related to their growth rate (electronic supplementary material, figure S3A) because cultures never achieved steady state, but the rate of change in silica content declined with the growth rate of diatoms (figure 1b). Overall, we obtained diatoms with a sixfold variation in silica content among species, and a nearly twofold variation within species.

(b) Grazing on diatoms with different degrees of silicification

Ingestion rates of the adult copepod T. longicornis decreased with increasing silica content of diatoms when compared across species, and varied approximately inversely with the silica content (hyperbolic relation) (figure 2a). A hyperbolic relationship implies that grazing mortality initially decreases rapidly with increasing silica content but as silica content increases further the mortality reduction diminishes. Overall, a sixfold variation in silica content from 1 to 6 fmol Si μm^{-3} lead to a fourfold decrease in grazing mortality. The same pattern of reduced grazing mortality with increasing silica content was also found within all species and was significant in four out of seven cases.

Cell size was a confounding factor since ingestion rates also varied with prey size, although much less than with silica content (figure 2b). Among the tested models for the ingestion rates, the AIC indicated that the best fit was with silica content per volume as a fixed factor, and diatom species as a random effect (table 2).

Similarly, ingestion rates of the copepod nauplii decreased with increasing silica content of diatoms both within and among the species (figure 2c). Overall, less than

a threefold increase in silica content led to almost negligible ingestion rates. Moreover, prey size variation within the examined range did not affect the ingestion rates of the nauplii (figure 2d).

Grazing of the dinoflagellate *Oxyrrhis marina* on diatoms and flagellates was unaffected by silica content, and was similar between the two prey groups (non-silicified nanoflagellates versus diatoms) (figure 2*e*). On the other hand, the grazing rates significantly varied with prey size (figure 2*f*).

(c) Survival of ingested diatoms

A small fraction of the ingested diatoms survived gut passage intact (up to approx. 0.5%); however, survival of ingested diatoms was independent of biogenic silica content (figure 3a) and/or size of diatoms (figure 3b).

4. Discussion

(a) Protective value of silicified cell walls

Here, we have shown that the siliceous shell provides partial protection against copepod grazing, and that the protective value increases with the silica content in the cells (figure 2a,c). The relations found here are, of course, specific to the grazers used, and smaller or larger copepod species may be more or less constrained by the silica shell than found here—just as the small nauplii are more impacted by the shell thickening than the larger adults are. The fact that very few cells pass the gut intact suggests that even small cells are broken during or after ingestion, and this is consistent with mechanical protection. This may also explain why the less powerful copepod nauplii, in particular, were unable to consume heavy silicified cells. Note that the differences in silicification among species can be higher than those obtained in our experiments [25], and therefore expected differences in predation mortality from copepods can be even more drastic than observed here. Similar protective value of silicification may be expected against euphausiids, which also need to crush diatoms before ingestion.

Other major metazoan grazers of diatoms in the ocean are tunicates. Tunicates are considered unselective feeders that ingest all prey within their food size spectrum. However, unlike copepods, consumed diatoms are not broken during ingestion and digestion, and thus the silica frustule will allow higher survival in tunicate faecal pellets [45].

This dependency of copepod grazing mortality on silica content is consistent with experimental measurements as well as finite-element method (FEM) simulations, demonstrating that the mechanical strength of diatom shells increases with increasing silica content [23,29,46,47]. It is also consistent with scaling arguments (see below) that both the fracture strength and the critical buckling load of a diatom shell increase approximately with the cell wall thickness raised to the power of 3. This scaling additionally explains, in a general sense, why the advantage of further thickening decreases as the silica content increases: the unbreakability of cells increases rapidly with their silica content.

While the diatom wall reduces copepod grazing mortality in our experiments, the wall seems to provide no protection against the protistan grazer *Oxyrrhis marina* (figure 2*e*). Phagotrophic dinoflagellates display a variety of feeding mechanisms, such as direct engulfment of prey, pallium

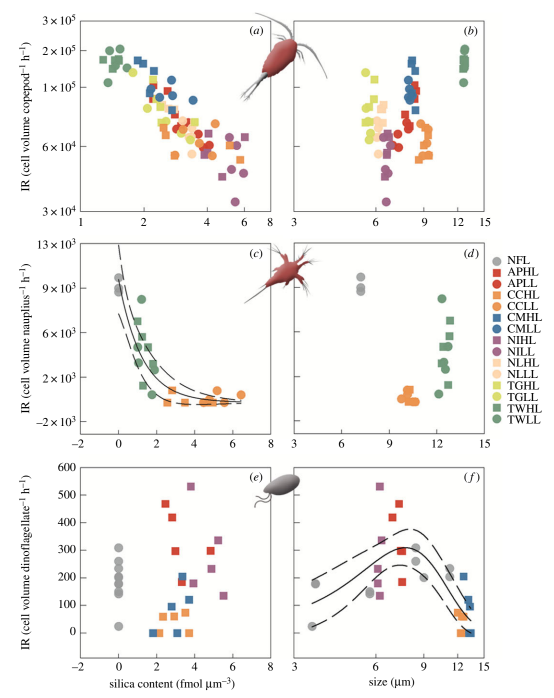


Figure 2. (*a*) Log-log relationship between copepod ingestion rate (IR; cell volume copepod⁻¹ h⁻¹) and biogenic silica content (Si; fmol μ m⁻³), and (*b*) between copepod ingestion rate and cell size (size; μ m). See table 2 for the model selection. (*c*) Relationship between nauplii ingestion rate (IR; cell volume nauplius⁻¹ h⁻¹) and biogenic silica content (Si; fmol μ m⁻³). A nonlinear regression was fitted to the silica range: IR = 19360 × exp(-0.81Si) [r^2 = 0.71; p < 0.05]; long-dashed lines are 95% confidence intervals. (*d*) Relationship between nauplii ingestion rate and cell size (size; μ m) (p > 0.05). (*e*) Relationship between dinoflagellate ingestion rate (IR; cell volume dinoflagellate⁻¹ h⁻¹) and biogenic silica content (fmol μ m⁻³) (p > 0.05). (*f*) Relationship between dinoflagellate ingestion rate and prey size (size; μ m). A log-normal distribution was fitted to the prey size spectrum: IR = 309.1 × exp[-0.5 × ((log(size) - 7.7)/2.9]²] (r^2 = 0.55; p < 0.05); long-dashed lines are 95% confidence intervals. Details about experimental organisms are provided in table 1. HL, high light conditions; LL, low light conditions. (Online version in colour.)

Table 2. Model comparisons of ingestion rates log(IR), where cell size log(size), silica content per volume log(Si) or silica content per volume and cell size log(Si) + log(size) are fixed factors, and species is a random effect. Selected model is presented in italics. *p < 0.05; **p < 0.01; and ***p < 0.001.

model	response variable	fixed structure	random structure	slope	confidence intervals	AIC
1	log(IR)	log(size)**	species***	1.67	[0.58, 2.76]	4.6
2	log(IR)	log(Si) ***	species**	<i>− 0.79</i>	[-0.98, -0.60]	<i>−23.7</i>
3	log(IR)	log(Si)*** + log(size)	species**	-0.70	[-0.91, -0.48]	-23.1
				0.32	[-0.32, 0.95]	

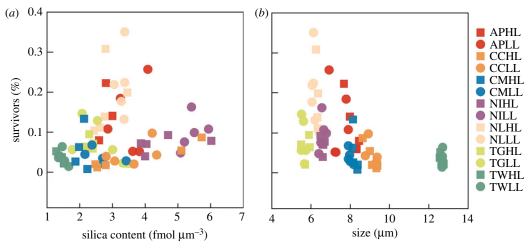


Figure 3. (a) Relationship between fraction of viable cells in copepod faecal pellets (survivors; %) and biogenic silica content of ingested cells (fmol μ m⁻³) (p > 0.05); and (b) between viable cells and size of diatoms (μ m) (p > 0.05). Details about experimental organisms are provided in table 1. HL, high light conditions; LL, low light conditions. (Online version in colour.)

feeding and tube-feeding mechanisms [48], but regardless of the strategy, the ability of dinoflagellates to use a specific prey is governed mainly by the size and shape of the prey [49]. For pallium feeders and those that engulf their prey, such as Oxyrrhis marina, crushing resistance of the silicified cell walls is of no importance, as these organisms digest the cell contents without damaging the walls [8]. It has been speculated that the wall may increase the prey handling time and hence decrease feeding rate [7], but we did not find any significant effects. On the other hand, a silicified wall may be an obstacle to tube-feeding dinoflagellates that pierce the prey by means of a peduncle and suck out its contents, and therefore it is not surprising that these dinoflagellates predominantly feed on naked cells [22,50]. However, there are some parasitoid peduncle-feeding dinoflagellates that successfully feed on diatoms, although their host range is generally restricted to closely related species [50], possibly indicating coevolution of silica walls and attack strategies based on the mode of accessing the frustule [22].

Thus, the defensive role of a silicified cell wall in diatoms appears to primarily provide partial protection against grazers that crush the cells. Below we discuss how the mechanical properties of silica shells vary with cell size and silica content in the context of copepod predation. We next discuss the cost of silicification and explore the trade-off function (i.e. the relation between the costs and benefits of the siliceous shell).

(b) Size scaling of the mechanical properties of the diatom shell

The material properties of the diatom shell are unique: it has the highest mechanical strength relative to its density among any known biological material, and for diatom shells this ratio is close to the theoretical maximum [51]. In addition to several other potential functions of the shell—to support the vacuole [52], enhance nutrient uptake rate [10] and facilitate light harvesting [53,54]—its outstanding strength strongly suggests that it also has a protective and defensive function, as demonstrated here. The silica content of diatoms scales approximately with cell volume (figure 1a), demonstrating that the thickness of the shell increases with cell size. As rationalized below, the observed scaling is further evidence that the

shell mainly serves a protective function. We can explore the scaling by asking how the silica content must increase with cell size to be able to resist the same force before it cracks or buckles. For this crude analysis, we ignore structural differences in the shell among species.

For a thin-walled structure (shell thickness h, size L and surface area $A \approx L^2$), the silica wall mass W scales with its volume as

$$W \approx L^2 h. \tag{4.1}$$

Several criteria may be derived for the mechanical strength, such as maximal (fracture) stress, or the critical buckling load. Assume first that failure is due to fracture. Consider a ring of radius R and cross section $h \times h$ that is subject to a centrally directed force F, and apply the Bernoulli–Euler beam equation. The ring would experience a maximum moment $M = RF/\pi$ and maximal stress $\sigma = 6RF/\pi h^3$, implying for $R \approx L$ that $h \approx L^{1/3}$. If this relation, on grounds of dimensional analysis, also applies to a sphere, spheroid or cylinder, then equation (4.2) leads to the scaling law

$$W \approx L^{2.33}. (4.2)$$

With similar arguments, the exact same scaling arises if one considers a plate that is subject to a central force applied by a bar, maybe better resembling the mandible acting on a diatom cell. Next, consider instead that failure is due to buckling. The buckling load P for a column (cross section $h \times h$ and length L) is $P = \pi 2EI/4L^2$, where E is Young's modulus, and the moment of inertia I scales as $I \approx h^4$ (as derived by Euler in 1755 [55]). This implies $I/L^2 \approx h^4/L^2 \approx \text{constant}$ or $h \approx L^{1/2}$. Attempting to approach the morphology of a diatom shell, assume the column has a cross-section $h \times b$ with $b \gg h$ (i.e. a plate of thickness h), then the buckling load would scale as $P \approx h^3b/L^2 \approx h^3$ just as suggested by the FEM simulation of [47]. For this case, using the criterion $P \approx h^3b/L^2 \approx \text{constant}$ (i.e. $h \approx L^{2/3}$), equation (4.2) now leads to the scaling law

$$W \approx L^{2.67} \approx V^{0.87},\tag{4.3}$$

where V is the cell volume (assumed proportional to the cube of linear dimension of the cell, L^3). Incidentally, this is identical to the observed size scaling of silica content in diatoms (figure 1a). Both arguments therefore suggest that shell

thickness should increase with size to retain constant resistance to external forces, thus further supporting the adaptive value of the shell as a defensive structure.

(c) Trade-offs

While our experiments demonstrate the benefits of the silica wall in terms of reduced predation mortality, the actual magnitude of the benefit of course depends linearly on the concentration of grazers. However, there may be costs of the defence. These costs are not immediately obvious from our experiments, but may nevertheless be explored from available information. The biochemical costs of forming a siliceous wall seem to be negligibly small; in fact, building a silicified cell wall appears to be much cheaper than the formation of ordinary cell walls [33]. There is evidence that shell thickening can be induced by grazer cues [5], and that thickening of the silica wall may require reduced cell division rate, suggesting a possible trade-off. This is because the incorporation of silica is linked to the duration of the cell wall synthesis phase in the cell division cycle. Since slow growth prolongs this phase, this will lead to higher incorporation of silica in the frustule [32]. As a consequence, silica content in diatoms will decline with growth rate under non-limiting silica conditions, as demonstrated repeatedly [32,35,37,56]. If the silica deposition rate was constant and independent of cell division rate, the silica content at steady state would vary inversely with growth rate (i.e. growth rate raised to the power of -1). Because the mortality rate due to copepod grazing in our experiments varies approximately inversely with silica content, this would imply a linear relationship between growth rate and grazing mortality (i.e. suggesting a linear trade-off). However, data from the literature suggest that silica incorporation rate does depend on cell division rate and that the steady-state silica content rather varies with the cell division rate raised to a power of b that is smaller than -1(figure 4). From the five experiments reported in figure 4, an average slope of -0.54 is estimated. This, in turn, implies that the grazing mortality increases with the growth rate raised to the power of 0.43 (when $IR \sim Si^{-0.79}$ from table 2). This suggests that as growth decreases, the reduction in predation mortality accelerates.

While the above demonstrates how shell thickness is a 'passive' function of cell division rate, there is evidence for one species, Thallasiosira weissflogii, that shell thickening may be induced by grazer cues [5]. If this observation can be generalized, it remains to be demonstrated whether and to what extent induced shell thickening requires a reduced growth rate, as suggested by the above, or whether the thickening is the result of an increased rate of silica incorporation in the frustule. However, inducible defences suggest that the defence is costly [57], and those costs must eventually lead to a reduced cell division rate. More experiments are required to clarify this and to quantify the trade-off function between the benefits and costs of induced thickening of cell walls in diatoms.

(d) Survival of diatoms during gut passage

Previous studies reported intact diatoms in copepod faecal pellets, suggesting that the silica wall may enable viable gut passage of ingested diatoms [18,19]. Our experiments show, however, that only a minute fraction of the cells survive and that the survival is independent of the silica content in the cells (figure 3a). However, the number of live cells may

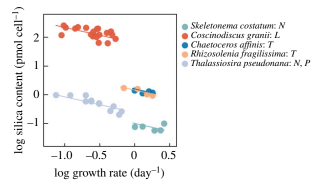


Figure 4. Steady-state silica content in diatoms as a function of cell division rate for diatoms that are limited by factors other than silica (nitrogen N, phosphorus P_r , light L or temperature T_r , as indicated in the legend). Data obtained from [32,35,37,56]. A fitted mixed linear model with one slope and random intercepts was fitted to the log-log relationships between silica content and growth rate. The estimated slope is -0.54 [-0.36, -0.72]. (Online version in colour.)

have been underestimated in our study, as some viable cells are not auto-fluorescent [58]. The few cells that do survive gut passage may sink to the seafloor together with the faecal pellet and thus be removed from the pelagic population. Sinking and overwintering in the sediment are an integral part of the diatom life-cycle strategy [34], and sedimented cells contribute to reseeding the pelagic the following year when brought back to the surface by upwelling or vertical mixing [9].

5. Concluding remarks

Diatoms are the major contributors to phytoplankton blooms because their mortality rates are lower than those of other groups with similar growth rates, making them central in aquatic ecosystems as well as the global carbon cycle [46]. But what makes diatoms unique are their silicified cell walls, whose mechanical strength relative to density is the highest among any known biological material [51]. Our results demonstrate that the silicified cell wall truly is an effective morphological defence mechanism against copepods, whose opal teeth have probably coevolved with their prey, but ineffective against protozoans with direct engulfment as feeding mechanisms. However, a specific trait does not need to perform perfectly to be maintained in a population, and if the grazing mortality rate is decreased by even a small factor, then as long as this benefit offsets the costs of employing the trait, it can be retained [8].

Data accessibility. Data available as part of the electronic supplementary material.

Competing interests. We declare we have no competing interests.

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