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34

35 **Abstract**

36 To broaden our understanding of pelagic ecosystem responses to environmental change, it is essential
37 that we improve the spatio-temporal resolution of *in situ* monitoring of phytoplankton communities. A
38 key challenge for existing methods is in classifying and quantifying cells within the nanophytoplankton
39 size range (2-20µm). This is particularly difficult when there are similarities in morphology, making
40 visual differentiation difficult for both trained taxonomists and machine learning based approaches.
41 Here we present a rapid fluoro-electrochemical technique for classifying nanophytoplankton, and using
42 a library of 52 diverse strains of nanophytoplankton we assess the accuracy of this technique based on
43 two measurements at the individual level: charge required to reduce per cell chlorophyll *a* fluorescence
44 by 50%, and cell radius. We demonstrate a high degree of accuracy overall (92%) in categorising cells
45 belonging to widely recognised key functional groups, however this is reduced when we consider the
46 broader diversity of “nano-phytoflagellates”. Notably, we observe that some groups, for example
47 calcifying *Isochrysidales*, have much greater resilience to electrochemically driven oxidative conditions
48 relative to others of a similar size, making them more easily categorised by the technique. The findings
49 of this study present a promising step forward in advancing our toolkit for monitoring phytoplankton
50 communities. We highlight that, for improved categorisation accuracy, future iterations of the method
51 can be enhanced by measuring additional predictor variables with minimal adjustments to the set-up. In
52 doing so, we foresee this technique being highly applicable, and potentially invaluable, for *in situ*
53 classification and enumeration of the nanophytoplankton size fraction.

54

55 **Introduction**

56 The essential role that phytoplankton play in sustaining marine ecosystems and driving key
57 biogeochemical cycles, notably the biological carbon pump, is unequivocal. As we progress through
58 the Anthropocene, the oceans are facing unprecedented rates of environmental change. In order to
59 improve our global understanding of the impacts that such change is having on phytoplankton
60 communities, and how this might impact key ecosystems services, we must advance our ability to
61 monitor phytoplankton community structure, both spatially and temporally. It is pertinent that *in situ*
62 measurements can capture and quantify the abundance of dominant functional groups of phytoplankton
63 that are present (*i.e.* groups with different biogeochemical and/or ecological functions), as this can
64 inform us about the likely implications of environmental change for marine food webs, biogeochemical
65 cycles, and the capacity of such communities to contribute to sequestration of dissolved CO₂. Widely
66 recognised functional groups, include (but are not limited to): coccolithophores, diatoms,
67 dinoflagellates, and cyanobacteria (Anderson 2005; Nair et al. 2008; Anderson et al. 2021).

68 An important consideration when attempting to quantitatively assess *in situ* phytoplankton communities
69 is that, as single unicellular organisms, phytoplankton can exhibit a large variation in size, spanning
70 over four orders of magnitude; ranging in length from picophytoplankton <2µm (including most
71 cyanobacteria and picoeukaryotes) to the largest diatoms >1000µm (Snoeijs et al. 2002; Finkel et al.
72 2010). From an ecological perspective, size is considered a master trait and is known to significantly
73 influence growth rates, nutrient requirements, grazing susceptibility, and sinking rates (Litchman and
74 Klausmeier 2008); all of which have a bearing on important biogeochemical cycles. In this respect, it
75 has been demonstrated that eukaryotic unicells in the nanophytoplankton size range (2-20µm) display
76 the greatest mass-specific metabolic rates, and thus growth rates, relative to larger microphytoplankton
77 (20µm- 2000µm) and smaller picophytoplankton (Marañón et al. 2013; López-Sandoval et al. 2014;
78 Ward et al. 2017), making them highly competitive and fast responding to environmental perturbations.
79 Indeed, the general success of nanophytoplankton and their ability to dominate phytoplankton biomass
80 has been widely demonstrated from field-based measurements in both the open-ocean (Tarran et al.
81 2006; Balzano et al. 2012; de Vargas et al. 2015; Bolaños et al. 2020) and coastal waters (Pinckney et
82 al. 2015; Barnes et al. 2015; Alves-De-Souza et al. 2017; Leblanc et al. 2018; Pivosz 2019).
83 Subsequently, as sentinels of the phytoplankton assemblage, nanophytoplankton are an insightful target
84 area for monitoring phytoplankton ecological and biogeochemical functioning in response to ocean
85 change. To date, however, there are numerous constraints and limitations to the available methods in
86 obtaining *in situ* time series measurements of nanophytoplankton diversity and abundance.

87 Arguably, the most accurate method for quantifying phytoplankton abundance is microscope taxonomy.
88 If specifically focussing on the nanophytoplankton size fraction, taxonomists can quantify abundance
89 to the genus and species level when there are easily identifiable cell characteristics or morphologies
90 (*e.g.* the unique extracellular calcite structures of coccolithophores, or the complex silica frustules of

91 diatoms). However, this size fraction also contains a large proportion of cell types (typically between 2
92 and 10 μ m) that are extremely hard to identify due to their similar morphology and lack of external
93 inorganic structure (Widdicombe et al. 2010; Piwosz 2019). As a result, a significant proportion of the
94 nanophytoplankton fraction is often given the blanket label of ‘nano-phytoflagellates’ (or similar) in
95 taxonomy surveys. For example, a long-term time series (> 15 years) of the pelagic phytoplankton
96 communities at the L4 station in the Western English Channel has consistently observed that these
97 ‘nano-phytoflagellates’ make up >80% of the total cell counts per unit volume (Widdicombe et al.
98 2010). The other main limitations of traditional microscopy are that it is time-consuming in nature,
99 requires highly skilled labour input, and live samples being fixed and preserved prior to analysis.

100 Flow cytometry is a higher throughput quantitative approach that is often used to distinguish
101 nanophytoplankton and picophytoplankton size fractions, of either live or fixed samples. Light
102 scattering and autofluorescence detection allow for clustering of cells with similar optical properties.
103 This gives a high degree of separation within the picophytoplankton size fraction, as the cyanobacteria
104 autofluorescence and scattering signal is distinct from that of picoeukaryotes (Collier 2000). Where
105 cytometry falls short is at being able to fully characterise ecologically relevant components of the
106 nanophytoplankton fraction; only coccolithophores (detectable due to their unique light scattering) and
107 cryptophytes (due to their phycoerythrin content) can be discriminated from other nanoeukaryotes
108 (Tarran et al. 2006; Tarran and Bruun 2015). Thus, in order to obtain a true representation of the species
109 present in the nanophytoplankton fraction, previous studies have often complemented microscope
110 taxonomy and cytometry analyses with genetics-based interpretations (Balzano et al. 2012; Leblanc et
111 al. 2018; Piwosz 2019; Bolaños et al. 2020; Stern et al. 2023), which generally only provide a value for
112 relative abundances.

113 Over the last decade or so, there has been an increased application of rapid throughput imaging flow
114 cytometry (Lombard et al. 2019). This technique demonstrates a high degree of accuracy in classifying
115 phytoplankton cell types from a combination of image-based machine learning and autofluorescence
116 measurements (Olson and Sosik 2007; Sosik and Olson 2007; Dugenne et al. 2014; Álvarez et al. 2014;
117 Camoying and Yñiguez 2016; Fragoso et al. 2019; Irisson et al. 2022; Fuchs et al. 2022; Kraft et al.
118 2022); thereby, in essence, merging the identification skills of a microscope taxonomist, with the speed
119 and tools of a flow cytometer. The FlowCam (Yokogawa Fluid Imaging Technologies, Inc) is a
120 laboratory-based device (Sieracki et al. 1998), that has been shown to produce highly comparable results
121 when validated alongside traditional microscopical estimates (Álvarez et al. 2014) and, depending upon
122 the FlowCam model, can analyse a broad range of particle sizes from 300nm to 1mm. As with
123 microscope techniques however, the current models of FlowCam still struggle to classify
124 nanophytoplankton cells that have similar sizes and morphologies. Like FlowCam, the CytoSense
125 (CytoBuoy b.v.) is a non-submersible imaging flow cytometry device that has also performed well
126 alongside traditional microscope taxonomy, but validation is poor for classifying cells that are <5 μ m in

127 size (Haraguchi et al. 2017). The Imaging FlowCytobot, or IFCB (McClane Research Laboratories,
128 Inc), and CytoBuoy (CytoBuoy b.v.) are devices that operate similarly to those above, but have been
129 developed to function autonomously at sea (Dubelaar et al. 1999; Olson and Sosik 2007; Fragoso et al.
130 2019). These are promising steps forward for the generation of *in situ* spatio-temporal data for
131 monitoring phytoplankton community structure. Indeed, a recent study has shown strong correlation
132 between IFCB measurements and microscopy data for quantifying blooms of filamentous cyanobacteria
133 in the Baltic Sea (Kraft et al. 2021). Nonetheless, whilst the imaging of these devices covers a broad
134 size range, with the IFCB imaging between 10 and 150 μ m and the CytoBuoy from 1 μ m to 778 μ m, both
135 devices are reported to have poor resolution at their lower detection limits necessary for categorising
136 and quantifying nanophytoplankton (Dugenne et al. 2014; Fragoso et al. 2019; Bolaños et al. 2020;
137 Chase et al. 2020; Fuchs et al. 2022). Furthermore, despite advancements in the machine learning
138 techniques used to improve the classification of data obtained from such devices (Fuchs et al. 2022;
139 Kraft et al. 2022), this is only as good as the resolution of the images acquired and the human
140 interpretation that drives the training (Irisson et al. 2022). This presents a substantial challenge in the
141 nanophytoplankton size range, where there are constraints on image resolution at the finer scale
142 (dependant on the objective lens and camera technology employed), along with human limitations in
143 validating (not all nanophytoplankton are easily distinguishable due to similar size and morphology).
144 We therefore identify that a key frontier in monitoring the health of pelagic ecosystems is in developing
145 novel high-throughput techniques that allow for higher resolution *in situ* discrimination and
146 quantification in the nanophytoplankton size range.

147

148 Moving forward, recent developments in the field of analytical chemistry have shown that
149 electrochemically induced oxidative stress destroys phytoplankton chlorophyll *a* fluorescence in a
150 manner that is idiosyncratic, allowing for differentiation of phytoplankton species from measurements
151 that are obtained within 10s of seconds (Yang et al. 2019; Yu et al. 2022). When a sufficiently high
152 potential is applied to an electrode that is immersed in seawater, a wide range of oxidants can form and
153 diffuse from the electrode surface, for example; oxidation of water to hydrogen peroxide and hydroxyl
154 radicals, bromide to hypobromous acid, and chloride to dichlorine (Yang et al. 2019; Yu et al. 2022).
155 The subsequent reaction of these oxidants with phytoplankton is seen to cause a rapid decay of the
156 cellular chlorophyll *a* fluorescence signal. The rate at which this happens is dependent on a number of
157 factors, including; the distance of the phytoplankton cells from the electrode, the potential applied (and
158 thus the species and concentration of oxidants generated), and more importantly for the work presented
159 here – the type of phytoplankton cell. To that end, we apply this novel fluoro-electrochemical method
160 to a much broader suite of ecologically relevant pico- and nanophytoplankton (52 cultured strains) to
161 produce a ‘susceptibility library’ based on two variables; chlorophyll *a* fluorescence ‘switch-off’ and
162 measured cell radius. This library is assessed using a random forest approach to determine how well the

163 electrochemical method can be used to classify cells into relevant groups. We specifically focussed on
164 strains of nanophytoplankton that (a) represent key functional groups and (b) represent some of the
165 traditionally hard to identify “nano-phytoflagellates”. We then explore a possible biological
166 underpinning to the method, as well as discussing its current limitations and suggested improvements.

167

168 **Methods and Procedures**

169

170 **Culturing for the ‘susceptibility library’**

171 Phytoplankton monocultures were selected so that we had a good representation of each of the key
172 functional groups (5 groups, represented by 37 strains in total), along with an assortment of strains that
173 we consider to be likely ‘nano-phytoflagellate’ candidates (6 groups, represented by 15 strains in total,
174 within the 2 to 10 μ m size range). The key functional groups that we analyse are herein labelled as:
175 “Calcifying Isochrysidales” (7 strains), “Coccolithophores” (8 strains), “Diatoms” (10 strains),
176 “Dinoflagellates” (8 strains), and “Picophytoplankton” (4 strains). In this instance, “Calcifying
177 Isochrysidales” includes strains of both *Emiliana huxleyi* (7 strains) and *Gephyrocapsa oceanica* (1
178 strain). Whilst they might both be considered coccolithophores, given their cosmopolitan nature and the
179 fact they are the most globally abundant calcifiers (particularly *E. huxleyi*) we considered them as a
180 separate group for this analysis. We also chose to include a range of *E. huxleyi* cell stages (both calcified
181 and non-calcified diploid strains, along with a haploid strain) and morphologies (representing different
182 extents of calcification), allowing us to robustly investigate the within species variance in the
183 electrochemical susceptibility. Picophytoplankton are included to allow us to test the lower size limits
184 for detection and classification of our method, including two strains of cyanobacteria (both
185 *Synechococcus* sp.) and two pico-eukaryotes (*Micromonas pusilla* and *Ostreococcus tauri*). Broadening
186 the dataset to encompass nanophytoplankton beyond the key functional groups, the additional groups
187 in our analysis were as follows: “Eustigmatales” (1 strain), “Green algae” (3 strains), “Non-calcifying
188 Isochrysidales” (4 strains), “Pavlovaes” (3 strains), “Phaeocystales” (1 strain), and Prymnesiales (3
189 strains). These strains were selected as it has previously been noted that the unrecognisable “nano-
190 phytoflagelletes”, which are a challenge for microscope taxonomy, could broadly be assigned to the
191 phyla of haptophytes (non-calcified types), chlorophytes (or green algae), and cryptophytes (Piwosz
192 2019). A full list of the 52 strains used in this study can be found in SI Table S1. In the case of
193 “Eustigmatales” we assume that a single strain of *Nannochloropsis granulata* is broadly representative,
194 and for “Phaeocystales” we worked with a single strain of *Phaeocystis globosa*.

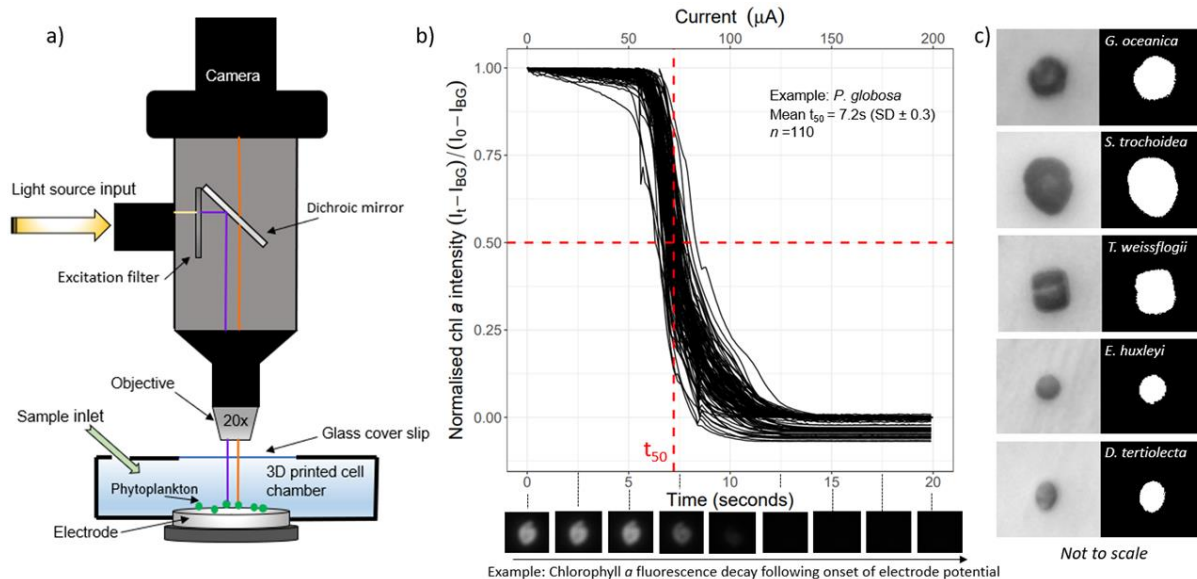
195 All strains were obtained from reputable culture collections: Roscoff Culture Collection (Roscoff,
196 France), Culture Collection of Algae and Protozoa (Oban, UK) and The Marine Biological Association
197 (Plymouth, UK). Following their arrival, they were maintained in exponential growth through regular

198 sub-culturing under sterile conditions on their advised growth medium (see Table S1, SI). All growth
199 media were prepared using synthetic ocean water (Morel et al. 1979), allowing greater control over the
200 main composition of the seawater (see Table S2, SI), and thus consistency in the probable oxidants
201 formed when measuring the electrochemical responses (note: a negligible effect of growth medium on
202 the susceptibility to the electrochemical stress was observed when compared alongside natural seawater,
203 see Fig. S3, SI). Cultures were incubated in PHCbi MLR-352-PE Incubators (PHC Europe B.V.) set to
204 17°C (or 20 °C for all diatoms), with a 14:10h light-dark regime at a PAR intensity of 20-40 $\mu\text{mol m}^{-2}$
205 s^{-1} , and were kept under these conditions for a minimum of two months prior to carrying out the
206 electrochemical susceptibility measurements. The growth of the cultures was tracked on a daily basis
207 using a TECAN Spark plate reader (Tecan Group Ltd.), where three technical replicates of 200 μl of
208 each culture were measured for chlorophyll *a* fluorescence as a proxy for culture biomass. We then
209 selected a time-point in mid-exponential growth phase, when each strain was not at saturation point (*i.e.*
210 carrying capacity) and therefore not nutrient limited, to conduct our experiments (see Figure S1, SI);
211 however, for a couple of strains, susceptibility measurements were taken at different growth phases
212 (and thus different nutrient availability) and only a minimal difference was observed across the phases
213 (see Fig. S4, SI). Immediately prior to the electrochemical experiments, to obtain a sufficient number
214 of cells per image series (as described below), all strains were concentrated by centrifugation
215 (Centrifuge 5702, Eppendorf UK Limited) at 1000 r.p.m for ten minutes, and resuspended to a
216 concentration typically ranging between 5 and 10x.

217

218 **The fluoro-electrochemical technique**

219 A more detailed description of the fundamental electrochemical principles and details of the underlying
220 methodology, including specifics of the equipment used, can be found in the previously published work
221 by Yu *et al.* (2022). For the purposes of this study, the step-by-step method described below outlines
222 the essential procedures that were followed to yield the underlying dataset for the susceptibility library.
223 In summary, we used a galvanostat based ramping linear current, applied to phytoplankton cells settled
224 onto the surface of a carbon electrode (thus controlling for distance from the electrode) and monitored
225 chlorophyll *a* fluorescence decay over time per individual cell (see Table S3 for number of individual
226 cell measurements per strain). The constant rate of current ramping (10 $\mu\text{A s}^{-1}$) means that the moles of
227 oxidant being generated electrochemically increases with time in a controlled fashion. This approach
228 allows for greater possible discrimination of phytoplankton cell types and over a shorter experimental
229 time frame (e.g. 10s of seconds), as opposed to running experiments at a single set potential.
230 Consequently, as cell types have different levels of resilience to such oxidative stress, this technique
231 enables us to quantify the differences in time that is required to drive the chlorophyll *a* quenching across
232 the phytoplankton investigated.



233

234 **Figure 1. (a)** a simple schematic diagram of the electrochemistry – fluorescence microscope set up adapted from
 235 Yu *et al.* (2022), **(b)** an example of normalised Chlorophyll *a* fluorescence transient data for the species
 236 *Phaeocystis globosa*, following onset of the potential (at time 0 seconds) and with a current ramping of $10\mu\text{A s}^{-1}$.
 237 Each black line represents a single cell measurement ($n = 110$), and the dashed red markings highlight the time
 238 point, t_{50} , where normalised fluorescence values have been reduced by 50%. The time-series of images below the
 239 plot illustrate the loss of fluorescence with time for one individual cell. **(c)** examples of brightfield images taken
 240 before the electrochemical experiments for a selection of strains (left column), and the respective estimate of
 241 projected pixel area that was subsequently used to determine the effective radius of each cell (after assuming a
 242 circular shape).

243

244 The fluoro-electrochemical measurements were made as follows (See Fig. 1a, adapted from Yu *et al.*
 245 2022).

246 1) A concentrated sample from a phytoplankton monoculture in exponential growth phase (as
 247 described in Culturing section above) was ‘drop cast’ onto the surface of the working electrode.
 248 After ~ 1 minute of allowing the cells to be deposited on the surface, excess solution was gently
 249 drawn-off using a tissue. Immediately following this, the 3D printed cell chamber was filled up
 250 to maximum capacity with culture growth medium via the sample inlet. Once the chamber was
 251 at capacity a glass cover slip was put in place.

252 2) Using the fluorescence microscope (Axio Examiner, Carl Zeiss Ltd., Cambridge U.K.), we
 253 focussed on a field of view whereby we had a high proportion of phytoplankton cells on the
 254 electrode surface (mean number per experiment = 23 ± 18 S.D., across 212 unique experimental
 255 image series).

256 3) With the microscope set in bright-field mode, we took an image of the starting positioning of
 257 the cells. These images were later used to obtain an estimate of effective cell radius (see Fig.
 258 1c).

259 4) Next, switching the microscope to fluorescence mode, the cells were excited using a $475 \pm$
 260 35nm excitation filter and emission signal passed through a dichroic mirror specific to

261 wavelengths >590nm for chlorophyll *a* fluorescence detection. Simultaneously, the
262 galvanostatic control was synchronised with the camera and data acquisition was started. For
263 the first 40s of the electrochemical experiment no potential was applied, allowing the
264 fluorescence signal to stabilise. Following this, the current was ramped from 0 μ A at a rate of
265 10 μ A s⁻¹ and images recorded at a capture rate of 10f.p.s. For each set of experiments the current
266 was ramped until the chlorophyll signal had completely ‘switched off’.

267 5) After the experiment, the 3D chamber was rinsed clean with DI water and the above steps were
268 repeated a minimum number of 3 times for each phytoplankton strain.

269 Following the experimental data collection, the raw fluorescence transient data was processed for each
270 individual cell within each experimental time-series ($n = 4884$, across all strains) using ImageJ software
271 (v1.53c, Fiji distribution), where “ n ” is the number of individual phytoplankton cell measurements. The
272 integrated fluorescence intensity values for each individual cell (I_t) were corrected by subtracting the
273 background signal of the electrode surface (I_{BG}) for all time points: $I_t - I_{BG}$, following this all values were
274 then normalised by dividing by the fluorescence intensity at the onset of the potential (I_0): $(I_t - I_{BG}) / (I_0 - I_{BG})$. We then used the normalised transient data to determine the time-point for each individual cell
275 where normalised intensity had decreased by 50% (t_{50}), see Fig. 1b. Due to the consistent linear ramping
276 of current at 10 μ A s⁻¹ and that all phytoplankton cells imaged were settled on the surface of the electrode
277 within the timescale of the experiments, we were able to determine the total charge required to be
278 injected to reach t_{50} for each cell, and thus we herein refer to our chlorophyll *a* susceptibility factor as
279 charge at t_{50} (in mC). In some instances where there was significant movement of individual cells it was
280 not possible to accurately measure the chlorophyll *a* fluorescence profile throughout the time series,
281 and subsequently data for these cells was considered erroneous and removed from the analysis.

282
283 For each cell specific chlorophyll *a* fluorescence transient, we used the corresponding bright-field image
284 collected prior to electrochemical experiments to derive a corresponding cellular area based on white
285 pixel area of the cell (Fig. 1c). This was achieved by using the auto-threshold function in ImageJ
286 freeware. From the total pixel area and using a predefined μ m to pixel ratio (0.31 μ m per pixel), we were
287 able to estimate an effective radius (in μ m) of each cell by assuming a circular 2D cell geometry (or a
288 spherical cell). For cells in the picophytoplankton size range, due to the lack of contrast with the
289 electrode surface it was not possible to accurately distinguish cell area, and therefore in these instances
290 we obtained a measurement of mean cell pixel area by manually measuring the area of a subset of at
291 least 10 cells per experimental time-series (using ImageJ).

292

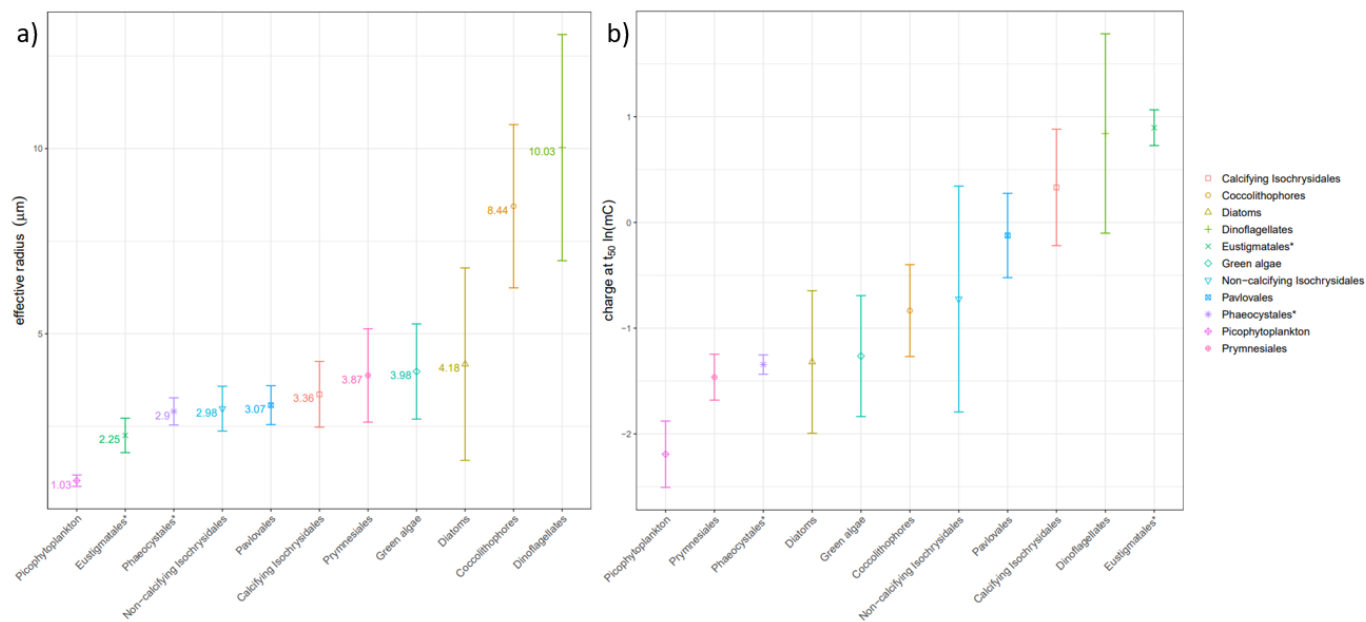
293

294

295 **Assessment of the susceptibility library**

296 With the two parameters of charge at t_{50} (mC) and effective radius (μm) defined for each individual
 297 cell, we assessed the applicability of the susceptibility library for distinguishing the cells into pre-
 298 defined groupings of relevance, as defined in the Culturing section above. Prior to this assessment it
 299 was necessary to balance the dataset for even strain representation within each pre-defined grouping.
 300 This was necessary as there was limited control over the number of individual cell transients obtained
 301 per experimental image series per strain (due to unpredictable numbers of cells depositing on the
 302 electrode surface), resulting in under/over-representation within groups (see Table S3, SI). To balance
 303 the data at the group level, we identified the strain within each group that had the lowest number of
 304 observations, and used that minimum number to randomly subset a sample of the same length for each
 305 of the other strains within the grouping level. This was achieved using the “*sample_n*” function in the
 306 R package “*dplyr*” (R version 4.2.2). The resultant balanced dataframe ($n = 2277$) was subsequently
 307 used to derive mean values at the group level and for the analyses described below (see Tables S4 and
 308 S5 for balanced data, SI).

309



310

311 **Figure 2. (a)** Mean effective radius (μm) at the level of each phytoplankton grouping defined in this
 312 study. The numbers alongside respective datapoints are the mean effective radius in μm . **(b)** Natural
 313 logarithm converted mean charge at t_{50} (mC) for each phytoplankton grouping. Data points represent
 314 the mean for each group, and error bars represent the standard deviation of the mean (see Table S5).
 315 Colour coding and shapes represent each group (see legend). Note: the ordering on the x-axis is in
 316 ascending order for each plot, demonstrating that across the groups larger cell radius does not
 317 necessarily result in greater charge at t_{50} .

318

319 First, looking at charge at t_{50} independently from radius, across all strains we see a large range spanning
 320 three orders of magnitude (see Table S4, SI), from a mean of 0.08mC (± 0.02 S.D.) for the pico-

321 eukaryote *Ostreococcus tauri*, up to 7.34mC (± 1.88 S.D.) for the dinoflagellate *Scrippsiella trochoidea*.
322 At the level of the pre-defined groupings described above we see the lowest mean charge at t_{50} within
323 the “Picophytoplankton” of 0.11mC (± 0.03 S.D.), and the greatest mean charge at t_{50} within the
324 “Eustigmatales” 2.45mC (± 0.41 S.D.), preceded by “Dinoflagellates” at 2.32mC (± 2.22 S.D.) (see Fig.
325 2b and Table S5, SI). The large range in these values indicate that there is a strong effect of cell grouping
326 on the resilience to electrochemically driven oxidative stress. Critically, however, there is also
327 considerable overlap in the values, as an example: “Calcifying Isochrysidales” have a mean charge at
328 t_{50} of 1.39mC (± 0.57 S.D.), which sits well within the large deviation range of the “Dinoflagellates”.
329 Subsequently, in order to further distinguish the groupings in such instances, use of the effective radius
330 can provide an additional dimension for separation where there is overlap. In the case of the previous
331 example, we see that “Calcifying Isochrysidales” have a mean effective radius of 3.36 μ m (± 0.89 S.D.),
332 whereas “Dinoflagellates” have a significantly larger mean effective radius of 10.03 μ m (± 3.05 S.D.)
333 (see Fig. 2a and Table S5, SI). Taking this forward, we next use a random forest analysis to test the
334 potential for single cell categorisation across all of the groups based on the variance in both charge at
335 t_{50} and effective radius.

336

337 Random forest analysis uses the pre-defined classes (in this case the phytoplankton groupings) to
338 construct a range of ‘decision trees’ for discrimination of the individual observations (in this case each
339 phytoplankton cell) based upon the predictor variables (in this case charge at t_{50} and effective radius) of
340 a dataset. To effectively test the accuracy of classification, this requires input of a ‘training’ dataset so
341 that the random forest algorithm can create the necessary discrimination functions for the pre-defined
342 classes. A ‘testing’ dataset can then be used to determine the accuracy of the discrimination on an
343 independent set of ‘blind’ observations. As an example of its application in a relevant field, random
344 forest algorithms have previously been used to successfully distinguishing individual populations of
345 phytoplankton strains from flow cytometry measurements on artificial communities (Bestion et al.
346 2020, 2021). Whilst a flow cytometer records a wide range of fluorescence and light scattering variables
347 per individual cell, here we are limited to testing the distinguishing power of our method with only the
348 two aforementioned predictor variables (though additional variables for future iterations of the method
349 are discussed later). Specifically, for our analysis, we used the “randomForest” function in the R
350 package “randomForest”; this function uses Breiman’s random forest algorithm for classification
351 (Breiman 2001). From this we could determine categorisation ‘decision trees’ for two subsets of the
352 susceptibility library data: (a) The key functional groups only (5 groups) with the balanced dataset (see
353 Fig. 3) and (b) all groups (11 groups) with the balanced dataset (see Fig. 5, Table S5). For each subset,
354 we randomly split the datasets using the “sample_n” function as described above, assigning 80% of
355 data for the training component, and 20% for the testing. The two input predictor variables of charge at
356 t_{50} and effective radius were natural log transformed prior to running the analysis. Following the training

357 and subsequent testing, confusion matrices were returned to demonstrate the predictions of the random
358 forest models (see Fig. 4 and 6, and Tables S6-S9). At the level of group for each testing subset, we
359 investigated the prediction success in terms of recall, precision, and F1-score metrics (Kraft et al. 2022),
360 see Tables 1 and 2.

361 Recall determines how well the random forest quantifies true positives (TP) for each class *i.e.* a higher
362 recall means fewer false negative predictions are made (FN):

$$363 \text{ Recall} = (\text{TP} / (\text{TP} + \text{FN}))$$

364 Precision determines how well incorrect false positives (FP) are rejected for each class *i.e.* a higher
365 precision means fewer false positive predictions are made, and more true negatives (TN) are correctly
366 identified:

$$367 \text{ Precision} = (\text{TP} / (\text{TP} + \text{FP}))$$

368 The F1-score combines both the recall and precision into a single metric to define the overall prediction
369 accuracy for each class:

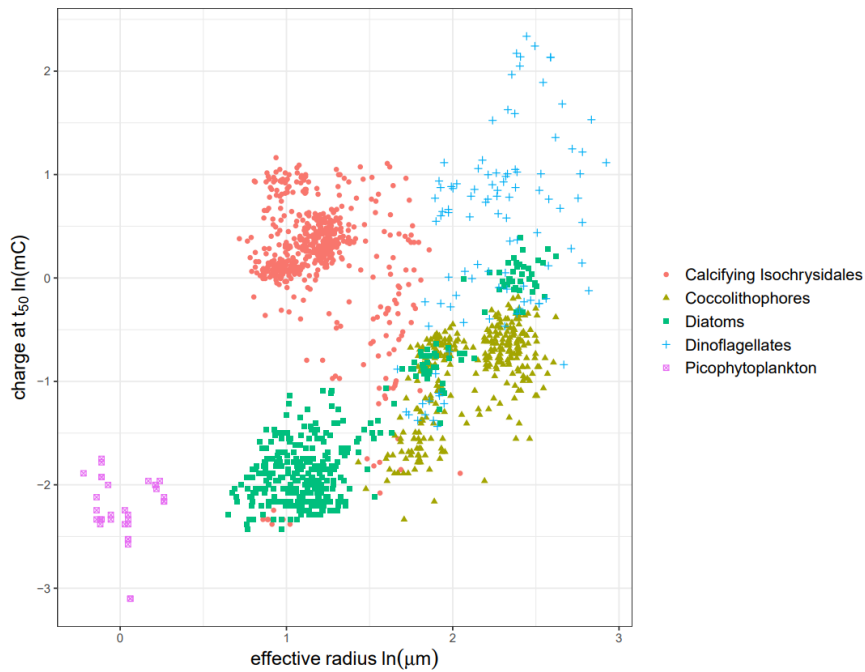
$$370 \text{ F1-score} = (2 * (\text{precision} * \text{recall}) / (\text{precision} + \text{recall}))$$

371 In addition to the above, overall accuracy of the random forest was returned to demonstrate the success
372 rate of categorisation across the testing dataset:

$$373 \text{ Accuracy} = (\text{TP} + \text{TN}) / (\text{TP} + \text{FP} + \text{TN} + \text{FN})$$

374

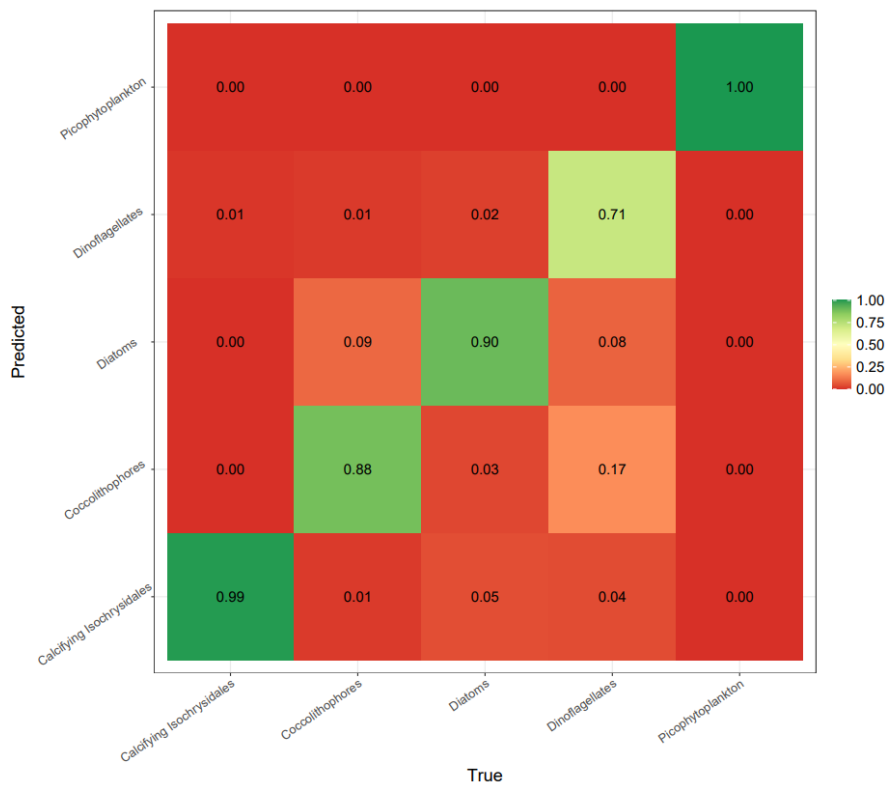
375



376

377 **Figure 3.** A scatterplot of natural log charge at t_{50} (mC) against natural log effective cell radius (μm) for the ‘key
 378 functional groups’ of cells measured in this study ($n = 1406$, see Table S5 for a summary of the balanced dataset).
 379 Whilst there is some overlap of the groupings, the random forest analysis on testing subset of this data returned
 380 an overall accuracy of 92% (see Fig. 4, Table 1, and Table S7).

381



382

383 **Figure 4.** A normalised confusion matrix demonstrating the prediction success of testing the random forest model
 384 with just the key groups included. To normalise, predicted number of cases were divided by the true total number
 385 of cases in each group (see Table S7 for original values).

386 **Table 1.** Evaluation metrics obtained from testing the random forest model trained for the key groups only.

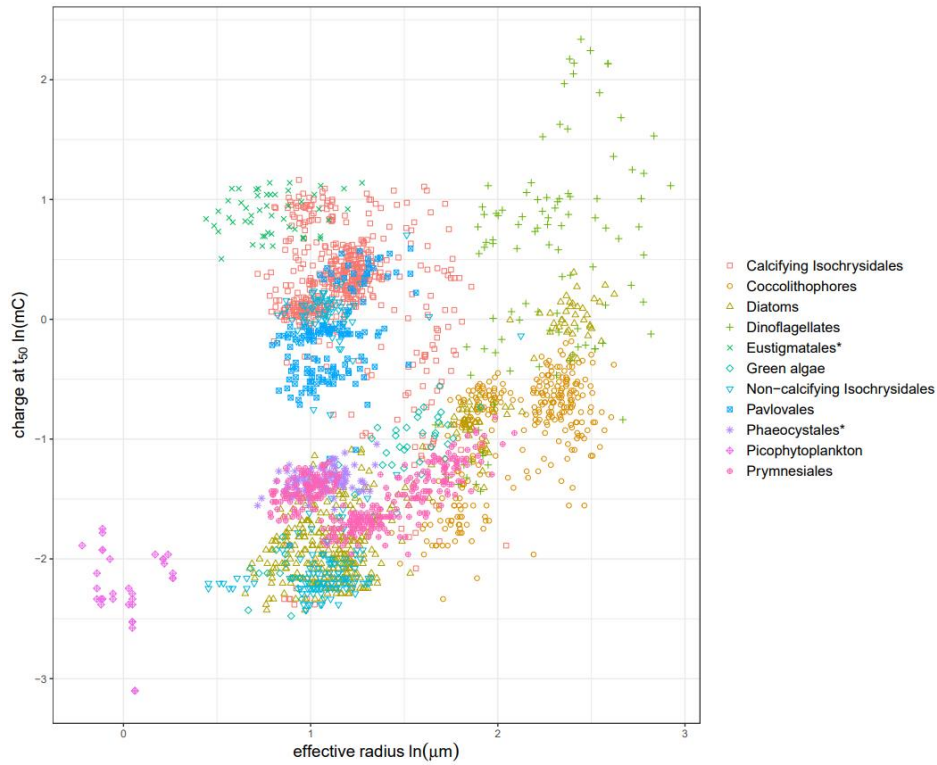
Group	Recall	Precision	F1-score
Calcifying Isochrysidales	0.99	0.95	0.97
Coccolithophores	0.88	0.91	0.89
Diatoms	0.9	0.89	0.9
Dinoflagellates	0.71	0.81	0.76
Picophytoplankton	1	1	1
Overall accuracy			0.92

387

388

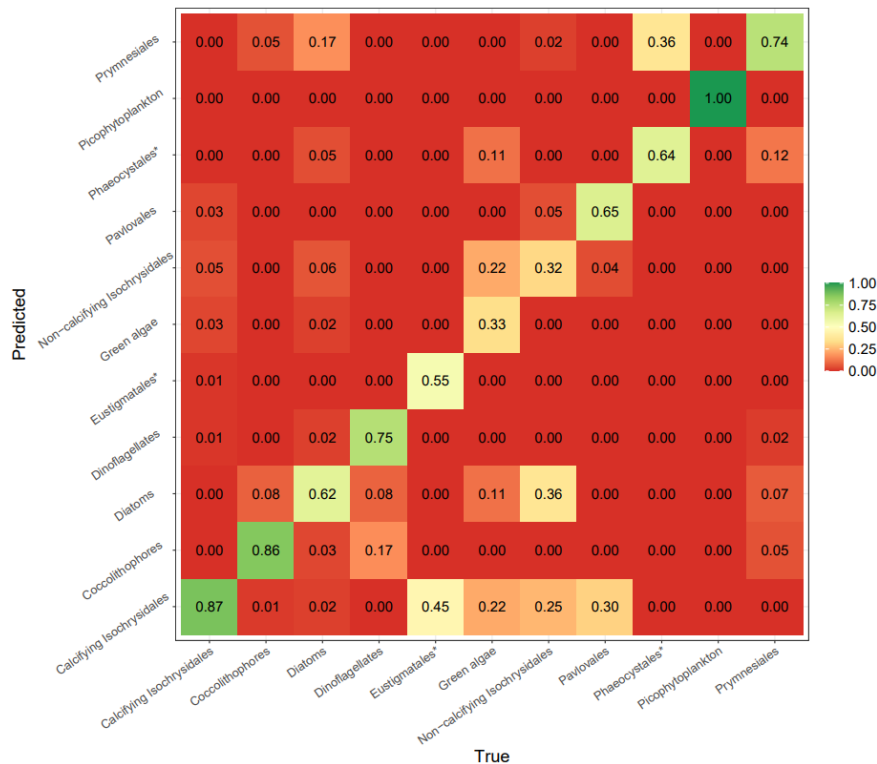
389 From training the random forest with the balanced dataset of the key groups only, we found there to be
390 an overall testing accuracy of 92% (see Table 1, and Table S7). The high level of accuracy is perhaps
391 unsurprising considering the visual clustering of the datapoints (see Fig. 3). Within the groupings
392 however, some classifications perform better than others. “Picophytoplankton” are correctly classified
393 in all instances (recall = 1), and this is primarily driven by their much smaller size relative to the other
394 groups. “Calcifying Isochrysidales” are the next best predicted group with a recall of 0.99, and this
395 appears to primarily be driven by their greater electrochemical resilience (*i.e.* greater charge at t_{50})
396 relative to cells of a similar small size. The poorest performance was in classifying “Dinoflagellates”,
397 with a recall of 0.71, albeit having a precision score of 0.81. This is likely due to considerable variation
398 in the charge at t_{50} within this group, meaning that some strains of dinoflagellate are misclassified as
399 larger diatoms and coccolithophore, contributing to a greater number of false negatives in this instance.
400 On the whole, the fact that all five of the key groups had F1-scores >0.75 is an indication that the
401 electrochemical sensitivity provides a good degree of separation, in combination with size. Nonetheless,
402 in order to be more reflective of the diversity of nanophytoplankton found in naturally occurring
403 communities, we next considered the level of performance once all of the other groups are included in
404 the random forest analysis (see Fig. 5).

405



406

407 **Figure 5.** A scatterplot of natural log charge at t_{50} (mC) against natural log effective cell radius (μm) for all groups
 408 of cells measured in this study ($n = 2277$, see Table S5 for a summary of the data presented here). Compared to
 409 Fig. 3, where just the key functional groups are presented, we now see more overlap of the groupings. The random
 410 forest analysis on the testing subset of this data returned an overall accuracy of 71% (see Fig. 6, Table 2, and
 411 Table S9). *denotes groups that were just represented by a single species.



412

413 **Figure 6.** A normalised confusion matrix demonstrating the prediction success of testing the random forest model
 414 with all groups included. To normalise, predicted number of cases were divided by the true total number of cases
 415 in each group (see Table S9 for original values). *denotes groups that were just represented by a single species.

416 **Table 2.** Evaluation metrics obtained from testing the random forest model trained for all groups. *denotes groups
 417 that were just represented by a single species.

Group	Recall	Precision	F1-score
Calcifying Isochrysidales	0.87	0.8	0.83
Coccolithophores	0.86	0.87	0.86
Diatoms	0.63	0.65	0.64
Dinoflagellates	0.75	0.82	0.78
Eustigmatales*	0.55	0.86	0.67
Green algae	0.33	0.33	0.33
Non-calcifying Isochrysidales	0.32	0.5	0.39
Pavlovaes	0.65	0.71	0.68
Phaeocystales*	0.64	0.54	0.58
Picophytoplankton	1	1	1
Prymnesiales	0.74	0.61	0.67
Overall accuracy			0.71

418

419 When all of the groupings are considered, following training, the overall accuracy on testing was 71%
 420 (see Table 2). The lower level of accuracy compared to the interpretation with only the key groups is
 421 also unsurprising, given that the number of potential classes in the random forest has more than doubled
 422 (5 to 11) and considering the greater extent of overlap in the group level clustering of the datapoints
 423 (see Fig. 5). Despite the overall reduction in accuracy, all of the key groups maintain a relatively high
 424 level of predictability with all five returning recall, precision and F1 scores of more than 0.6. Of the key
 425 groups, the biggest reduction in performance is in the “Diatom” group, with recall being reduced from
 426 0.9 in the key groups subset (see Table 1) to 0.63 when all groups are included. An element of confusion
 427 in this instance is being caused by cell types belonging to the other groups, such as “Non-calcifying
 428 Isochrysidales” and “Prymnesiales”, which have similar size and electrochemical susceptibility
 429 distributions. Of the additional groups, the best performing were the “Pavlovaes” with an F1-score of
 430 0.68, closely followed by “Prymnesiales” and “Eustigmatales”, both with F1-scores of 0.67. The worst
 431 performing were, “Green algae” and “Non-calcifying Isochrysidales” with F1-scores of 0.33 and 0.39,
 432 respectively. In both cases, considerable confusion was coming from “Diatoms” and “Calcifying
 433 Isochrysidales”, and specifically for “Green algae” there was substantial confusion with “Non-
 434 calcifying Isochrysidales”. As dinoflagellates are often the group most commonly associated with
 435 harmful algal blooms, it is interesting to note that we see a higher precision (0.82) than recall (0.75) for
 436 this group. This indicates that whilst the random forest model is better at rejecting false positives, it
 437 could be improved in terms of reducing false negatives (i.e. true cases of dinoflagellate that are being
 438 missed). From an applicability point of view, this could suggest that as things stand, improvement is
 439 needed if such a technique was to be employed to monitor for potentially harmful algae.

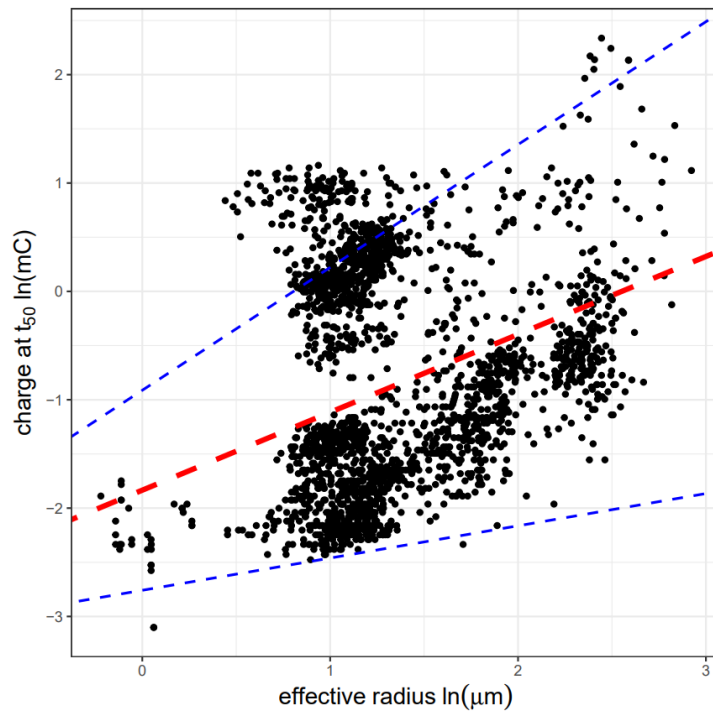
440 Significantly, of the key groups, there were some notable high performers following testing, despite the
 441 inclusion of the other groupings: “Calcifying Isochrysidales” with a recall of 0.87, “Coccolithophores”

442 a recall of 0.86, and for “Picophytoplankton” recall remained at 1. In the case of the “Coccolithophores”,
443 their relatively strong predictability is likely due to their low electrochemical resilience relative to their
444 large size, in contrast to “Dinoflagellates” of a comparable size which generally have a greater
445 resilience. “Calcifying Isochrysidales” on the other hand are highly distinguishable for their remarkable
446 electrochemical resilience relative to other groups of a similar smaller size. This poses some key
447 questions about the underlying mechanism of this technique for distinguishing cell types across the
448 groups: How much is cell size driving the electrochemical resilience? Are there any contradicting
449 exceptions to any potential size scaling? With this in mind, we next scrutinise the effect of size on the
450 susceptibility factor to see how much this is driving the overall resilience to the oxidative stress across
451 the groups.

452

453 **Understanding the mechanism: the size scaling of the relationship and size adjusted** 454 **resilience**

455 Using the balanced dataset of all groupings, we investigated if there was a significant linear relationship
456 between natural log transformed charge at t_{50} and effective radius (see Fig. 7). This was achieved using
457 the ‘*lme4*’ package in R for linear mixed effects modelling, whereby we fitted and compared linear
458 models to the data both with and without the random effect of grouping on both the intercept and slope
459 of the response. Of the two models, the model including the random effect of grouping on both the slope
460 and intercept scored more favourably than the model without (see Table S10, SI), indicating that there
461 was an overall significant positive linear scaling between charge at t_{50} and effective radius ($p < 0.001$),
462 but that this scaling was highly variable across the groupings. The overall model returned a slope value
463 of 0.72 (95% CIs: 0.30 – 1.13). Out of the 11 ‘groups’, 7 of them had significant positive within group
464 size dependence, notably “Green algae”, “Diatoms”, and “Dinoflagellates” had particularly strong
465 scaling of more than 1 (See Fig. S7 and Table S11, SI). There was no significant size scaling within the
466 groups of “Picophytoplankton”, and “Eustigmatales”, “Phaeocystales”; this is perhaps unsurprising
467 given that the latter two were only represented by one single strain. Notably, the only group that had a
468 significant negative size scaling were the “Calcifying Isochrysidales”, -0.48 (95% CIs: -0.66 – -0.31).

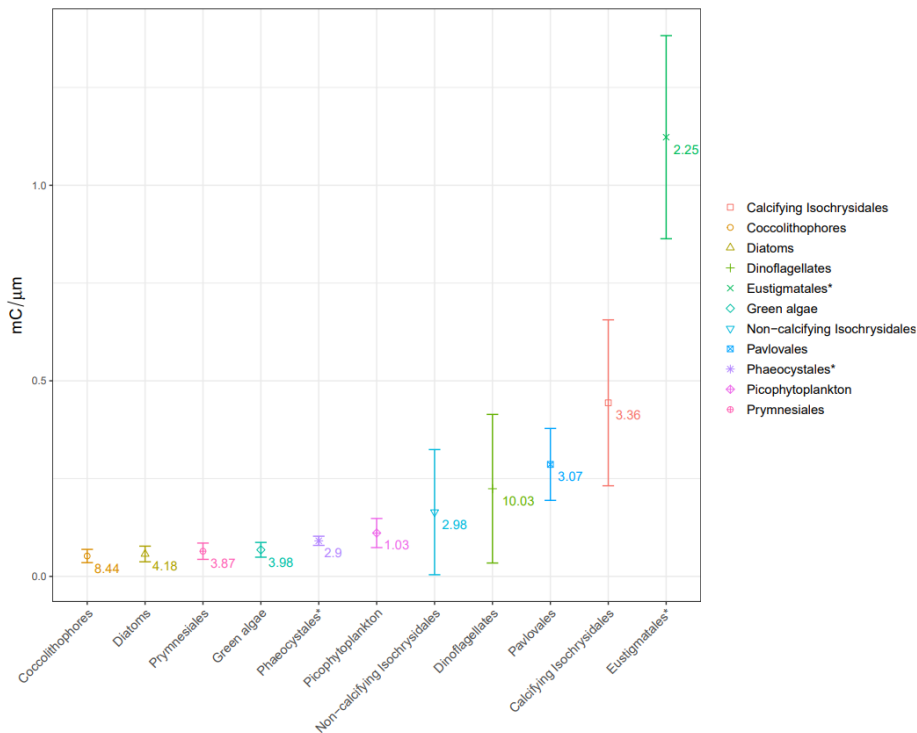


469

470 **Figure 7.** A scatterplot of natural log charge at t_{50} (mC) against natural log effective cell radius (μm) of cells
 471 measured in this study, following the balancing of strain representation per group ($n = 2277$, see Table S4 SI for
 472 a summary of the data presented here). The red line indicates the overall slope (0.72) of the allometric relationship
 473 modelled using linear mixed effects with the random effect of grouping on the slope and intercept factored for,
 474 and the blue dashed lines indicate the 95% confidence of this model fit (slopes of 0.30 and 1.13 for lower and
 475 upper, respectively), see Tables S10 and S11 for statistics.

476

477 On the whole, this demonstrates that across most of the groupings there is an intrinsic allometric scaling
 478 of charge required to “switch-off” the chlorophyll *a* signal. This was previously found at the species
 479 level in a recent study by Yu *et al.* (2023), which demonstrated that differing electrochemical
 480 susceptibility of life stages of *Chlamydomonas concordia*, were primarily driven by size variation in
 481 the different cell types, and not necessarily an underlying biological factor (Yu et al. 2023).
 482 Consequently, we could postulate that any significant differences in electrochemical susceptibility
 483 following a size normalisation might therefore indicate which of the groupings in this study have an
 484 ‘unknown’ underlying biological feature that results in higher or lower resilience. We investigated this
 485 by normalising all charge at t_{50} values for size, by simply dividing charge (mC) by effective radius (μm)
 486 for each of the individual measurements in the balanced dataset (see Fig. 8).



487

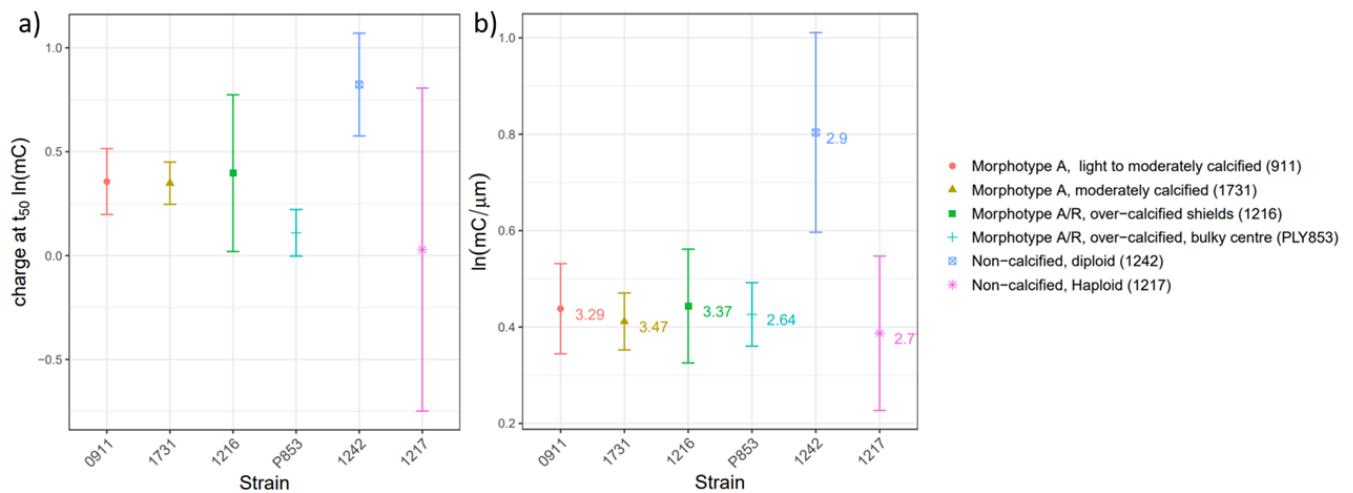
488 **Figure 8.** Comparison of per group means of size normalised charge at t_{50} , following the balancing of strain
 489 representation per group. Data points represent the mean value for each group and the error bars are for standard
 490 deviation. The labelled numbers alongside the data points represent the mean effective radius for each of the
 491 groups (in μm), illustrating that in some instances there is a disproportionate resilience to the electrochemical
 492 charge relative to cell size – notably for “Calcifying Isochrysidales” and “Eustigmatales”. For pairwise
 493 comparisons see Table S12, SI. *denotes groups that were just represented by a single species.

494

495 Following size normalisation of the charge at t_{50} , we carried out pairwise comparisons across the groups
 496 using Pairwise Wilcoxon Rank Sum testing (due to the non-parametric distribution of data in most of
 497 the groups), for this we used the function “*pairwise.wilcox.test*” in the R package “*stats*”, with
 498 “*p.adjust.method*” set to the “Bonferroni” correction of p values see Table S12, SI. We see no
 499 significant difference between a number of the key groupings, notably “Diatoms” and
 500 “Coccolithophores” ($p = 0.388$), and “Picophytoplankton” and “Dinoflagellates” ($p = 0.325$). Given
 501 that “Diatoms” in this dataset have a mean effective radius of $4.18\mu\text{m}$ (± 2.60 S.D.) and
 502 “Coccolithophores” of $8.44\mu\text{m}$ (± 2.20 S.D.), this indicates that most distinguishing between these two
 503 groups within the random forest must primarily be driven by size, given that their size normalised charge
 504 values are indistinguishable. Likewise, “Picophytoplankton” have a mean effective radius of $1.03\mu\text{m}$
 505 (± 0.15 S.D.) and “Dinoflagellates” a mean effective radius of $10.03\mu\text{m}$ (± 3.05 S.D.), demonstrating
 506 extreme ends of the size spectrum within this dataset, yet after size normalisation of their respective
 507 charge values they are indistinguishable in terms of their electrochemical resilience. Contrary to these
 508 observations, it is evident that there are some clear outliers, whereby following size normalisation they
 509 are more distinguishable from the rest of the groups, notably “Calcifying Isochrysidales” and
 510 “Eustigmatales”. In both cases, their size normalised charge was statistically greater than all the other

511 groups, with “Eustigmatales” having the greatest overall per unit size resilience (see Fig. 8, and Table
 512 S12). Whilst we only have one strain representing “Eustigmatales” in this dataset, *Nannochloropsis*
 513 *granulata*, we might infer that there is something about the particular biology of these two groups that
 514 is driving their greater resistance to the high levels of oxidative stress. This is of relevance to the method,
 515 because across all our assessments using the random forest analysis it was the “Calcifying
 516 Isochrysidales” group that consistently retained one of the highest levels of prediction accuracy of the
 517 key functional groups. Taking our investigation further, we next take a closer look at within species
 518 variability to disentangle if any of the particular strains of *E. huxleyi* (from the “Calcifying
 519 Isochrysidales” group), representing different life stages and calcification morphologies (Green et al.
 520 1996; Young et al. 2003; Bendif et al. 2023), can help to identify any further trends that may lead to
 521 understanding the biological mechanism.

522



523

524 **Figure 9. (a)** Comparison of natural log transformed mean charge at t_{50} across the different strains of *E. huxleyi*
 525 representing different coccosphere morphologies (0911 (RCC911), 1731 (RCC1731), 1216 (RCC1216),
 526 (PLY853)), and non-calcified diploid and haploid life stages (1242 (RCC1242), (1217 (RCC1217)). For details
 527 on the strains see Table S1, **(b)** Comparison of natural log transformed size normalised charge across the same
 528 strains. The labelled numbers alongside the data points represent the mean effective radius for each of the strains
 529 (in μm), illustrating that across the strains there is minimal difference in size. For pairwise comparisons see Table
 530 S13, SI. For both plots, data points represent the mean value for each group and the error bars are for standard
 531 deviation.

532

533 Despite their differing coccosphere morphologies (ranging from light-, moderate-, and over- calcified
 534 features), in most cases across the four calcified (diploid) strains there was negligible difference in the
 535 size normalised charge at t_{50} (see Fig. 9, and Table S13). This result is reassuring in terms of the
 536 proposed sensing technology, as it suggests that the within strain variance of “Calcifying
 537 Isochrysidales” is not too substantial to cause confusion with other groups. Of particular curiosity, the
 538 ‘1242’ non-calcified diploid strain (RCC 1242), had a significantly greater charge per unit size relative
 539 to all other strains, and furthermore the ‘1217’ non-calcified haploid strain (RCC 1217) was generally

540 indistinguishable from the calcified strains (except for its calcified diploid version RCC 1216). This
541 observation is of relevance to understanding any potential underlying mechanism to the sensor
542 technology for two reasons. Firstly, it indicates that there is minimal effect of the extracellular inorganic
543 calcite layers in either increasing or reducing the resilience of the cell type. This is also more broadly
544 demonstrated when comparing the “Calcifying Isochrysidales” group to the “Coccolithophores”, the
545 latter in many instances (*e.g. Coccolithus braarudii*) having much larger extracellular calcium carbonate
546 shell volumes (Yang et al. 2022), yet relatively much faster “switch-off” times and lower size
547 normalised charge tolerance under this method (see Fig. 2b, and Fig. S9, SI). Secondly, it suggests that
548 there is something biologically different between the non-calcified and calcified diploid cells that is
549 driving the discrepancy in the resilience. Disentangling this is beyond the scope of this study, but
550 presents an avenue for further investigation as to why such cell types have greater resilience. Indeed,
551 along a similar vein of thinking, exploring the biological differences of each of the groups against the
552 highly resilient *Nannochloropsis granulata* could help to understand what is driving such resilience
553 irrespective of the cell size. As a tentative suggestion, greater resilience could simply be determined by
554 the membrane structure of the cell, whereby the presence of more membrane layers surrounding the
555 cytoplasm and/or the plastids could potentially present a barrier acting to slow down the transmission
556 of the oxidative radicals to the chlorophyll molecules (Yu et al. 2022). It has previously been reported
557 that *Isochrysidales* (and specifically *E. huxleyi* and *G. oceanica*) are distinctive from other
558 coccolithophorids due to a number of different periplast and membrane features, including:
559 unmineralized outer cell scales, more rigid double membrane structures of the peripheral endoplasmic
560 reticulum, and unique long-chain membranous alkenones (Fujiwara et al. 2001). Perhaps it is features,
561 such as these, slowing the transmission of radical oxidants to the chloroplasts of “Calcifying
562 Isochrysidales”, resulting in apparent resilience relative to other groups in this study.

563

564 **Discussion**

565 Overall our method has demonstrated a good degree of accuracy when it comes to making
566 classifications of cell types into ecologically relevant groups. The groups that consistently had the
567 greatest accuracy were “Calcifying Isochrysidales”, “Coccolithophores”, and “Picophytoplankton”; all
568 of which maintained a recall of >0.85, and precision and F1-scores of >0.8, across the random forest
569 predictions (both key group and all group libraries). The inclusion of the suspected “nano-
570 phytoflagellate” groups did cause some overall reduction in the accuracy of the technique, from 92% to
571 71%, but given that we have just two variables to make predictions (charge at t_{50} and effective radius),
572 this is something that we anticipate can be improved if the technique was adjusted to capture additional
573 predictor variables. The findings of the susceptibility library assessment also demonstrate that following
574 size normalisation there must be some underlying biological feature that is enhancing the strong
575 classification ability of some of the groups, notably “Calcifying Isochrysidales” and “Eustigmatales”,

576 and thus this presents an important avenue for further investigative studies into the potential mechanistic
577 underpinnings.

578

579 **Current applicability of the method and recommendations**

580

581 Whilst we demonstrate the ability of our technique to classify nanophytoplankton into groups of
582 relevance, it is critical that the method can quantify the abundance of different groups in natural
583 seawater samples. This will require field testing alongside more traditional techniques *e.g.* microscope
584 taxonomy and cytometry to validate the applicability of using a predefined susceptibility library based
585 on a limited selection of 52 monocultures. Methodological and engineering advancements will need to
586 be made to take this method to such a stage. As things stand, in obtaining the susceptibility
587 measurements presented here, samples of an uncontrolled volume were drop cast following
588 concentration by centrifugation and left to settle on the electrode beforehand (Kumar et al. 2020).
589 Therefore, given the uncertainties around the volume of sample used, it was not possible to quantify the
590 original abundance of cells in the samples measured. A prototype instrument that implements the fluoro-
591 electrochemical technique with a flow-cell type system could help to overcome this issue, as both
592 sample volume and flow rate could be quantifiable and controllable. To achieve the ‘current ramping’
593 approach as demonstrated in this study it is likely that a series of in-flow ring electrodes with different
594 applied currents, separated at periodic intervals, would be required in a future flow-type device. This
595 may allow for a similar level of classification, but with the advantage of the high throughput of a flow-
596 cell setup where number of events can be quantified per unit time, providing a measurement of actual
597 abundance for the different classified cell types. Not only could this make the laboratory-based
598 measurements higher throughput (akin to imaging flow cytometry), it would also be an essential step
599 towards the long-term goal of the technique being used on autonomous platforms, whereby the
600 collection of *in situ* spatio-temporal data of nanophytoplankton community structure would be
601 invaluable to monitoring the effects of environmental change. From a practical point of view, our
602 technique could lend itself well to long-term *in situ* monitoring via an autonomous platform. The
603 electrochemical technique is reagent free (whereby seawater serves as the electrolyte), thus minimising
604 the need for regular retrieval of the device. Furthermore, the production of oxidants on the electrode
605 surface provides an intrinsic anti-fouling mechanism; indeed, electrochemical generation of hydroxyl
606 radicals is a technique that has been successfully used to rapidly kill invasive phytoplankton species in
607 ship’s ballast water within seconds, not dissimilar to what we see in the experiments presented here
608 (Zhitao et al. 2005; Bai et al. 2010, 2012; Zhang et al. 2013). Given the relatively simple optics set-up
609 of our technique, compared to others, it is also likely the key components be scaled down from the
610 current lab set-up (see Fig.1a), making it more feasible for deployment in the field.

611 As mentioned previously, devices such as FlowCam, CytoSense, CytoBuoy and IFCB use combinations
612 of flow cytometry fluorescence measurements with rapid imaging of cells. Whilst highly progressive in
613 helping us to understand planktonic communities, they fall short in their ability to distinguish at the
614 much smaller size ranges – especially when many cell types in the nanophytoplankton range can have
615 a similar apparent morphology and size. Understandably, there is a consequent trade-off between
616 magnification, image quality, and the size range of phytoplankton being measured. This is likely to
617 make any machine learning algorithms poorer at the lower limits of the nanophytoplankton size range.
618 The key novelty of our classifying technique at the nanophytoplankton level is the extra tool for
619 distinguishing phytoplankton cell types that has not been previously applied: the electrochemical
620 susceptibility of the chlorophyll *a* fluorescence signal, which is less dependent on magnification and
621 image resolution. We can get a good degree of categorisation overall from simply combining the charge
622 at t_{50} value with a simple effective radius estimate. We also demonstrate that our design has potential
623 to distinguish picophytoplankton, as well as larger nanophytoplankton, spanning three orders of
624 magnitude.

625 It was evident from our assessment that some groups were more easily predicted than others, especially
626 when all the groups were considered. Of the key functional groups, our technique currently falls short
627 when it comes to distinguishing diatoms (F1-score of 0.64, see Table 2), and of the other groups, the
628 lowest accuracy was in predicting “Non-calcifying Isochrysidales” and “Green algae” (F1-scores of
629 0.39 and 0.33 respectively, see Table 2). Therefore, due to the differing levels of classification ability
630 across the groups as things stand, it is likely the set-up would yield more promising *in situ* measurements
631 from nanophytoplankton communities dominated by taxa belonging to the more easily classifiable
632 groups presented here. To advance our method further, and with minimal increase in cost, a couple of
633 adjustments to both the apparatus and method could give us additional variables that will likely improve
634 predictive ability. Firstly, with the addition of more excitation and emission pathways, a measurement
635 of secondary chlorophyll *b* pigment fluorescence could be obtained. In marine phytoplankton, this
636 pigment is unique to chlorophytes (or green algae). In the nanophytoplankton range, a large degree of
637 confusion for our method was between the “Diatoms” and “Green algae”, and therefore this addition
638 would help to reduce this. Such an advancement could also assist at the picophytoplankton scale, where
639 the majority of pico-eukaryotes are either prasinophytes containing chlorophyll *b*, or cyanobacteria
640 containing phycocyanin (a chlorophyll accessory pigment, also with distinguishable autofluorescence
641 properties).

642 Secondly, where there is overlap between calcified cell groups and others, we might be able to use the
643 intrinsic dissolution of the calcite during the electrochemical experiments to observe changes in the
644 apparent radius before and after the experiment. In brief, the electrochemical oxidation of water means
645 that H^+ is generated in the vicinity of the electrode, decreasing the pH around the cells. Consequently,
646 it has been observed that during the short time span of the experimental measurements presented here,

647 the smaller extracellular calcite coccospheres (*e.g. E. huxleyi*) can be completely dissolved, such that a
648 before and after measure of cell radius could indicate the calcification of a cell. Previous work has
649 demonstrated that such a method can also be applied to estimate the mass of extra-cellular calcium
650 carbonate of entire coccospheres (Yang et al. 2022; Fan et al. 2022), which is another relevant
651 measurement for understanding marine biogeochemical cycles, notably the ‘rain ratio’ (Hutchins 2011).
652 Having an additional predictor variable of the ratio of cellular radius before and after the fluoro-
653 electrochemical experiment would thereby provide an entirely affordable and achievable additional
654 dimension for improved differentiation of calcified and non-calcified cell types in the random forest
655 algorithms, as well as generating an estimate of cellular calcite which could be of great value to ocean
656 biogeochemists.

657 Lastly, additional predictor variables can be gained by taking advantage of the full range of data that is
658 harvested from the experiments presented here. In terms of the chlorophyll *a* fluorescence measurement,
659 we only use a single parameter derived from each transient profile: charge at t_{50} . If the full transient
660 profiles were to be assessed then it is likely this could improve the predictive power of the susceptibility
661 library. Indeed, the shape of the transients appear to be idiosyncratic at a group-specific level (see Fig.
662 S2, SI). With this in mind, by extracting the time (and thus charge) data for additional stages of
663 normalised chlorophyll *a* fluorescence intensity (*e.g.* at 75%, 25%, 0%) we are more likely to capture
664 the variation in the shape, or gradient, of the ‘switch-off’ profile. Likewise, in terms of the bright-field
665 imaging, we only make a relatively crude interpretation of the cell radius. There are other variables that
666 can be characterised from the images, such as the minor and major axis lengths to determine aspect
667 ratio (and thus giving an indication on how spherical or elongated a cell is).

668 Taking things forward, whilst we demonstrate relatively good accuracy with just using the two predictor
669 variables used in this study (charge at t_{50} and effective radius), a greater wealth of data could be yielded
670 with only minor methodological tweaks (as suggested above). By training the random forest with more
671 variables (such as: secondary pigment fluorescence, change in cell radius before and after experiment,
672 chlorophyll *a* fluorescence at different stages in the ‘switch-off’, and various cell characteristics from
673 2D bright-field images) this is likely to improve the classification accuracy across more of the relevant
674 groups.

675 Overall, following on from the promising results presented here, we foresee that our technique (subject
676 to the improvements discussed) could be extremely complimentary to broader efforts of ocean
677 monitoring. The focus of our technique on the nanophytoplankton size fraction can add detail that is
678 currently missing from existing techniques. As we demonstrate, there is potential for distinguishing
679 cells into relevant groupings, going beyond the limited capacity of standard flow cytometry where, as
680 things stand, only a select few of the functional groups can be differentiated within nanoeukaryotic
681 communities; coccolithophores and cryptophytes (Tarran et al. 2006; Tarran and Bruun 2015).

682 Furthermore, as discussed above, the application of our technique would help to unlock detail at the
683 finer size scale that is also poorly characterised by current *in situ* imaging flow cytometry. Through
684 developing our ‘susceptibility library’ with a diverse range of taxa, we also expect that our technique
685 could help to tease out more information from previously labelled unknown ‘nano-phytoflagellate’
686 populations in natural communities, which are notoriously problematic for taxonomists due to their
687 similar sizes and morphology when viewed under the microscope (Widdicombe et al. 2010; Piwosz
688 2019). For example, both “Pavloales” and “Prymnesiales” haptophyte groups in this study had F1-
689 scores of >0.65, indicating that whilst they might be of a similar size and morphology (and thus are
690 likely “nano-phytoflagellate” candidates), they are somewhat distinguishable by their susceptibility to
691 the electrochemically driven oxidative conditions.

692

693 **Summary**

694

695 We have presented a novel fluoro-electrochemical technique for classifying marine nanophytoplankton,
696 and critically assessed this by testing its ability to predict phytoplankton groupings from two simple
697 parameters: charge required to reduce per cell chlorophyll *a* fluorescence by 50% and effective cell
698 radius. This returned an excellent degree of accuracy when only considering taxa belonging to key
699 functional groups (5 groups), but a reduced degree of accuracy when a broader range of groups,
700 encompassing likely “nano-phytoflagellates”, were considered (11 groups). We demonstrate that the
701 technique relies on the general positive size scaling of the susceptibility across the groups to provide
702 additional distinguishing power, and when size is normalised for there are some groups that demonstrate
703 exceptional resilience to the highly oxidative conditions of our technique, notably “Calcifying
704 Isochrysidales” and “Eustigmatales”. This presents an avenue for further investigation into the
705 biological underpinnings of this new method. Whilst the technique currently has its limitations as we
706 report, with advancement of the set-up to make complimentary measurements, the predictive power of
707 the method could be enhanced. Critically, if the technique is to eventually be used for real world *in situ*
708 measurements, the next step of assessment is to test its ability in quantifying abundance of different
709 groupings in natural samples alongside more traditional techniques. If such further testing yields
710 positive results, we anticipate that our technique could be adapted to work in conjunction with
711 autonomous platforms, with the potential to greatly enhance our ability in monitoring
712 nanophytoplankton community structure.

713

714

715

716 **Author Contributions**

717 S.B., M.Y., C.B.-M., R.G.C., H.A.B., and R.E.M.R. conceptualized the study; S.B. (phytoplankton),
718 M.Y., and C.B.-M (electrochemistry), contributed to the methodology; S.B. conducted the experiments;
719 S.B. and M.Y., analysed the data, S.B. wrote the manuscript, M.Y., H.C., C.B.-M., R.G.C., H.A.B., and
720 R.E.M.R., reviewed and edited the manuscript.

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724

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889 **Supplemental Information for:**

890 **A novel fluoro-electrochemical technique for classifying diverse marine**
891 **nanophytoplankton**

892

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895

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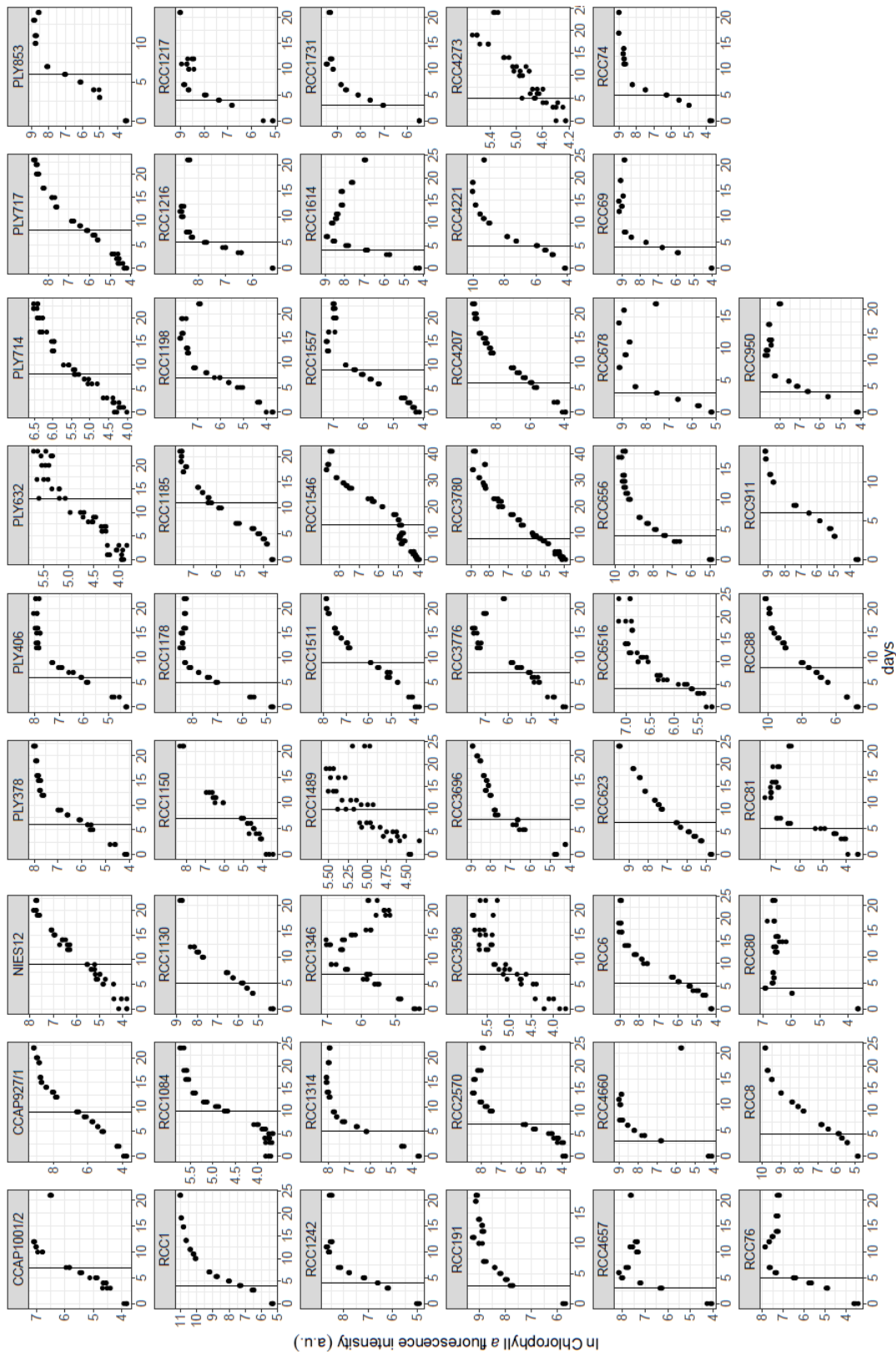
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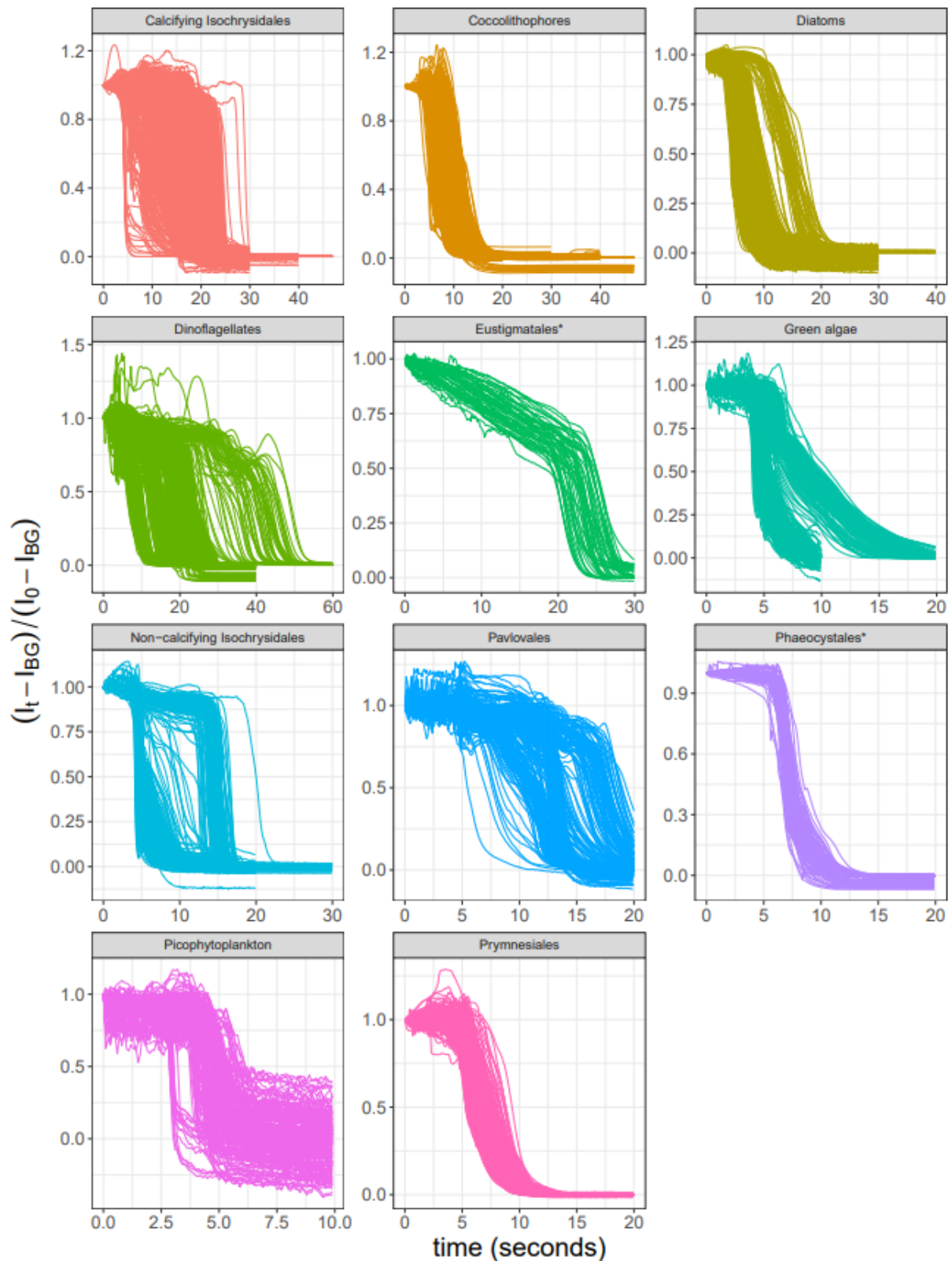
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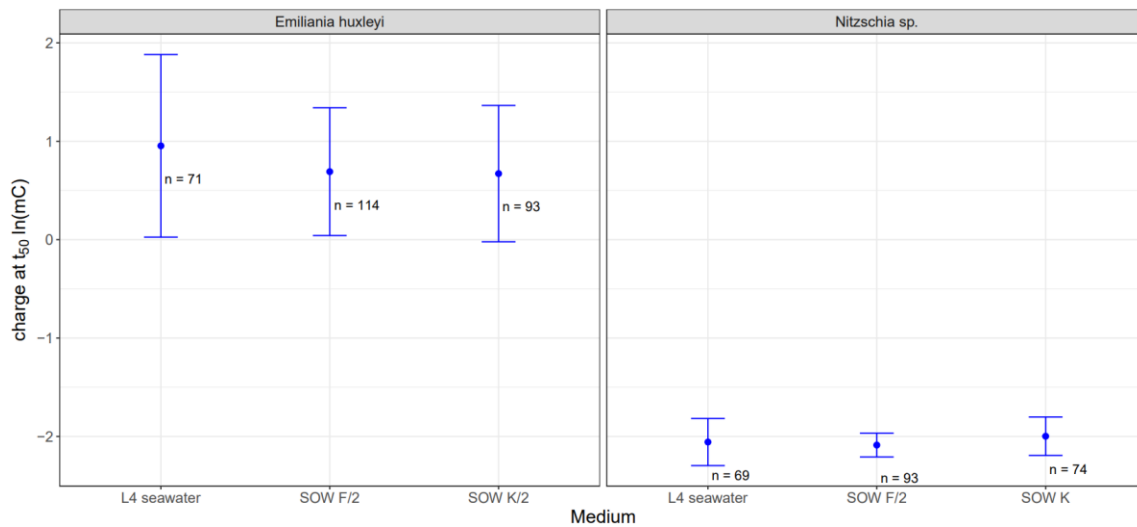
908 **Figure S1:** Culture growth curves for each of the strains listed in Table S1, as natural log transformed
 909 Chlorophyll *a* fluorescence intensity against days. The data points represent technical replicated
 910 measurements for each culture on each day. The vertical black line corresponds to the time-point when
 911 the electrochemical measurements were made, as described in the main text.

912



913

914 **Figure S2:** Normalised Chlorophyll *a* fluorescence transients for each of the relevant phytoplankton
 915 groups in this study. Each line represents a single cell measurement (4884 individual cell measurements
 916 in total, across the groups). Time ‘0 seconds’ is the time-point when the current ramping of the
 917 electrochemical experiments commenced. *denotes groups that were just represented by a single
 918 species.

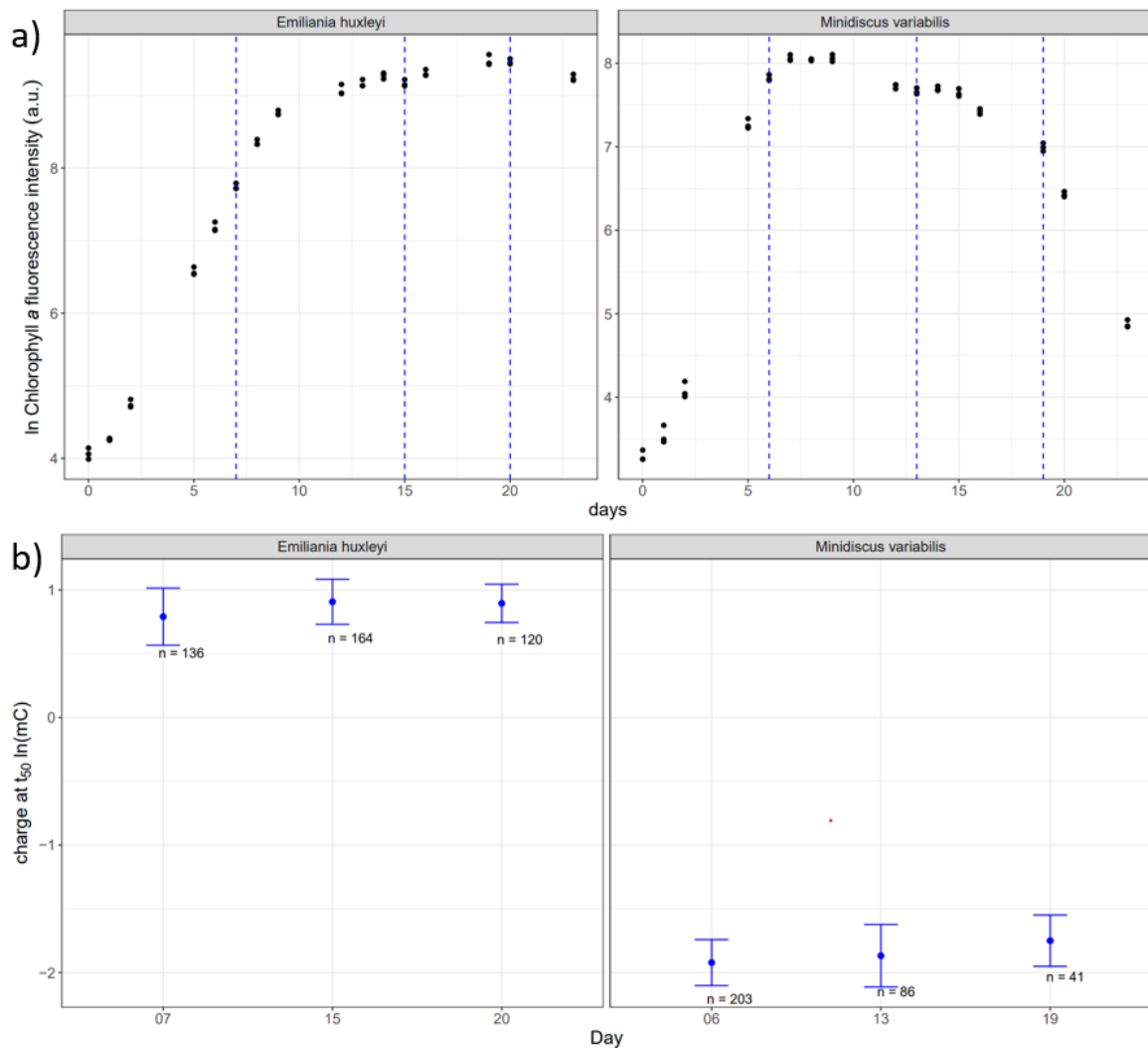


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921 **Figure S3:** Mean and standard deviation of natural log charge at t_{50} (in mC) measured for each of the
 922 strains when using different mediums as electrolyte. In the case of the calcifying isochrysidale
 923 *Emiliana huxleyi* (RCC1216), SOW K/2 (synthetic ocean water enriched with K/2 medium stocks) was
 924 the original growth medium. For the diatom *Nitzschia sp.* (RCC80), SOW K (synthetic ocean water
 925 enriched with K medium stocks) was the original growth medium. For each of the above, 15mL of
 926 culture in exponential growth phase was centrifuged at 1000 r.p.m for 10 minutes, supernatant (original
 927 growth medium) was removed, and pelleted cells were resuspended in either L4 seawater or SOW F/2
 928 (synthetic ocean water enriched with F/2 medium stocks). L4 seawater is natural seawater (collected
 929 at 10m depth from the L4 station, Western Channel Observatory, English Channel), which had been
 930 aged and refrigerated in the dark for > 1 year and 0.22 μ m filtered prior to making measurements.
 931 Numbers alongside data points represent the number of cells measured with each medium. Considering
 932 the deviation in the values, there is a negligible effect of each medium (or electrolyte) on the
 933 susceptibility of the strains. It is also clear that the variance for each medium response is within the
 934 broader range of values obtained for the strains' respective classified groups in the 'susceptibility library'
 935 (see Fig. 3 and Fig. S5).

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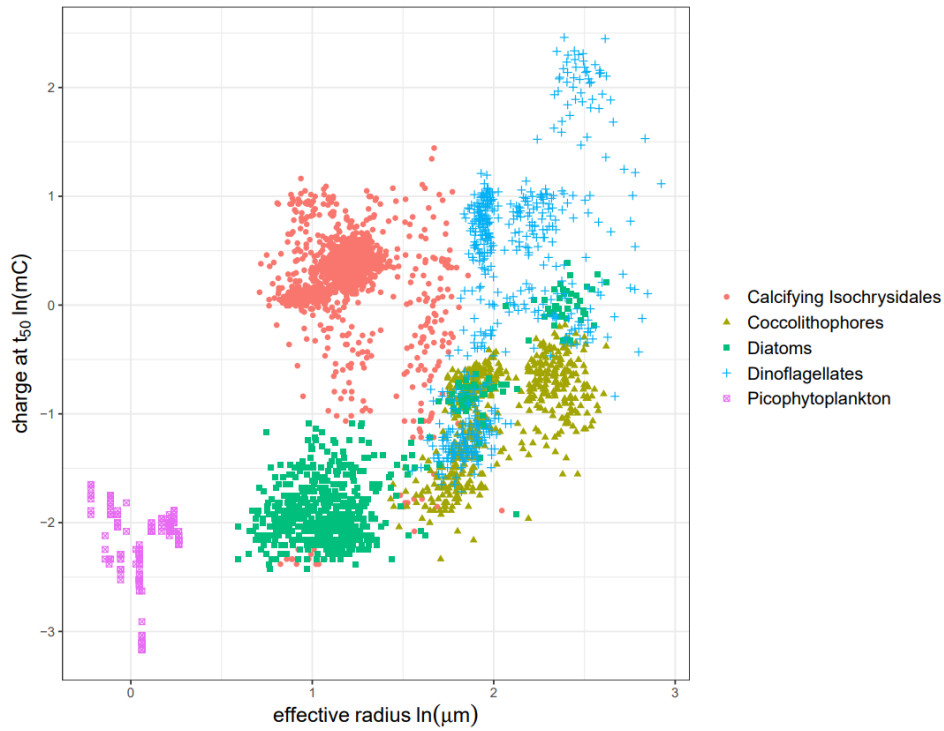
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939 **Figure S4:** (a) Culture growth curves for two strains: the calcifying isochrysidale *Emiliana huxleyi*
 940 (RCC 1731) and the diatom *Minidiscus variabilis* (RCC 4657), as natural log transformed Chlorophyll
 941 *a* fluorescence intensity against days. The data points represent technical replicated measurements for
 942 each culture on each day. The vertical dashed blue lines correspond to the time-points when the
 943 electrochemical measurements were made (b) Mean and standard deviation of natural log charge at t_{50}
 944 (in mC) measured for each of the strains, at each of the labelled time-points in (a). Numbers alongside
 945 data points represent the number of cells measured. As can be seen, there is a negligible effect of
 946 population growth stage on the susceptibility of the strains. Furthermore, it is clear that the variance of
 947 data from each point of the growth curve is within the range of values obtained for the strains' respective
 948 classified groups in the 'susceptibility library', where all measurements were made in the exponential
 949 growth phase (see Fig. 3 and Fig. S5).

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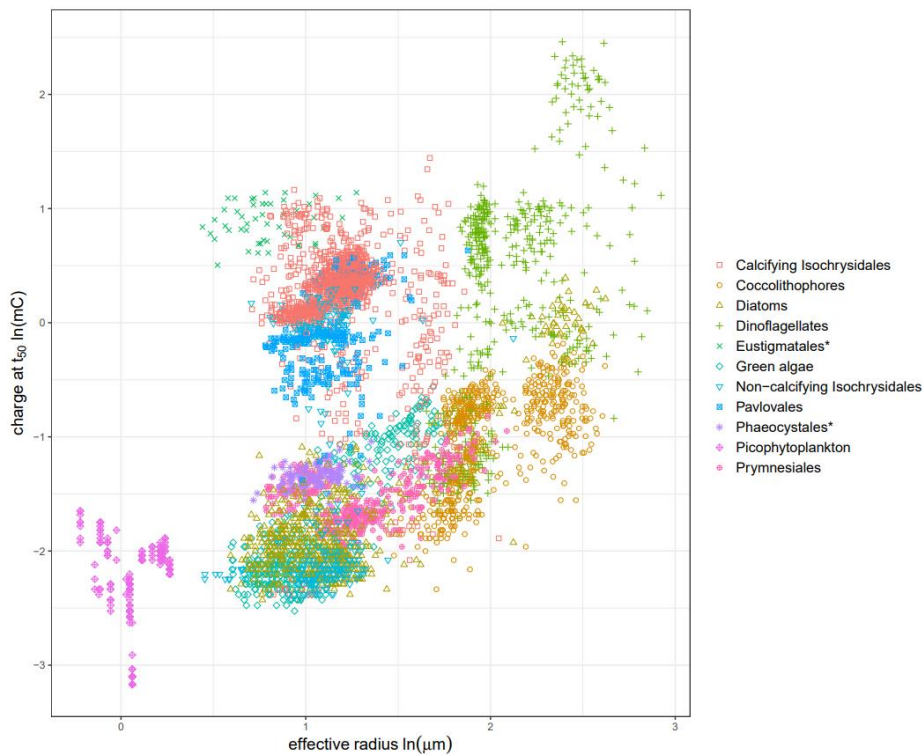
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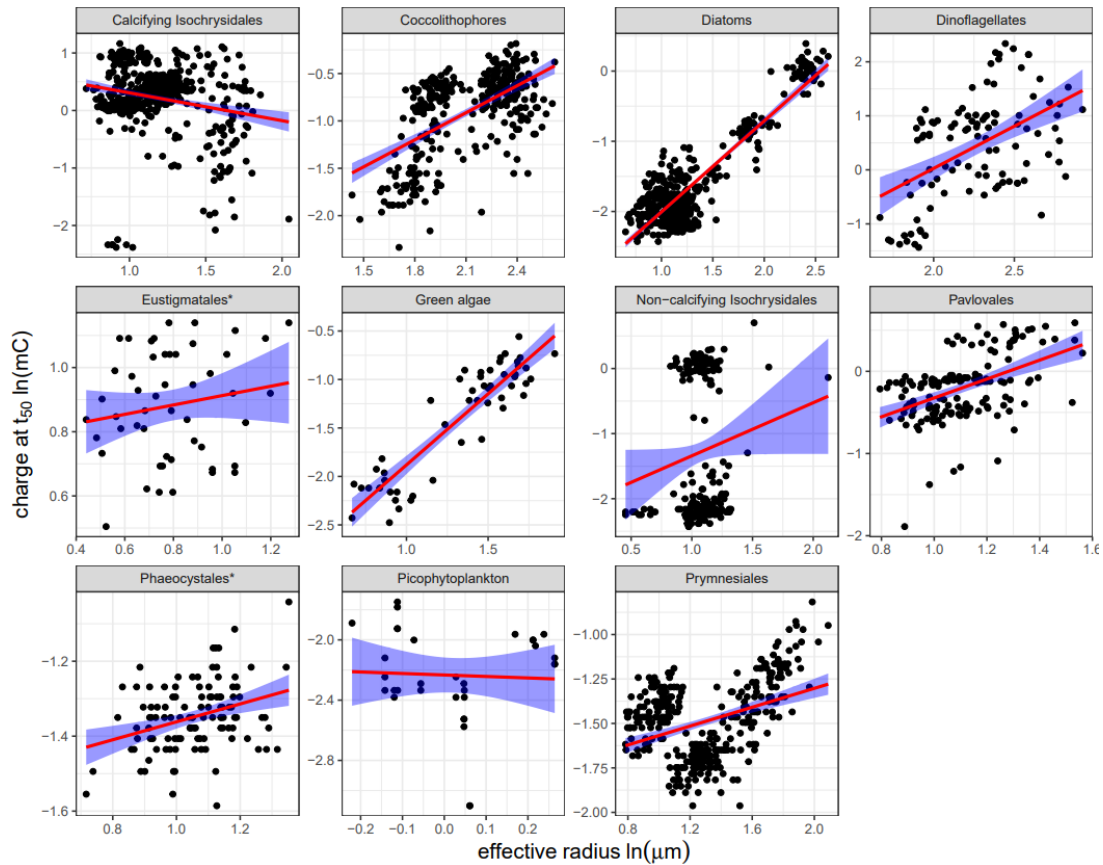
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954 **Figure S5:** A scatterplot of natural log charge at t_{50} (mC) against natural log effective cell radius (μm)
 955 for the ‘key functional groups’ of cells measured in this study, using the unbalanced dataset ($n = 3880$).



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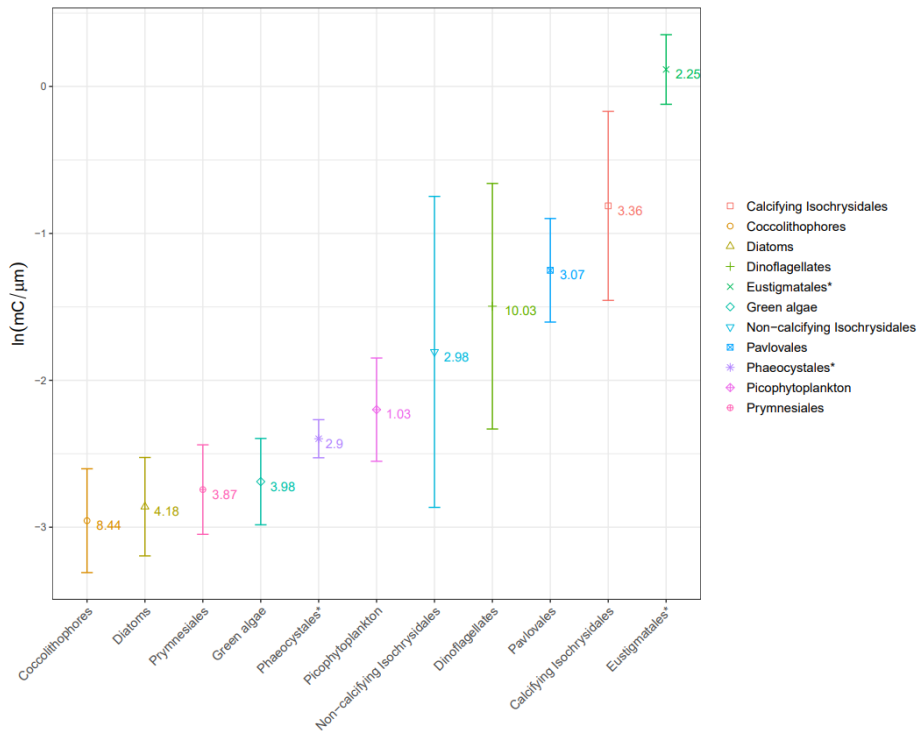
957 **Figure S6:** A scatterplot of natural log charge at t_{50} (mC) against natural log effective cell radius (μm)
 958 for all groups of cells measured in this study, with the unbalanced dataset ($n = 4884$, see Table S3 for
 959 a summary of the data presented here). *denotes groups that were just represented by a single species.



960

961 **Figure S7:** Scatterplots of natural log charge at t_{50} (mC) against natural log effective cell radius (μm)
 962 of cells measured for each phytoplankton group in this study, following the balancing of strain
 963 representation per group ($n = 2277$). The red line indicates the overall slope of the allometric
 964 relationship modelled for each level of grouping, and the blue shading is a visual representation of the
 965 95% confidence of this fit. See Table S11 for the coefficients of each individual model fit, and respective
 966 p values. *denotes groups that were just represented by a single species.

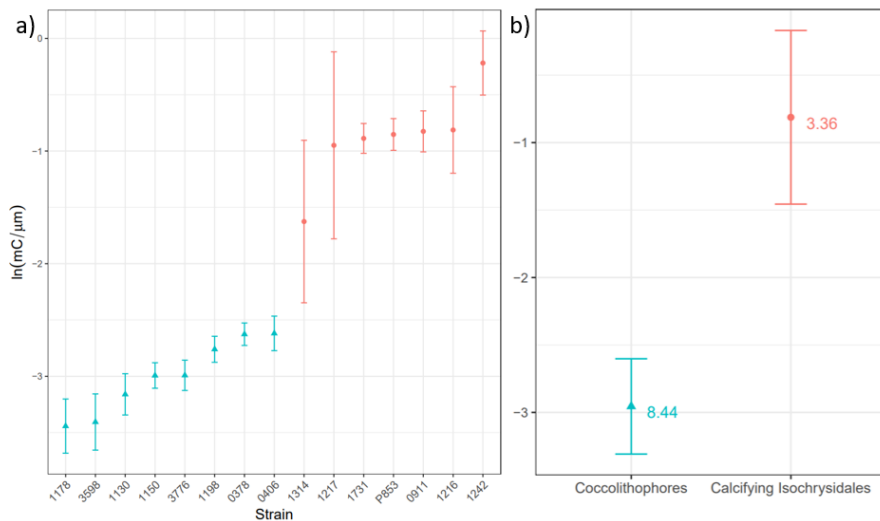
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969 **Figure S8:** Comparison of per group natural log transformed mean of size normalised charge at t_{50} .
 970 Data points represent the mean value for each group and the error bars are for standard deviation,
 971 following balancing the representation of the individual strains within each group. The labelled numbers
 972 alongside the data points represent the mean effective radius for each of the groups. *denotes groups
 973 that were just represented by a single species.

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976 **Figure S9: (a)** Comparison of natural log size normalised charge at t_{50} across the different strains of
 977 “Coccolithophore” (blue) and “Calcifying Isochrysidales” (red) **(b)** Comparison of natural log size
 978 normalised mean charge at t_{50} the group level for “Coccolithophores” and “Calcifying Isochrysidales”.
 979 Data points represent the mean values and the error bars are for standard deviation. The labelled
 980 numbers alongside the data points represent the mean effective radius. These plots clearly demonstrate
 981 that despite their much smaller size, “Calcifying Isochrysidales” are significantly more resilient in terms
 982 of their chlorophyll *a* “switch-off”.

983 **Table S1:** A list of all the 52 strains used, including their grouping (in this study), their culture collection
984 strain numbers, and their respective growth medium enrichment (see Table S2). Strain prefixes relate
985 to the respective culture collections where the strain was obtained (RCC = Roscoff Culture Collection
986 (Roscoff, France), CCAP = Culture Collection of Algae and Protozoa (Oban, UK), and PLY = The
987 Marine Biological Association (Plymouth, UK)

Group	Species detail	Strain	Synthetic seawater-based growth medium
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , non-calcified haploid	RCC 1217	K/2
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A, light -moderately calcified	RCC 911	K/2
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A, moderately calcified	RCC 1731	K/2
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A/R, over-calcified with bulky centre	PLY 853	K/2
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> morphotype A/R, over-calcified shields	RCC 1216	K/2
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , non-calcified diploid	RCC 1242	K/2
Calcifying Isochrysidales	<i>Gephyrocapsa oceanica</i>	RCC 1314	K/2
Coccolithophores	<i>Calcidiscus leptoporus</i> (1)	RCC 1130	K/2
Coccolithophores	<i>Calcidiscus leptoporus</i> (2)	RCC 1150	K/2
Coccolithophores	<i>Calyptosphaera sphaeroidea</i>	RCC 1178	K/2
Coccolithophores	<i>Chrysotila dentata</i> (1)	PLY 378	K/2
Coccolithophores	<i>Chrysotila dentata</i> (2)	PLY 406	K/2
Coccolithophores	<i>Coccolithus braarudii</i>	RCC 1198	K/2
Coccolithophores	<i>Coccolithus pelagicus</i>	RCC 3776	K/2
Coccolithophores	<i>Scyphosphaera apsteinii</i>	RCC 3598	L1
Diatoms	<i>Nitzschia closterium</i>	RCC 81	K
Diatoms	<i>Nitzschia</i> sp.	RCC 80	K
Diatoms	<i>Phaeodactylum tricoratum</i>	RCC 69	K
Diatoms	<i>Coscinodiscus</i> sp.	RCC 4273	K
Diatoms	<i>Halamphora coffeaeformis</i>	CCAP 1001/2	K
Diatoms	<i>Minidiscus comicus</i>	RCC 4660	K
Diatoms	<i>Minidiscus variabilis</i>	RCC 4657	K
Diatoms	<i>Skeletonema japonicum</i>	RCC 74	K
Diatoms	<i>Thalassiosira pseudonana</i>	RCC 950	K
Diatoms	<i>Thalassiosira weissflogii</i>	RCC 76	K
Dinoflagellates	<i>Amphidinium carterae</i>	RCC 88	F/2
Dinoflagellates	<i>Karenia papilionacea</i>	RCC 6516	K
Dinoflagellates	<i>Lepidodinium chlorophorum</i>	RCC 1489	F/2
Dinoflagellates	<i>Heterocapsa triquetra</i>	PLY 717	F/2
Dinoflagellates	<i>Prorocentrum micans</i>	NIES-12	F/2
Dinoflagellates	<i>Prorocentrum minimum</i>	PLY 714	F/2
Dinoflagellates	<i>Scrippsiella trochoidea</i>	PLY 632	F/2
Dinoflagellates	<i>Thoracosphaera heimii</i>	RCC 1511	K/2
Eustigmatales*	<i>Nannochloropsis granulata</i>	RCC 8	F/2
Green algae	<i>Chlamydomonas concordia</i>	RCC 1	F/2
Green algae	<i>Dunaliella tertiolecta</i>	RCC 6	K
Green algae	<i>Bigelowiella natans</i>	RCC 623	F/2
Non-calcifying Isochrysidales	<i>Isochrysis galbana</i>	CCAP 927/1	K
Non-calcifying Isochrysidales	<i>Isochrysis litoralis</i>	RCC 1346	F/2

Non-calcifying Isochrysidales	<i>Isochrysis</i> sp.	RCC 4207	K
Non-calcifying Isochrysidales	<i>Ruttnera</i> sp.	RCC 3696	F/2
Pavloales	<i>Diacronema vlkianum</i> (1)	RCC 1546	F/2
Pavloales	<i>Diacronema vlkianum</i> (2)	RCC 3780	F/2
Pavloales	<i>Pavlova granifera</i>	RCC 1557	F/2
Phaeocystales*	<i>Phaeocystis globosa</i>	RCC 678	K
Picophytoplankton	<i>Micromonas pusilla</i>	RCC 1614	K
Picophytoplankton	<i>Ostreococcus tauri</i>	RCC 4221	F/2
Picophytoplankton	<i>Synechococcus</i> sp. (1)	RCC 1084	F/2
Picophytoplankton	<i>Synechococcus</i> sp. (2)	RCC 2570	F/2
Prymnesiales	<i>Chrysochromulina camella</i>	RCC 1185	K/2
Prymnesiales	<i>Chrysochromulina</i> sp.	RCC 656	K
Prymnesiales	<i>Prymnesium parvum</i>	RCC 191	K

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991 **Table S2:** The final concentration of the components of Aquil Synthetic Ocean Water (Morel et al.
992 1979), used to make all of the culture growth mediums (See Table S1). For medium enrichment applied
993 to each see the following references: K (Keller et al. 1987), F/2 (Guillard and Ryther 1962), L1 (Guillard
994 and Hargraves 1993).

		Molar Concentration in final SOW based medias (mol dm ⁻³)
Synthetic Ocean Water (SOW) based on the Aquil medium recipe	NaCl	4.20 x 10 ⁻¹
	Na ₂ SO ₄	2.88 x 10 ⁻²
	KCl	9.39 x 10 ⁻³
	NaHCO ₃	2.38 x 10 ⁻³
	KBr	8.40 x 10 ⁻⁴
	H ₃ BO ₃	4.85 x 10 ⁻⁵
	NaF	7.15 x 10 ⁻⁵
	MgCl ₂ ·6H ₂ O	5.46 x 10 ⁻²
	CaCl ₂ ·2H ₂ O	1.05 x 10 ⁻²
	SrCl ₂ ·6H ₂ O	6.38 x 10 ⁻⁵

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Table S3. A summary of the predictor variables at the strain level, prior to balancing the dataset (i.e. variable n across the strains within each group).

Group	Species detail	Strain	n	t_{50} (s)	SD t_{50}	Charge at t_{50} (mC)	SD Charge at t_{50}	effective radius (μm)	SD effective radius
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , non-calcified haploid	RCC 1217	115	13.60	3.82	1.00	0.42	2.76	0.59
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A, light - moderately calcified	RCC 911	198	16.92	1.40	1.44	0.25	3.31	0.33
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A, moderately calcified	RCC 1731	311	16.94	0.93	1.44	0.17	3.44	0.32
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A/R, over-calcified with bulky centre	PLY 853	158	14.97	0.91	1.12	0.14	2.64	0.22
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> morphotype A/R, over-calcified shields	RCC 1216	277	16.74	2.81	1.44	0.42	3.35	0.23
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , non-calcified diploid	RCC 1242	80	21.21	2.44	2.28	0.49	2.90	0.52
Calcifying Isochrysidales	<i>Gephyrocapsa oceanica</i>	RCC 1314	132	13.65	4.98	1.05	0.78	5.12	0.52
Coccolithophores	<i>Calcidiscus leptoporus</i> (1)	RCC 1130	111	7.21	0.81	0.26	0.06	6.26	0.56
Coccolithophores	<i>Calcidiscus leptoporus</i> (2)	RCC 1150	55	10.08	0.68	0.51	0.07	10.10	0.78
Coccolithophores	<i>Calyptrosphaera sphaeroidea</i>	RCC 1178	93	6.17	0.71	0.19	0.05	5.87	0.81
Coccolithophores	<i>Chrysotila dentata</i> (1)	PLY 378	69	9.89	0.58	0.49	0.06	6.79	0.54
Coccolithophores	<i>Chrysotila dentata</i> (2)	PLY 406	106	9.83	0.58	0.48	0.06	6.60	0.62
Coccolithophores	<i>Coccolithus braarudii</i>	RCC 1198	52	11.29	0.76	0.64	0.08	10.20	0.87
Coccolithophores	<i>Coccolithus pelagicus</i>	RCC 3776	40	10.15	0.78	0.52	0.08	10.33	0.97
Coccolithophores	<i>Scyphosphaera apsteinii</i>	RCC 3598	67	8.58	0.99	0.37	0.08	11.14	1.24
Diatoms	<i>Nitzschia closterium</i>	RCC 81	59	5.28	0.78	0.14	0.05	3.22	0.34
Diatoms	<i>Nitzschia sp.</i>	RCC 80	79	5.27	0.33	0.14	0.02	3.24	0.27
Diatoms	<i>Phaeodactylum tricornutum</i>	RCC 69	95	4.77	0.27	0.11	0.01	3.28	0.34
Diatoms	<i>Coscinodiscus sp.</i>	RCC 4273	40	14.23	1.15	1.02	0.17	11.03	1.15
Diatoms	<i>Halamphora coffeaeformis</i>	CCAP 1001/2	39	6.05	0.59	0.18	0.04	3.58	0.73
Diatoms	<i>Minidiscus comicus</i>	RCC 4660	95	5.28	0.68	0.14	0.04	2.51	0.35
Diatoms	<i>Minidiscus variabilis</i>	RCC 4657	126	5.49	0.44	0.15	0.02	2.56	0.73
Diatoms	<i>Skeletonema japonicum</i>	RCC 74	69	6.67	0.77	0.23	0.05	2.94	0.39
Diatoms	<i>Thalassiosira pseudonana</i>	RCC 950	60	5.36	0.59	0.15	0.03	2.94	0.32
Diatoms	<i>Thalassiosira weissflogii</i>	RCC 76	49	9.15	0.71	0.42	0.06	6.52	0.73
Dinoflagellates	<i>Amphidinium carterae</i>	RCC 88	157	7.65	0.80	0.30	0.06	6.31	0.61
Dinoflagellates	<i>Karenia papilionacea</i>	RCC 6516	35	12.81	1.17	0.83	0.15	12.56	1.92
Dinoflagellates	<i>Lepidodinium chlorophorum</i>	RCC 1489	34	18.11	3.74	1.71	0.70	10.20	1.32
Dinoflagellates	<i>Heterocapsa triquetra</i>	PLY 717	62	21.21	1.82	2.27	0.38	9.29	0.80
Dinoflagellates	<i>Prorocentrum micans</i>	NIES-12	13	23.07	3.98	2.73	0.93	15.05	1.81
Dinoflagellates	<i>Prorocentrum minimum</i>	PLY 714	51	12.69	1.87	0.82	0.23	7.63	1.47
Dinoflagellates	<i>Scrippsiella trochoidea</i>	PLY 632	51	38.90	5.05	7.69	1.89	11.91	1.10
Dinoflagellates	<i>Thoracosphaera heimii</i>	RCC 1511	138	20.45	2.57	2.12	0.51	6.99	0.31

Eustigmatales*	<i>Nannochloropsis granulata</i>	RCC 8	50	22.06	1.85	2.45	0.41	2.25	0.46
Green algae	<i>Chlamydomonas concordia</i>	RCC 1	107	8.38	0.90	0.35	0.08	4.32	0.66
Green algae	<i>Dunaliella tertiolecta</i>	RCC 6	17	8.85	0.84	0.39	0.08	5.24	0.62
Green algae	<i>Bigelowiella natans</i>	RCC 623	269	4.84	0.43	0.12	0.02	2.41	0.33
Non-calcifying Isochrysidales	<i>Isochrysis galbana</i>	CCAP 927/1	96	9.27	4.34	0.52	0.42	3.05	0.36
Non-calcifying Isochrysidales	<i>Isochrysis litoralis</i>	RCC 1346	75	5.10	1.76	0.15	0.17	2.68	0.51
Non-calcifying Isochrysidales	<i>Isochrysis</i> sp.	RCC 4207	48	14.20	2.94	1.05	0.33	3.13	0.90
Non-calcifying Isochrysidales	<i>Ruttnera</i> sp.	RCC 3696	70	5.30	2.23	0.16	0.21	3.04	0.37
Pavlovales	<i>Diacronema vlkianum</i> (1)	RCC 1546	103	13.22	0.32	0.87	0.04	2.82	0.39
Pavlovales	<i>Diacronema vlkianum</i> (2)	RCC 3780	49	10.90	1.11	0.60	0.10	2.81	0.33
Pavlovales	<i>Pavlova granifera</i>	RCC 1557	117	15.03	2.99	1.17	0.42	3.55	0.52
Phaeocystales*	<i>Phaeocystis globosa</i>	RCC 678	110	7.22	0.33	0.26	0.02	2.90	0.37
Picophytoplankton	<i>Micromonas pusilla</i>	RCC 1614	121	5.08	0.18	0.13	0.01	1.23	0.06
Picophytoplankton	<i>Ostreococcus tauri</i>	RCC 4221	93	4.01	0.53	0.08	0.02	1.05	0.01
Picophytoplankton	<i>Synechococcus</i> sp. (1)	RCC 1084	42	5.27	0.63	0.14	0.03	0.90	0.05
Picophytoplankton	<i>Synechococcus</i> sp. (2)	RCC 2570	8	4.49	0.20	0.10	0.01	0.92	0.07
Prymnesiales	<i>Chrysochromulina camella</i>	RCC 1185	114	7.31	0.74	0.27	0.05	5.41	0.79
Prymnesiales	<i>Chrysochromulina</i> sp.	RCC 656	107	6.96	0.39	0.24	0.03	2.66	0.28
Prymnesiales	<i>Prymnesium parvum</i>	RCC 191	172	6.00	0.29	0.18	0.02	3.58	0.39
TOTAL			4884						

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Table S4. A summary of the predictor variables at the strain level, with the balanced dataset (i.e. standardised *n* across the strains within each group, see Methods).

Group	Species detail	Strain	<i>n</i>	<i>t</i> ₅₀ (s)	SD <i>t</i> ₅₀	Charge at <i>t</i> ₅₀ (mC)	SD Charge at <i>t</i> ₅₀	effective radius (µm)	SD effective radius
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , non-calcified haploid	RCC 1217	80	13.89	3.60	1.03	0.40	2.77	0.68
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A, light - moderately calcified	RCC 911	80	16.85	1.37	1.43	0.24	3.29	0.32
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A, moderately calcified	RCC 1731	80	16.81	0.91	1.42	0.16	3.47	0.31
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , morphotype A/R, over-calcified with bulky centre	PLY 853	80	14.91	0.85	1.12	0.13	2.64	0.22
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> morphotype A/R, over-calcified shields	RCC 1216	80	17.04	2.65	1.49	0.39	3.37	0.24
Calcifying Isochrysidales	<i>Emiliana huxleyi</i> , non-calcified diploid	RCC 1242	80	21.21	2.44	2.28	0.49	2.90	0.52
Calcifying Isochrysidales	<i>Gephyrocapsa oceanica</i>	RCC 1314	80	13.30	4.77	1.00	0.71	5.11	0.55
Coccolithophores	<i>Calcidiscus leptoporus</i> (1)	RCC 1130	40	7.26	0.81	0.27	0.06	6.25	0.50
Coccolithophores	<i>Calcidiscus leptoporus</i> (2)	RCC 1150	40	10.06	0.58	0.51	0.06	10.14	0.70
Coccolithophores	<i>Calyptrosphaera sphaeroidea</i>	RCC 1178	40	6.08	0.84	0.19	0.06	5.91	0.82
Coccolithophores	<i>Chrysotila dentata</i> (1)	PLY 378	40	9.95	0.57	0.50	0.06	6.88	0.57
Coccolithophores	<i>Chrysotila dentata</i> (2)	PLY 406	40	9.85	0.60	0.49	0.06	6.74	0.79
Coccolithophores	<i>Coccolithus braarudii</i>	RCC 1198	40	11.31	0.77	0.64	0.09	10.15	0.84
Coccolithophores	<i>Coccolithus pelagicus</i>	RCC 3776	40	10.15	0.78	0.52	0.08	10.33	0.97
Coccolithophores	<i>Scyphosphaera apsteinii</i>	RCC 3598	40	8.52	1.15	0.37	0.10	11.13	1.32
Diatoms	<i>Nitzschia closterium</i>	RCC 81	39	5.17	0.66	0.14	0.04	3.24	0.31
Diatoms	<i>Nitzschia sp.</i>	RCC 80	39	5.29	0.34	0.14	0.02	3.27	0.26
Diatoms	<i>Phaeodactylum tricornutum</i>	RCC 69	39	4.76	0.28	0.11	0.01	3.30	0.40
Diatoms	<i>Coscinodiscus sp.</i>	RCC 4273	39	14.26	1.15	1.02	0.16	11.04	1.16
Diatoms	<i>Halamphora coffeaeformis</i>	CCAP 1001/2	39	6.05	0.59	0.18	0.04	3.58	0.73
Diatoms	<i>Minidiscus comicus</i>	RCC 4660	39	5.19	0.68	0.14	0.04	2.49	0.36
Diatoms	<i>Minidiscus variabilis</i>	RCC 4657	39	5.47	0.42	0.15	0.02	2.46	0.25
Diatoms	<i>Skeletonema japonicum</i>	RCC 74	39	6.62	0.81	0.22	0.06	2.93	0.36
Diatoms	<i>Thalassiosira pseudonana</i>	RCC 950	39	5.35	0.59	0.14	0.03	2.96	0.35
Diatoms	<i>Thalassiosira weissflogii</i>	RCC 76	39	9.13	0.72	0.42	0.06	6.51	0.70
Dinoflagellates	<i>Amphidinium carterae</i>	RCC 88	13	7.72	0.67	0.30	0.05	6.31	0.54
Dinoflagellates	<i>Karenia papilionacea</i>	RCC 6516	13	12.64	1.46	0.81	0.18	12.44	2.12
Dinoflagellates	<i>Lepidodinium chlorophorum</i>	RCC 1489	13	19.71	3.86	2.01	0.74	10.01	1.08
Dinoflagellates	<i>Heterocapsa triquetra</i>	PLY 717	13	21.78	1.57	2.38	0.34	9.58	0.79
Dinoflagellates	<i>Prorocentrum micans</i>	NIES-12	13	23.07	3.98	2.73	0.93	15.05	1.81
Dinoflagellates	<i>Prorocentrum minimum</i>	PLY 714	13	13.06	1.68	0.87	0.21	8.14	1.77
Dinoflagellates	<i>Scrippsiella trochoidea</i>	PLY 632	13	38.01	5.03	7.34	1.88	11.63	1.42
Dinoflagellates	<i>Thoracosphaera heimii</i>	RCC 1511	13	20.36	2.98	2.11	0.55	7.06	0.29
Eustigmatales*	<i>Nannochloropsis granulata</i>	RCC 8	50	22.06	1.85	2.45	0.41	2.25	0.46
Green algae	<i>Chlamydomonas concordia</i>	RCC 1	17	8.12	1.01	0.33	0.08	4.28	0.58
Green algae	<i>Dunaliella tertiolecta</i>	RCC 6	17	8.85	0.84	0.39	0.08	5.24	0.62
Green algae	<i>Bigeloviella natans</i>	RCC 623	17	4.85	0.41	0.12	0.02	2.43	0.32

Non-calcifying Isochrysidales	<i>Isochrysis galbana</i>	CCAP 927/1	48	10.09	4.19	0.60	0.41	3.04	0.38
Non-calcifying Isochrysidales	<i>Isochrysis litoralis</i>	RCC 1346	48	4.87	0.78	0.12	0.05	2.71	0.54
Non-calcifying Isochrysidales	<i>Isochrysis</i> sp.	RCC 4207	48	14.20	2.94	1.05	0.33	3.13	0.90
Non-calcifying Isochrysidales	<i>Ruttnera</i> sp.	RCC 3696	48	5.36	2.36	0.17	0.22	3.03	0.36
Pavloales	<i>Diacronema vlkianum</i> (1)	RCC 1546	49	13.29	0.32	0.88	0.04	2.84	0.37
Pavloales	<i>Diacronema vlkianum</i> (2)	RCC 3780	49	10.90	1.11	0.60	0.10	2.81	0.33
Pavloales	<i>Pavlova granifera</i>	RCC 1557	49	14.96	3.15	1.17	0.44	3.56	0.47
Phaeocystales*	<i>Phaeocystis globosa</i>	RCC 678	110	7.22	0.33	0.26	0.02	2.90	0.37
Picophytoplankton	<i>Micromonas pusilla</i>	RCC 1614	8	5.08	0.21	0.13	0.01	1.26	0.04
Picophytoplankton	<i>Ostreococcus tauri</i>	RCC 4221	8	3.89	0.59	0.08	0.02	1.05	0.01
Picophytoplankton	<i>Synechococcus</i> sp. (1)	RCC 1084	8	5.26	0.55	0.14	0.03	0.90	0.05
Picophytoplankton	<i>Synechococcus</i> sp. (2)	RCC 2570	8	4.49	0.20	0.10	0.01	0.92	0.07
Prymnesiales	<i>Chrysochromulina camella</i>	RCC 1185	107	7.32	0.74	0.27	0.05	5.40	0.80
Prymnesiales	<i>Chrysochromulina</i> sp.	RCC 656	107	6.96	0.39	0.24	0.03	2.66	0.28
Prymnesiales	<i>Prymnesium parvum</i>	RCC 191	107	6.02	0.26	0.18	0.02	3.56	0.36
TOTAL			2277						

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1024 **Table S5.** A summary of the predictor variables at the group level, with the balanced dataset (i.e.
1025 standardised n across the strains within each group, see Methods). *denotes groups that were just
1026 represented by a single species.

Group	n	t_{50} (s)	SD t_{50}	Charge at t_{50} (mC)	SD Charge at t_{50}	effective radius (μm)	SD effective radius
Calcifying Isochrysidales	560	16.29	3.66	1.39	0.57	3.36	0.89
Coccolithophores	320	9.15	1.80	0.43	0.16	8.44	2.20
Diatoms	390	6.73	2.86	0.27	0.27	4.18	2.60
Dinoflagellates	104	19.54	9.10	2.32	2.22	10.03	3.05
Eustigmatales*	50	22.06	1.85	2.45	0.41	2.25	0.46
Green algae	51	7.27	1.92	0.28	0.14	3.98	1.29
Non-calcifying Isochrysidales	192	8.63	4.75	0.48	0.47	2.98	0.60
Pavloales	147	13.05	2.55	0.88	0.35	3.07	0.52
Phaeocystales*	110	7.22	0.33	0.26	0.02	2.90	0.37
Picophytoplankton	32	4.68	0.68	0.11	0.03	1.03	0.15
Prymnesiales	321	6.76	0.74	0.23	0.05	3.87	1.26
TOTAL	2277						

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1033 **Table S6.** Results from training the random forest using a random subset of 80% of the balanced data
 1034 for the key groups only ($n = 1406$). The green highlighted grid squares indicate the number of successful
 1035 categorisations for each group within the training.

		TRUE (Training)				
		Calcifying Isochrysidales	Coccolithophores	Diatoms	Dinoflagellates	Picophytoplankton
PREDICTED (Training)	Calcifying Isochrysidales	423	1	9	1	0
	Coccolithophores	4	227	24	17	0
	Diatoms	7	10	262	6	0
	Dinoflagellates	0	6	7	56	0
	Picophytoplankton	0	0	0	0	26
	RECALL	0.97	0.93	0.87	0.7	1
PRECISION	0.97	0.83	0.92	0.81	1	
F1	0.97	0.88	0.89	0.75	1	
					Overall accuracy	0.92

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1037 **Table S7.** Results from testing the trained random forest using a subset of 20% of the balanced data
 1038 for the key groups only ($n = 1406$). The green highlighted grid squares indicate the number of successful
 1039 categorisations for each group within the training.

		TRUE (Testing)				
		Calcifying Isochrysidales	Coccolithophores	Diatoms	Dinoflagellates	Picophytoplankton
PREDICTED (Testing)	Calcifying Isochrysidales	125	1	4	1	0
	Coccolithophores	0	67	3	4	0
	Diatoms	0	7	79	2	0
	Dinoflagellates	1	1	2	17	0
	Picophytoplankton	0	0	0	0	6
	RECALL	0.99	0.88	0.9	0.71	1
PRECISION	0.95	0.91	0.9	0.81	1	
F1	0.97	0.89	0.9	0.76	1	
					Overall accuracy	0.92

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1050 **Table S8.** Results from training the random forest using a random subset of 80% of the balanced data
 1051 for all groups ($n = 2277$). The green highlighted grid squares indicate the number of successful
 1052 categorisations for each group within the training.

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		TRUE (Training)										
		Calcifying Isochrysidales	Coccolithophores	Diatoms	Dinoflagellates	Eustigmatales*	Green algae	Non-calcifying Isochrysidales	Pavlovaes	Phaeocystales*	Picophytoplankton	Prymnesiales
PREDICTED (Training)	Calcifying Isochrysidales	372	0	5	0	9	7	36	32	1	0	0
	Coccolithophores	4	212	22	13	0	1	0	0	0	0	13
	Diatoms	3	10	176	8	0	13	49	2	5	0	31
	Dinoflagellates	0	5	7	54	0	0	1	0	0	0	0
	Eustigmatales*	8	0	0	0	30	0	0	0	0	0	0
	Green algae	5	1	6	1	0	11	0	0	1	0	3
	Non-calcifying Isochrysidales	19	0	29	0	0	2	42	3	2	0	0
	Pavlovaes	18	0	1	0	0	0	12	84	0	0	2
	Phaeocystales*	0	0	10	0	0	0	2	1	42	0	32
	Picophytoplankton	0	0	0	0	0	0	0	0	0	29	0
	Prymnesiales	5	16	46	4	0	8	6	2	37	0	182
	RECALL	0.86	0.87	0.58	0.68	0.77	0.26	0.28	0.68	0.48	1	0.69
PRECISION	0.81	0.8	0.59	0.81	0.79	0.39	0.43	0.72	0.48	1	0.59	
F1	0.83	0.83	0.59	0.73	0.78	0.31	0.34	0.7	0.48	1	0.64	
Overall accuracy											0.69	

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1055 **Table S9.** Results from testing the trained random forest using a subset of 20% of the balanced data
 1056 for all groups ($n = 2277$). The green highlighted grid squares indicate the number of successful
 1057 categorisations for each group within the training.

		TRUE (Testing)										
		Calcifying Isochrysidales	Coccolithophores	Diatoms	Dinoflagellates	Eustigmatales*	Green algae	Non-calcifying Isochrysidales	Pavlovaes	Phaeocystales*	Picophytoplankton	Prymnesiales
PREDICTED (Testing)	Calcifying Isochrysidales	110	1	2	0	5	2	11	7	0	0	0
	Coccolithophores	0	65	3	4	0	0	0	0	0	0	3
	Diatoms	0	6	55	2	0	1	16	0	0	0	4
	Dinoflagellates	1	0	2	18	0	0	0	0	0	0	1
	Eustigmatales*	1	0	0	0	6	0	0	0	0	0	0
	Green algae	4	0	2	0	0	3	0	0	0	0	0
	Non-calcifying Isochrysidales	6	0	5	0	0	2	14	1	0	0	0
	Pavlovaes	4	0	0	0	0	0	2	15	0	0	0
	Phaeocystales*	0	0	4	0	0	1	0	0	14	0	7
	Picophytoplankton	0	0	0	0	0	0	0	0	0	3	0
	Prymnesiales	0	4	15	0	0	0	1	0	8	0	43
	RECALL	0.87	0.86	0.63	0.75	0.55	0.33	0.32	0.65	0.64	1.00	0.74
PRECISION	0.80	0.87	0.65	0.82	0.86	0.33	0.50	0.71	0.54	1.00	0.61	
F1	0.83	0.86	0.64	0.78	0.67	0.33	0.39	0.68	0.58	1.00	0.67	
Overall accuracy											0.71	

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1059 **Table S10.** Summary of the model comparison for fitting linear models to $\ln(\text{charge at } t_{50})$ against
 1060 $\ln(\text{effective radius})$, across the full balanced dataset ($n = 2277$), see Fig. 7 (Main text). The best model
 1061 (lowest AIC) is the mixed model, including the random effect of grouping on the slope and intercept of
 1062 the response. This indicates that there is an overall significant positive size scaling, but that the
 1063 relationship differs across the groups (see Fig. S7).

	npar	AIC	BIC	logLik	deviance	Chisq	Df	Pr(>Chisq)
lm_size_dep	3	6398.24	6415.43	-3196.12	6392.24			
lmer_size_dep2	6	3394.90	3429.29	-1691.45	3382.91	3009.33	3	<2.2 x 10⁻¹⁶

Models:
 lm_size_dep: $\ln_{mC} \sim \ln_{rad}$
 lmer_size_dep2: $\ln_{mC} \sim \ln_{rad} + (\ln_{rad} | \text{eco_group})$

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1065 **Table S11:** Linear model output for each of the group specific relationships (see Fig. S7). Bold
 1066 highlights indicate a significant allometric scaling, using p values for the slope coefficient only. ‘Lower’
 1067 and ‘Upper’ represent the 95% confidence intervals of the intercept and slope (\ln_{rad}) coefficients.
 1068 The overall size scaling is visualised in Fig. 7 (Main text). *denotes groups that were just represented
 1069 by a single species.

	(Intercept)	lower	upper	\ln_{rad}	lower	upper	p (slope only)
Calcifying Isochrysidales	0.79	0.58	1.00	-0.48	-0.66	-0.31	0.000000
Coccolithophores	-2.91	-3.35	-2.48	0.95	0.75	1.16	0.000000
Diatoms	-3.30	-3.45	-3.15	1.30	1.19	1.40	0.000000
Dinoflagellates	-3.10	-3.83	-2.36	1.56	1.24	1.88	0.000000
Eustigmatales*	0.77	0.19	1.35	0.15	-0.56	0.85	0.680259
Green algae	-3.35	-3.88	-2.81	1.47	1.07	1.86	0.000000
Non-calcifying Isochrysidales	-2.16	-2.58	-1.73	0.82	0.43	1.20	0.000040
Pavloales	-1.47	-2.03	-0.92	1.15	0.65	1.64	0.000005
Phaeocystales*	-1.60	-2.39	-0.82	0.24	-0.50	0.98	0.524655
Picophytoplankton	-2.23	-2.41	-2.06	-0.10	-1.31	1.11	0.872391
Prymnesiales	-1.83	-2.07	-1.60	0.27	0.09	0.44	0.002990
Overall (lmer model, see Table S10)	-1.83	-2.76	-0.91	0.72	0.30	1.13	NA

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1071 **Table S12.** A summary of the Pairwise Wilcoxon Rank Sum testing, to complement Fig. 8 (Main text).
 1072 Bold text indicates a significant difference ($p < 0.05$) in the $mC/\mu m$ values between the pairs.

Group1	Group2	p value (Bonferroni adjusted for nonparametric data)
Coccolithophores	Calcifying Isochrysidales	0.000000
Diatoms	Calcifying Isochrysidales	0.000000
Dinoflagellates	Calcifying Isochrysidales	0.000000
Eustigmatales*	Calcifying Isochrysidales	0.000000
Green algae	Calcifying Isochrysidales	0.000000
Non-calcifying Isochrysidales	Calcifying Isochrysidales	0.000000
Pavloales	Calcifying Isochrysidales	0.000000
Phaeocystales*	Calcifying Isochrysidales	0.000000
Picophytoplankton	Calcifying Isochrysidales	0.000000
Prymnesiales	Calcifying Isochrysidales	0.000000
Diatoms	Coccolithophores	0.388386

Dinoflagellates	Coccolithophores	0.000000
Eustigmatales*	Coccolithophores	0.000000
Green algae	Coccolithophores	0.000012
Non-calcifying Isochrysidales	Coccolithophores	0.009010
Pavloales	Coccolithophores	0.000000
Phaeocystales*	Coccolithophores	0.000000
Picophytoplankton	Coccolithophores	0.000000
Prymnesiales	Coccolithophores	0.000000
Dinoflagellates	Diatoms	0.000000
Eustigmatales*	Diatoms	0.000000
Green algae	Diatoms	0.005731
Non-calcifying Isochrysidales	Diatoms	1.000000
Pavloales	Diatoms	0.000000
Phaeocystales*	Diatoms	0.000000
Picophytoplankton	Diatoms	0.000000
Prymnesiales	Diatoms	0.000124
Eustigmatales*	Dinoflagellates	0.000000
Green algae	Dinoflagellates	0.000000
Non-calcifying Isochrysidales	Dinoflagellates	0.001037
Pavloales	Dinoflagellates	0.000007
Phaeocystales*	Dinoflagellates	0.000007
Picophytoplankton	Dinoflagellates	0.325385
Prymnesiales	Dinoflagellates	0.000000
Green algae	Eustigmatales*	0.000000
Non-calcifying Isochrysidales	Eustigmatales*	0.000000
Pavloales	Eustigmatales*	0.000000
Phaeocystales*	Eustigmatales*	0.000000
Picophytoplankton	Eustigmatales*	0.000000
Prymnesiales	Eustigmatales*	0.000000
Non-calcifying Isochrysidales	Green algae	1.000000
Pavloales	Green algae	0.000000
Phaeocystales*	Green algae	0.000000
Picophytoplankton	Green algae	0.000001
Prymnesiales	Green algae	1.000000
Pavloales	Non-calcifying Isochrysidales	0.000000
Phaeocystales*	Non-calcifying Isochrysidales	0.056749
Picophytoplankton	Non-calcifying Isochrysidales	1.000000
Prymnesiales	Non-calcifying Isochrysidales	1.000000
Phaeocystales*	Pavloales	0.000000
Picophytoplankton	Pavloales	0.000000
Prymnesiales	Pavloales	0.000000
Picophytoplankton	Phaeocystales*	0.010620
Prymnesiales	Phaeocystales*	0.000000
Prymnesiales	Picophytoplankton	0.000000
Overall Kruskal-Wallis for significant effect of group		<2.2 x 10 ⁻¹⁶

1074 **Table S13.** A summary of the Pairwise Wilcoxon Rank Sum testing, to complement Fig. 9b (Main
 1075 text), showing the difference in natural $\ln(\text{mC}/\mu\text{m})$ across the different *Emiliana huxleyi* strains in this
 1076 study. Bold text indicates a significant difference ($p < 0.05$) in the $\text{mC}/\mu\text{m}$ values between the pairs.
 1077 The non-calcified diploid strain of *E. huxleyi* is significantly different to all other strains, with a higher
 1078 average $\text{mC}/\mu\text{m}$.

Group 1	Group 2	<i>p</i> value (Bonferroni adjusted for nonparametric data)
Morphotype A, moderately calcified (1731)	Morphotype A, light to moderately calcified (911)	0.601538
Morphotype A/R, over-calcified shields (1216)	Morphotype A, light to moderately calcified (911)	0.051449
Morphotype A/R, over-calcified, bulky centre (PLY853)	Morphotype A, light to moderately calcified (911)	1.000000
Non-calcified, diploid (1242)	Morphotype A, light to moderately calcified (911)	0.000000
Non-calcified, Haploid (1217)	Morphotype A, light to moderately calcified (911)	0.581945
Morphotype A/R, over-calcified shields (1216)	Morphotype A, moderately calcified (1731)	0.000078
Morphotype A/R, over-calcified, bulky centre (PLY853)	Morphotype A, moderately calcified (1731)	0.526381
Non-calcified, diploid (1242)	Morphotype A, moderately calcified (1731)	0.000000
Non-calcified, Haploid (1217)	Morphotype A, moderately calcified (1731)	1.000000
Morphotype A/R, over-calcified, bulky centre (PLY853)	Morphotype A/R, over-calcified shields (1216)	0.000945
Non-calcified, diploid (1242)	Morphotype A/R, over-calcified shields (1216)	0.000000
Non-calcified, Haploid (1217)	Morphotype A/R, over-calcified shields (1216)	0.001045
Non-calcified, diploid (1242)	Morphotype A/R, over-calcified, bulky centre (PLY853)	0.000000
Non-calcified, Haploid (1217)	Morphotype A/R, over-calcified, bulky centre (PLY853)	1.000000
Non-calcified, Haploid (1217)	Non-calcified, diploid (1242)	0.000000
Overall Kruskal-Wallis for effect of <i>Emiliana huxleyi</i> strain type		<2.2 x 10⁻¹⁶

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