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Variable inter- and intra-species alkaline phosphatase activity within single cells of revived dinoflagellates

Mathias Girault¹, Raffaele Siano², Claire Labry², Marie Latimier², Cécile Jauzein², Thomas Beneyton¹, Lionel Buisson¹, Yolanda Del Amo³, Jean-Christophe Baret^{1,4}

1) CNRS, Univ. Bordeaux, CRPP, UMR 5031, 33600 Pessac, France.

2) Ifremer, DYNECO, F-29280 Plouzané, France.

3) Université de Bordeaux, UMR CNRS 5805 EPOC, Station Marine d'Arcachon, 33120 Arcachon, France.

4) Institut Universitaire de France, 75005 Paris.

Corresponding authors: raffaele.siano@ifremer.fr, jean-christophe.baret@u-bordeaux.fr

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Abstract:

Adaptation of cell populations to environmental changes is mediated by phenotypic variability at the single cell level. Enzyme activity is a key factor in cell phenotype and the expression of the alkaline phosphatase activity (APA) is a fundamental phytoplankton strategy for maintaining growth under phosphate-limited conditions. Our aim was to compare the APA among cells and species revived from sediments of the Bay of Brest (Brittany, France), corresponding to a pre-eutrophication period (1940's) and a beginning of a post-eutrophication period (1990's) during which phosphate concentrations have undergone substantial variations. Both toxic marine dinoflagellate *Alexandrium minutum* and the non-toxic dinoflagellate *Scrippsiella acuminata* were revived from ancient sediments. Using microfluidics, we measured the kinetics of APA at the single-cell level. Our results indicate that all *S. acuminata* strains had significantly higher APA than *A. minutum* strains. For both species, the APA in the 1990's decade was significantly lower than in the 1940's. For the first time, our results reveal both inter-and intra-specific variabilities of dinoflagellate APA and suggest that, at a half-century time-scale, two different species of dinoflagellate may have undergone similar adaptative evolution to face environmental changes and acquire ecological advantages.

Introduction

Microphytoplankton abundances, community structure and species ecological niches are commonly reported to change at ocean surfaces in response to modified environmental conditions [1]. These changes can be associated with the ability of the microorganisms to adapt their physiology in order to maintain their resilience or spread in the environment [2-4]. Physiological responses of phytoplankton to environmental changes are generally estimated by the analysis of relatively few biological parameters in a limited number of cultivated species and strains. For a single strain in standard culture and analytical conditions, these estimates integrate the combined responses of hundreds of thousands of cells within a phytoplankton culture. This approach is limited because it overlooks cellular adaptations at the single cell level, which are crucial for explaining changes in species distribution and phenology [5-9]. The single phytoplankton cell adaptation is a critical and a complex problem, especially given that several studies report that long-term acclimation and/or genetic selection can contribute to the maintenance of

population fitness under environmental change scenarios [10-11]. In addition, other studies suggest that both physiological adaptations and phenotypic variability of the cells can be insufficient for survival through environmental changes [12-13].

Physiological adaptations of phytoplankton species are classically studied by monitoring variations in biological parameters (*e.g.* growth rate, photosystem) under culture conditions reproducing environmental change scenarios [14]. Even though these approaches can demonstrate how phytoplankton respond to changes in temperature, nutrients and heavy metals, they remain limited to short-term incubations (*e.g.*, weeks or months) because phytoplankton experiments are cumbersome to be carried out for longer periods [14-17]. For studying long-term physiological variations within phytoplankton species, an innovative approach consists of comparing biological functions among strains of species revived from sediments of different ages. These strains are obtained from the germination of cysts that were stored in ancient sediments [18, 19]. Protected by a resistant biopolymer wall, the genomes and the physiological constants of cells are theoretically well preserved [20]. Consequently, these sediment archives become “time capsules” that contain populations from the past [21, 22]. The revived cells can be cultivated under different environmental conditions, including nutrient starvation scenarios, and physiological indicators of the biological function of interest (*e.g.* nutrient uptake) are monitored and compared among strains [20]. The differences in biological function indicators within strains of different ages can reflect adaptations of the corresponding populations from distinct time periods.

Phosphate concentration is known to shape the structure of marine planktonic communities [23]. A low concentration of phosphate is generally reported to limit cell abundances and species distributions in many marine or coastal ecosystems [24-26]. To survive phosphate depletion, microorganisms such as diatoms and dinoflagellates can activate a set of enzymes, such as the alkaline phosphatases (AP), to use dissolved organic phosphorus instead of phosphate [27, 28]. Induction of extracellular AP is linked to the physiological conditions of the cells and is used as an indicator of phosphate stress in phytoplankton [29]. Here we focus on extracellular AP activity (APA) because a part of the intracellular AP can be constitutive, and substrate used for the determination of enzyme activity can be in competition with acid phosphatases involved in cellular processes not linked to cellular P-stress status [27, 30, 31]. In general,

APA is determined using fluorescent substrates in bulk experiments [32, 33] and the fluorescence measurements are not species-specific. Although size fractionation through filtration separates free-bacteria from microphytoplankton, both living and dead cells are integrated in such enzymatic assays, which leads to an underestimation of the APA measured in the sample. Moreover, the presence of free AP, which is released by cell lyses, can overestimate APA in the sample. To limit these potential biases, microfluidics and image processing algorithms have been recently combined in order to measure APA at single phytoplankton cell levels [34].

In this study, we demonstrate a new microfluidic platform that micro-compartmentalizes micron-sized droplets of cells from suspension. By sorting the water-in-oil droplets containing dinoflagellates, the APA assay is performed at the single cell level and only counts cells of interest. To consider the potential buffer effect of the intracellular phosphorus (P) pool on the APA, we monitor the expression of APA during consecutive days, up to 7 days, and compare APA at the maximum expression. We used this new approach to analyse extracellular APA for single living phytoplankton cells and compare strains of different species and ages. We used two dinoflagellate species (i.e., *Alexandrium minutum* and *Scrippsiella acuminata*) revived from two distinct sediment layers of a core sampled in the coastal environment of the Bay of Brest (France). These two sampling layers corresponded to a pre-eutrophication period with low human P loadings (1940's) and a start of post-eutrophication period for P supplies (1990's). We explored the variability of APA among cells of a same strain (intra-clonal), among strains of the same species (intra-species), and between species (inter-species). Finally, we explored APA modification at a timescale of ~50 years, a period of important changes in agricultural activities on the coastal land of the Bay of Brest. From our results, we hypothesize a co-evolving adaptations scenario of two different dinoflagellate species to the environmental variations in terms of phosphorus availability occurring in the Bay of Brest during the 1940–1990 period.

Materials and methods

Phytoplankton culture establishment and sample preparation

By using a Kullenberg corer, a sediment core (KS06) was sampled on-board of the R/V *Thalia* on the April 25th, 2017 in the Bay of Brest (48°22'52.74"N–04°26'54.60"W Brittany, France). The core was

collected near the Brest harbor in a non-dredged area with a low sedimentation rate, at a water depth of 7.10 m, (Fig. 1). The sediment core (3.44 m long) was immediately extruded from the Plexiglas tube and sliced into 1-cm layers. Aliquots of sediments were used for dating as previously described [19, 21]. Materials for dinoflagellate cyst germination were stored in 50 mL Nalgene tubes at 4°C in the dark and without oxygen. Aliquots of the sediment-sieved fraction 25–100 µm were suspended in a culture K medium (EDMK/5; ref. 35) and cell germinations were monitored daily with light microscopy. Single germinated cells were quickly isolated in order to obtain monoclonal cultures and these were characterized genetically (LSU 18S rRNA gene sequencing) as described elsewhere [21]. The date of each culture was determined according to the sediments dating from which cells were revived as described previously [19, 21].

For this study, we selected six revived strains (i.e., AM-47-1, AM-92-1, AM-92-2, SC-47-1, SC-47-2, SC-92-1) corresponding to dinoflagellates *Alexandrium minutum* (Halim) and *Scrippsiella acuminata* (Ehrenberg) from two selected dates (1947±11 years and 1992±4 years). Dinoflagellate strains were acclimated for two weeks into sterile flasks (30 mL) with F/2 medium in artificial seawater at 17°C [36]. The F/2 medium was chosen for the presence of one single source of inorganic P (phosphate). This step enabled rapid growth before transfer into phosphate-deplete F/2 medium. The transfer into P-deplete medium was performed in order to obtain a cellular stress to phosphate limitation. In this study, day zero was considered as the day when cultures started to grow on the P-depleted F/2 medium (artificial seawater). The low concentrations of phosphate measured on day zero corresponded to the residual phosphate from the inoculum (Table 1). Cultures were exposed to a 12:12h day/night cycle at 80 µE m⁻² s⁻¹. Before sampling, the cultures were filtered by gravity on a 10 µm nucleopore filter (Whatman) and were gently washed with fresh P-deplete F/2 medium under laminar flow hood. This step was performed to concentrate cells in the samples, prevent bacteria contamination and ensure comparable nutrient concentrations during the APA assay on-a-chip.

Lab-on-a-chip device and experimental setup

The microfluidic chips were made of poly-(dimethylsiloxane) (PDMS, Sylgard 184) from SU8-3000 negative photoresist (MicroChem Corp) moulds produced using standard soft-lithography procedures

[37]. The surfaces of the microfluidic channels were treated using fluorosilane (Aquapel) in order to increase the hydrophobic properties of the chip. The microfluidic chip was placed on a microfluidic platform dedicated to APA measurement [34]. In this study, we added a controlled X/Y motorized platform, a microfluidic module to sort droplets containing single cells. We optimized hydrodynamic trapping by increasing the height of the channel from 30 μm to 60 μm (Supplementary materials Fig. 1). These three improvements allowed us to monitor a total of 100 droplets trapped in the incubation channel for each APA assay.

Alkaline phosphatase activity (APA) assay on a chip

A first series of water-in-oil droplets (500 pL) were generated by flow-focusing the culture stream containing phytoplankton with two streams of FC40 oil containing a surfactant (5% w/w, Fluosurf, Emulseo). The droplets of interest were sorted according to the morphology of the cell, as previously described [38, 39]. The droplets of interest slightly compressed in the channel were rerouted to the series of high height areas corresponding to the traps. By using the second droplet module on the chip, a train of droplets (80 pL) was generated by flow focussing a stream containing 100 μM of the ELF-97 phosphatase substrate (E6588, ThermoFisher, referred after ELF-P) with two streams of FC40 oil containing a surfactant (5% w/w). According to the concentrations of organic phosphate saturating the AP site of the dinoflagellate in the literature, we assumed that all the active AP sites were saturated by 100 μM of ELF-P. This is a prerequisite for a meaningful comparison of APA between cells [40]. Actually, the maximal velocity rate assumed to be measured is proportional to the quantity of AP. As a consequence, our APA measurements are indicative of the enzymatic equipment of cells. By under pressurizing the outlet, the small ELF-P droplets were rerouted to a series of small hydrodynamic traps and stored in contact with the droplets containing a single dinoflagellate cell. Then, the positions of droplets of interest were saved in the time-lapse sequence program (LabVIEW). A short width pulse of voltage (100 ms, 100 V) is applied to the electrodes located on each side of the collection channel in order to simultaneously fuse each droplet containing a phytoplankton with a droplet containing the ELF-P substrate. The fusion of the droplets initiates the APA assay and the images were automatically captured according to droplet locations. The APA was determined by monitoring the fluorescence signal

of droplets containing a single living cell. In this study, a living cell was defined as a cell swimming in the droplet when the APA assay started. The fluorescence signal was due to the hydrolysis of ELF-P and subsequent precipitation of phenol form of ELF alcohol into a fluorescent water-insoluble form ELF-A [41]. The measurement of the fluorescent product was made according to the method described previously [34]. Blank experiments were conducted to test the APA response of strains cultivated under P-replete conditions. No ELF-A fluorescence signals were detected in the blank experiments. The presence of natural pigments, bleaching of the ELF-A and bacterial contamination were also considered in order to decrease potential bias in the fluorescence measurement (Supplementary materials Figs. 2, 3).

Image processing systems

In addition to the droplet sorting system based on the morphology of cells, a second image processing algorithm was developed to automatically detect the droplets and measure the fluorescence intensity in each time-lapse image (Supplementary materials Fig. 4; refs. 37, 42). Because the fluorescent images were captured as a function of time, it was possible to precisely follow the variability of the fluorescence intensity of each droplet trapped in the collection channel.

Phosphate measurement in the medium

Samples for the determination of phosphate were carefully filtered through pre-combusted (12 h at 480°C) 25 mm Whatman GF/F. Pre-combustion of the filters is used to avoid any potential contamination. Phosphate was analyzed immediately after filtration on a spectrophotometer (Shimadzu UV 160) with a 5 cm optical path cell. The phosphate concentration was determined according to the method of Murphy and Riley [43], with the detection limit of 0.02 μM .

Statistical analysis

To perform a one-way analysis of variance (ANOVA) test, we first determined all APA labelling kinetics at a single cell level. Then, for each strain, we selected the experiment that had maximum APA

values. From these maximum values, we used ANOVA to test significant differences of APA among strains of different species and ages.

Results

Automated microfluidic platform for the APA assay

The APA has been determined on three strains of *Alexandrium minutum* and three strains of *Scrippsiella acuminata* (Fig. 1). The strains SC-47-1, SC-47-2 and AM-47-1 corresponded to the 1940's decade (1947 ± 11 years; blue symbols in this study). The strains AM-92-1, AM-92-2 and SC-92-1 corresponded to the 1990's decade (1992 ± 4 years; green symbols). By using the microfluidic platform, a total of 25,230 images were captured and analysed during the experiments. For each captured image, the fluorescence intensity of water-in-oil droplets containing single cells was determined as an index of APA. Six examples of the increase in fluorescence of single living dinoflagellate encapsulated in the droplets are shown in Fig. 2. The APA fluorescence curves show a typical enzyme labelling pattern with the maximum of the APA observed when the cell started to be labelled. The microphotographs showed the increase in fluorescent ELF-A product (green) on the surface of the cells as a function of time. The results showed that all strains of *A. minutum* had high numbers of small ELF-A spots located between the plates. All strains of *S. acuminata* also showed relatively high numbers of ELF-A spots distributed along the plate sutures, with a brighter fluorescence signal than *A. minutum* (Supplementary materials Fig. 3)

Temporal evolution of the APA labelling

The simultaneous fusion of every droplet containing the phytoplankton with the droplet containing the ELF-P substrate initiated the APA assay. We found an intra-clonal variability between the start of the APA assay and the observation of the first fluorescent spots at the surface of the dinoflagellates. For *S. acuminata*, we observed a fast labelling with a short lag-time (less than 5 s). In contrast, the labelling of *A. minutum* was obtained 10 min to 30 min after droplet fusion. All strains induced the synthesis of APA when the cells were cultivated in phosphate-deplete F/2 medium (Table 1). However, the timing of induction varied between the two species. *S. acuminata* strains induced the synthesis of AP between

3 and 5 days since the experiment started, whereas *A. minutum* strains induced the synthesis of the AP between six and seven days (Supplementary materials Fig. 5). For the strains SC-47-1, SC-92-1, AM-92-1, the APA and the number of cells labelled increased as a function of time and revealed the progressive increase in phosphate stress that the cells incurred (Table 1, Fig. 3). For strains SC-47-2, AM-47-1 and AM-92-2, the APA was found only once after 3, 6.5, and 6 days, respectively and was considered as the maximum for this study. The proportion of labelled cells ranged from 6% (AM-92-1, day 6) to 23% (AM-47-1, day 6.5), with the highest values found at the APA maximum. The results indicated that a period of 6–7 days was required to reach the maximum APA value for *A. minutum*, whereas 3–5 days were needed for *S. acuminata*. The day after the APA maxima, two main patterns were observed in the cultures. A first group of strain (AM47-1; SC92-1) had some cells still alive but not labelled. A second group of strains (SC47-1; SC47-2; AM92-2) had no living cells the day after APA maxima detection.

Intraclonal, intra- and inter-specific variability of the APA

The higher average values of APA were measured for two strains of *S. acuminata* corresponding to 1940's period (SC-47-1: 441 ± 51 fmol min⁻¹ cell⁻¹; SC-47-2: 421 ± 55 fmol min⁻¹ cell⁻¹). These values were higher than average APA values of the *A. minutum* strain of the same period (Table 1; Fig. 4; ANOVA, $p < 0.001$). The average value of APA of the *S. acuminata* strain of the 1990's (241 ± 56 fmol min⁻¹ cell⁻¹) is significantly higher than those of *A. minutum* strains of the same period (ANOVA, $p < 0.001$). These two *A. minutum* strains had the lowest average values of APA measured among all strains and ages (AM-92-1: 17 ± 7 fmol min⁻¹ cell⁻¹; AM-92-2: 21 ± 4 fmol min⁻¹ cell⁻¹). No significant differences were found between the maximum values of the APA of the strains AM-92-1 and AM-92-2 and between the strains SC-47-1 and SC-47-2. In summary, all *S. acuminata* strains had higher APA than all *A. minutum* strains and, for both species, strains from the 1940's had a significantly higher activity than the corresponding strains from the 1990's (ANOVA, $p < 0.001$).

Discussion

Extra/intra APA and cellular P stress status:

We developed an integrated microfluidic platform to automatically sort droplet containing cells and perform APA assays for dinoflagellates based on fluorescence at a single cell level. In our study, we mainly focus on external APA as an indicator of cellular P-stress status. This choice has been supported because dissolved organic phosphorus (DOP) compounds must be hydrolyzed extracellularly before being assimilated as a P source [44]. Moreover, the most important dissolved organic utilizing enzyme is AP and is reported to be mainly located at the cell surface and its activity is mostly linked to cellular P-stress status [27]. In the literature, intracellular AP is sometimes counted for the determination of cellular P-stress status. However, there are several major issues when considering intracellular AP as a P-stress indicator. Firstly, a part of the intracellular APA has been reported to be constitutive [30]. Secondly, intracellular APA activity seems to not have an access to some external DOP compounds, because no phosphoester transporters from the cellular membrane to the intracellular contents have been discovered yet [27]. Thirdly, the internal pH of the cell can vary within the day, leading to a range of pH values more suitable for acid phosphatases that have been described as more constitutive enzymes and ensuring different cellular functions [30, 31, 44-46]. By considering the extracellular APA as a key indicator for cellular P stress, we reported the first measurements of the regulation of the extracellular APA for two dinoflagellate species, the toxic *A. minutum* and non-toxic *S. acuminata*. These species were obtained from germination of cysts from ancient sediments of two contrasting periods of phosphorus loadings of the Bay of Brest. Our results show significant intraspecific and interspecies APA heterogeneities and a coherent pattern of APA variability between the two tested periods of time.

Temporal evolution of the APA at the single-cell level

The microfluidic approach enabled us to enumerate APA labelled living cells. To limit false-positive APA-labelled cells, we considered the swimming capability of the phytoplankton as a screening process for the detection of cell living status. This step allowed us to discard cysts and dead cells that have been reported to overestimate APA [47]. Because ELF substrate did not enter through a compromised cellular membrane and only living dinoflagellates marked with fluorescent ELF-A spots were counted as a positively labelled cell, the microfluidic approach enabled us to improve the measurement accuracy of the APA [48]. Furthermore, cells were not fixed with either ethanol, paraformaldehyde nor HgCl₂, which

are fixatives commonly reported to permeabilize cell membranes and affect cellular integrity [49-51]. The permeabilization of the cell membrane overestimates the proportion of positive cells because substrates can enter through the cell and label intracellular enzyme activity. Indeed, we obtained lower proportions of labelled cells in P-depleted samples (6–22%) than reported by other studies testing dinoflagellate fixed cells (*e.g.*, ~79% for *A. catenella* and *A. tamarensis* [52]; 68% for *Alexandrium* spp., 82–84% for *Protoperidinium* spp. and *Karenia mikimotoi* [29]; 50–90% for *S. acuminata* (there named *S. trochoidea*) [53]). Yet, we consider that the number of mobile labelled cells is sufficient for identifying significant intra- and inter-specific variabilities among strains and species, and to study this variability across strains of different ages. A higher number of cells would likely decrease the standard deviation but not significantly change our conclusions.

Alkaline phosphatase (AP) synthesis is commonly reported to be induced by phytoplankton during phosphorus stress [54]. However, the regulation of AP synthesis appears not to be ubiquitous among phytoplankton species because several species, including the dinoflagellate *Alexandrium catenella* (named as *A. fundyense*), were observed to constantly express variation in AP whatever the P content in the media [50, 55, 56]. Moreover, due to the lack of clear ecological relationships between APA and phosphate concentrations, Flynn *et al.*, reported that APA was not a reliable indicator of the P-status for the species *A. minutum* [57]. In our study, the absence of labelled cells at the beginning of monitoring and the progressive increase in number of labelled cells and APA through time suggested that almost all extracellular AP enzymes were not constitutive but were induced under P-deplete conditions. Moreover, our results suggest that APA is directly regulated by the cells through time. The temporal variability of APA observed here highlights the need for time series monitoring of enzyme regulation over several days in order to measure and compare the highest APA as possible among strains. Two different dynamics of APA increase have been found. First, a progressive increase of APA has been observed at the timescale of several days in *S. acuminata* strains (2–5 days). Second, a fast regulation of APA (<1 day) was found for the *A. minutum* strains AM-47-1 and AM-92-2. This fast regulation of the APA suggests that, despite our sampling efforts, the maximum values of APA determined in this study were probably not the highest that the strains can reach. Because the sorting and labelling steps in our

microfluidic platform took ~3 h, further improvements of the APA assay and microfluidic procedures may be required for studying fast regulation of APA at shorter timescales.

No living cells of AM-47-1 and SC-92-1 strains were labelled the day after a maximum of APA. Although the repression of APA can take three days for some dinoflagellate species, fast repression of APA has also been observed in on-a-field experiments and in cultures growing under P-deplete conditions [58-60]. The fast repression of AP synthesis observed in this study tends to confirm previous analyses showing a decrease in APA after 4–5 days of incubation, concomitant with the presence of APA in the dissolved fraction [34, 56, 58]. To explain such a quick repression of the AP synthesis while dissolved inorganic phosphate concentrations are still under the detection limit in the culture medium, we previously suggested that free-AP enzyme released from living cells or cell lysates in the dissolved fraction of the culture allowed new P-availability for the cell and consequently repressed intracellular AP synthesis [34]. The potential effect of phosphate released by free-AP enzymes in the medium on the cellular repression of AP synthesis has not been tested in this study. However, according to our culture conditions, the lyses of P-limited cells might have released free-AP enzyme in the batch culture and might have caused fast repression of AP synthesis, especially given that free-AP enzymes have been reported to have a long lifetime (>2 weeks) in laboratory-incubated environmental samples [61].

Intra-clonal and Intra-specific variabilities of the APA

Our results revealed a particularly high intraclonal variability of the APA (SD: from $\pm 12\%$ to $\pm 42\%$, Table 1). The important variability of APA at the intraclonal level adds evidence for high phenotypic heterogeneity of cells growing under the same environmental conditions. This result helps to confirm a recent single-cell labelling study that reported order of magnitude differences between the lowest and highest APA cell measurements for cultivation in phosphate-limited conditions [34]. High metabolic heterogeneity was also observed in nitrate and nitrogen uptakes and was suggested to be an investment of some cells in advance of nutrient availability changes [8, 62]. To cope with fluctuating nutrient conditions, some cells develop a set of enzymes with different affinities to the substrate. For example, Dyhrman and Palenik suggested that the haptophyte *Emiliania huxleyi* has a set of cell-surface AP enzymes with different substrate affinities [63]. By using AP enzymes with high or low affinities,

Emiliana huxleyi cells show different kinetics as a function of the P concentration in the environment. Due to the high phenotypic heterogeneity of APA, and the relatively low number of positive cells observed in this study, identification of different kinetic patterns of the APA is difficult. Indeed, observations of ELF-A spots with different fluorescent intensities between the armour plates on the same cell can be the result of a locally higher spatial density of enzymes or the presences of enzymes with different affinities, as reported for the species *Emiliana huxleyi* [62].

Although the intraclonal APA heterogeneity found here was high, APA kinetic differences between the two pairs of strains from the same species and same age cannot be discriminated significantly (i.e. SC47-1 and SC47-2; AM-92-1 and AM-92-2, Fig. 4). This result indicates that APA variability among strains of one species is hidden by the high intraclonal heterogeneity of APA within a strain. The enzymatic responses of the two pairs of strains to phosphate stress are particularly homogeneous and suggest that the strains of a species of same age had a same enzymatic answer to a P-limited environment. Therefore, among the population of dinoflagellates tested in this study and living in the same environment, the variation of APA appears to be more linked to the phenotypic heterogeneity than the intraspecies variability.

Inter-specific variability of the APA

The range of the APA measured by cell in the present study ($16\text{-}441 \text{ fmol min}^{-1} \text{ cell}^{-1}$; Table 1) was in agreement with APA measured in other dinoflagellate species using different protocols (e.g. $160\text{-}830 \text{ fmol min}^{-1} \text{ cell}^{-1}$ for *A. tamarensis*; $12.5 \text{ fmol min}^{-1} \text{ cell}^{-1}$ for *A. catenella*, [64, 65]. In this range, a constantly higher APA was observed for *S. acuminata* compared to *A. minutum* strains. This difference of level of activity, associated with a rapid trigger of the APA regulation found in *S. acuminata*, suggests that P-requirement was higher for *S. acuminata* than for *A. minutum*. The long delay time to trigger the APA in *A. minutum* cultures could be related to the capacity of this species to store phosphate in the cell and a lower P requirement needed for growth. The intracellular phosphorus content is well known to be a factor able to more directly control the APA than external phosphate concentration [66-68]. To estimate the P-storage capacities of a species, the “luxury coefficient” has been commonly used in the literature since its original publication [69]. This indicator is the ratio of the P-content of cell cultivated

in P-replete conditions to the P-content of cell living in P-deplete conditions. Literature records tend to support the first hypothesis, stating a larger storage capacity of *A. minutum* in comparison with *S. acuminata*. Indeed, a luxury coefficient of 8.1 can be determined for *A. minutum* from the compilation of different studies conducted on the same strain [70-72], whereas the ratio reported of *S. acuminata* ranges from 1.4 to 2.9 [73-75]. Therefore, abilities of *A. minutum* to store P in the intracellular phosphate pool may help this species to maintain the functional cell machinery even when phosphate becomes depleted in the medium and may delay or limit the synthesis of alkaline phosphatases under P-stress. The high storage capacity of *A. minutum* associated with a low P requirement may also result in difficulty to observe APA under low phosphate concentration in on-a-field studies [54].

Ecological implications of APA variabilities

Few studies addressed the effect of the cellular metabolism heterogeneity on the dynamics of phytoplankton population [76-78]. However, the emergence of single-cell methods brings growing evidence that among a population composed of theoretically “genetically identical” cells, heterogeneity of metabolism can modify the cell distribution and resilience [6, 79, 80]. Our results have shown that, for a same species, the average values at maximum APA in the old strains (1947±11 years) are significantly higher than the activities of more recent ones (1992±4 years). This inter-age variability of the APA observed suggests that two different dinoflagellate species have co-adapted to the changing environmental conditions occurred in the Bay of Brest at half-century timescale. However, it remains difficult to precisely link the regulation of APA and the hydrological parameters such as phosphate concentrations and nutrient stoichiometry measured in the Bay of Brest. Indeed, nutrient concentrations were not routinely measured at the end of the 1940’s and one of the first standard protocols for the measurement of phosphate in marine water has only been published by 1962 [44]. Despite these technical limitations, several hydrological patterns can be reconstructed based of the historical measurements (Supplementary materials). Although other environmental parameters, such as light, could limit the growth of dinoflagellates in turbid coastal environment or during winter, the historical trends suggest that, within all nutrients, phosphate appeared to be a potential limiting factor for phytoplankton development since the early 1990’s (Supplementary materials, Ref. 80). This limitation

is mainly linked to the higher increase of nitrate concentrations than phosphate in rivers from the end of 1940's to the 1990's decade and the consequent dystrophic unbalancing of N/P ratios in the environment. Strains that lived in 1990's, in potentially more important P-limited conditions than 1940's, would be expected to show high APAs. Surprisingly, even if the N/P ratio ranged from 170 to 426 in the early 1990's decade, the APA assays performed on *A. minutum* and *S. acuminata* strains indicated, for both species, an opposite trend, showing lower APA level for 1990's decade strains.

Given that the progressive P limitation of phytoplankton cannot be demonstrated during the 1940-1990 period because of the absence of nutrient data, we can advance two different hypotheses to explain low APAs observed for recent strains. A first hypothesis would be that dinoflagellate populations have not been P limited during the 1940-1990 period. The regular high P riverine concentrations and sediment inputs associated with the storage ability of some species could have ensured P needs of both dinoflagellate species. Therefore, these populations were not selected across time on the efficiency to use P and the increase of the APA because the P inputs were sufficient for living cells. This hypothesis is plausible if phosphate concentrations have actually increased from the 1940's to the 1990's in parallel with nitrate concentrations and the P-storage ability did not significantly change at the half century time scale. A second hypothesis supposes a lower concentration of P in the environment during the 1940-1990 period, to which some dinoflagellate sub-populations had progressively adapted. In this scenario, dinoflagellates of the 1990's period would have optimized the utilisation of intracellular stock of P, relying on this storage when limiting conditions occurred in the environment. Being adapted to lower P concentrations in the environment, dinoflagellates of the 1990's decade would have lower APA because the P cell requirement would be ensured by intracellular pool storage. This hypothesis is consistent with the high phenotypic plasticity of phytoplankton to adapt its cellular requirement under P starvation [82, 83]. Additional experiments including a higher number of strains are required to better understand the ecological importance of the APA in phosphate limiting conditions, including modern/actual strains from this same ecosystem. A major issue would be to understand if the energy cost for a cell to express and maintain the APA constitutes a significant advantage over other survival strategies at a half-century timescale. A concluding evidence of our observations is however that dinoflagellates can modify some physiological constants in order to survive in a changing environment [84].

Conclusions

We determined the regulation of the APA at the single cell level for two dinoflagellate species, *A. minutum* and *S. acuminata*, which were revived from ancient sediment core and cultivated under P-deplete conditions. The inter-species comparison of the APA indicated that *S. acuminata* had a significantly higher APA than *A. minutum* strains. This inter-species variability of the APA may result from a relatively high storage capacity of phosphate reported in *A. minutum*. The intra-specific comparison of strains with different ages revealed that recent strains (1992±4 years) had a significantly lower APA than the old strains (1947±11 years). Although genotypic analysis should be conducted to confirm our observations, our results suggested that the significant modification in the expression of the APA can take place at the half-century timescale in order to cope with the P concentration modifications in the environment. Adaptation to variations in P concentrations had probably contributed to generate similar evolution pattern in APA of these dinoflagellate strains. The microfluidic technology demonstrated here provides an efficient approach for exploring and measuring physiological changes at the single-cell level to modifications of the nutrient conditions. These results highlight the need to consider the plasticity of cells in ecological models, especially if the phytoplankton species can modify its own physiology faster than the long-term projection of the climate change scenario.

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Author contributions: MG, JCB and RS conceived and designed the study (MG, JCB microfluidics; MG, RS microbiology). Experiments were performed by MG (APA experiments, microfluidics), CL (phosphate concentrations measurements) and ML (cultivation and preparation of cells) under the supervision of RS and JCB. MG, LB and TB contributed analytical tools (MG, LB microfluidic platform and instrumentation; MG, TB microfluidic chips, MG image processing). MG, CL, CJ, YDA and JCB contributed to data analysis. MG, JCB and RS wrote the manuscript with contributions from all authors.

Conflict of interest: JCB is a co-founder of Emulseo whose surfactant formulation was used in this study.

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List of table and figures:

Table 1: Alkaline phosphatase activity (APA) of six dinoflagellate strains as a function of time. Number of living cells and percentage of labelled cells are also indicated in the table. The average of the APA in $\text{fmol min}^{-1} \text{cell}^{-1}$ is calculated using the living labelled cells. (*na*) is the absence of APA measure due to lack of living cell found in the droplets.

Figure 1: Sampling area from the sediment core PALM-KS-06 (Bay of Brest, Brittany, France) and microphotographs of the *Scrippsiella acuminata* (SC) and *Alexandrium minutum* (AM) strains revived from sediments (Scale bar 10 μm).

Figure 2: Examples of the six strains of dinoflagellates having alkaline phosphatase activity (APA). The top panel displays the photomicrographs of the cells encapsulated in the water-in-oil droplets progressively labelled with the ELF substrate (green fluorescence). The bottom panel shows the increase in fluorescence of the cells of each strain measured by the microfluidic platform during APA assays. Scale bar 20 μm .

Figure 3: Variability of the alkaline phosphatase activity (APA) for two strains of *Scrippsiella acuminata* (SC-47-1; SC-92-1) as a function of time. The hairlines show the APA of each cell which induce alkaline phosphatase (AP) synthesis, the thick lines are the average of APA and shade areas are the standard deviations. ‘No living cell captured’ means that no living cell was trapped in the microfluidic chip. ‘No living cell labelled’ means that all living cells captured in the microfluidic did not induce AP synthesis.

Figure 4: Difference in the maximum alkaline phosphatase activity (APA) of the *Scrippsiella acuminata* (SC) and *Alexandrium minutum* (AM) strains. The left panels show the increase of the fluorescent product of the enzymatic assay (ELF-A) as a function of time. Each hairline shows the ELF-A measured of a single cell. The thick lines are the average of the ELF-A and shade area is the standard deviation of the ELF-A measured during the experiment. The data collected within the 20 min are used to determine the maximum of the APA. The corresponding APA data is plotted on the right panel. Using all the data set, the one-way ANOVA test was performed to compare APA among the different strains.

Table 1: Alkaline phosphatase activity (APA) of six dinoflagellate strains as a function of time. Number of living cells and percentage of labelled cells are also indicated in the table. The average of the APA in $\text{fmol min}^{-1} \text{cell}^{-1}$ is calculated using the living labelled cells. (*na*) is the absence of APA measure due to lack of living cell found in the droplets.

	Time (day)	[PO ₄ ³⁻] (μM)	Number of living cells captured	Number of labelled cells	Labelled cells (%)	APA ($\text{fmol min}^{-1} \text{cell}^{-1}$)	APA SD (%)
SC-47-1	3	0.07	58	7	12	128 ± 33	± 26%
	4	<0.02	135	14	10.4	235 ± 60	± 26%
	5	<0.02	78	17	21.8	441 ± 51	± 12%
	6	<0.02	0	<i>na</i>	<i>na</i>	<i>na</i>	<i>na</i>
SC-47-2	3	0.03	83	12	14.5	421 ± 55	± 13%
	4	<0.02	0	<i>na</i>	<i>na</i>	<i>na</i>	<i>na</i>
AM-47-1	5	0.44	55	0	0	0	<i>na</i>
	6	<0.02	70	0	0	0	<i>na</i>
	6.5	<0.02	95	22	23.0	77 ± 27	± 35%
	7	<0.02	100	0	0	0	<i>na</i>
SC-92-1	3	0.38	30	0	0	0	<i>na</i>
	4	<0.02	86	6	7	36 ± 15	± 42%
	5	<0.02	153	20	13.1	241 ± 56	± 23%
	6	<0.02	61	0	0	0	<i>na</i>
AM-92-1	6	0.06	106	7	5.7	16 ± 5	± 31%
	7	<0.02	116	17	14.7	17 ± 7	± 41%
AM-92-2	5	2.85	92	0	0	0	<i>na</i>
	6	0.05	162	20	12.3	21 ± 4	± 19%
	7	<0.02	0	<i>na</i>	<i>na</i>	<i>na</i>	<i>na</i>

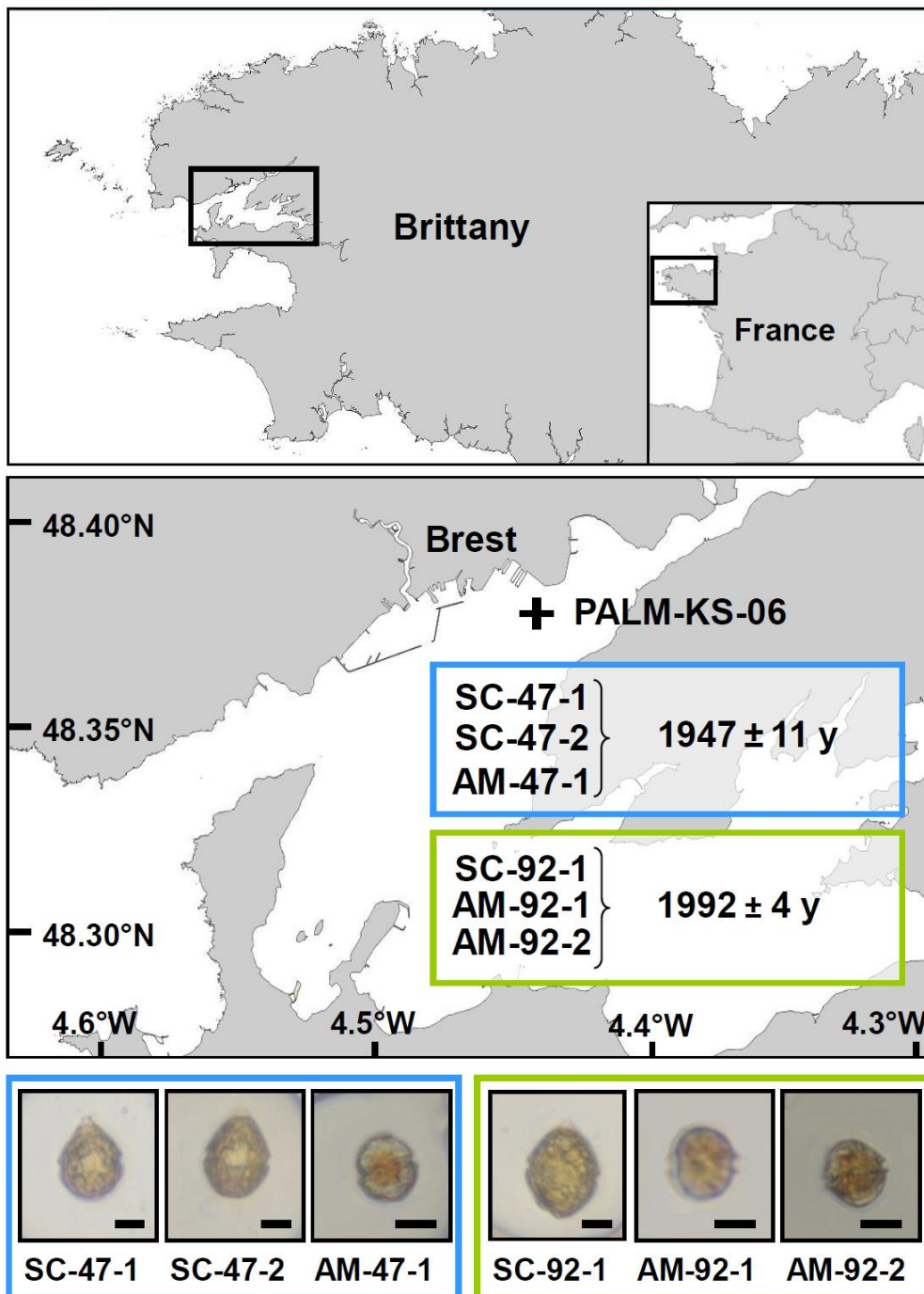


Figure 1: Sampling area from the sediment core PALM-KS-06 (Bay of Brest, Brittany, France) and microphotographs of the *Scrippsiella acuminata* (SC) and *Alexandrium minutum* (AM) strains revived from sediments (Scale bar 10 μ m).

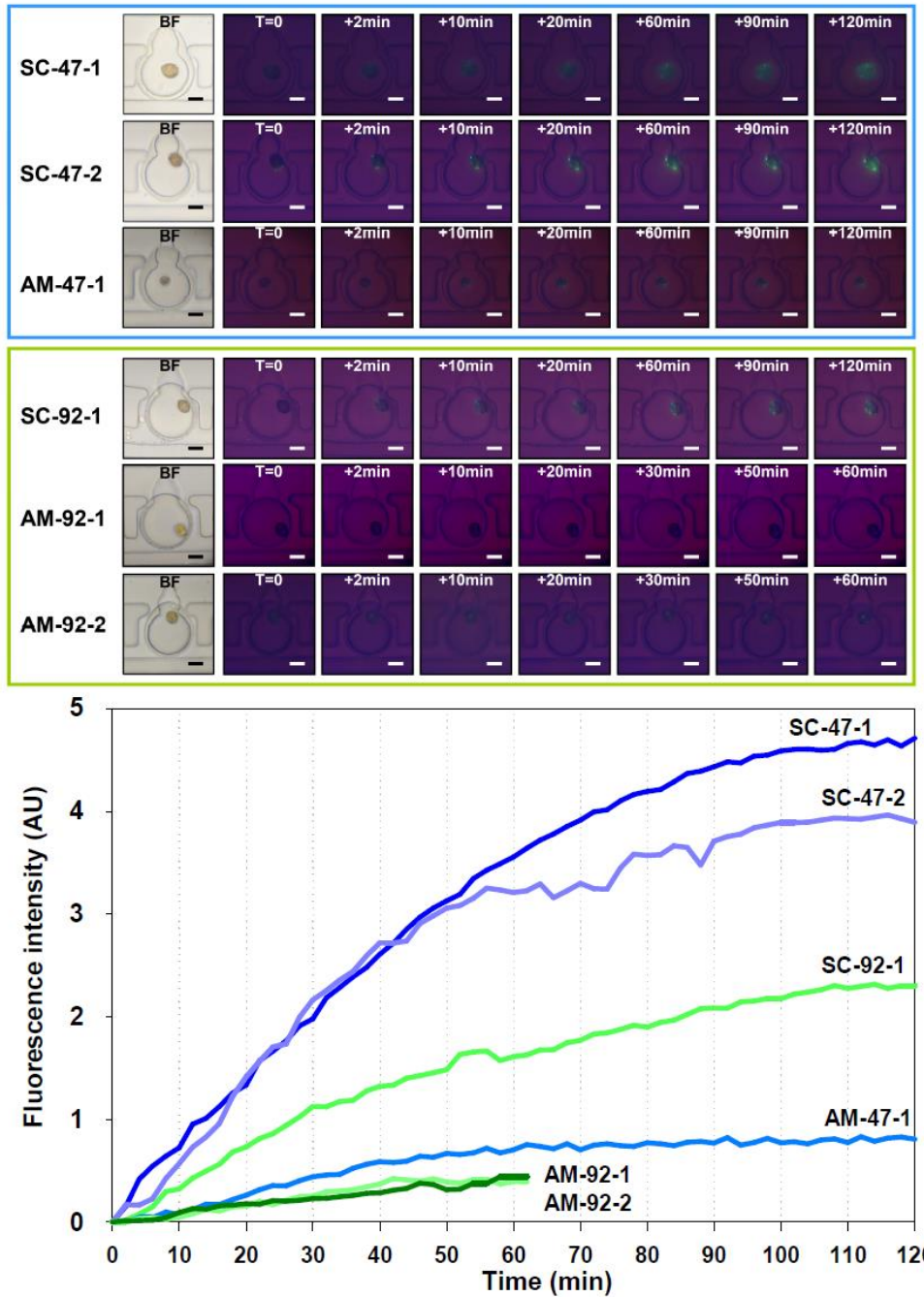


Figure 2: Examples of the six strains of dinoflagellates having alkaline phosphatase activity (APA). The top panel displays the photomicrographs of the cells encapsulated in the water-in-oil droplets progressively labelled with the ELF substrate (green fluorescence). The bottom panel shows the increase in fluorescence of the cells of each strain measured by the microfluidic platform during APA assays. Scale bar 20 μm .

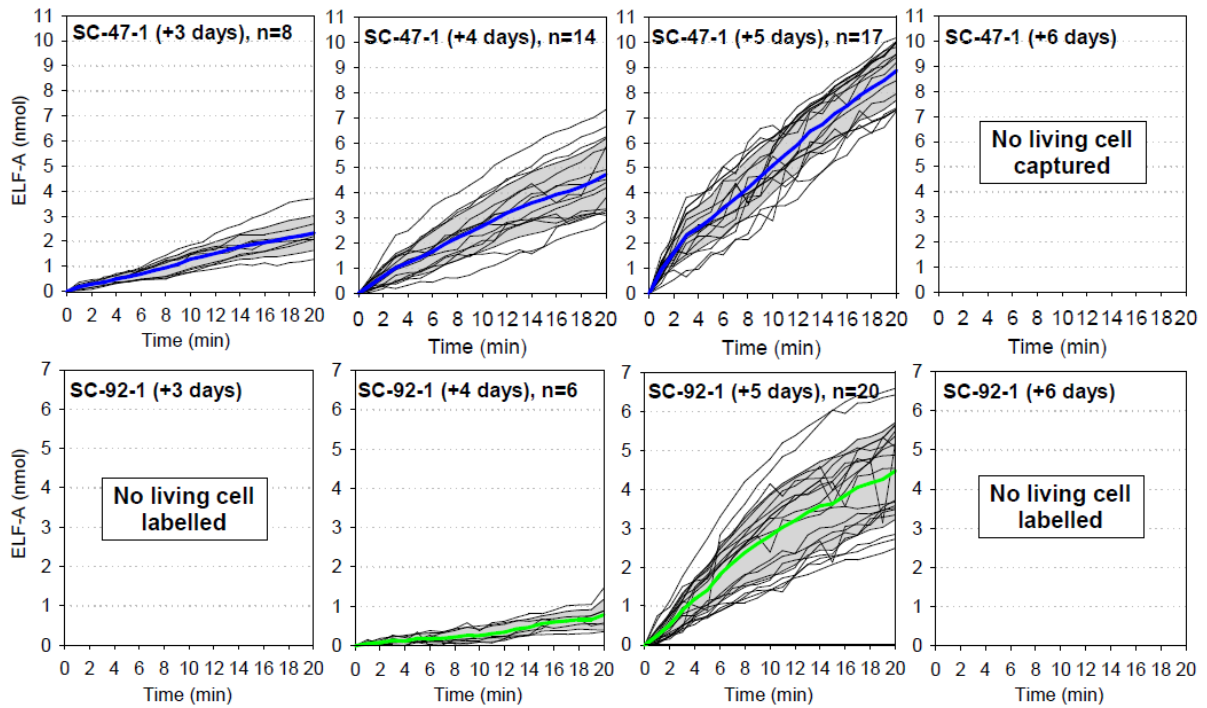


Figure 3: Variability of the alkaline phosphatase activity (APA) for two strains of *Scrippsiella acuminata* (SC-47-1; SC-92-1) as a function of time. The hairlines show the APA of each cell which induce alkaline phosphatase (AP) synthesis, the thick lines are the average of APA and shade areas are the standard deviations. ‘No living cell captured’ means that no living cell was trapped in the microfluidic chip. ‘No living cell labelled’ means that all living cells captured in the microfluidic did not induce AP synthesis.

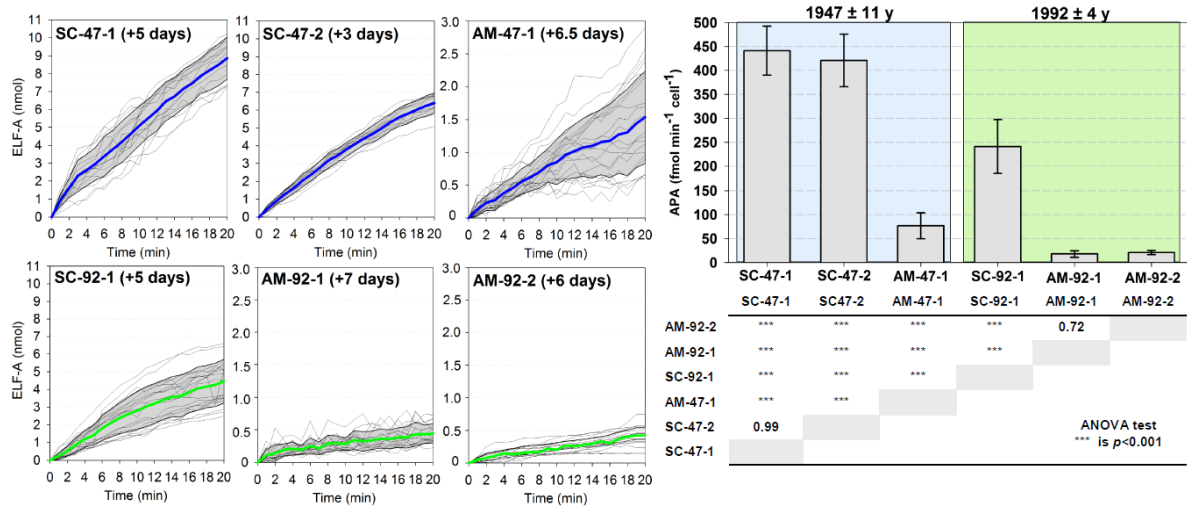


Figure 4: Difference in the maximum alkaline phosphatase activity (APA) of the *Scrippsiella acuminata* (SC) and *Alexandrium minutum* (AM) strains. The left panels show the increase of the fluorescent product of the enzymatic assay (ELF-A) as a function of time. Each hairline shows the ELF-A measured of a single cell. The thick lines are the average of the ELF-A and shade area is the standard deviation of the ELF-A measured during the experiment. The data collected within the 20 min are used to determine the maximum of the APA. The corresponding APA data is plotted on the right panel. Using all the data set, the one-way ANOVA test was performed to compare APA among the different strains.

Variable inter- and intra-species alkaline phosphatase activity within single cells of revived dinoflagellates

Supporting Information

Mathias Girault¹, Raffaele Siano², Claire Labry², Marie Latimier², Cécile Jauzein², Thomas Beneyton¹, Lionel Buisson¹, Yolanda Del Amo³, Jean-Christophe Baret^{1,4}

1) CNRS, Univ. Bordeaux, CRPP, UMR 5031, 33600 Pessac, France.

2) Ifremer, DYNECO, F-29280 Plouzané, France.

3) CNRS, Laboratoire d'Environnements et Paléoenvironnements Océaniques et Continentaux (EPOC), UMR 5805, 33615 Pessac, France.

4) Institut Universitaire de France, 75005 Paris, France

Corresponding authors: raffaele.siano@ifremer.fr, jean-christophe.baret@u-bordeaux.fr

Key words: Alkaline phosphatase activity, single cell, microfluidic, ELF, dinoflagellate.

Supplementary information Figure 1: Schema of the microfluidic platform setup dedicated to the alkaline phosphatase assay (APA) at the single cell level. The top left panel show the optical setup and the different devices to control the path of the droplets in the microfluidic chip. The detail of the microfluidic chip is shown on the top right panel. The photomicrographs (bottom panel) illustrated the mains steps of the APA assay performed in the microfluidic.

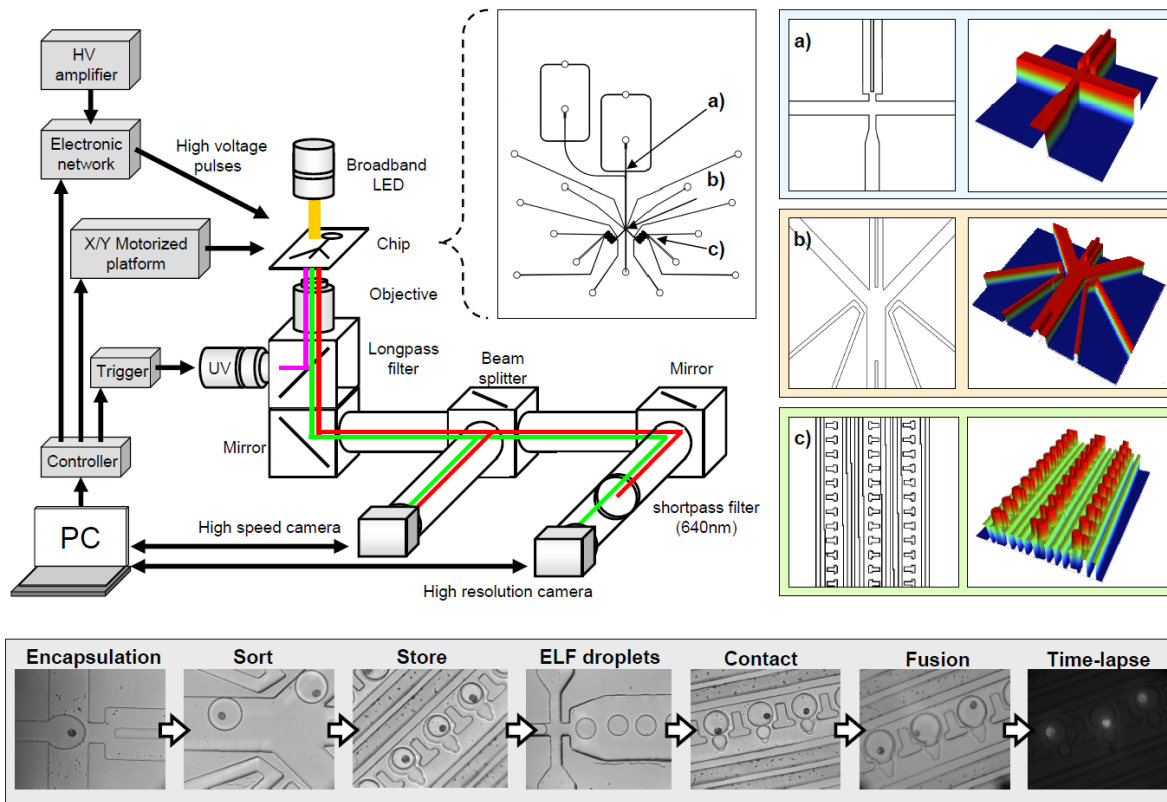
Supplementary information Figure 2: Fluorescence spectra of four dinoflagellate strains before and after the alkaline phosphatase assay (circles and triangles, respectively; ex: 365nm). The blue and the red dashed lines are the long and shortpass filters (470-640nm) used to block blue light and the fluorescence signal of the chlorophyll pigments captured by the camera.

Supplementary information Figure 3: Photomicrographs of the double staining labelling protocol performed on the six strains of dinoflagellates. The DAPI fluorescences are located inside the cell (i.e. nucleus) while the ELF fluorescence (green dots) is located at the surface of the dinoflagellates. Scale bars are 10 μ m.

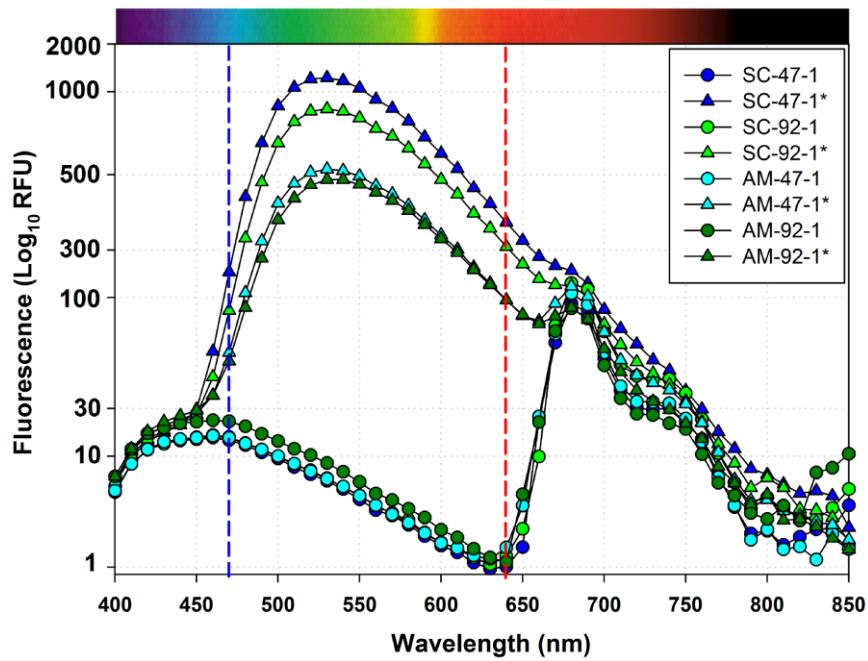
Supplementary information Figure 4: Flow chart of the image processing algorithm developed in this study. The main part of the software was to efficiently detect the edge of the droplets in the brightfield image. To optimize their detections, a median filter along the x axis was used to delete the salt and pepper noises. Then, a gamma correction (0.6) was applied in order to increase the difference of intensity between the objects and the background of the image. By using a threshold of 10 in pixel intensity and a distance of 5 between two interrogation pixels, an eight directions wavelet decomposition method was processed to the image in order to detect the edge of objects. From the binary images showing the edges of objects, a flood fill algorithm was computed to fill the objects. Then, circles were detected using the Hough transform. The locations of the center of the gravity and radius of droplets were saved in an array. Then, the array including the location and radius of the droplets was used to create a series of masks which perfectly fit the droplet areas. For each mask, intensity of each pixel was summed and the results were normalized by the area of the droplet.

Supplementary information Figure 5: Kinetics of the alkaline phosphatase activity of the *Scrippsiella acuminata* and *Alexandrium minutum* strains depending on the number of day where cells are cultivated in a phosphate deplete medium.

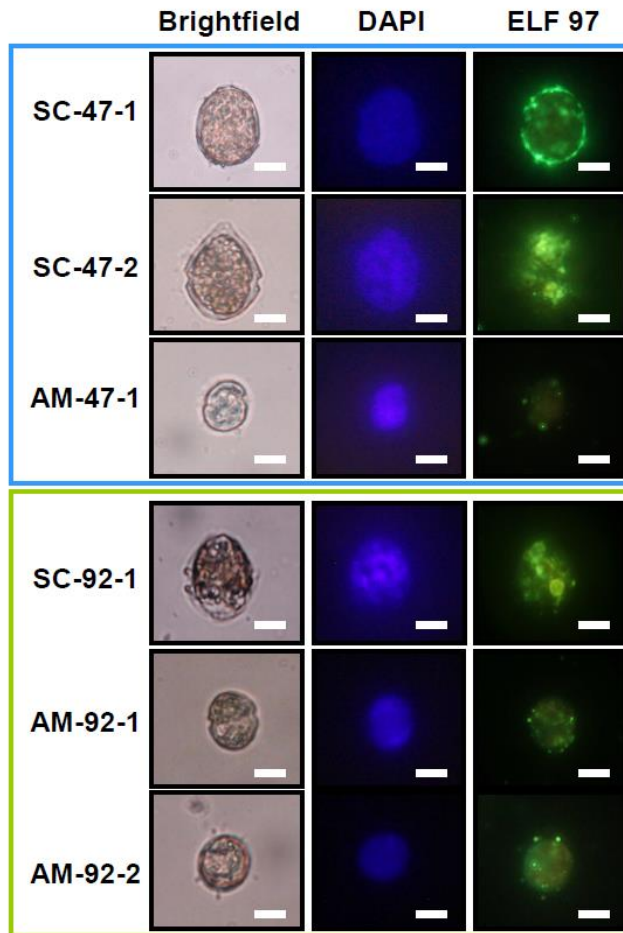
Supplementary information Figure 6: Nitrate and phosphate concentrations in the Elorn river (Pont ar Bled and Kerigeant sampling stations, respectively).



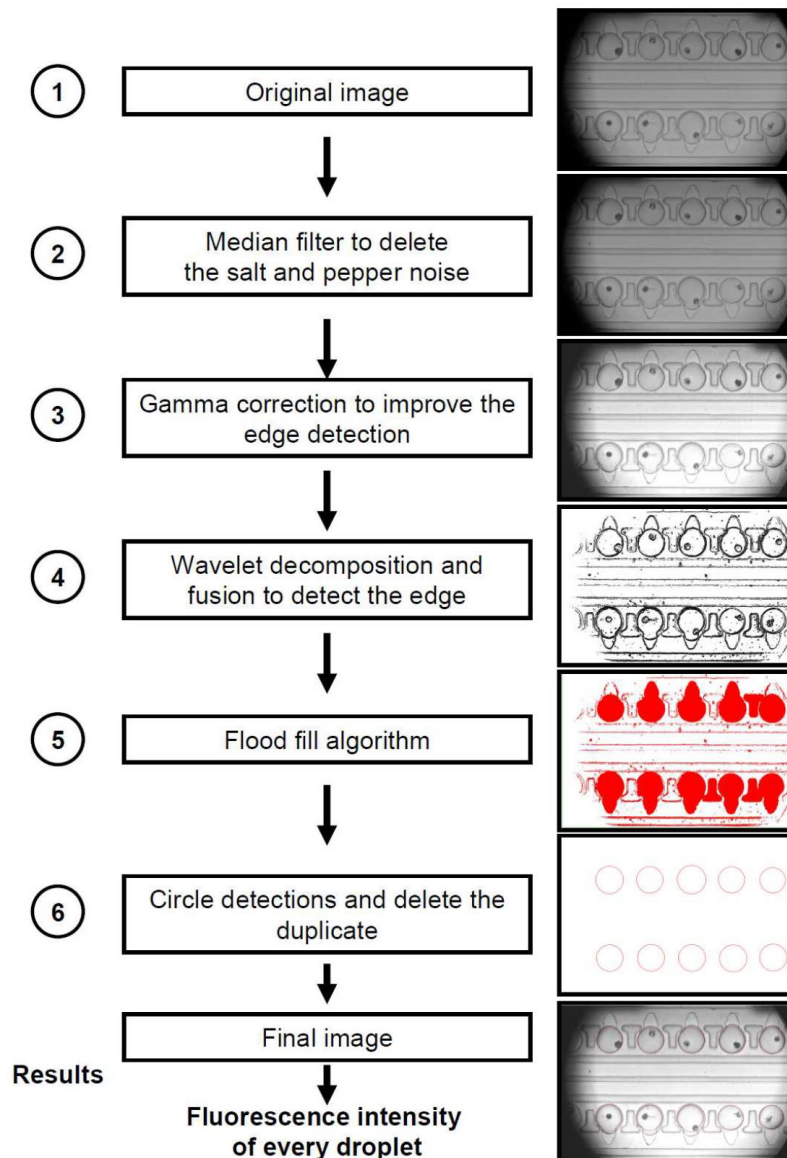
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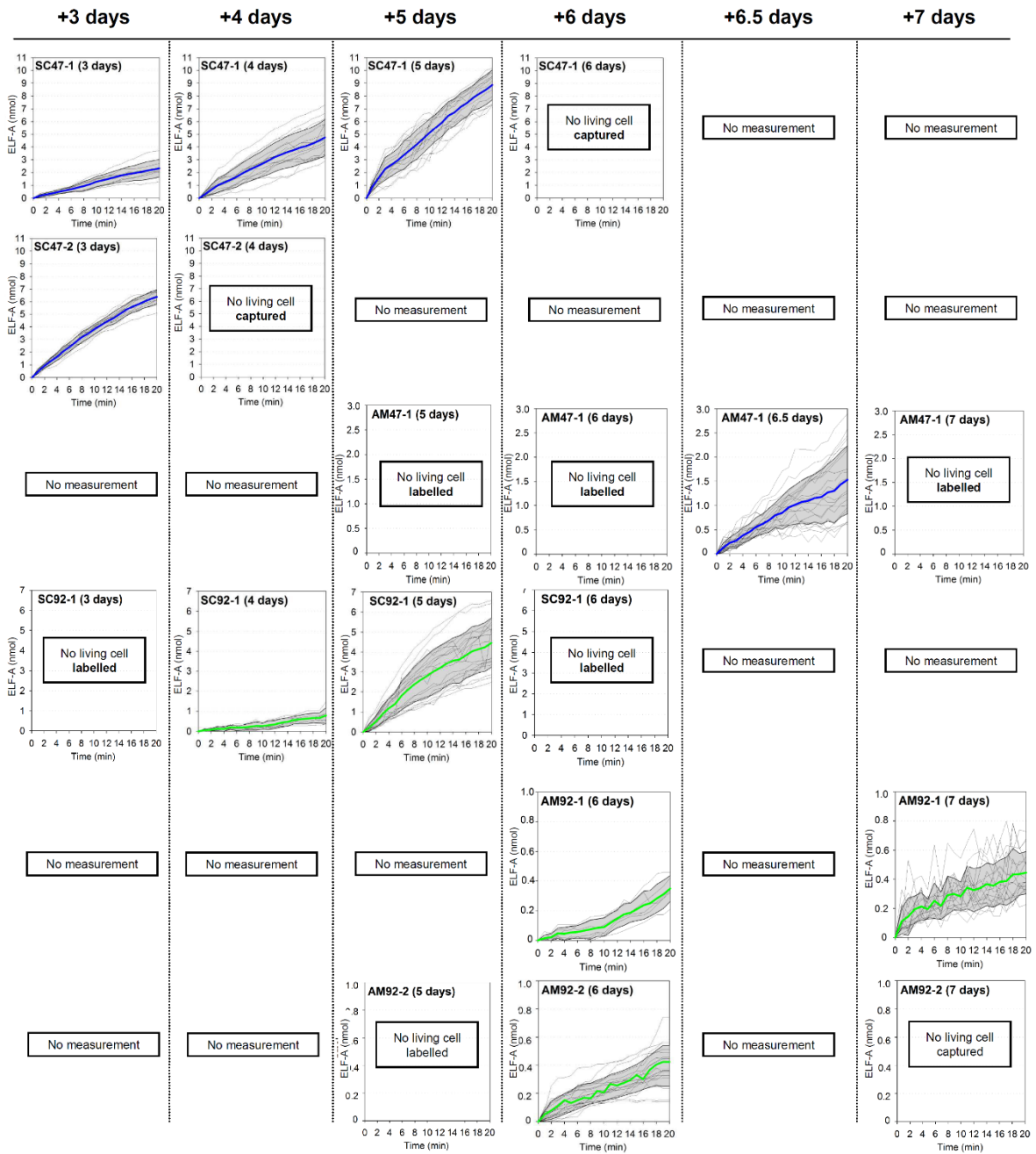
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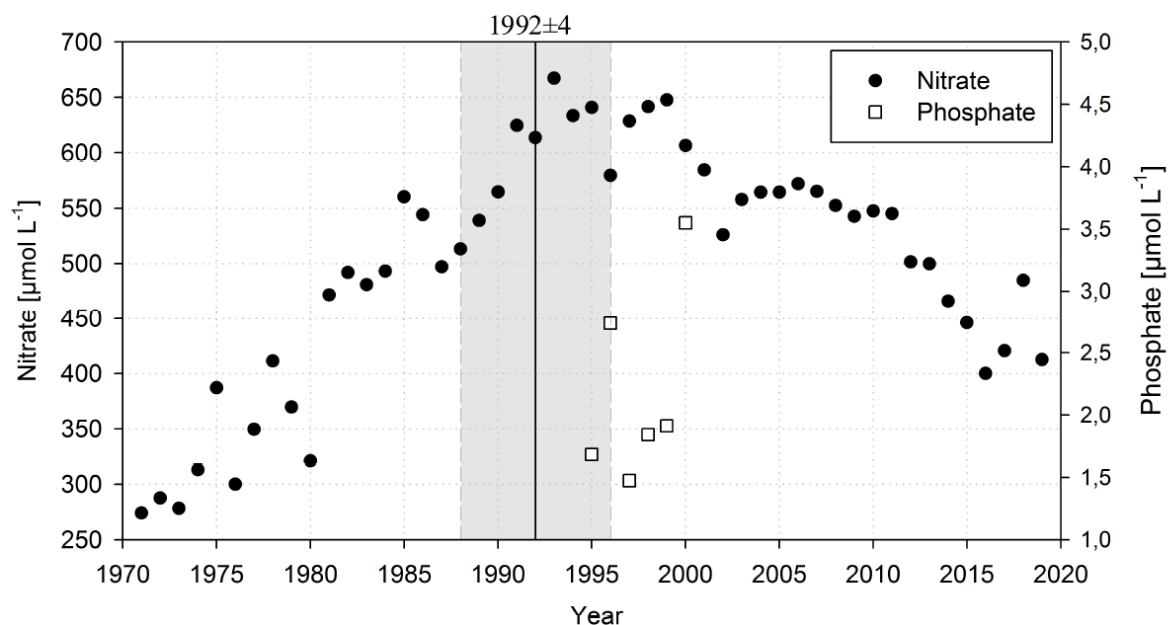
Control of the fluorescence signal during the APA assay

To test the fluorescence intensity of the natural pigments between 470nm and 640nm, an APA assay has been performed on four cultures of dinoflagellates in microtiter plate (SC-47-1; SC-92-1; AM-47-1; AM-92-1; Supplementary materials Fig. 2). As the fluorescence of the ELF-A product is located between the armour plates of the dinoflagellates, the presence of the bacteria harboured in these spaces have also been investigated in the cultures. A double staining protocol (4',6-diamidino-2-phenylindole DAPI and ELF) was assayed when the cell expressed the APA. The results indicated that only the nucleus of the dinoflagellate is stained by the DAPI (Supplementary materials Fig. 3). The absence of DAPI product at the same location as the ELF showed that bacteria were not responsible for the fluorescence of the ELF in the samples. Finally, to minimize the exposure time of cells to the harmful short wavelength, the ultraviolet LED was connected to a diaphragm shutter with a controller (SHB1T, Thorlabs). Typically, an exposition of 500 ms every 2 min was observed to not induce lyses of cells and not significantly bleach the intensity of the fluorescent product (*i.e.* ELF-A)

Historical nutrient data in the Bay of Brest

The Bay of Brest is a semi-enclosed basin fertilized by two main rivers (Aulne and Elorn) and wastewaters from human activities. According to the location of the sediment core used in this study, the hydrology of the Elorn river was considered in order to describe the changes of the environmental conditions. At the end of WWII, the industry and agriculture were particularly weakened by years of wars. Under these circumstances, nitrate and phosphate loads to the Bay through riverine inputs were probably particularly low. Because of the important circulation in the Bay of Brest (driven by tidal movement and residual currents) nutrient accumulation is limited (the residence time of seawater ranged from 3 days to 1 month; [85-86]). From the end of the WWII, a broad period of worldwide economic expansion, mainly initiated by the Marshall and Monnet plans tended to modernize both the industry and agriculture in France. Within this new context, the use of fertilizers (mainly nitrate and in a lesser extent phosphate) was largely promoted and used from the 1960's in order to increase the agricultural yield. Due in part to the high affinity of nitrate to leach from the soil, the nitrate inputs from the rivers strongly fertilized the roadstead of Brest. For example, one of the first studies conducted after the WWII pointed out that nitrate concentration in the estuary of Elorn river in the winters 1979-1981 was particularly high ($400 \mu\text{mol.L}^{-1}$; same order as the Seine river; a major river near Paris metropole, France) and nitrate charge in summer was so high that the phytoplankton growth in the estuary did not significantly decrease the nitrate concentration in summer [87, 88]. At the end of the 1970's, the growth of phytoplankton was strongly correlated to the nutrient pulses from rivers where several phytoplankton blooms were observed within the same year [89]. In the early 1980's, the ratio N/P reached 30 (*i.e.* >16 Redfield ratio) before the spring bloom of phytoplankton but the main limiting factor for phytoplankton was still reported to be the nitrate [89, 90]. From the early 1980's to the 1990's, the N/P ratio keeps increasing because of the long delay in efficiently regulating nitrate in agriculture together with the laws which severely limited phosphate discharges from anthropogenic activities (European directive 91/271, May 21st 1991; CJEU June 13th 2013: 62012CJ0193). As a consequence, the 1990's decade was characterized by the highest nitrate concentrations ever measured in the Elorn river (annual average is $613 \mu\text{M}$ in 1992 and up to $667 \mu\text{M}$ in 1993, Supplementary materials Fig. 6). In the same decade, phosphate concentrations in the Elorn river ranged from $1.4 \mu\text{M}$ to $3.6 \mu\text{M}$ leading to N/P ratios spread from 170 to 426 (Supplementary materials, Fig. 6). Despite the potential increase of phosphate concentrations within the bay waters from 1947 to 1992, these values indicated that the inputs of nutrients in the Bay of Brest were progressively unbalanced in relation to the higher nitrate concentrations. A possible visible consequence was that diatom blooms observed in 1993 and 1994 appeared to be firstly Si- and P-limited and then, in the lesser extend N-limited within the productive period from spring to summer [91]. Fifteen years before, N was the main limiting factor for phytoplankton in the Bay of Brest [92].

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Supplementary materials Figure 6: Nitrate and phosphate concentrations in the Elorn river (Pont ar Bled and Kerigeant sampling stations, respectively).

