

# Pyrenoidal sequestration of cadmium impairs carbon dioxide fixation in a microalga

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1	Pyrenoidal sequestration of cadmium impairs carbon dioxide
2	fixation in a microalga <sup>#</sup>
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## 35 ABSTRACT

36 Mixotrophic micro-organisms are able to use organic as well as inorganic carbon sources and, 37 thus, play an essential role in the biogeochemical carbon cycle. In aquatic ecosystems, the alteration of CO<sub>2</sub> fixation by toxic metals such as cadmium - classified as a priority pollutant -38 39 could contribute to the unbalance of the carbon cycle. In consequence, the investigation of 40 cadmium impact on carbon assimilation in mixotrophic micro-organisms is of high interest. 41 We exposed the mixotrophic microalga *Chlamydomonas reinhardtii* to cadmium in a growth medium containing both CO<sub>2</sub> and labeled <sup>13</sup>C-[1,2] acetate as carbon sources. We showed that 42 43 the accumulation of cadmium in the pyrenoid, where it was predominantly bound to sulfur 44 ligands, impaired CO<sub>2</sub> fixation to the benefit of acetate assimilation. TEM/X-EDS and 45 µXRF/µXANES at Cd L<sub>III</sub>-edge indicated the localization and the speciation of cadmium in the cellular structure. In addition, NanoSIMS analysis of the  ${}^{13}C/{}^{12}C$  ratio in pyrenoid and starch 46 47 granules revealed the origin of carbon sources. The fraction of carbon in starch originating from CO<sub>2</sub> decreased from 73% to 39% during cadmium stress. For the first time, the 48 49 complementary use of high-resolution elemental and isotopic imaging techniques allowed 50 relating the impact of cadmium at the subcellular level with carbon assimilation in a 51 mixotrophic microalga.

## 52 **KEYWORDS**

Biogeochemical carbon cycle, CO<sub>2</sub> fixation, toxic metals, cadmium stress, subcellular imaging,
 NanoSIMS, synchrotron µXRF, µXANES, TEM/X-EDS, isotopic labeling, isotope ratio

55

## 56 INTRODUCTION

57 The increase of carbon dioxide concentration in the atmosphere over the past decades has been 58 associated with global climate change. Therefore, investigation and understanding of the 59 biogeochemical carbon cycle is of primordial importance. The terrestrial and oceanic biosphere 60 plays a major role in the global carbon cycle by the production and conversion of organic matter 61 (Field, Behrenfeld, Randerson & Falkowski, 1998). Plants have developed a process for CO<sub>2</sub> 62 fixation into organic carbon using the enzyme ribulose-1,5-bisphosphate 63 carboxylase/oxygenase (RuBisCO) which is regarded to be the most abundant protein in the biosphere (Ellis, 1979). Microalgae are small eukaryotic organisms where RuBisCO is 64 65 concentrated in the specific organelle called the pyrenoid (Badger et al., 1998). Pyrenoid containing algae have been identified as being responsible for 28 - 44% of the global carbon 66 67 fixation which underlines the key role of microalgae in the biogeochemical carbon cycle 68 (Mackinder et al., 2016).

Eukaryotic algae are classically considered as photoautotrophic using light and inorganic compounds for energy and carbon fixation, or heterotrophic using only organic compounds. Recently, mixotrophy has been integrated into aquatic biogeochemical cycle modeling (Caron, 2016; Ward & Follows, 2016). Mixotrophic organisms are able to combine photoautotrophic growth conditions as well as heterotrophic nutrition. They are dominant among micro-algal communities in oligotrophic and eutrophic environments (Burkholder, Glibert & Skelton, 2008; Hartmann et al., 2012). To date, the consideration of mixotrophy in biogeochemical
carbon cycle models is still at the beginning (Caron, 2016). Recently it has been shown that
mixotrophy globally lead to an increase of the size of organisms in the aquatic food chain and
thus enhances carbon sinking into the deep ocean by 35% (Ward & Follows, 2016).

79 Cadmium is a highly toxic metal and is considered as a priority pollutant in ecosystems 80 (Campbell, 2006). In high polluted natural waters micromolar cadmium concentrations were 81 found such as in the Tinto river in Spain (40 - 620 µM) (Aguilera & Amils, 2005) while for 82 other natural waters lower Cd concentration are described, e.g. 2.4 - 17.8 nM in polluted 83 Macedonian rivers (Ramani et al., 2014) and only 0.2 - 2 nM in a Swiss river (Xue & Sigg, 84 1998). Also, in polluted soils and soil solutions high cadmium concentrations were measured, 85 for example, 5.4 - 45.5 µM Cd in soil solutions of soils close to a chemical engineering plant 86 in Northern Taiwan (Lee, Lai & Chen, 2004) and 284 mg Cd per kg in paddy soils from 87 Thailand (Simmons, Pongsakul, Saiyasitpanich & Klinphoklap, 2005). Moreover, Cd has been 88 shown to be biomagnified in some aquatic food webs (Croteau, Luoma & Stewart, 2005; 89 Ikemoto et al., 2008; Rouleau, Gobeil & Tjlve, 2006; Signa, Mazzola, Tramati & Vizzini, 90 2017). At the microbial level, this accumulation occurs in communities between bacteria and 91 eukaryote organisms (Worden et al., 2015). In view of the importance of mixotrophy in the 92 aquatic carbon cycle, the impact of toxic metals such as Cd on carbon assimilation in 93 microalgae is of growing interest.

*Chlamydomonas reinhardtii* is a unicellular green alga which is found worldwide in soils and
fresh water as its natural habitat. This alga is used as a model for many cellular processes (Dent,
Han & Niyogi, 2001; Goto & Johnson, 1995; Harris, 1989; Merchant et al., 2006; Merchant,
Kropat, Liu, Shaw & Warakanont, 2012; Merchant et al., 2007; Rochaix, M. GoldschmidtClermont & Merchant, 1998; Silflow & Lefebvre, 2001) including metallic stress (Hanikenne,

99 2003). Cadmium is known to inhibit its sexual reproduction (Goodenough et al., 1993), to limit 100 photosynthesis activity (Faller, Kienzler & Krieger-Liszkay, 2005; Nagel & Voigt, 1989, 1995; 101 Voigt & Nagel, 2002; Voigt, Nagel & Wrann, 1998) and to induce oxidative stress (Aksmann 102 et al., 2014; Vega, Garbayo, Domínguez & Vigara, 2006). In order to reduce free intracellular 103 Cd<sup>2+</sup>, the microalga synthesizes thiol peptides such as phytochelatins (Bräutigam, 104 Schaumlöffel, Preud'Homme, Thondorf & Wesenberg, 2011; Gekeler, Grill, Winnacker & 105 Zenk, 1988) and sequesters Cd bound to polyphosphate granules into vacuoles which were 106 compared to acidocalcisomes (Penen et al., 2017; Ruiz, Marchesini, Seufferheld, Govindjee & 107 Docampo, 2001).

C. reinhardtii can grow photoautotrophically in the presence of light and CO<sub>2</sub>, 108 109 heterotrophically in the dark when acetate is available as an organic carbon source or 110 mixotrophically with acetate, CO<sub>2</sub> and light (Johnson & Alric, 2013). In the latter case, CO<sub>2</sub> is 111 photosynthetically fixed by RuBisCO (Johnson & Alric, 2012, 2013), while non-112 photosynthetic acetate assimilation begins by its incorporation into acetyl coenzyme A (acetylCoA) (Johnson & Alric, 2012, 2013). Then, acetylCoA is metabolized in the 113 114 tricarboxylic acid (TCA) cycle in mitochondria or in the glyoxylate cycle in cytosol (Johnson 115 & Alric, 2012, 2013; Singh, Shukla, Chary & Rao, 2014). Both acetate assimilation and 116 photosynthesis lead to starch formation (Ball, Dirick, Decq, Martiat & Matagne, 1990; Heifetz, 117 Förster, Osmond, Giles & Boynton, 2000; Johnson & Alric, 2012, 2013; Park et al., 2015). In 118 spite of an abundant literature about carbon assimilation under mixotrophic conditions in C. reinhardtii, the impact of toxic metals on the carbon metabolism has not been investigated yet. 119

Ion beam and X-ray beam imaging techniques are powerful tools to study the isotopic and
elemental composition of single-cells (Ortega, Devès & Carmona, 2009; Penen et al., 2016;
Roschzttardtz et al., 2011; Sarret et al., 2013; Schaumlöffel et al., 2016). Nanoscale secondary

123 ion mass spectrometry (NanoSIMS) allows elemental and isotopic mapping with a resolution 124 of about 50 nm. In biology, it is used to localize and to determine at a subcellular level the turnover of proteins and metabolites using stable isotopic tracers (<sup>13</sup>C and <sup>15</sup>N) (Gao, Huang & 125 Tao, 2016; Lechene et al., 2006). Moreover, when NanoSIMS is combined with transmission 126 127 electron microscopy (TEM) in correlative imaging, isotopic tracers can be localized in the cell 128 structure (Clode, Stern & Marshall, 2007; Hoppe, Cohen & Meibom, 2013; Kopp et al., 2015; 129 Penen et al., 2016). NanoSIMS mapping has been also used to determine metal distributions in 130 plants and microalgae (Hong-Hermesdorf et al., 2014; Moore et al., 2010, 2011, 2012). As 131 complementary techniques, synchrotron micro X-ray fluorescence (µXRF) and micro X-ray 132 absorption spectroscopy (µXAS) can be combined and directly applied on frozen hydrated 133 samples to locate metals and decipher their structural environment, respectively, allowing the 134 determination of metal distribution and speciation in subcellular regions of microalgae (Adams 135 et al., 2016; Leonardo et al., 2014; Leonardo et al., 2016; Penen et al., 2017; Wang, Lv, Ma & Zhang, 2016). 136

137 Here, we investigated the impact of cadmium on carbon assimilation in C. reinhardtii grown 138 under mixotrophic conditions. The objective was to relate subcellular cadmium localization and speciation with acetate and  $CO_2$  assimilation. Microalgae were thus exposed up to 70  $\mu$ M 139 Cd and cultivated in the presence of labeled  ${}^{13}C$ -[1,2] acetate to distinguish carbon originating 140 141 from acetate assimilation or from photosynthesis. A multimodal approach at subcellular level 142 allowed the localization of cadmium in the algal structure by TEM/X-EDS (X-ray energy dispersive spectroscopy) and the determination of carbon origins by measuring the  ${}^{13}C/{}^{12}C$  ratio 143 144 in the pyrenoid and starch using NanoSIMS. In addition, Cd was localized by µXRF, which 145 was combined to µXANES (micro X-ray absorption near edge structure spectroscopy) at the Cd L<sub>III</sub>-edge to identify Cd speciation. Our study showed that in C. reinhardtii cadmium 146 147 sequestration in the pyrenoid impaired CO<sub>2</sub> fixation to the benefit of acetate assimilation.

## 148 MATERIALS AND METHODS

### 149 **Biological material and growth conditions**

150 Chlamydomonas reinhardtii wild type strain (SAG 11/32b, Experimental Phycology and 151 Culture Collection of Algae at Goettingen University (EPSAG), Germany) was grown 152 heterotrophically in a tris-acetate-phosphate medium (Harris, 1989) where 153 ethylenediaminetetraacetic acid (EDTA) and FeSO4.7H2O were replaced by iron 154 ethylenediamine-N,N'-bis(2-hydroxyphenylacetate) (Fe-EDDHA) to increase the availability of cadmium in solution. All chemicals and reagents were purchased from Sigma-Aldrich 155 156 (Saint-Quentin Fallavier, France) unless stated otherwise. The composition of this TAPEDDHA medium is described in Table S1. For carbon fixation experiments, the  $TAP_{EDDHA}$  (<sup>13</sup>C) medium 157 was prepared with <sup>13</sup>C-labeled acetate (sodium <sup>13</sup>C-[1,2] acetate, 99% <sup>13</sup>C, Sigma Aldrich) 158 159 instead of acetic acid and pH was equilibrated to 7 with a HCl solution. Microalgae grew at 22 160 °C under constant illumination (by one Osram 827 and two Osram 840 fluorescent tubes) and 161 constant agitation (120 rpm). Fresh medium was inoculated with stock culture at the end of 162 exponential phase to reach an initial optical density at 730 nm (OD<sub>730nm</sub>) of 0.040. Micro-algae 163 were exposed to 10, 20, 30, 40, 50 and 70 µM of CdCl<sub>2</sub> added 24 h after the inoculation while control cultures were grown in parallel without Cd exposure (control). For controls and each 164 165 Cd concentration three independent algal cultures were grown (triplicates) and used for replicated determination of the growth rate, chlorophyll and starch concentrations, and 166 167 cadmium uptake.

As the main aim of this study was the investigation of the cadmium stress response at cellular level during the exponential growth phase, cells were exposed to Cd after 24h. Therefore changes in the lag phase, the delay before the start of exponential growth, were not assayed since this delay allowed the adaptation required for cells to begin the experience of new 172 environmental conditions, i.e. cadmium exposure in this study. The cell cycle of C. reinhardtii 173 is about 24 h which is confirmed with the start of the exponential phase at this time point. This 174 means that under these conditions the cells of the second (and third) cell cycle are exposed. 175 Cells were harvested after 72 h for different analyses, i.e. at the end of the third cell cycle when these cells experienced similar exposure conditions. It is a good approximation although 176 177 cultivation was not synchronized. This procedure also applied in our previous studies (Bräutigam, Schaumlöffel, Preud'Homme, Thondorf, & Wesenberg, 2011; Bräutigam, 178 179 Schaumlöffel, Krauss, & Wesenberg, 2009; Penen et al., 2017).

180 The concentrations of free metals in the medium were simulated (Table 1) with the freeware 181 Visual MINTEQ 3.0 (vminteq.lwr.kth.se/visual-minteq-ver-3-0/) (Gustafsson, 2013) using 182 Lindsay's database containing thermodynamic equilibrium constants compiled by Lindsay 183 (1979). Before use, glassware was washed with 5% HNO<sub>3</sub>. All solutions were prepared with 184 ultrapure water (18 M $\Omega$  cm at 25 °C) obtained from a Milli- Q system (Millipore, Bedford, 185 MA).

### 186 Growth rate determination

187 Optical density at 730 nm (OD<sub>730nm</sub>) was measured every 12 h in each control and Cd-exposed 188 algal cultures, grown in triplicate, using a spectrophotometer. Growth rates µ were calculated 189 as follows:  $\mu = (\ln N_f - \ln N_i)/\Delta t$ . N<sub>f</sub> was the OD<sub>730nm</sub> of the culture measured at the end of the 190 exponential phase. N<sub>i</sub> was the OD<sub>730nm</sub> of the culture at the beginning of the exponential phase 191 and  $\Delta t$  was the duration of the exponential phase (in days). Relative growth rate was expressed 192 as the ratio  $\mu/\mu_0$  determined during the exponential phase of growth ( $\mu$ : growth rate during 193 cadmium exposure,  $\mu_0$ : growth rate in control conditions). The Excel macro REGTOX 194 (http://www.normalesup.org/~vindimian/en\_index.html) was used to calculate the half 195 maximal efficient concentrations ( $EC_{50}$ ) applying the Hill model.

### 196 Chlorophyll concentration measurement

197 The chlorophyll concentration assay was adapted from Arnon et al. (Arnon, 1949). After 72 h 198 cultivation, 1 ml of each control and Cd-exposed algal cultures (grown in triplicate) was 199 centrifuged at  $13000 \times g$  for 1 min. Each algal pellet was resuspended in 1 mL of 80% acetone 200 and vortexed for 2 min. Then, solubilized chlorophyll was separated from cell debris by a 201 centrifugation at  $13000 \times g$  for 4 min. After that, the concentration (C) of chlorophyll a and b 202 in the culture medium was determined spectrophotometrically at 652 nm and calculated as follows:  $C = Abs_{652nm} \times 1000 / 34.5$  with C in mg mL<sup>-1</sup>. Finally, in order to relate the chlorophyll 203 204 content to the biomass, the determined chlorophyll concentration was normalized to the optical 205 density at 730 nm whose value is proportional to the number of cells in the medium (unit:  $\mu g$  $mL^{-1} OD_{730nm}^{-1}$ ). 206

### 207 Starch concentration measurement

208 After 72 h cultivation, microalgae were harvested from in triplicate grown control and Cd-209 exposed (70  $\mu$ M) algal cultures by centrifugation at 3000  $\times$  g for 3 min and each pellet was resuspended in 1.5 mL water. In order to estimate the dry biomass, a 0.5 mL aliquot from each 210 211 sample was dried at 60 °C for 24 h and the resulting pellet was weighted. The remaining 1 mL 212 samples were washed once with water and once with 80% ethanol, and centrifuged  $(3000 \times g)$ 213 for 3 min). Then, each pellet was boiled in 2 mL water for 10 min. Finally, starch concentrations 214 were determined in these extracts, using the enzymatic starch assay kit HK (Sigma Aldrich) 215 described in Delrue et al. (1992).

### 216 Cadmium uptake

217 After 48 h cadmium exposure (70  $\mu$ M), microalgae were harvested from in triplicate grown 218 cultures by centrifugation at 3000  $\times$  g for 3 min. In order to obtain the intracellular Cd 219 concentration, each algal pellet was subjected to the following washing protocol according to 220 Macfie et al. (1994): a first time in water, then twice in a mix of Na<sub>2</sub>-EDTA (1 mM) / CaCl<sub>2</sub> 221 (3.77 mM) and finally in water with centrifugation between in (3000  $\times$  g for 3 min). The 222 remaining pellets were dried at 60°C for 24 h and weighted. This dry algal biomass was digested in a mixture of 1 mL 70% (v/v) HNO<sub>3</sub> and 1 mL 30% (v/v) H<sub>2</sub>O<sub>2</sub> at 80 °C for 3 h. 223 224 Samples were then diluted to reach a 2% HNO<sub>3</sub> concentration. The cadmium concentration was determined by ICP-MS (Inductively Coupled Plasma Mass Spectrometry, 7500 model, 225 226 Agilent Technologies).

227

### 228 Sample preparation for TEM/X-EDS and NanoSIMS

229 Microalgae from a control and a Cd-exposed (70 µM) culture containing <sup>13</sup>C-labeled acetate in 230 the medium were harvested after 72 h of inoculation by centrifugation at  $3,000 \times g$  for 3 min. A protocol including high pressure freezing, and freeze substitution followed by resin inclusion 231 232 was used to preserve cell integrity. For high-pressure freezing, each microalgae pellet was 233 resuspended and incubated for 1 h in 150 mM mannitol. After a centrifugation at  $3000 \times g$  for 234 3 min, the harvested cells were included in agarose 2%. Then, the inclusion was plunged in 235 cryo-protectant 1-hexadecene and high-pressure freezing was performed immediately (EM 236 HPM 100, Leica Microsystems, Vienna, Austria). Samples were maintained in liquid nitrogen 237 until the freeze-substitution steps.

Freeze-substitution was adapted from O'Toole's protocol (O'Toole, 2010). Under liquid nitrogen, algal samples were post-fixed in 1% osmium tetroxide, 0.1% uranyl acetate and 0.25% glutaraldehyde in anhydrous acetone and transferred into the Automatic Freeze Substitution System AFS2 (Leica microsystems). Temperature was maintained at -90 °C for 3 d and then warmed to -30 °C for 8 h. Samples were removed from freeze-substitution system,
kept at 0 °C for 1 h and washed with water-free acetone.

Resin inclusion was carried out at room temperature. Samples were embedded in epoxy resin
(EPON 812, Delta Microscopies, Mauressac, France) using a graded resin and acetone series.
The following steps were applied for infiltration: 3:1 acetone:resin, 1:1 acetone:resin, 1:3
acetone:resin, and 2 baths with 100% resin. Finally, resin embedded samples were polymerized
for 48 h at 60 °C.

Samples were cut in 70 nm and 300 nm sections for TEM and NanoSIMS, respectively, using a diamond knife (Diatome, Biel-Bienne, Switzerland) on an ultra-microtome [EM Ultracut-UC7, Leica Microsystems) and placed on the respective sample holder. In order to carry out correlative imaging by TEM and NanoSIMS, a first 70 nm section was placed on a copper grid for TEM and then, an adjacent 300 nm section was placed on a silicon wafer (Wafer Solution, Le Bourget du lac, France) for NanoSIMS as previously described (Penen et al., 2016). Cells were localized and observed first by TEM and then, relocalized and analyzed by NanoSIMS.

### 256 **TEM/X-EDS analysis**

A transmission electron microscope FEI TECNAI 12 (Eindhoven, The Netherlands) using an accelerating voltage 120 kV and equipped with an X-Energy Dispersive Spectroscopy (X-Flash 6T 60 BRUKER-SYNERGIE 4, Evry, France) was used. The parallel electron beam of the nanoprobe was focused on 0.25-1.2  $\mu$ m areas. Under these conditions X-EDS spectra of 1-2k counts s<sup>-1</sup>, a deadtime of 15-20% and analytical time of 120 - 240 s per spectrum were generated.

### 263 NanoSIMS analysis

264 A nanoscale secondary ion mass spectrometer NanoSIMS 50L (CAMECA, Gennevilliers, France) was used to perform SIMS analysis. This device was equipped with a primary Cs<sup>+</sup> ion 265 source (lateral resolution down to 50 nm) for the mapping of electronegative elements, and 266 267 with a novel O<sup>-</sup> RF plasma primary ion source enabling a high sensitivity for electropositive elements combined to a high lateral resolution (40 nm lateral resolution) (Malherbe et al., 268 269 2016). The seven parallel electron multiplier detectors equipped on the NanoSIMS 50L 270 instrument allowed detection of seven elements at the same time. In order to remove major 271 interferences on the chosen element masses, the mass resolution (M/ $\Delta$ M) was tuned to be about 5000. Initially, ion maps (256×256 pixels, 64-144  $\mu$ m<sup>2</sup>) of <sup>12</sup>C<sup>-</sup>, <sup>13</sup>C<sup>-</sup>, <sup>12</sup>C<sup>14</sup>N<sup>-</sup> (for N detection) 272 and  ${}^{32}S^{-}$  were carried out using the primary Cs<sup>+</sup> ion source. Mapping of  ${}^{31}P^{+}$  and  ${}^{40}Ca^{+}$  was then 273 274 performed using the O<sup>-</sup> RF plasma primary ion source.

### 275 NanoSIMS data processing

276 Image processing was performed using ImageJ software (v.1.48, Wayne Rasband, National 277 Institutes of Health (NIH), Bethesda, MD, USA). Processing of NanoSIMS images required 278 additionally the openMIMS plugin developed at the National Resource for Imaging Mass 279 Spectrometry (NRIMS, Cambridge, MA, USA). Ion maps were corrected for detector dead time (44 ns) with the openMIMS plugin and the  ${}^{13}C/{}^{12}C$  ratio maps were corrected for the mass 280 bias as a function of the  ${}^{13}C/{}^{12}C$  ratio measured on microalgae not enriched in  ${}^{13}C$ . The  ${}^{13}C/{}^{12}C$ 281 ratio was measured in regions of interest (ROI) defined on pyrenoid and starch granules of six 282 control cells and six Cd-exposed cells as detailed in Figure S1. The isotopic enrichment of <sup>13</sup>C 283 was expressed as its relative abundance  $\delta^{13}$ C compared to the <sup>13</sup>C natural isotopic abundance 284 285 and calculated using the following formula:

$$\delta^{13}C = \frac{R_{\text{sample}}}{R_{\text{nat}}} - 1 \tag{1}$$

12

where  $R_{sample}$  is the measured  ${}^{13}C/{}^{12}C$  ratio in the ROI of the cells and  $R_{nat}$  is the natural carbon isotope ratio.

The origin of carbon in starch granules from either acetate or  $CO_2$  as carbon source was determined by data processing adapted from Terrado et al. (2017). First, the fractional abundance  $F_{\text{starch}}$  of the carbon isotope <sup>13</sup>C in starch granules of the sum of both carbon isotopes is given as (in percentage):

293 
$$F_{\text{starch}} = \frac{{}^{13}\text{C}}{{}^{13}\text{C} + {}^{12}\text{C}} \times 100$$
 (2)

which can also be expressed as

295 
$$F_{\text{starch}} = \frac{R_{\text{starch}}}{1 + R_{\text{starch}}} \times 100$$
(3)

where  $R_{starch}$  is the <sup>13</sup>C/<sup>12</sup>C ratio measured in the ROI within starch granules. Then, the origin of starch carbon was determined using the following formulas:

298 
$$F_{\text{starch}} = F_{\text{acetate}} \times f_{\text{acetate}} + F_{\text{CO2}} \times f_{\text{CO2}}$$
(4)

299 with: 
$$f_{\text{acetate}} + f_{\text{CO2}} = 1$$
 (5)

300 
$$f_{\text{acetate}} = \frac{F_{\text{starch}} - F_{\text{CO2}}}{F_{\text{acetate}} - F_{\text{CO2}}} \times 100 \text{ and } f_{\text{CO2}} = \frac{F_{\text{starch}} - F_{\text{acetate}}}{F_{\text{CO2}} - F_{\text{acetate}}} \times 100$$
(6)

301 The formula (6) results from (4) and (5) where  $F_{acetate}$  and  $F_{CO2}$  are the fractional abundance of 302 <sup>13</sup>C in the carbon sources acetate (99%) and CO<sub>2</sub> (1.09%), respectively, and  $f_{acetate}$  and  $f_{CO2}$ 303 represent the percentage of the origin of carbon within the ROI (starch granule) from the 304 respective source.

### 305 Samples preparation for synchrotron-based techniques

306	Microalgae from a control and a Cd-exposed (70 $\mu$ M) culture were harvested after 72 h of
307	inoculation by centrifugation at $3000 \times g$ for 3 min. They were rapidly washed in ultrapure
308	water three times to remove metals and organic compounds weakly adsorbed on the cell wall.
309	Two sample preparations were used.

310 For µXRF and Cd L<sub>III</sub>-edge µXANES, droplets of microalgae suspension were deposited on 4 311 µm-thick Ultralene film (SPEX sample prep Metuche, NJ). After decantation for 10 min, 312 microalgae were frozen into liquid nitrogen-chilled isopentane (-160 °C).

313 For Cd L<sub>III</sub>-edge XANES bulk analysis, the algae pellet was frozen in liquid nitrogen, 314 homogenized and pressed as a 5 mm frozen pellet. All the samples were kept frozen at -80 °C 315 until measurements and transferred to the sample stage in their hydrated frozen state.

316

#### 317 **µXRF** imaging

318 Measurements were performed on the ID21 beamline at the European Synchrotron Radiation 319 Facility (ESRF, Grenoble, France) equipped with a fixed exit Si(111) two-crystal 320 monochromator (Cotte et al., 2017). The X-ray photons were focused by a KB mirror system 321 providing a sub-micron resolution of 0.6  $\mu$ m (H)  $\times$  0.3  $\mu$ m (V) FWHM on the sample. The Xray fluorescence signal was recorded with a large 100 mm<sup>2</sup> SDD detector (Bruker) while the 322 323 samples were scanned to obtain elemental maps. All the measurements were carried out under 324 cryogenic conditions at -160 °C using a liquid N<sub>2</sub> cryostat to limit beam radiation damage. The 325 fluorescence signal was normalized by the incident photon intensity  $(I_0)$  measured with a 326 photodiode. Phosphorus, sulfur, chloride, and cadmium maps were recorded using an incident 327 energy of 3570 eV, below the absorption edge of potassium, while potassium and calcium maps 328 were recorded at 4100 eV as described in Isaure et al. (2006). The fluorescence signal was

deconvoluted from fluorescence background and fluorescence elemental overlapping to obtain
elemental maps using the PYMCA software (Solé, Papillon, Cotte, Walter & Susini, 2007).

331 Cd LIII-edge µXANES and XANES. µXANES spectra were recorded with the same lateral 332 resolution as for  $\mu XRF$  in fluorescence mode on points of interest visualized on the  $\mu XRF$ 333 maps. For Cd model compounds and bulk samples, XANES spectra were recorded in 334 fluorescence mode with a defocused beam (100  $\mu$ m  $\times$  100  $\mu$ m). Measurements were also done 335 under cryogenic conditions to limit radiation damage and speciation change. µXANES and 336 XANES spectra were collected in the energy range 3520-3590 eV. Approximately 30 scans 337 were calibrated with a metallic Cd foil, averaged, and normalized according to standard 338 methods using ATHENA software (Ravel & Newville, 2005). Then, experimental normalized 339 spectra were compared to a library of standard spectra previously collected (Huguet et al., 340 2012; Isaure et al., 2006, 2015; Penen et al., 2017). A fingerprint approach was used to simulate 341 the unknown spectra by linear combination fits (LCFs) of Cd model compounds as described 342 by Isaure et al. (2006). The quality of the fits was estimated by the normalized sum-squares 343 residuals NSS =  $\Sigma$ (Xanes<sub>experimental</sub>-Xanes<sub>fit</sub>)<sup>2</sup>/ $\Sigma$ (Xanes<sub>experimental</sub>)<sup>2</sup>×100, in the 3520-3580 for 344 µXANES spectra and 3520-3590 eV range for XANES spectra. Linear combination fits with 345 one, two and three components were tested and the combination with n+1 components was 346 retained if the NSS parameter decreased more that 20% in comparison to the fit with n 347 components (Figure S2). The uncertainty of the proportion of each compound was estimated 348 to 10% (Isaure et al., 2015).

### 349 Statistics

350 One-way ANOVA followed by a post-hoc Tuckey test Statistical analyses were performed on 351 growth rate, chlorophyll concentration, starch concentration and <sup>13</sup>C/<sup>12</sup>C ratio measured on 352 NanoSIMS maps to identify significant differences among the different conditions. Results were considered significant for a p < 0.05. Kruskal-Wallis one-way analysis of variance on ranks (p<0.05) was performed when the condition of homogeneity of variances was not respected.

## 356 **RESULTS**

### 357 Availability of cadmium and essential trace elements in TAPEDDHA medium

In a first approach, cadmium availability in TAP<sub>EDDHA</sub> medium, i.e. free Cd<sup>2+</sup> concentration, was simulated using Visual Minteq software (Table 1). Concentrations of 10, 20, 30, 40, 50 and 70  $\mu$ M of CdCl<sub>2</sub> added to the medium correspond to free ionic Cd concentrations of 1, 4, 7, 10, 13 and 20  $\mu$ M. Cd availability was thus increased by 10 compared to the classic TAP medium for the higher Cd concentration.

The free ion concentrations of several essential trace elements in the TAP<sub>EDDHA</sub> medium were markedly different from that in the original TAP medium (Table 1). Most important, the concentration of free  $Zn^{2+}$  increased by 10000 and of free  $Cu^{2+}$  by 100 whereas  $Co^{2+}$  availability decreased by 10000 compared to the classic TAP medium.

### 367 Cadmium exposure leads to starch accumulation

C. reinhardtii was exposed to a range of 10-70 µM CdCl<sub>2</sub> for 48 h in TAP<sub>EDDHA</sub> medium and 368 369 relative growth rates were measured in order to determine Cd efficient concentrations (EC<sub>x</sub>) 370 with the Hill model (Figure 1A). Up to 20 µM Cd, no significant effect on the algae growth was observed (EC<sub>5</sub> = 22  $\mu$ M Cd [15–31  $\mu$ M Cd Cl<sub>95%</sub>]) but increasing Cd concentrations 371 372 induced a strong decrease of the relative growth rate (EC<sub>50</sub> = 74  $\mu$ M Cd [67–86  $\mu$ M Cd Cl<sub>95%</sub>]). 373 In order to investigate the impact of cadmium on C. reinhardtii in stress conditions, the Cd exposure concentration of 70  $\mu$ M (i.e. 20  $\mu$ M free Cd<sup>2+</sup>) was applied in this study since it was 374 375 the nearest to  $EC_{50}$  value. These concentrations can seem quite high compared to other studies

focused on Cd concentrations in the 10-100 nM range (Stoiber et al. 2012, Lavoie et al. 2012)
. However, contrary to these studies, the present experiments were carried out in mixotrophic
conditions, which likely modified the cell response. Our observations support that under
mixotrophic conditions *C. reinhardtii* was more resistant against Cd stress.

When exposed to 70  $\mu$ M Cd, the intracellular cadmium concentration was 1530  $\pm$  132  $\mu$ g g<sup>-1</sup> 380 381 (dry weight). The global impact of this cadmium stress on C. reinhardtii vitality was examined 382 by the determination of growth rate, chlorophyll concentration and starch concentration (Figure 383 1B, C and D). Compared to the control condition, Cd induced a decrease of the growth rate 384 (from  $0.76 \pm 0.05$  to  $0.39 \pm 0.06$  d<sup>-1</sup>) and the chlorophyll concentration (from  $8.3 \pm 0.4$  to 6.4 $\pm$  0.3 µg mL<sup>-1</sup> OD<sub>730nm</sub><sup>-1</sup>) while starch concentration was increased (from 47.6  $\pm$  4.7 to 87.9  $\pm$ 385 13.2 mg  $g^{-1}$ ). These results showed an important stress in *C. reinhardtii* when it exposed to 70 386 387 µM Cd leading to an impaired photosynthesis and an accumulation of starch. Carbon 388 metabolism in C. reinhardtii was thus strongly impacted by Cd toxicity.

### 389 Cadmium sequestration in pyrenoid and vacuoles

390 *C. reinhardtii* cells exposed to 70  $\mu$ M Cd for 48 h were examined by TEM/X-EDS and 391 NanoSIMS (Figure 2). Figure 2 A and B show that cell structure was altered by Cd exposure, 392 particularly the chloroplast where the pyrenoid and its starch shell were destructured. The 393 starch plaques were also more abundant in the Cd-exposed cells corroborating the increase of 394 starch measured by enzymatic digestion.

Elemental composition of organelles observed on TEM micrographs (Figure 2C, F) was determined on their corresponding NanoSIMS elemental maps (Figure 2E, H). Vacuoles were rich in calcium, phosphorus and sulfur. Pyrenoids were characterized by a composition rich in sulfur and nitrogen, which could be attributed to the high concentration in proteins (RuBisCO in particular) inside this organelle. Moreover, pyrenoids were surrounded by a halo of 400 phosphorus which did not necessarily correspond to the starch shell. Finally, starch shells and401 starch plates were rich in nitrogen.

402 Additional X-EDS analysis (Figure 2D, G) revealed that cadmium concentration was high in 403 the pyrenoid. This pyrenoidal sequestration of Cd could thus be responsible for Cd toxicity in 404 *C. reinhardtii*. Furthermore, Cd was localized beside Ca and P in electron dense bodies in 405 vacuoles suggesting a cadmium sequestration in calcium polyphosphate granules as confirmed 406 in our previous study by  $\mu$ XANES analysis (Penen et al., 2017). It is important to note that the 407 elemental composition of vacuoles and pyrenoid determined by NanoSIMS was confirmed by 408 the X-EDS measurements.

### 409 Cadmium localized in pyrenoid is preferentially bound to sulfur ligands

Elemental  $\mu$ XRF maps (Figure 3A) of *C. reinhardtii* cells exposed to 70  $\mu$ M Cd for 48 h corroborated the elemental distribution obtained by TEM/X-EDS and NanoSIMS analysis (Figure 3A). A sulfur spot was present in each cell which corresponded to the pyrenoid. Although cadmium was detected at low level in the whole cells, it was highly concentrated in one spot in each cell co-localized with sulfur, confirming its pyrenoidal sequestration. In addition, phosphorus, calcium, and to a lesser extent cadmium were co-localized in spots of 1  $\mu$ m size representing Ca polyphosphate granules in vacuoles.

In combination with  $\mu$ XRF,  $\mu$ XANES gave information about Cd binding ligands in areas of interest evidenced by  $\mu$ XRF. In contrast, XANES collected in bulk samples allowed to determine the average *in situ* cadmium binding in the microalgae. Cd L<sub>III</sub>-edge XANES was demonstrated powerful to distinguish sulfur from oxygen ligands due to absence of a peak at 3539 eV in sulfur binding atoms (Isaure et al., 2006). Moreover, as indicated by arrows in Cd reference XANES spectra in figures 3B and 3C, the second oscillation was shifted to higher energy values for Cd-acetate and Cd-malate where Cd is bound to carboxyl groups compared to Cd bound to phosphate groups in Cd-phosphate and Cd-phytate (Penen et al., 2017). This enabled a differentiation between the oxygen donor ligands in Cd-carboxyl and Cd-phosphate. In contrast, Cd-phytochelatin 2 (Cd-PC<sub>2</sub>), Cd-glutathione (Cd-GSH) and Cd-cysteine spectra had similar pattern, thus hampering the exact determination of the nature of sulfur donor ligands; however, they can be encompassed as thiol groups. Therefore, the distribution of Cd ligands, as results of linear combination fits (LCFs) of Cd L<sub>III</sub>-edge XANES and  $\mu$ XANES spectra, was described as ratios of Cd-S, Cd-O-P, and Cd-O-C compounds (Figure 3D).

In the S and Cd-enriched areas attributed to the pyrenoid,  $\mu$ XANES analysis revealed that Cd was mainly coordinated by thiol ligands (67 ± 10% and 56 ± 10% for the two presented pyrenoids), but also by carboxyl groups (33 ± 10% and 44 ± 10%). No phosphate ligand was identified in these spots. A pyrenoid consists principally of proteins, mainly RuBisCO, thus  $\mu$ XANES results could indicate cadmium binding to protein thiol groups. Finally, Cd bound to carboxyl groups could be attributed to Cd sequestered to the starch shell around the pyrenoid since  $\mu$ XANES here probed a three-dimensional volume.

438 XANES measurements of bulk samples showed that Cd- thiol ligands in algae accounted for 439  $31 \pm 10\%$  only while Cd- carboxyl groups amounted to  $25 \pm 10\%$ . The main part of Cd was 440 associated with phosphate ligands (44 ± 10%) suggesting that the calcium polyphosphate 441 granules sequestered an important part of the metal in *C. reinhardtii*. Indeed, these granules 442 were previously probed by µXANES and revealed the sequestration of Cd by phosphate ligands 443 (Penen et al., 2017).

### 444 Cadmium promotes carbon incorporation from acetate in starch

445 In order to investigate the impact of Cd on carbon fixation and to distinguish inorganic and 446 organic carbon sources of *C. reinhardtii*, i.e.  $CO_2$  and acetate, <sup>13</sup>C labeled acetate was employed for tracer experiments. *C. reinhardtii* control cells and cells exposed to 70  $\mu$ M Cd for 48 h in TAP<sub>EDDHA</sub> medium containing <sup>13</sup>C labeled acetate were examined by NanoSIMS (Figure 4). <sup>13</sup>C/<sup>12</sup>C ratio maps highlight intracellular areas of carbon resulting from acetate assimilation (high <sup>13</sup>C/<sup>12</sup>C ratio) (Figure 4A and B).

Two different subcellular regions of interest (ROI) were selected in <sup>13</sup>C/<sup>12</sup>C ratio maps (Figure 451 S1) to investigate carbon assimilation: the pyrenoid as organelle of CO<sub>2</sub> fixation and the starch 452 453 plates as a final storage form of carbon in C. reinhardtii. Based on data from ROI in 6 individual control cells and 6 exposed cells (Figure S1), the <sup>13</sup>C isotopic enrichment  $\delta^{13}$ C was calculated 454 455 as represented in Figure 4C. Hence, <sup>13</sup>C enrichment was found in the pyrenoid, but it did not change significantly under Cd exposure. In contrast, starch plates showed a 5-fold higher <sup>13</sup>C 456 457 enrichment under Cd exposure than in unexposed cells. In particular, the study of carbon 458 isotopic distribution in starch as the final carbon storage form allowed the quantification of 459 carbon assimilation from CO<sub>2</sub> and acetate sources.

460 The determination of  $f_{\text{acetate}}$  and  $f_{\text{CO2}}$  in starch granules revealed that, under control conditions, 73% of starch carbon came from CO<sub>2</sub> (<sup>12</sup>C) and 27% from acetate (<sup>13</sup>C) (RSD  $\pm$  9%). In 461 contrast, Cd stress seemed to disturb CO<sub>2</sub> (<sup>12</sup>C) assimilation which decreased to 39% while 462 acetate ( $^{13}$ C) assimilation contributed for 61% of starch carbon (Figure 4D) (RSD  $\pm$  13%). 463 464 Moreover, when the isotopic distribution of starch carbon was combined with the starch 465 concentration previously analyzed (Figure 1D), it appeared that overall  $CO_2$  carbon ( $^{12}C$ ) assimilation remained limited (1.28  $\pm$  0.18 mmol (<sup>12</sup>C) g<sup>-1</sup> in control condition; 1.26  $\pm$  0.43 466 mmol (<sup>12</sup>C) g<sup>-1</sup> under Cd exposure) (Figure 4E) relative to the highly increased starch 467 468 biosynthesis under Cd exposure. In contrast, acetate carbon (<sup>13</sup>C) assimilation was about 4-fold higher during Cd exposure (0.48  $\pm$  0.15 mmol (<sup>13</sup>C) g<sup>-1</sup> under control conditions; 1.99  $\pm$  0.47 469 mmol  $({}^{13}C)$  g<sup>-1</sup> under Cd exposure). These results suggest that Cd exposure interfered with CO<sub>2</sub> 470

471 assimilation due to the Cd fixation in the pyrenoid possibly due to the Cd binding to protein472 thiol groups.

## 473 **DISCUSSION**

### 474 Effect of the culture medium on cell vitality

475 C. reinhardtii was cultivated in a modified version of TAP medium where EDTA was replaced 476 with EDDHA (TAP<sub>EDDHA</sub>) in order to increase Cd availability and, at the same time, Cd 477 toxicity. However, under control conditions, this modification induced already a decrease of 478 cell vitality. Indeed, in our previous work (Penen et al., 2017), C. reinhardtii grown in classic TAP medium showed a higher growth rate  $(1.15 \pm 0.03 \text{ d}^{-1})$  and a higher production of 479 chlorophyll (10.78  $\pm$  0.38 µg ml<sup>-1</sup> OD<sub>730nm</sub><sup>-1</sup>) compared to the present study where microalgae 480 481 have been cultivated in TAP<sub>EDDHA</sub> medium (Figure 1B, C). These effects of TAP<sub>EDDHA</sub> medium 482 on cell vitality in control conditions could be explained by changes in the availability of some 483 trace elements compared to the classic TAP medium (Table 1). Indeed, higher availability 484 mainly of zinc but also, to a lesser extent, of copper, and in addition deficiency of cobalt was 485 found in the control TAPEDDHA medium.

### 486 Cell response to cadmium stress

Cell vitality results showed that a 70 μM Cd exposure for 48 h induced a decrease of the chlorophyll concentration, which is a well-known sign of photosynthesis impairment (Nagel & Voigt, 1989; Collard & Matagne, 1990; Nagel & Voigt, 1995; Prasad, Drej, Skawińska & Strałka, 1998). The degradation of the chlorophyll pool was concomitant with a growth decrease and an increase of carbon storage in the form of starch. Usually, this stress reaction has been described in *C. reinhardtii* during nitrogen (Ball et al., 1990; Cakmak et al., 2012), phosphorus (Ball et al., 1990) and sulfur (Ball et al., 1990; Cakmak et al., 2012) starvation.

494 Moreover, Juergens et al. (2015, 2016) showed that starch was accumulated to maintain a 495 carbon pool during N-starvation instead of delaying photoinhibition and oxidative stress due 496 to energetic overflows from photosynthesis. In case of metallic stress, starch storage was 497 reported in Chlamydomonas acidophila (Nishikawa, Yamakoshi, Uemura & Tominaga, 2003) 498 exposed to 20 µM Cd where the size of starch plates observed microscopically increased 2.43-499 fold. Metal stress could thus induce, in an indirect way, symptoms similar to those of nutrient 500 starvation. Working with C. reinhardtii, Bräutigam et al. (2011) showed that Cd stress induced 501 the synthesis of thiol polypeptides like phytochelatins, resulting in a decrease in the 502 intracellular concentration of its precursors like glutathione,  $\gamma$ -EC and cysteine leading 503 presumably to a sulfur starvation. Although, sulfur supply by the medium should be sufficient 504 (Bräutigam et al., 2011), an intracellular sulfur deficiency is conceivable. Furthermore, a 505 proteomic study (Gillet, Decottignies, Chardonnet & Maréchal, 2006) revealed that enzymes 506 involved in sulfur metabolism were modified under Cd stress in order to promote the synthesis 507 of phytochelatins.

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### 510 Cadmium is bound to RuBisCO in pyrenoid and to phosphate in acidocalcisome

For the first time, a pyrenoidal sequestration of Cd and its binding to sulfur ligands were clearly observed in *C. reinhardtii*. In addition, high sulfur and nitrogen content indicated a high protein concentration in the pyrenoid. A model of the impact of Cd sequestration on carbon fixation in C. reinhardtii is proposed in Figure 5. Pyrenoids are essentially composed of the protein RuBisCO (Borkhsenious, Mason & Moroney, 1998). They are the place of the carbon fixation from CO<sub>2</sub> via the Calvin cycle (Harris, 1989). Beside RuBisCO, the enzyme RuBisCO-activase is involved in this process (McKay, Gibbs & Vaughn, 1991). Thus, it can be hypothesized that 518 Cd binding to protein thiol groups interferes with CO<sub>2</sub> fixation in *C. reinhardtii*, for example 519 due to conformational changes caused by cadmium-thiol bonds. Indeed, it has been reported 520 that modifications at cysteines 449 and 459 of the RuBisCO large subunit induce a 521 conformational change and a decrease of its carboxylase activity (Marín-Navarro & Moreno, 522 2006). Moreover, the RuBisCO large subunit protein is down-regulated under Cd exposure 523 (Gillet et al., 2006). For higher plants, it has been shown that Cd impairs the CO<sub>2</sub> assimilation 524 rate by RuBisCO activity decrease, e.g. in the crop Lactuca sativa (Dias et al., 2012) and in 525 Nicotiana rustica (Afef, Leila, Donia, Houda & Chiraz, 2011).

In addition, TEM/X-EDS,  $\mu$ XRF and Cd L<sub>III</sub>-edge XANES measurements have shown that Cd was also coordinated by phosphate ligands and localized in 0.5-1  $\mu$ m sized vacuoles containing phosphorus and calcium. In our previous work (Penen et al., 2017), cadmium was shown to be localized in vacuoles which were compared to acidocalcisomes due to their specific elemental composition in phosphorus and calcium (Goodenough, Heiss, Roth, Rusch & Lee, 2019; Hong-Hermesdorf et al., 2014; Komine, Eggink, Park & Hoober, 2000; Komine, Park, Wolfe & Hoober, 1996; Ruiz et al., 2001).

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### 534 CO<sub>2</sub> fixation is impaired by cadmium stress

535 Cadmium has been shown to disturb the light-dependent part of the photosynthesis (Faller et 536 al., 2005) but no study has previously focused on the impact of Cd on carbon assimilation. *C*. 537 *reinhardtii* is able to grow photoautotrophically using CO<sub>2</sub>, heterotrophically using acetate in 538 the dark and mixotrophically using both carbon sources in the light. In the present study, 539 mixotrophic growth conditions under constant illumination induced carbon storage as starch 540 (Figure 1D). Moreover, NanoSIMS analysis revealed that, carbon from acetate was 541 accumulated in the pyrenoid and in starch granules (Figure 4B). The pathways involved in acetate assimilation as well as the impact of cadmium on these pathways are discussed in thenext paragraphs and are represented in Figure 5.

544 The accumulation of carbon from acetate in the pyrenoid can be explained by the assimilation 545 of acetate through the tricarboxylic acid cycle (TCA). Indeed, acetate assimilation starts with 546 its incorporation into acetyl coenzyme A (acetyl-CoA), either directly converted by acetyl-CoA 547 synthetase, or in two steps with the successive action of acetate kinase and phosphate 548 acetyltransferase. Acetyl-CoA is used as substrate in the TCA cycle in mitochondria. The TCA 549 cycle produces ATP as well as NADH and releases the carbon of acetyl-CoA as CO<sub>2</sub> (Johnson 550 & Alric, 2013). Then, the mitochondrial carbonic anhydrase CaH4/5 (Raven, 2001) converts 551  $CO_2$  in bicarbonate which is carried to the pyrenoid by the  $CO_2$  concentrating mechanism 552 (CCM) (Meyer & Griffiths, 2013). This latter is a combination of active carbon transporters 553 (Burow, Chen, Mouton & Moroney, 1996; Im & Grossman, 2002; Miura et al., 2004; Pollock, 554 Prout, Godfrey, Lemaire & Moroney, 2004), carbonic anhydrases (Moroney et al., 2011) 555 specific to cellular compartments and the soluble proteins (LCIB/LCIC) (Yamano et al., 2010) 556 allowing CO<sub>2</sub> trapping inside the pyrenoid. In the present work, the isotopic ratio of carbon 557 (<sup>13</sup>C enrichment) inside the pyrenoid was similar under control and exposure conditions (Figure 558 4C) showing that cadmium did not seem to disturb TCA cycle and CCM.

Starch biosynthesis involves the successive actions of ADP-glucose pyrophosphorylase (AGPP) and starch synthase (SS) on the final product of the gluconeogenesis pathway (glucose-1-phosphate (G1P)). Carbon from acetate enters gluconeogenesis in two different ways, once incorporated into AcetylCoA. On the one hand, as mentioned in the previous paragraph, AcetylCoA feeds the TCA cycle releasing CO<sub>2</sub>. In the chloroplast, the two carbon sources thus share the CO<sub>2</sub> fixation step by RuBisCO into the Calvin cycle pathway. The final product of the Calvin cycle (glyceraldehyde-3-phosphate (G3P)) can enter the gluconeogenesis (Johnson & Alric, 2013). On the other hand, Acetyl-CoA feeds the glyoxylate cycle which is a TCA cycle where CO<sub>2</sub> releasing steps are by-passed (Johnson & Alric, 2012, 2013) in the cytosol. Indeed, using AcetylCoA and isocitrate as substrates, isocitrate lyase produces glyoxylate and succinate. This latter is successively converted in mitochondria to fumarate, malate, oxaloacetate then phosphoenolpyruvate (PEP) which is exported to cytosol and enters the gluconeogenesis pathway.

*C. reinhardtii* is able to assimilate inorganic carbon via PEP carboxylation by the two PEP carboxylases CrPpc1/CrPpc2, leading to the production of oxaloacetate (Mamedov, Moellering & Chollet, 2005). However, the algal PEP carboxylase activity is mainly anaplerotic and not photosynthetic (Giordano, Norici, Forssen, Eriksson & Raven, 2003). Although the impact of cadmium on PEPC activity is not established in *C. reinhardtii*, it has been shown that Cd inhibits PEPC activity in higher plants like Miscanthus species (Guo et al., 2016) and Zea mays (Wang, Zhao, Liu, Zhou, & Jin, 2009).

579 Despite <sup>13</sup>C-[1,2] labeling of acetate, our experiments do not allow us to distinguish the 580 pathways taken by acetate carbons to be finally incorporated in starch. However, unlabeled 581 carbon assimilation from CO<sub>2</sub> can only pass by the Calvin cycle. In consequence, after a 48 h Cd exposure, the relative decrease in <sup>12</sup>C from 73 to 39% of starch carbon (Figure 4D) shows 582 583 that carbon fixation through Calvin cycle pathway is limited by Cd stress but not completely 584 inhibited. Hence, the high increase of starch biosynthesis under Cd stress (Figure 1D) is 585 achieved in mixotrophic conditions by an about 4-fold higher carbon assimilation from acetate. 586 However, the overall amount of carbon assimilation from CO<sub>2</sub> into starch remains limited in 587 relation to the increase of starch production (Figure 4E). Moreover, the decrease in chlorophyll 588 concentration (figure 1C) coupled with a massive accumulation of Cd in the pyrenoid (Figures

2D, G and 3A) seems to disturb not only the light-dependent reactions (Faller et al., 2005) but
also the light-independent reactions of photosynthesis.

591 In unexposed C. reinhardtii grown under mixotrophic conditions, 78% of total carbon (Heifetz 592 et al., 2000) has been found to be from  $CO_2$  assimilation which is similar to the 73% of starch 593 carbon originating from CO<sub>2</sub> found in this work. Although Cd exposure and nutrient starvation 594 have a similar impact on starch accumulation, the limitation of CO<sub>2</sub> fixation is specific to Cd 595 stress, since during nitrogen starvation, 65% of starch carbon (Juergens et al., 2016) as well as 596 80% of total carbon (Juergens et al., 2016) are taken from an inorganic carbon source. In 597 addition, enzymes involved in Calvin cycle have been shown to be down-regulated under Cd 598 stress (phosphoglycerate kinase, ribose-5phosphate isomerase) while enzymes involved in 599 acetate assimilation were up-regulated as for the isocitrate lyase (Gillet et al., 2006) involved 600 in the glyoxylate cycle or the phosphoglycerate mutase (Gillet et al., 2006) taking part in 601 gluconeogenesis.

Using NanoSIMS in combination with <sup>13</sup>C-labeled acetate it was possible to detect, localize, and quantify <sup>13</sup>C-labeled starch as end product within the cells. Although these are important results, the limitation of this approach is, that between the acetate substrate and the end product no other metabolites of the carbon metabolism pathway during cadmium stress were measured. A metabolomic or a fluxomic approach would be necessary to complete our results with regard to the overall carbon fixation and metabolism pathway involved during cadmium stress in *C*. *reinhardtii*.

In conclusion, the present work shows that cadmium clearly impairs carbon assimilation in *C. reinhardtii* grown in mixotrophic conditions. Due to the pyrenoidal sequestration of Cd, CO<sub>2</sub> fixation is limited during Cd exposure whereas acetate assimilation is favored. Mixotrophy could thus allow phytoplankton surviving toxic metal pollution events. Although mixotrophy 613 seems to render aquatic ecosystems more resistant, the alteration of  $CO_2$  fixation by toxic 614 metals could contribute to the unbalance of the biogeochemical carbon cycle.

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## 628 CONFLICT OF INTEREST

629 The authors declare no conflict of interests.

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- 990 *Cell Physiology*, *51*(9), 1453–1468.

#### **TABLES**

F	ree ion conce	entrations (I	VI)	
		Free ion concentrations (M)		
TAP		TAPEDDHA		
Control	70 µM Cd	Control	70 µM Co	
2.5 10 <sup>-5</sup>	2.6 10 <sup>-5</sup>	2.6 10 <sup>-5</sup>	2.6 10 <sup>-5</sup>	
3.1 10 <sup>-4</sup>	3.1 10 <sup>-4</sup>	3.1 10 <sup>-4</sup>	3.1 10 <sup>-4</sup>	
5.6 10 <sup>-9</sup>	5.0 10 <sup>-7</sup>	-	-	
-	-	1.1 10 <sup>-22</sup>	1.2 10 <sup>-22</sup>	
2.4 10 <sup>-10</sup>	3.0 10 <sup>-6</sup>	3.1 10 <sup>-6</sup>	3.1 10 <sup>-6</sup>	
6.5 10 <sup>-14</sup>	6.1 10 <sup>-10</sup>	6.9 10 <sup>-12</sup>	7.5 10 <sup>-12</sup>	
1.5 10 <sup>-11</sup>	2.0 10 <sup>-7</sup>	5.6 10 <sup>-15</sup>	4.7 10 <sup>-15</sup>	
10.7 10 <sup>-10</sup>	8.0 10 <sup>-10</sup>	8.3 10 <sup>-10</sup>	8.4 10 <sup>-10</sup>	
5.6 10 <sup>-6</sup>	1.1 10 <sup>-8</sup>	5.6 10 <sup>-6</sup>	1.1 10 <sup>-9</sup>	
-	2.1 10 <sup>-6</sup>	-	2.0 10 <sup>-5</sup>	
	Control 2.5 10 <sup>-5</sup> 3.1 10 <sup>-4</sup> 5.6 10 <sup>-9</sup> - 2.4 10 <sup>-10</sup> 6.5 10 <sup>-14</sup> 1.5 10 <sup>-11</sup> 10.7 10 <sup>-10</sup>	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	$\begin{array}{c c c c c c c c } \hline Control & 70 \ \mu M \ Cd & Control \\ \hline 2.5 \ 10^{-5} & 2.6 \ 10^{-5} & 2.6 \ 10^{-5} \\ \hline 3.1 \ 10^{-4} & 3.1 \ 10^{-4} & 3.1 \ 10^{-4} \\ \hline 5.6 \ 10^{-9} & 5.0 \ 10^{-7} & - \\ \hline & & & & & & \\ \hline & & & & & & \\ \hline & & & &$	

**Table 1**: Comparison of ion availability in classic TAP medium and TAP<sub>EDDHA</sub> medium.

# 1000 FIGURE CAPTIONS

**Figure 1**: Impact of cadmium exposure to *C. reinardtii* grown in TAP<sub>EDDHA</sub> medium. (A) 1002 Relative growth rate was expressed as the ratio  $\mu/\mu 0$  determined during the exponential phase

1003 of growth after addition of 0, 10, 20, 30, 40, 50 and 70  $\mu$ M of CdCl<sub>2</sub>. (n=3 ± sd).  $\mu$  = growth 1004 rate during cadmium exposure,  $\mu_0$  = growth rate in control conditions. Black curve represents 1005 the dose-response effect calculated with the Hill model. Impact of a 70  $\mu$ M Cd exposure on 1006 growth (B) and on chlorophyll concentration (C), and starch concentration (D) after a 48 h 1007 exposure. Significant differences with respect to the control condition at a significance level of 1008 0.05 (p<0.05) are indicated by an asterisk (\*).

1009 Figure 2: Structure and elemental composition of *C. reinhardtii* cells exposed to cadmium. (A) 1010 Electron micrographs of two C. reinhardtii cells in control condition and (B) of two cells 1011 exposed to 70  $\mu$ M Cd for 48 h, scale bar = 1  $\mu$ m. Abbreviations: N, nucleus; P, pyrenoid; S, 1012 starch; Th, thylakoid; CW, cell wall. (C to H) Correlative TEM/X-EDS and NanoSIMS 1013 imaging of two cadmium exposed C. reinhardtii cells: (C and F) Electron micrographs of C. 1014 reinhardtii cells exposed to 70 µM Cd for 48 h, white arrows show the area analyzed by X-1015 EDS analysis, scale bar = 1  $\mu$ m. Abbreviations: N, nucleus; P, pyrenoid; S, starch; Th, 1016 thylakoid; V, vacuole; CV, contractile vacuole; CW, cellwall. (E and H) Corresponding NanoSIMS elemental maps obtained by  $Cs^+$  ion source ( ${}^{12}C^{14}N^-$  for N detection,  ${}^{32}S^-$ ) and O<sup>-</sup> 1017 RF plasma ion source  $({}^{31}P^+, {}^{40}Ca^+)$ , 10×10 µm<sup>2</sup> (E) and 12×12 µm<sup>2</sup> (H) fields of 256×256 1018 1019 pixels, scale bar =  $2 \mu m$ . (D and G) X-EDS spectra of vacuoles and pyrenoid located on electron 1020 micrographs, copper signal is produced by the copper grid

**Figure 3:** In situ cadmium localization and speciation in *C. reinhardtii* exposed to 70  $\mu$ M Cd for 48 in TAP<sub>EDDHA</sub> medium. (A) False color  $\mu$ -XRF elemental maps of Cd, S, P and Ca, arrows show points of interest where  $\mu$ -XANES analyses were performed, scale bar = 2  $\mu$ m. Step size = 0.4  $\mu$ m, dwell-time = 300 ms/pixel at 3570 eV for S, P, Cd and at 4100 eV for Ca. (B) Two or three components fits (dotted line) of Cd L<sub>III</sub>-edge  $\mu$ -XANES spectra (solid line) of points of interest marked by arrows and of bulk sample; (C) Derivative of Cd L<sub>III</sub>-edge  $\mu$ XANES; (D) 1027 Distribution of Cd ligands after normalization of the percentages to 100%. The uncertainty is 1028 estimated to  $\pm 10\%$ .

**Figure 4:** Carbon assimilation in *C. reinhardtii* grown in TAP<sub>EDDHA</sub> medium (labeled <sup>13</sup>C-[1,2] 1029 1030 acetate). (A and B)  ${}^{13}C/{}^{12}C$  isotope ratio images obtained by NanoSIMS using Cs<sup>+</sup> source. The 1031 analyzed fields of 256×256 pixels were, from left to right, 12×12 µm<sup>2</sup>, 8×8 µm<sup>2</sup>, 10×10 µm<sup>2</sup> 1032 for control conditions (A) and  $12 \times 12 \ \mu m^2$ ,  $10 \times 10 \ \mu m^2$ ,  $10 \times 10 \ \mu m^2$  for cadmium exposure 1033 conditions (70  $\mu$ M Cd for 48 h) (B), scale bar = 2  $\mu$ m, ROI and annotations for pyrenoid and starch plates are shown in Figure S1. (C) Impact of 70 µM Cd exposure for 48 h on <sup>13</sup>C isotopic 1034 1035 enrichment in the pyrenoid and in starch plates ( $n=6 \pm sd$ ). Isotopic distribution of carbon in 1036 starch plates obtained from  $f_{\text{acetate}}$  and  $f_{\text{CO2}}$  (D) and in starch carbon equivalent (E), (n = 6 ± sd). 1037 Significant differences with respect to the control condition according to Kruskal-Wallis one-1038 way analysis of variance on ranks (p<0.05) are indicated by an asterisk (\*).

Figure 5: Proposed model of cadmium impact on the assimilation of carbon in 1039 1040 Chlamydomonas reinhardtii in mixotrophic conditions. The description of the general 1041 metabolic pathways for carbon assimilation from organic (acetate) and inorganic (CO<sub>2</sub>) sources 1042 are adapted from Johnson & Alric (2013). Under exposure conditions, cadmium is highly 1043 sequestered in the pyrenoid by sulfur ligands, impairing the Calvin cycle. Thus, CO<sub>2</sub> and 1044 acetate assimilation involving the Calvin cycle is limited (thin grey arrows) whereas acetate 1045 assimilation through glyoxylate cycle is favored (thick blue arrows). In addition, the 1046 anaplerotic PEPC activity could hypothetically participate in inorganic carbon fixation. Starch 1047 is accumulated in the chloroplast to prevent cellular troubles due to Cd stress. Finally, Cd is 1048 also sequestered in vacuoles, coordinated by phosphate ligands. Abbreviations: SDH, Succinate dehydrogenase; FUM, Fumarase; MDH, Malate Dehydrogenase; PCK, 1049

1050 Phosphoenolpyruvate Carboxykinase; *PEPC*, Phosphoenolpyruvate carboxylase; *CAH*,
1051 Carbonic anhydrase.

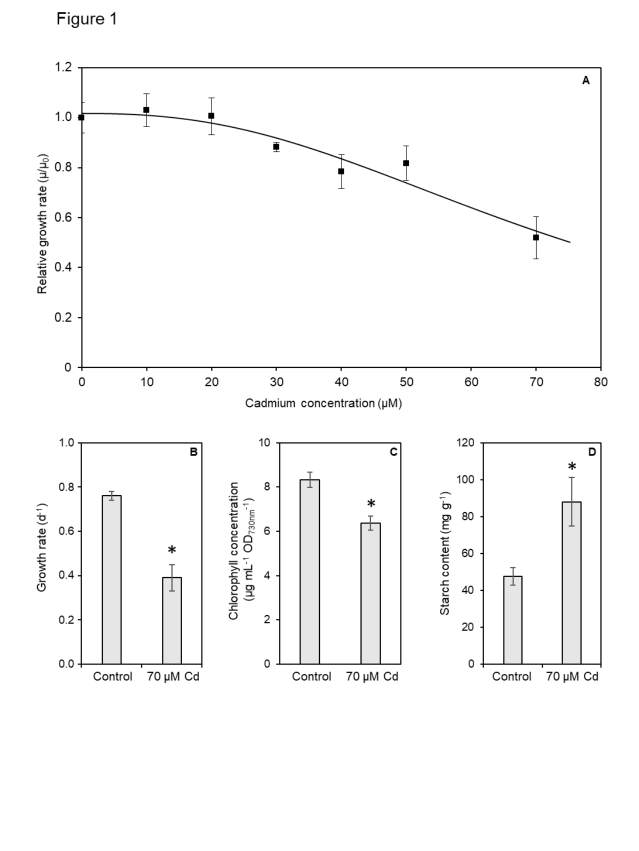
#### 1053 SUPPORTING INFORMATION

1054 Supporting information is available at the publisher's web-site. Supporting information is given

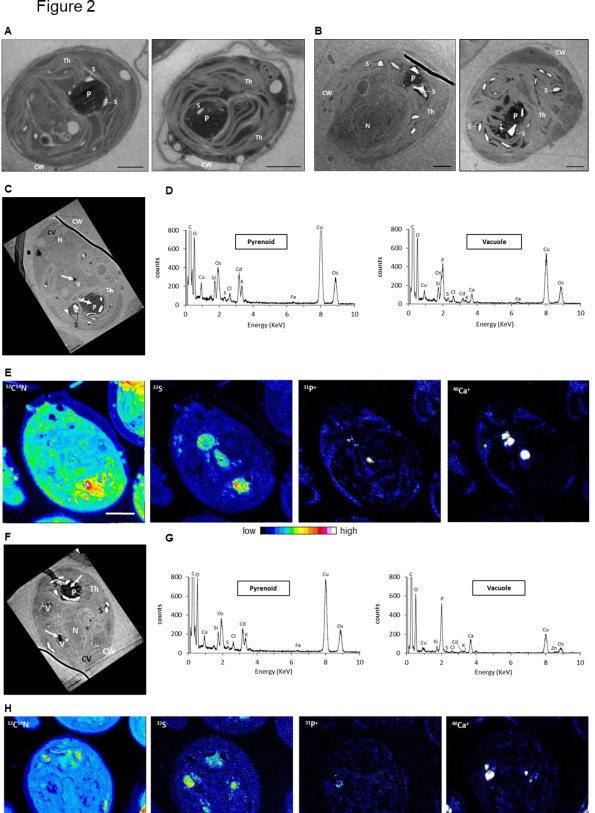
1055 to the composition of the TAP<sub>EDDHA</sub> medium (**Table S1**), to the definition of regions of interest

1056 (ROI) in NanoSIMS <sup>13</sup>C/<sup>12</sup>C images for data analysis (**Figure S1**), and to linear combination

- 1057 fits (LCFs) of XANES spectra (Figure S2).







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