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The singular properties of photosynthetic cytochrome c_{550} from the diatom *Phaeodactylum tricornutum* suggest new alternative functions

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Cytochrome c_{550} is an extrinsic component in the luminal side of photosystem II (PSII) in cyanobacteria, as well as in eukaryotic algae from the red photosynthetic lineage including, among others, diatoms. We have established that cytochrome c_{550} from the diatom *Phaeodactylum tricornutum* can be obtained as a complete protein from the membrane fraction of the alga, although a C-terminal truncated form is purified from the soluble fractions of this diatom as well as from other eukaryotic algae. Eukaryotic cytochromes c_{550} show distinctive electrostatic features as compared with cyanobacterial cytochrome c_{550} . In addition, co-immunoseparation and mass spectrometry experiments, as well as immunoelectron microscopy analyses, indicate that although cytochrome c_{550} from *P. tricornutum* is mainly located in the thylakoid domain of the chloroplast – where it interacts with PSII – , it can also be found in the chloroplast pyrenoid, related with proteins linked to the CO₂ concentrating mechanism and assimilation. These results thus suggest new alternative functions of this heme protein in eukaryotes.

Introduction

Photosynthetic cytochrome c_{550} (Cc₅₅₀) is a c-type heme protein (molecular mass around 15 kDa, encoded by the PSBV gene) with an unusual bis-histidinyl axial coordination (Frazão et al. 2001) that acts as an extrinsic subunit of photosystem II (PSII) by binding stoichiometrically to the luminal surface of this photosynthetic complex (Shen 2015, Ago et al. 2016). In spite of its redox character, the role of Cc_{550} in PSII appears to be just structural, by stabilizing the Mn_4CaO_5 cluster and binding to Cl^- and Ca^{2+} ions, as a redox function of the Cc_{550} heme cofactor in PSII has not yet been

established (Shen and Inoue 1993, Shen et al. 1998, Enami et al. 2003, Nagao et al. 2010). Cc_{550} is present in cyanobacteria and in eukaryotic algae from the red photosynthetic lineage, which comprises, among others, red algae and diatoms, but is absent in the green lineage, which includes green algae and plants, that have replaced Cc_{550} for the Ca^{2+} binding PsbP subunit, which lacks any cofactor (revised in Roncel et al. 2012).

In many organisms, Cc_{550} can be easily purified as a soluble protein (Navarro et al. 1995, Kerfeld and Krogmann 1998, Bernal-Bayard et al. 2017), and the existence of two different populations of Cc_{550} (bound to the PSII or free in the lumen) has been postulated (Kirilovsky

Abbreviations – CA, carbonic anhydrase; Cc_{550} , cytochrome c_{550} ; Cc_6 , cytochrome c_6 ; CCM, CO_2 -concentrating mechanism; DSS, disuccinimidyl suberate; FBA, fructose-bisphosphate aldolase; PSI, photosystem I; PSII, photosystem II; RubisCO, ribulose-1,5-bisphosphate carboxylase/oxygenase.

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et al. 2004). Soluble Cc_{550} has a very low midpoint redox potential (ranging from -190 to $-314\,\mathrm{mV}$; Alam et al. 1984, Navarro et al. 1995, Roncel et al. 2003, Bernal-Bayard et al. 2017) incompatible with a redox function in PSII. However, much more positive potential values were determined when bound to PSII (from -80 to $+200\,\mathrm{mV}$; Roncel et al. 2003, Guerrero et al. 2011). Several alternative roles have been proposed for the soluble Cc_{550} in cyanobacteria, as in anaerobic carbon and hydrogen metabolism (Krogmann 1991, Morand et al. 1994), cyclic photophosphorylation (Kienzel and Peschek 1983) or in the reduction of nitrate to ammonium (Alam et al. 1984), but none of these possible functions have been clearly established.

Diatoms belong to the red lineage of algae that diverged from the green lineage that finally evolved to higher plants. In addition to Cc_{550} (or PSBV), the assembly of extrinsic proteins at the lumenal side of diatom PSII includes the cyanobacterial-like subunits PsbO and PsbU, as well as the PsbQ' subunit, also present in red algae (Enami et al. 2003, Nagao et al. 2010), and an additional extrinsic protein, named as Psb31 (Okumura et al. 2008, Nagao et al. 2017). However, although the structure of diatom PSII has not yet been solved, the crystal structure of PSII from the red alga *Cyanidium caldarium* has shown an overall structure similar to the cyanobacterial complex, including the position of Cc_{550} (Ago et al. 2016).

In order to optimize photosynthetic efficiency, the chloroplasts of diatoms display a refined thylakoid architecture (Flori et al. 2017), which includes the presence of the pyrenoid, an electron-dense semicrystalline protein aggregate present in the chloroplast of most unicellular eukaryotic algae. It is considered that CO₂ fixation occurs in the pyrenoid, where the enzyme ribulose-1,5-bisphosphate carboxylase/oxygenase (RubisCO) is preferentially localized, accounting for over 90% of the pyrenoid's protein content (reviewed in Badger et al. 1998, Meyer et al. 2017). In addition, the presence of carbonic anhydrase (CA) enzymes in the pyrenoid has been reported (Tachibana et al. 2011, Sinetova et al. 2012, Hopkinson et al. 2016, Kikutani et al. 2016). Thus, the role of the pyrenoid is to sustain efficient carbon fixation by increasing the CO₂ concentration [CO₂ concentrating mechanism (CCM)] in the vicinity of RubisCO (Giordano et al. 2005, Matsuda et al. 2017). Moreover, in most unicellular eukaryotic algae, including diatoms, the pyrenoid is penetrated by one or more thylakoid lamellae (Bedoshvili et al. 2009, Engel et al. 2015, Meyer et al. 2017). In the marine diatom Phaeodactylum tricornutum, in particular, the pyrenoid matrix is crossed by a single thylakoid lamella that contains a luminal θ -type CA, that adds to the two ß-type CAs, PtCA1 and PtCA2, located inside the pyrenoid (Tachibana et al. 2011, Kikutani et al. 2016). Components of the photosystem I (PSI) complex have also been found in pyrenoids, indicating that intrapyrenoid thylakoids could supply ATP to this compartment via cyclic electron transport, thus covering the energetic demands of carbon fixation at the same time as avoiding the evolution of inhibitory O2 (McKay and Gibbs 1991, Meyer and Griffiths 2013). Consequently, it was generally accepted that intrapyrenoid thylakoids are enriched in PSI and lack PSII, although in the green alga Chlamydomonas reinhardtii, the presence of PSII components has been recently reported (Mackinder et al. 2017). However, the presence in this space of either the soluble photosynthetic cytochrome c_6 (Cc₆) or the Cc_{550} , has not been yet addressed.

Recently, the photosynthetic Cc₅₅₀ from *P. tricornu*tum has been purified from algal cells and characterized (Bernal-Bayard et al. 2017). The purified protein was described as being truncated in the last hydrophobic residues of the C-terminus. Moreover, a weaker affinity of Cc₅₅₀ for the diatom PSII complex was also described (Bernal-Bayard et al. 2017). Here, we provide additional insights into the peculiar properties of Cc₅₅₀ from the diatom P. tricornutum. First, we have determined that it is possible to purify Cc₅₅₀ as the complete non-truncated form. In addition, P. tricornutum Cc₅₅₀ has been immunolocalized in the pyrenoid of the chloroplast, in close contact with proteins previously described as located in this microcompartment. Thus, in P. tricornutum, the Cc₅₅₀ could play a role related to the pyrenoid function that should yet be defined.

Materials and methods

Cell cultures

Cells of the coastal pennate diatom *P. tricornutum* CCAP 1055/1 were grown in artificial seawater medium (12 µM Fe, iron-replete culture; McLachlan 1964, Goldman and McCarthy 1978) in a rotatory shaker (50 rpm) at 20°C, with regular transfer of the cells into fresh media. The cultures were illuminated by white led lamps giving an intensity of 4.35 W m⁻² (T8-150MWBL led lamps; Wellmax, Shanghai, China) under a light/dark cycle of 16/8 h. *Chaetoceros muelleri, Nannochloropsis gaditana* and *Isochrysis galbana* cells were obtained from indoor cultures from the Centro Público Integrado de Formación Profesional Marítimo Zaporito (San Fernando, Spain).

Protein purification

Purification of truncated Cc_{550} from *P. tricornutum*, *C. muelleri*, *N. gaditana* and *I. galbana* cells was carried

out essentially as recently described in Bernal-Bayard et al. (2017), in the presence of the protease inhibitors phenylmethylsulfonyl fluoride (PMSF) (1 mM), benzamidine (1 mM), aminocaproic acid (100 mM), D-phenylalanine (10 mM) and hydrocinnamic acid (1 mM). Purification of the complete Cc₅₅₀ from *P. tricor*nutum membrane fractions was performed essentially following the protocol previously described to obtain PSII-enriched samples (Bernal-Bayard et al. 2017) with some modifications. Basically, P. tricornutum cells were disrupted three times in a French press at 15 000 psi, the \(\begin{aligned} \text{-dodecyl-maltoside} \) (\(\beta - DM \) incubation was done for 45 min, and the solubilized samples were loaded onto a 0.17–0.3 M sucrose continuous density gradient. Finally, the complete Cc₅₅₀ was collected from the top gradient band, corresponding to the free protein (Bernal-Bayard et al. 2017). For the MS analysis, samples were exhaustively washed and concentrated by ultrafiltration with 5 mM Tris-HCl buffer, pH 8.5.

Protein samples for in-gel peptide fingerprint analysis of P. tricornutum Cc_{550} were obtained from crude cell extracts resolved on polyacrylamide gel electrophoresis. Fresh P. tricornutum cells were washed twice in ice-cold PBS 20 mM buffer (pH 8.0), and resuspended in 25 mM Tris—HCl (pH 8.5), 50 mM NaCl buffer, supplemented with 0.5% Triton X-100, DNase and the protease inhibitors PMSF, benzamidine and aminocaproic acid. Samples were incubated for 5 min at 70° C followed by a cycle of French-press disruption (20 000 psi). Samples were then centrifuged at $170\,000\,g$ for $15\,\text{min}$, and the resultant supernatant was resolved on 20% (w/v) sodium dodecyl sulfate-polyacrylamide gel electrophoresis and visualized by Coomassie blue staining.

For the co-immunoseparation and immunodetection of Cc_{550} , polyclonal antibodies raised against this cytochrome (Bernal-Bayard et al. 2017) were purified by their affinity to pure Cc_{550} , by using 4 mg of Cc_{550} (Bernal-Bayard et al. 2017) and 2 ml of the AminoLink Plus Immobilization Kit (Thermo Scientific, Waltham, MA) at pH7.2, according to the instructions of the supplier. Antibodies against Cc_6 (Roncel et al. 2016) and the large and small plant RubisCO subunits (Agrisera, Vännäs, Sweden) were also used in control experiments. For immunodetection of *P. tricornutum* RubisCO subunits, initial crude extracts were in some cases enriched in RubisCO by precipitation with PEG 4000 (20% w/v), as previously described (Haslam et al. 2005).

DNA analysis

Genomic DNA extraction was carried out using a simplified CTAB-extraction procedure (Lukowitz et al. 2000). Total extracted DNA from *C. muelleri* and *I. galbana* was

used as template to amplify by polymerase chain reaction (PCR) the corresponding PSBV genes, using adequate oligonucleotide pairs (Fig. S1, Supporting Information). Thus, whereas C. muelleri gene amplification was completed by using oligonucleotide primers designed from the Chaetoceros gracilis sequence (C1-F and C2-R), I. galbana PSBV gene amplification and sequencing was done in two steps. First, using degenerate oligonucleotide primers designed from a consensus sequence from C. gracilis, P. tricornutum and the five closest PSBV genes (according to a BLAST search), which allowed the amplification and sequencing of the 3'-end (242 pb), that showed a 83% identity with a similar sequence of the PSBV gene from the Isochrysidal alga Emiliania huxleyi. Thus, a second PCR amplification was carried out to obtain the *PSBV* 5'-end by using oligonucleotide primers designed from E. huxleyi. In all cases, the PCR fragments were cloned for subsequent sequencing in the pGEM-T vector (Promega, Madison, WI) according to the protocol of the supplier. DNA sequencing was carried out in the sequencing service of the company Secugen (Madrid, Spain).

Co-immunoseparation analysis

different samples were used co-immunoseparation experiments. First, fresh P. tricornutum cells, resuspended in 25 mM Tris-HCl (pH 7.5), 50 mM NaCl and 0.1% Triton X-100, supplemented with DNase and the protease inhibitors PMSF, benzamidine and aminocaproic acid, were disrupted in a French press at 20 000 psi (three cycles). Samples were then centrifuged at 170 000 g for 30 min, and the resultant supernatant was considered as the soluble fraction. Second, P. tricornutum cells resuspended in 50 mM Tris-HCl (pH 7.5), 50 mM NaCl and 5 mM EDTA (buffer A) supplemented with proteases inhibitors, DNase and 1 M betaine, were disrupted at a lower pressure (7500 psi, two cycles). Unbroken cells were separated by centrifugation at 5000 g for 5 min and the supernatant was centrifuged at 170 000 g for 30 min. The resultant pellet was resuspended in buffer A at 1 mg Chl ml⁻¹ and later diluted to 0.5 mg Chl ml⁻¹ with the same volume of ß-DM 3% (w/v) prepared in buffer A. The solution was then incubated 30 min in the dark at 4°C under gentle stirring followed by centrifugation at 170 000 g for 30 min. The resulting detergent-solubilized supernatant was diluted with buffer A to a ß-DM concentration of 0.1% and concentrated in an Amicon pressure cell (10 K cutoff filter; Merck Millipore, Darmstadt, Germany). The final concentrated solution was considered as the membrane-extracted fraction. Finally, crosslinked samples, with the crosslinking reagent disuccinimidyl

suberate (DSS; Thermo Scientific), were obtained from fresh P. tricornutum cells by following the instructions of the supplier. Cells were resuspended in PBS 20 mM (pH 8.0) buffer and washed three times with ice-cold PBS (pH 8.0) to remove amine-containing agents from the culture media. Cells were then treated with DSS to 5 mM final concentration, followed by 30 min incubation at room temperature. Crosslinking reactions were stopped by the addition of Quench Solution (1 M Tris-HCl, pH 7.5) to a final concentration of 20 mM Tris, followed by 15 min incubation at room temperature. Samples were then centrifuged at 5000 g for 5 min and pellets were resuspended in 20 mM Tris-HCl (pH 7.5), 50 mM NaCl and 5 mM EDTA buffer, supplemented with protease inhibitors and DNase, and disrupted in a French press at 20 000 psi (three cycles). Unbroken cells were discarded by centrifugation, and cell lysates were diluted to 1 mg Chl ml⁻¹ and treated with \(\beta - DM \) as described previously for the membrane-extracted fraction. The resulting final concentrated supernatant was considered as the crosslinked fraction. In all cases, the protein content of the samples was quantified using the Lowry method, whereas chlorophyll concentration was determined according to Arnon (1949).

Immunopurifications using μ columns and μ MACS separator (Miltenyi Biotec, Bergisch Gladbach, Germany) were carried out following the instructions of the supplier. An amount of 5 mg of protein mixed with 5 μ l of purified polyclonal antibodies against Cc₅₅₀ were used, and the eluted fraction was analyzed by liquid chromatography tandem-mass spectrometry (LC-MS/MS). Control immunoseparation experiments, using the preimmune serum, were also carried out and the identified proteins were subtracted from those obtained when using the immune serum.

MS analysis

Matrix-assisted laser desorption/ionization time-of-flight MS (MALDI-TOF MS) and LC-MS/MS analyses were performed at the Instituto de Bioquímica Vegetal y Fotosíntesis Proteomic Service (Sevilla, Spain). Tryptic digestion of in-gel *P. tricornutum* proteins for peptide fingerprint analysis and MALDI-TOF MS were carried out as previously described (Bernal-Bayard et al. 2017). Tryptic digestion in solution of immunoseparated samples, as well as peptide analysis by LC and MS, were performed basically as previously described (Vowinckel et al. 2013) on a Tandem Quadrupole TOF MS (AB Sciex TripleTOF 5600) coupled to a NanoSpray III Ion Source (AB Sciex) and nano-HPLC (Eksigent ekspert nanoLC 425). Peptide separation was first carried out on a precolumn (Acclaim PepMap 100 C18, 5 μm, 100 Å, 100 μm

id \times 20 mm, Thermo Fisher Scientific) and then eluted onto the analytical column (New Objective PicoFrit column, 75 µm id \times 250 mm, emitter included and packed with ReproSil-Pur 3 µm). Data acquisition was achieved as previously described (Vowinckel et al. 2013). The ion accumulation time was set to 250 ms (MS) and to 65 ms (MS/MS), resulting in a total duty cycle of 2.89 s.

LC-MS/MS data acquired in DDA mode were analyzed and processed basically as described in Vowinckel et al. (2013), by using the Paragon algorithm (Protein-Pilot Software, AB Sciex, v. 5.0.1) and the reference proteome of *P. tricornutum* in the UniProt database of protein sequences. AB Sciex contaminants and rabbit proteins from the UniProt database were discarded. Only peptides with a confidence score > 0.05 were considered for further analysis.

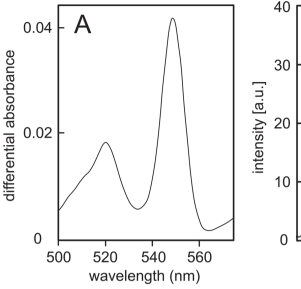
Immunoelectron microscopy

Phaeodactylum tricornutum cells pellets were fixed in 2% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.3, for 2 h at 4°C, washed several times in the same buffer at 4°C and dehydrated through a graded ethanol series. Finally, cells were embedded in LR White medium grade resin (Sigma-Aldrich, Saint Louis, MO) as described in Lichtlé et al. (1992).

For immunolocalization, paled gold sections were cut with an ultramicrotome (Reichert-Jung Ultracut, Wien, Austria) with a diamond knife and mounted on nickel grids. Ultrathin sections were treated with the rabbit purified antiserum anti- Cc_{550} (Bernal-Bayard et al. 2017) for 1 h, and the secondary antibody (goat anti-rabbit 5-nm gold particles, GAR G5 EM; Janssen Life Sciences, Beerse, Belgium) was used at a dilution of 1:100 for 1 h. Finally, the grids were contrasted with 2% aqueous uranyl acetate for 8 min and observed with a Zeiss Libra 120 Electron Microscope at 80 kV. Two controls were performed: (1) replacing the primary antibody by a drop of PBS buffer; and (2) incubation with the primary antibody anti- Cc_6 (luminal soluble carrier) at a dilution of 1:1000 for 1 h.

Structural models

Structures of Cc₅₅₀ from *C. muelleri, N. gaditana* and *I. galbana* were modeled using the SWISS MODEL Workspace platform (https://swissmodel.expasy.org/interactive; Guex and Peitsch 1997), using as templates the crystal structures of Cc₅₅₀ from the cyanobacterium *Synechocystis* sp. PCC 6803 (pdb 1E29) and the red alga *C. caldarium* (pdb 4YUU). The representation of protein surface electrostatic potentials was performed by using the Swiss-Pdb Viewer program (https://spdbv



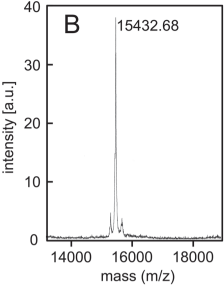


Fig. 1. (A) Differential absorption spectra between reduced and oxidized Cc₅₅₀ (via dithionite or ascorbate treatments) from *Phaeodactylum tricornutum* extracted by solubilization from membrane fraction. The concentration of cytochrome was 3 μM. (B) Molecular weight MS analysis of the complete holocytochrome.

.vital-it.ch/). Although interactions of the amino acid side chains – that can change their pKas and therefore their charge in solution – are not taken into account, this program is a useful tool for comparative purposes.

Data deposition

The MS proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD008763. The *PSBV* gene sequences from *C. muelleri* and *I. galbana* are deposited in the NCBI databank, GenBank accession numbers MG779498 and MG779497, respectively.

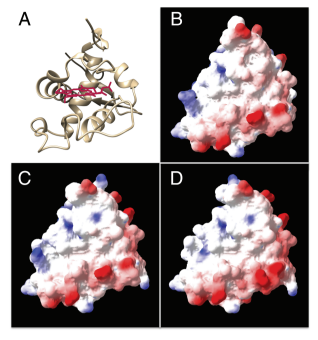
Results and discussion

In a previous paper, we showed that P. tricornutum Cc550 is purified from the soluble fraction as a truncated protein in the two last tyrosine residues of the C-terminus (Bernal-Bayard et al. 2017). Similar results were obtained (data not shown) even though the purification was carried out in the presence of broad-spectrum proteases inhibitors (5 mM EDTA and 1 mM PMSF), as well as specific inhibitors of either serine proteases (1 mM benzamidine and 100 mM aminocaproic acid) or carboxypeptidases competitive inhibitors (10 mM D-phenylalanine and 1 mM hydrocinnamic acid; Elkins-Kaufman and Neurath 1949, Hartsuck and Lipscomb 1971). In this work, we have, however, developed a protocol for the purification of the complete protein by detergent solubilization from the membrane fraction and the further isolation of the free protein from the top part of sucrose gradients (Fig. 1A). Thus,

an MALDI-TOF MW analysis of the solubilized Cc_{550} showed an MW (approximately 15 433 Da) that fits with the expected MW based on the *PSBV* gene sequence (15 438 Da; Fig. 1B). The occurrence of the complete holoprotein was also confirmed by tryptic digestion and MS peptide fingerprint analysis (data not shown).

A procedure was designed to check if the soluble or the membrane-associated Cc₅₅₀ correspond to different physiological forms (a soluble truncated protein and a complete protein bound to PSII), and not to the exposition to cell proteases in the soluble fraction during the purification course. Phaeodactylum tricornutum crude cell extracts were heated at 70°C and directly resolved on polyacrylamide gel electrophoresis, in order to inactivate possible proteolytic enzymes and shorten the process of proteins separation. Extracted gel spots in the MW range corresponding to Cc550 were analyzed by tryptic digestion and MS peptide fingerprint (Fig. S2). The fingerprint peptide analysis accurately covered 100% of the amino acid sequence corresponding to the complete non-truncated protein, whereas the theoretical peptide fingerprint for the truncated protein could not be correctly fitted to the obtained data (Fig. S2), which indicates that truncation is a non-physiological process. Remarkably, the enzymatic activity responsible of Cc₅₅₀ truncation seems to be widespread among several lines of the red lineage of eukaryotic algae, as truncated Cc₅₅₀ was purified from C. muelleri (a marine centric diatom), N. gaditana (Eustigmatophyte) and I. galbana (Haptophyte, Isochrysidales). However, this activity seems to be absent in cyanobacteria, as Cc₅₅₀ from Synechocystis sp. PCC 6803, used as a control, showed the expected MW for the complete protein (data not shown).

Fig. 2. (A) Backbone model of Cc_{550} from *Chaetoceros muelleri* obtained using the crystal structures of Cc_{550} from the cyanobacterium *Synechocystis* sp. PCC 6803 (PDB entry 1E29) and the red alga *Cyanidium caldarium* (PDB 4YUU) as main templates. (B–D) Surface electrostatic potential distribution of the structural models of Cc_{550} from (B) *C. muelleri*, (C) *Nannochloropsis gaditana* and (D) *Isochrysis galbana*. The view displays the heme groups in the same orientation as in (A), showing in front the cofactor exposed area, and in the top the protein C-terminal hydrophobic protuberance. Simulations of surface electrostatic potential distribution were performed using the Swiss-PDB Viewer Program assuming an ionic strength of 500 mM at pH 7.0. Positively and negatively charged regions are depicted in blue and red. respectively.



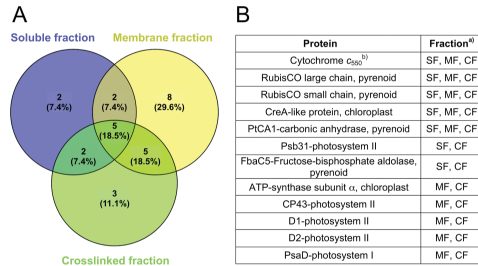


Fig. 3. Potential Cc₅₅₀ interacting proteins as identified by co-immunoseparation and LC–MS/MS in soluble, membrane and crosslinked fractions of the diatom *Phaeodactylum tricornutum*. (A) Venn diagram of the Cc₅₅₀-protein interaction dataset showing intersections of the three co-immunoseparated fractions. (B) Chloroplastic protein targets selected when appearing both in the crosslinked fraction and at least in one of the other two fractions (soluble or membrane). ^{a)} SF, co-immunoseparated in the soluble fraction; MF, co-immunoseparated in the membrane fraction; CF, co-immunoseparated in the crosslinked fraction. ^{b)} Cc₅₅₀ was identified in all the samples.

In the case of *C. muelleri* and *I. galbana*, previous sequencing of their *PSBV* genes was required in order to establish the MW of both complete proteins. The *C. muelleri PSBV* gene showed only 11 nucleotide variations as compared with the gene of *C. gracilis*, but these resulted only in two amino acid changes in the protein transit peptide (data not shown). The *I. galbana PSBV* gene showed 83% identity with the equivalent gene of the also Isochrysidal alga *E. huxleyi*, and the protein

alignment (94% identity) is shown in Fig. S1. Modeled structures of *C. muelleri*, *I. galbana* and *N. gaditana* Cc_{550} were obtained from the available sequences (Fig. 2). As previously described in the case of the *P. tricornutum* protein, the three Cc_{550} show a common folding similar to that described in cyanobacteria and red algae (Fig. 2) and conserve the hydrophobic northern finger (according to the orientation presented in Fig. 2) previously described (Frazão et al. 2001). However,

the surface of eukaryotic Cc₅₅₀ also shows exclusive electrostatic features as compared with cyanobacterial Cc₅₅₀. Thus, whereas in the prokaryotic protein the cofactor exposed area holds a negatively charged electrostatic character (Frazão et al. 2001, Bernal-Bayard et al. 2017), in the eukaryotic Cc₅₅₀, the area around the heme group is mainly hydrophobic, and the negative electrostatic potential is restricted to the southern area, opposite to the hydrophobic northern protuberance (Fig. 2). According to the PSII known structures, the facing surface of Cc_{550} (Fig. 2), that includes the cofactor exposed area, is involved in the binding to the photosystem, maintaining close contacts with the PSII surface (Ago et al. 2016). Consequently, the distinctive surface charge distribution of eukaryotic Cc550 could be significant when establishing the binding affinity to PSII, as previously suggested for the *P. tricornutum* Cc₅₅₀ (Bernal-Bayard et al. 2017).

Although Cc₅₅₀ can be obtained from soluble cell extracts in different organisms (Evans and Krogmann 1983, Navarro et al. 1995, Kerfeld and Krogmann 1998, Bernal-Bayard et al. 2017), this fact is particularly significant in P. tricornutum, where about 60-85% of total Cc550 is solubilized during the process of cell disruption in the absence of added detergents (Bernal-Bayard et al. 2017). A similar result has been here observed during the purification of this protein from C. muelleri, N. gaditana and I. galbana cells (data not shown). These results could be justified by a combination of a weaker affinity for PSII and an enhanced PSII turnover, as previously suggested in diatoms (Lavaud et al. 2016), both resulting in a higher fraction of unbound Cc₅₅₀. In this sense, most of the protein still attached to membrane fractions in P. tricornutum has been shown to be released by relatively weak detergent extraction procedures, thus suggesting a comparatively weaker affinity of Cc₅₅₀ for PSII (Bernal-Bayard et al. 2017). A less intense affinity for PSII may open the possibility of new functions for an increased fraction of unbound Cc₅₅₀. We have explored this possibility by performing a co-immunoseparation (from the endogenous Cc₅₅₀ already present in the different samples) and MS analysis, in order to identify possible novel protein interactors of P. tricornutum Cc₅₅₀. Three different cellular fractions were immunoseparated with purified Cc₅₅₀-specific antibodies and analyzed by LC-MS/MS: (1) the supernatant obtained after cell disruption at high pressure in a non-osmotically stabilized medium followed by sample ultracentrifugation (soluble fraction); (2) the detergent-extracted fraction obtained from membrane pellets after cell breaking at low pressure in an osmotically stabilized medium (membrane fraction) and (3) the supernatant obtained after DSS treatment of whole

cells followed by cell disruption and ultracentrifugation in the presence of detergent (crosslinked fraction).

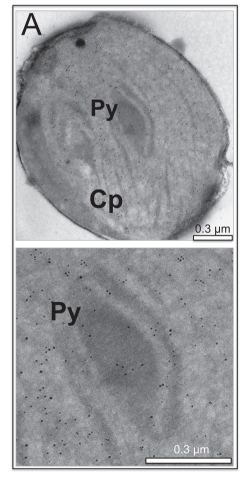
A relatively low number of proteins were detected by LC-MS/MS in the three co-immunoseparated samples, more particularly in the crosslinked fraction (data available via ProteomeXchange identifier PXD008763). Moreover, in order to minimize the occurrence of false positive or redundant interactions, LC-MS/MS data were filtered according to the following criteria: (1) possible protein partners have to be described as chloroplastic proteins and/or predicted to have a transit peptide targeting to this organelle; and (2) undefined predicted proteins, ribosomal proteins and fucoxanthin-chlorophyll light-harvesting antenna proteins were discarded. From these criteria, a preliminary list of potential targets of Cc₅₅₀ was initially obtained (Figs 3A and S3), which includes RubisCO. Although this enzyme is a highly abundant chloroplast protein complex in plants, it has been shown that RubisCO has a much lower abundance in microalgae, representing less than 6% of total protein (Losh et al. 2013). In addition, the use of the purified Cc₅₅₀-specific antibodies in western-blot experiments with cell extracts showed a major recognition of Cc₅₅₀, and no bands corresponding to the predominant components of the pyrenoid (i.e. RubisCO subunits or CA enzymes) were detected (Fig. S4). Thus, we consider RubisCO as a reliable co-immunoseparated target of P. tricornutum Cc₅₅₀.

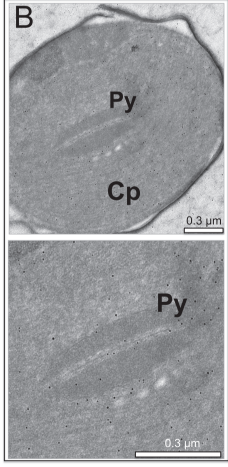
Co-immunoseparation in the soluble or membrane fractions does not firmly demonstrate a specific physiological interaction, since the method involves putting together proteins that are actually in different cellular compartments, which can lead to describing artefactual interactions. Thus, protein crosslinking in whole cells has been used to demonstrate close contacts between Cc₅₅₀ and other proteins as well as co-localization in the same subcellular compartment. Ultimately, validated protein targets where selected as appearing both in the crosslinked fraction and at least in one of the other two (soluble or membrane) fractions (Fig. 3A). This restrictive criterion discarded two PSII-associated proteins that appeared both in the soluble and membrane fractions, but not in the crosslinked one (Fig. S3), as they are the PSII extrinsic PsbQ' subunit and a FK506-binding protein with peptidylprolyl isomerase activity, that in plants is related to the PSII assembly (Gollan et al. 2012). The final list of co-immunoseparated proteins fitting the restrictive stablished criteria is shown in Fig. 3B. It is important to note that most proteins that co-immunoseparated in the crosslinked fraction have been already annotated as being located in the chloroplast (Figs 3 and S3), validating the reliability of the method used. In addition to several PSII subunits (Psb31,

CP43, D1 and D2), expected from the Cc₅₅₀ location in the red algal PSII structure (Ago et al. 2016), only a very limited number of proteins co-immunoseparated in the three types of samples analyzed (Fig. 3), including RubisCO (large and small subunits) and PtCA1 (a ß-CA), both located in the pyrenoid compartment (Satoh et al. 2001, Tachibana et al. 2011, Kikutani et al. 2016). Only another protein of unknown function, classified as a CreA-like protein, also co-immunoseparated in the three samples. The CreA family is a group of carbon metabolism transcription regulators described in other organisms (Ries et al. 2016). However, although distantly related to CreA proteins, this CreA-like protein of *P. tricornutum* lacks the typical sequence signatures corresponding to zinc fingers for DNA-binding, typically found in canonical CreA proteins, and its transit peptide indicates a chloroplast targeting. Moreover, a sequence-based structural analysis predicted the existence of a transmembrane domain in this protein (data not shown). On the other hand, another pyrenoid protein, the fructose-bisphosphate aldolase (FBA; Allen et al. 2012), was detected both in the crosslinked and soluble fractions (Fig. 3B). Finally, two proteins were detected both in the crosslinked and membrane fractions: the PSI PsaD subunit and the ATP-synthase α subunit, and these two results can be attributed to non-specific interactions of luminal Cc_{550} with the two thylakoid membrane complexes (ATP-synthase and PSI) to which these subunits belong. Actually, other ATP-synthase and PSI subunits were detected, but only in the immunoseparated membrane fractions (Fig. S3). Nevertheless, all the potential new targets of Cc_{550} here identified are located in the pyrenoid (RubisCO, PtCA1, FBA) or may be related to carbon metabolism (CreA-like protein).

To confirm the possible presence of Cc_{550} in the pyrenoid we have carried out an immunoelectron microscopy analysis of the location of Cc_{550} in *P. tricornutum* cells, by using purified polyclonal antibodies specific against this protein. As a control, the location of the luminal soluble Cc_6 carrier has been also studied. Fig. 4 shows the electron microscopy results for the immunodetection experiments. As expected, Cc_6 seems to be located both in the stromal and the intrapyrenoid thylakoids but not in the pyrenoid matrix

Fig. 4. Immunoelectron microscopy images of *Phaeodactylum tricornutum* cells (upper panel) showing localization of (A) photosynthetic Cc_{550} , and (B) the soluble luminal Cc_6 . The lower panel shows the expansion of the pyrenoid area. Cp, chloroplast; Py, pyrenoid.





(Fig. 4B), in agreement with its functional association with PSI. However, although Cc_{550} showed the expected chloroplastic localization, it appears not only in the stromal and intrapyrenoid thylakoids, but also in the pyrenoid matrix (Fig. 4A). Thus, the association of Cc_{550} with the pyrenoid in *P. tricornutum* is supported both by co-immunoseparation and immunoelectron microscopy analysis.

It is important to note that our results do not imply that Cc₅₅₀ functionally interacts with all the pyrenoid targets included in Fig. 3 (RubisCO, PtCA1 or FBA), as possible cross interactions between these proteins can result in indirect co-immunoseparations with Cc₅₅₀. However, our results strongly support the pyrenoid localization of Cc₅₅₀, confirmed by both the in vivo crosslinking with pyrenoid proteins and the pyrenoid immunolocalization (Figs 3 and 4). The alternative could be the existence of cross reactions between the purified antibodies against Cc₅₅₀ with any of the observed pyrenoid targets, which can result in parallel immunoseparations and immunodetections. However, we consider that this possibility can be ruled out, as western blot experiments have not shown signs of the occurrence of such cross reactions (Fig. S4).

The association of Cc₅₅₀ with the pyrenoid has not been previously reported and undoubtedly represents an intriguing result. Eukaryotic Cc550 is a chloroplast-encoded protein that has a transit peptide, similar to those from cyanobacteria, for targeting into the thylakoid lumen, and therefore the interaction with RubisCO (or other pyrenoid proteins) that is located in the stromal compartment is difficult to justify. In addition, PSII activity has been initially described to be restricted to stromal thylakoids (McKay and Gibbs 1991). However, very recently, the presence of both intrinsic and extrinsic PSII subunits in the pyrenoid of the green alga C. reinhardtii has been reported (Mackinder et al. 2017, Zhan et al. 2018), including the localization of the PsbQ extrinsic component of PSII in intrapyrenoid thylakoids (Mackinder et al. 2017). Thus, the presence of the PSII extrinsic PSBV subunit (Cc₅₅₀) in this compartment could not be totally unexpected.

We only can speculate in order to explain a location of Cc₅₅₀ in the pyrenoid of *P. tricornutum*. In *C. reinhardtii*, the described presence of PSII components in the pyrenoid could be related with the existence of a network of pyrenoid-penetrating tubules from the intrapyrenoid thylakoids. However, although an equivalent network has been described in the pyrenoid of red algae, it has not been found in diatoms (Engel et al. 2015, Meyer et al. 2017). On the other hand, in unicellular algal species, it has been proposed that the pyrenoid plays an important role in defining the starting point of thylakoidal

maturation and structural nucleation (reviewed in Rast et al. 2015). In particular, in *C. reinhardtii*, the pyrenoid is connected to biogenesis centers involved in the assembly of PSII (but not PSI), as ribosomes and mRNAs encoding PSII subunits are localized in the pyrenoid periphery (Uniacke and Zerges 2007). In addition, PSII mutants impaired in PSII assembly showed an accumulation of early PSII intermediates at the pyrenoid (Uniacke and Zerges 2007). A similar process in diatoms could maybe explain the presence of PSII subunits into or in close contact with the pyrenoid of *P. tricornutum*.

On the other hand, although the role (if any) that Cc₅₅₀ could play in the pyrenoid matrix may be a matter of discussion, it should be related to carbon fixation or the CCM located in this microcompartment. Interestingly, the existence of two PSII putative Ca2+ binding PsbP-type proteins, PSBP3 and PSPP4, has been reported in the pyrenoid of *C. reinhardtii* (Mackinder et al. 2017). However, there is no evidence of the presence of equivalent PsbP-like proteins in diatoms. Actually, PsbP is considered to be evolutionarily recruited as a replacement of Cc₅₅₀ in the photosynthetic green lineage, playing a similar role stabilizing the binding of Ca²⁺, and thus increasing PSII affinity for this ion (Frazão et al. 2001, Roncel et al. 2012). Moreover, a Ca2+-binding protein (CAS) has been also recently shown to specifically localize into the intrapyrenoid thylakoids of C. reinhardtii (Wang et al. 2016). Thus, the presence of Cc_{550} in the pyrenoid of *P. tricornutum* could be related to its role as a calcium-binding stabilizing protein.

Author contributions

J.A.N. and M. H. conceived the project and carried out the protein purification experiments; P. B.-B., C. A. and C. C. carried out the DNA analysis and the co-immunoseparation and immunodetection experiments; P.C. carried out the immunoelectron microscopy experiments with inputs from M. R.; J. A. N., M. H. and M. R. discussed the results and wrote the manuscript, which was corrected, revised and approved by all authors.

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Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

- **Fig. S1.** Sequences of oligonucleotides used in this work and alignment of sequences of *Isochrysis galbana* and *Emiliania huxleyi* eukaryotic cytochromes c_{550} .
- **Fig. S2.** Summary for peptide fingerprint analysis of *Phaeodactylum tricornutum* Cc_{550} directly resolved on polyacrylamide gel electrophoresis.
- **Fig. S3.** Co-immunoseparated proteins identified by LC–MS/MS in soluble, membrane and crosslinked fractions of *Phaeodactylum tricornutum*.
- **Fig. S4.** Western blot analysis of crude cellular fractions of *Phaeodactylum tricornutum* with antibodies against either Cc_{550} or the RubisCO large and small subunits.