Iron–Sulfur Cluster Biosynthesis in Algae with Complex Plastids

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Abstract

Plastids surrounded by four membranes harbor a special compartment between the outer and inner plastid membrane pair, the so-called periplastidal compartment (PPC). This cellular structure is usually presumed to be the reduced cytoplasm of a eukaryotic phototrophic endosymbiont, which was integrated into a host cell and streamlined into a plastid with a complex membrane structure. Up to date, no mitochondrion or mitochondrion-related organelle has been identified in the PPC of any representative. However, two prominent groups, the cryptophytes and the chlorarachniophytes, still harbor a reduced cell nucleus of symbiont origin, the nucleomorph, in their PPCs. Generally, many cytoplasmic and nucleus-located eukaryotic proteins need an iron–sulfur cofactor for their functionality. Beside some exceptions, their synthesis is depending on a so-called iron–sulfur complex (ISC) assembly machinery located in the mitochondrion. This machinery provides the cytoplasm with a still unknown sulfur component, which is then converted into iron–sulfur clusters via a cytosolic iron–sulfur protein assembly (CIA) machinery. Here, we investigated if a CIA machinery is present in mitochondrion-lacking PPCs. By using bioinformatic screens and in vivo-localizations of candidate proteins, we show that the presence of a PPC-specific CIA machinery correlates with the presence of a nucleomorph. Phylogenetic analyses of PPC- and host specific CIA components additionally indicate a complex evolution of the CIA machineries in organisms having plastids surrounded by four membranes.

Key words: iron–sulfur cluster, cytosolic iron–sulfur protein assembly (CIA), periplastidal compartment, Guillardia theta, Bigelowiella natans, Phaeodactylum tricornutum.

Introduction

Cryptophytes and chlorarachniophytes, both eukaryotic phototrophic unicellular organisms, provide extraordinary examples in respect to the cellular minimization of a previously free-living organism. By merging two eukaryotic cells, a phototrophic symbiont was reduced within a host cell to a complex organized plastid surrounded by four membranes. Extraordinary minimization can be observed by the presence of a remnant cytoplasm of the eukaryotic symbiont, which is located between the outer and inner plastid membrane pair (Maier et al. 2015). In cryptophytes and chlorarachniophytes, this remnant cytoplasm (periplastidal compartment [PPC]) harbors a tiny cell nucleus, as the only well-defined PPC-located compartment (Grosche et al. 2014). This so-called nucleomorph is reduced in coding capacity as well as overall size (Douglas et al. 2001; Gilson et al. 2006; Tanifuji et al. 2011; Moore et al. 2012; Tanifuji et al. 2014a; Suzuki et al. 2015). Although being, as mentioned, reduced, (e.g., Moore and Archibald 2009; Grosche et al. 2014), nucleomorphs are functional in respect to transcription and RNA processing (Tanifuji et al. 2014b; Suzuki et al. 2016), and the nucleomorph-encoded proteins are expected to be translated at PPC-specific 80S ribosomes. Thus, cryptophytes and chlorarachniophytes are unicellular organisms with two different cell nuclei located in two different eukaryotic cytoplasms, each one containing its own type of 80S ribosomes. However, many phylogenetically related organisms, such as diatoms, minimized their PPC more drastically, as here a nucleomorph and a second set of ribosomes are absent (Hempel et al. 2007; Bolte et al. 2009).

Assembly of ribosomes is highly coordinated (Maier et al. 2013) and requires, besides components present in mature
ribozymes, several accessory factors. One of these, the iron–sulfur cluster (Fe–S) containing protein Rli1, is involved in ribosome biogenesis in addition to translation initiation and termination (Kispal et al. 2005; Strunk et al. 2012). A central cellular role of Rli1 in this process can also be deduced from the fact that homologs are present in archaean and eukaryotes (Nurenenberg and Tampe 2013). Accordingly, the presence of a gene encoding the Fe–S cluster protein Rli1-homolog in the nucleomorph of cryptophytes is not surprising (first identified by Douglas et al. 2001; but, as a member of ATP binding cassette [ABC] ATPases protein family, annotated as an ABC transporter). Nevertheless, the need for factors taking part in ribosome functions might be different in chlorarachniophytes, as neither a nucleomorph- nor a nucleus-encoded PPC-located Rli1 has yet been identified in silico. However, more likely Rli1 might be encoded in two versions in chlorarachniophytes as well, but the genes are not detectable in the genome sequence of B. natans. In any case, the presence of a RLI1 gene encoded by the cryptophyte nucleomorph demands Fe–S clusters to be present in the PPC of at least cryptophytes.

One of the major cell biological highlights of the last decades was the finding that typical aerobic eukaryotes depend on mitochondria for the synthesis of Fe–S clusters (see: Lil et al. 2014, 2015; Mühlendorf et al. 2015). This makes mitochondria essential for the maturation of all cellular Fe–S proteins except for those of plastids, which possess a SUF (sulfur component indispensable for cytosolic and nuclear Fe–S cluster assembly). This component is transported via the ABC transporter Atm1 (yeast, Atm3/Starik1 in S. cerevisiae) into the cytoplasm (Kispal et al. 1999; Kushnir et al. 2001; Bernard et al. 2009; Lil et al. 2014). Due to its so far unknown nature this component is referred to as X-S (Lil et al. 2015). In the cytoplasm X-S, together with monothiol glutaredoxins, is processed by a so-called cytosolic iron–sulfur protein assembly (CIA) machinery to generate cytosolic and nuclear Fe–S proteins (Netz et al. 2014). Most components of both the ISC- and CIA-machineries are generally essential for aerobic eukaryotes and defects therefore affect the whole cell leading to so-called Fe–S diseases (Stehling et al. 2014).

Since at least the genetically active PPC of cryptophytes seems to require Fe–S clusters a mitochondrion or a MRO was never identified in the PPC of at least cryptophytes. Especially for the latter, at least remnants of a CIA machinery (Lil et al. 2015) for the assembling of PPC-specific Fe–S proteins might be present.

In order to search for the here postulated PPC-specific CIA machineries, we screened the genomes of a cryptophyte, a chlorarachniophyte and a diatom for putative CIA factors. Our results suggest that, in parallel to a CIA machinery in the host cytoplasm, PPC-specific CIA factors are encoded in the genomes of cryptophytes and most likely chlorarachniophytes, whereas diatoms lack a PPC-specific CIA machinery. Our phylogenetic studies on the central CIA factors Cfd1 and Nbp35 (both are central P-loop NTPases serving as scaffold proteins for cytosolic iron–sulfur protein assembly) indicated different origins of host- and symbiont-specific Nbp35 proteins in cryptophytes, chlorarachniophytes and the host version of diatoms. Moreover, the presence of Cfd1, generally not known from members of the SAR supergroup (Stramenopiles, Alveolates and Rhizaria), was confirmed for a host version for cryptophytes (Tsaousis et al. 2014). As a PPC-version of a mitochondrion or a MRO was never identified in cryptophytes, we finally hypothesize that a X-S component for ISC assembly in the PPC of cryptophytes is supplied by the plastid SUF system and imported into the PPC.

Materials and Methods

Culture Conditions

Phaeodactylum tricornutum was cultivated in f/2 medium (Guillard 1975), adjusted to pH 7.2, under constant illumination (80 μmol photons m⁻² s⁻¹) at 22 °C. Liquid cultures were grown with agitation (150 rpm) in a volume of 300 ml.

Guilardia theta was cultured in h/2 medium, a modified f/2 medium with a final NH₄Cl concentration of 500 μM, without agitation at 15 °C in a 12h/12h day/night cycle in a volume of 100–200 ml.

In silico Analyses and Factor Identification

Factors of the CIA machinery were identified using the known factors of S. cerevisiae in a reciprocal BLAST (Altschul et al. 1990) approach against the genomes of B. natans, G. theta, and P. tricornutum (Bowler et al. 2008; Curtis et al. 2012), the nucleomorph genomes of B. natans and G. theta were also included (Douglas et al. 2001; Gilson et al. 2006). Gene models of identified candidates were manually checked for, for example, EST or RNASeq coverage and analyzed for presence of N-terminal targeting signals using SignalP 3.0 (Bendtsen et al. 2004), TargetP 1.1 (Emanuelsson et al. 2000) and alignments of the N-terminus (supplementary fig. S1, Supplementary Material online). Cia1 and Cia2 candidates were analyzed with help of HmmerWeb Version 2.23.0 and Gene3D prediction (supplementary figs. S3 and S4, Supplementary Material online) (Finn et al. 2015; Dawson et al. 2017; Lewis et al. 2018; Potter et al. 2018).
Phylogenetic Reconstruction

To verify ortholog status of identified factors, yeast and before (see above) identified query protein sequences were searched with BLASTP (Altschul et al. 1990) against a plant-specific, custom-made protein database that included genomes of the species listed in supplementary table S2, Supplementary Material online. Results were filtered according to Rost (1999) to keep homologous sequences only. Additionally, sequences for Nbp35 and Cfd1 were added from the results of Tsouosis (Tsouosis et al. 2014) and from a NCBI Blast to add additional sequences from especially SAR and fungal species (see supplementary table S2, Supplementary Material online). Resulting sequences were aligned using MUSCLE version v3.8.31 (Edgar 2004) in automatic mode, and resulting alignments were inspected manually, duplicated sequences were removed and alignment was trimmed using Jalview version 2.8 (Clamp et al. 2004). Based on these alignments, neighbor-joining (NJ) guide trees were built using quicktree_sd (http://hdl.handle.net/10013/epic.33164.d001; Last accessed March 27, 2018) with 1,000 bootstrap samples. Sequences with very long branches, potentially representing flawed gene models, were removed upon inspection of initial trees. Afterwards, the appropriate models were selected based on AIC/BIC using ProtTest 3.4 (Guindon and Gascuel 2003; Darriba et al. 2011). Final phylogenies were constructed by Bayesian inference (BI) using Mr. Bayes 3.2.5 (Ronquist et al. 2012). BI analysis was run with two hot and two cold chains, discarding 25% of trees as burn-in, for 8,000,000 generations (standard deviation of split frequencies 0.027225). The resulting tree was displayed, colored, and midpoint-rooted with FigTree version 1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/; Last accessed March 27, 2018), schematic tree representation was done with iTOL (Letunic and Bork 2016).

In vivo Localizations of CIA Proteins

The sequences of PPC localized Dre2 (ProtID: 199508), Nar1 (ProtID: 138617), Nbp35 (ProtID: 186667), and Tah18 (ProtID: 199508) can be retrieved from the genome database (https://genome.jgi.doe.gov/Guith1/Guith1.home.html; Last accessed March 27, 2018). RNA was isolated from 100 ml culture of G. theta to the host-derived cytoplasms of Phaeodactylum tricornutum (diatom) (table 1 and supplementary table S1, Supplementary Material online).

In case a symbiont-specific CIA machinery is present in the PPC in parallel to a host version that is cytoplasmic, their factors should either be encoded by the nuclear Fe–S proteins (Kispal et al. 2005; Srinivasan et al. 2014). Not surprisingly, our inspection of the available genome sequences highlighted genes for several cytosolic CIA factors, most likely targeted to the host-derived cytoplasm of Guillardia theta (cryptophyte), Bigelowiella natans (chlorarachniophytes), and Phaeodactylum tricornutum (diatom) (table 1 and supplementary table S1, Supplementary Material online).

For heterologues in vivo localization studies in P. tricornutum, the amplificates were cloned 5’- to eGFP (5’-amplificate-eGFP-3’) into the nitrate inducible pPha-NR vector (GenBank: JN180663) after intermediate cloning into pJET vector (Thermo Scientific). Apart from Tah18 full length sequences were used for localization studies. In case of Tah18, we used the N-terminal part (first 972 of 2,286 bp/first 324 of 761 amino acids) for localization.

Fluorescence Microscopy

For in vivo localization the expression of the eGFP fusion proteins was induced with 0.9 mM of NaNO3 for 2 days. The localization was visualized with a confocal laser scanning microscope Leica TCS SP2 using a HCX PL APO 40×/1.25 – 0.75 Oil CS objective. eGFP and chlorophyll fluorescence was excited at 488 nm. eGFP fluorescence was detected at a bandwidth of 500–520 nm and the chlorophyll (plastid autofluorescence) at a bandwidth of 625–720 nm, respectively. Pictures were processed with Fiji ImageJ with usage of BioFormats Importer plug-in (Linkert et al. 2010; Schindelin et al. 2012; Schneider et al. 2012; Rueden et al. 2017).

Results

Screening the Genomes of Cryptophytes, Chlorarachniophytes, and Diatoms for Components of a CIA Machinery

Cryptophytes, chlorarachniophytes, and diatoms possess a mitochondrial within the cytoplasm. As it was shown for important model systems such as yeast, this organelle exports a sulfur component which is used by the CIA machinery for the maturation of cytosolic and nuclear Fe–S proteins (Kispal et al. 2005; Srinivasan et al. 2014). Not surprisingly, our inspection of the available genome sequences highlighted genes for several cytosolic CIA factors, most likely targeted to the host-derived cytoplasm of Guillardia theta (cryptophyte), Bigelowiella natans (chlorarachniophytes), and Phaeodactylum tricornutum (diatom) (table 1 and supplementary table S1, Supplementary Material online).
the TPL is neither an aromatic amino acid nor a leucine (Gruber et al. 2007; Patron and Waller 2007). Although a BTS is present in the presence of nucleus-encoded, PPC-directed proteins of chlorarachniophytes as well, a highly conserved amino acid at the first position of the TPL was not detected (Patron and Waller 2007). Using the absence/presence of a predicted BTS as a criterion, we identified CIA factors with a putative PPC destination for cryptophytes and chlorarachniophytes, as listed in table 1. Interestingly, we did not detect genes for PPC-directed Cia1 (a WD40 protein) and Cia2 (a MIP18 family-like motif harboring protein) factors in the nuclear genome of Guillardia theta. Instead, a reinspection of the nucleomorph genome sequence of the cryptophyte highlighted the two orfs, orf357 and orf143, respectively, most likely encoding the missing factors. The nucleomorph-encoded Cia1 and Cia2 candidate proteins show either a WD40-like motif and “Putative cytosolic iron–sulfur protein” prediction (Cia1, supplementary fig. S3, Supplementary Material online) or a MIP18 family-like motif and Fe–S cluster assembly (FSCA)’ prediction (Cia2, supplementary fig. S4, Supplementary Material online), which strongly indicate the here proposed functions in the CIA targeting complex. For the distinction of Cfd1 and Nbp35 homologs a ferredoxin-like domain is indicative (e.g., Netz et al. 2016) which is present in the N-terminal regions of Nbp35- but not Cfd1-homologs. Using this criterion, we annotated the encoded proteins. Most notably, our screening confirms the finding of Tsaousis et al. (2014) in respect to the presence of a Cfd1-homolog in G. theta.

In case of the diatom Phaeodactylum tricornutum, the proteome of the PPC was already investigated in vivo (Moog et al. 2011). In this data set no potential Fe–S protein was identified. As a nucleomorph is absent in diatoms, no indications exist for the need of Fe–S proteins acting in the PPC. In agreement with the previous data, no CIA-factors with a PPC-specificity could be identified in silico screens in the P. tricornutum genome (table 1).

For further analyses (see below), we reinspectied the predicted gene models identified in the G. theta genome and included, if present, EST-data for correcting the gene models in some cases. This led to a promising data set of putative CIA factors for the host and the symbiont (table 1 and supplementary table S1, Supplementary Material online). However, the BTS of the PPC-localized Tah18 of G. theta (supplementary fig. S1, Supplementary Material online) is predicted to be not very pronounced. The identification of CIA factors in the B. natans genome was more complicated due to the annotation of the genome. This was further complicated by the high intron load of the coding regions, missing EST-data covering the coding region and high levels of splice variants including both intron retention and exon skipping. Nevertheless, we identified several CIA factors (Nbp35, Nar1, Cia2, Dre2, and Tah18; table 1 and supplementary table S1, Supplementary Material online) in the B. natans genome, which are potentially PPC-localized, one of each with a putative BTS. However, in case of Dre2, Nbp35, and Tah18 the putative BTS is again not very pronounced.

Taken together, the in silico analyses indicated the presence of a CIA machinery in the PPCs of cryptophytes and most likely chlorarachniophytes, whereas evidence for such a machinery in the PPC of diatoms is absent.

Under the assumption that organisms having a transcriptionally active nucleomorph need a CIA machinery in their PPC, which is devoid of any traces of a mitochondrion, either a sulfur-component has to be imported for Fe–S cluster assembly or a NIF (nitrogen fixation)- or SUF-system (Lill and Muhlenhoff 2006; Stairs et al. 2014; Karnkowska et al. 2016; Freibert et al. 2017) should be present in this compartment. We have already postulated the presence of a SUF-system in cryptophytes, located in the stroma of their complex plastid (Hjorth et al. 2005), but found no indication of any SUF- or NIF-related factors with a PPC targeting signal in the genome of G. theta. Therefore, import of an activated sulfur component into the PPC, which is then used by a PPC-specific CIA machinery is likely. It is known from other eukaryotes that the mitochondria-localized ABC transporter Atm1 provides the cytoplasm with a sulfur component synthesized in the mitochondrion (e.g., Srinivasan et al. 2014). By scanning the genome of G. theta several potential ABC transporters were identified. One candidate shows, as expected, homology to Atm1 and a potential targeting signal for mitochondria (protein ID 188917). Aside from this canonical, most likely mitochondria localized version, other potential Atm1-like ABC transporters can be detected, but these have no striking similarity to Atm1. In addition, none of these candidates has a targeting signal for the outermost or second outermost membrane of the complex plastid. However, one of the putative ABC transporters from G. theta is predicted to be encoded as a preprotein having an N-terminal BTS with the specificity for the innermost plastid membrane (protein ID 164017).

### In vivo Localization

For G. theta and B. natans transformation protocols are not established, preventing homologous expression of proteins in
these organisms. However, in earlier studies concerning the targeting signals of cryptophytes, we have shown that PPC-specific targeting sequences of G. theta perfectly direct proteins into the remnant cytoplasm of the secondary symbiont of the diatom P. tricornutum (Gould et al. 2006). Therefore, we localized the cryptophytic CIA factors heterologously, expressing a PPC targeting signal, fused to eGFP, in the diatom. This approach is especially important for the predicted PPC version of Tah18 from G. theta, which shows, as mentioned, a less-pronounced BTS-like structure at the N-terminus. We have expressed eGFP fusion proteins for the predicted CIA factors Nbp35, Nar1, Tah18, Dre2, and Grx3 (fig. 1). These fusion proteins localized within the PPC, as shown by a so-called blob-like structure of the eGFP signal (Kilian and Kroth 2005). Thus, the here predicted, PPC-specific CIA factors of cryptophytes are equipped with a targeting signal directing proteins into the PPC when heterologously expressed in the diatom system.

**Phylogeny**

In 2014, Tsaoisis and colleagues published their results on the phylogeny of the CIA machinery in the stramenopile Blastocystis (Tsaoisis et al. 2014). As an important outcome they demonstrated that the CIA machinery is uniquely found in eukaryotes. Although sequences from chlorarachniophytes, cryptophytes and diatoms were included into their study, only one copy of each CIA factor was integrated into the phylogenetic trees in the case of cryptophytes and chlorarachniophytes. Since we have shown here that a cytosolic and a PPC-located version of CIA factors are encoded in the genomes of G. theta and B. natans, we reinvestigated the phylogeny of the CIA factors Cfd1 and Nbp35. In addition, we incorporated our novel data on the localization of the CIA representatives when evaluating those results (fig. 2 and supplementary fig. S2, Supplementary Material online). Although Cfd1 and Nbp35 are homologous proteins, their sequences can be undeniably discerned due to a ferredoxin-like domain only present in the N-terminal region of Nbp35 proteins.

The phylogeny of Cfd1 and Nbp35 sequences including Hcf101 representatives (fig. 2 and supplementary fig. S2, Supplementary Material online) indicated, similar to the work of Tsaoisis et al. (Tsaoisis et al. 2014), a separation of the Cfd1 and Nbp35 branches. Sequences for the chloroplast-localized Hcf101 were integrated into the analyses since this protein, similar to Cfd1 and Nbp35, harbors a P-loop NTPase domain; Nar1 sequences served as outgroup to root the tree.

In case of the Nbp35 phylogeny, four clades were identified, an Opisthokonta-type, a plant-type and two SAR clades. Nbp35 of diatoms was found in one of the SAR clades (SAR-I), whereas in SAR-II the PPC and host Nbp35 versions of B. natans are grouping. Interestingly, the Nbp35 versions of G. theta branch differently, namely in case of the host version (nucleus-encoded, cytoplasmic) in the Opisthokonta clade and, in case of the PPC version, together with red algal Nbp35 proteins in the plant-like clade. In case of Cfd1, we only found a host version for G. theta, which also has been shown before (Tsaoisis et al. 2014), but none for B. natans. Similar to the host Nbp35 version, Cfd1 clusters in the Opisthokonta clade.

**Discussion**

Several ecologically and medically important organisms such as diatoms or the malaria causing protist Plasmodium falciparum harbor complex organized plastids that evolved from a phototrophic eukaryotic endosymbiont (e.g., Gould et al. 2008). Most of the compartments of the symbiont, such as the mitochondrion, were eliminated during coevolution of host and symbiont. However, intermediates in the extent of intracellular reduction of the phototrophic symbiont that has become a complex plastid are known. This is the case for chlorarachniophytes and cryptophytes, both of which harbor a plastid surrounded by four membranes (Maier et al. 2000). Here, the cytoplasm of the eukaryotic symbiont (PPC), is reduced in such a way that the nucleus of the former symbiont (nucleomorph) is in respect to size and coding capacity minimized (Douglas et al. 2001; Gilson et al. 2006; Lane et al. 2007; Tanifuji et al. 2011; Moore et al. 2012; Tanifuji et al. 2014a; Suzuki et al. 2015), but still present and active. However, the cellular reduction in others such as diatoms went further, so that for these organisms no cellular traces of a symbiotic nucleus were preserved. The morphological differences between organisms harboring a plastid surrounded by four membranes are manifested in several cell biological/biochemical functions; as shown here, this is also the case for CIA machineries, which are essential to assemble cytoplasmic iron–sulfur clusters for Fe–S proteins (fig. 3).

We have previously studied the PPC-proteome of the diatom P. tricornutum (Moog et al. 2011). This organism, as mentioned, has no nucleomorph and, as known so far, has only to manage a small set of cellular functions in the PPC, for which no Fe–S proteins seem to be essential. Consequently, no CIA machinery is necessary in the PPC of diatoms, which is supported by the in silico analyses presented here. Thus, our report indicates a eukaryotic (remnant) cytoplasm devoid of a CIA machinery. As known from other eukaryotes, several nucleus-located Fe–S proteins are acting in DNA replication, DNA repair, chromosome segregation and telomere length regulation (Rudolf et al. 2006; Netz et al. 2011; Gari et al. 2012; Stehling et al. 2012; Wu and Brosh 2012; Stehling et al. 2013; Paul and Lill 2015). Therefore, essentiality of Fe–S clusters in at least some of these diatom proteins is expected and a CIA machinery should be expressed in the cytosol for the incorporation of Fe–S clusters into apo-proteins. Our screen for CIA factors supports this hypothesis. In addition, the CIA machinery of diatoms shows, by the lack of the factor Cfd1, a
typical feature for a member of the SAR group (Tsaousis et al. 2014).

By knowing the importance of Fe–S proteins in genome maintenance and expression (Paul and Lill 2015), one has to postulate that nucleomorph-containing PPCs are depending on Fe–S proteins. In fact, according to our analyses, B. natans as well as G. theta might import putative Fe–S cluster apoproteins into the PPC. In addition, an important Fe–S protein, Rli1, is encoded in the nucleomorph of G. theta (Douglas et al. 2001). Thus, cryptophytes and chlorarachniophytes should depend on a PPC-located CIA machinery. Our screen of the genomes of G. theta and B. natans genomes uncovered CIA factors equipped with a BTS besides those active in the cytosol. To verify that these factors are directed into a PPC, we performed in vivo localizations. As for G. theta and B. natans no transformation protocols are elaborated and therefore no in vivo localization experiments are possible, we studied targeting of the putative PPC-specific CIA factors in heterologous localization experiments in the diatom system. This approach resulted in a localization of the eGFP-fusion proteins within the remnant cytoplasm of the eukaryotic symbiont in P. tricornutum, thereby confirming our in silico predictions in a heterologous system (fig. 1).

Altogether, the here described PPC-specific CIA factors encoded in the nuclear genome of G. theta make up a nearly complete machinery known from studies in other organisms (fig. 3). Missing factors apparently not encoded in the nuclear genome of the cryptophyte are Cia1 and Cia2, which are part

**Fig. 1.**—Heterologous localization of G. theta CIA factors in P. tricornutum. Potential PPC localized CIA factors of G. theta were heterologously expressed in P. tricornutum as GFP fusion proteins. Brightfield, GFP, plastid autofluorescence and overlay (from left to right) are shown. eGFP fluorescence shows a localization in a so-called blob-like structure which indicates a localization in the PPC. In the case of Dre2, Nar1 and Tah18 cells show two blob-like structures, indicating dividing cells. PAF: plastid autofluorescence. Scale Bar: 10 μm.
of the so-called targeting complex responsible for the transfer of Fe–S clusters on target apo-proteins (Balk and Lobreaux 2005; Srinivasan et al. 2007; Stehling and Lill 2013). However, in *G. theta* these factors seem to be encoded in the nucleomorph genome, previously annotated as Orf357 and Orf143 (Douglas et al. 2001). Although the sequence similarity of these Orfs to nuclear-encoded Cia1 and Cia2 versions of other organisms is low, the presence of indicative domains supports their identity as factors of the CIA machinery (supplementary figs. S3 and S4, Supplementary Material online). This is not restricted to *G. theta*, because the nucleomorph of *Chroomonas mesostigmatica*, *Cryptomonas paramecium* and *Hemiselmis anderseni* (Lane et al. 2007; Tanifuji et al. 2011; Moore et al. 2012) encode these orfs as well. Therefore, the here presented data support the conclusion that a complete set of CIA factors is present in the PPC of the cryptophyte *G. theta* (fig. 3). Although not proven in respect to the localization of the factors, our data indicate the presence of a second, PPC-located CIA machinery for *B. natans* as well.

These results, together with earlier findings on the absence of a PPC-located mitochondrion, raise the question on the origin of a sulfur component necessary for Fe–S cluster assembly by the PPC-specific CIA machinery in cryptophytes. Fe–S clusters can be synthesized not only by an ISC machinery, but also by a SUF- or NIF-machinery. A SUF-machinery was indeed described for cryptophytes, but it is localized within the stroma (Hjorth et al. 2005). To pursue the hypothesis of a putative PPC-located NIF-machinery an in silico screening for NIF-specific factors was performed. However, no matches were found (data not shown). Thus, we predict that an activated sulfur component for ISC synthesis is imported into the PPC, which is in principle possible either from the host-derived cytoplasm or the stroma. In the case that the activated sulfur component stems from the host, two membranes have to be crossed, similar to an origin of the component in the stroma, where an active transport across the innermost and second innermost membrane of the plastid is required.

In order to investigate the origin of the sulfur component for Fe–S cluster synthesis in cryptophytes, we searched for
Atm1 homologs. This mitochondrial ABC transporter was shown to export a sulfur component used by the cytosolic CIA machinery (e.g., Kispal et al. 1999; reviewed in Lill et al. 2014). Although several ABC transporters are encoded by the G. theta genome, the ones with the highest sequence similarities are a predicted mitochondrial version of Atm1 and another predicted to be localized in a plastid membrane. The nature of the latter is not really indicative due to a generally high primary sequence conservation of ABC transporters. Nevertheless, we propose that the sulfur component used by the PPC-located CIA machinery of G. theta is synthesized by a SUF system of the complex plastid and exported from the plastid stroma into the PPC via an ABC transporter in the inner plastid membrane (fig. 3). Although the origin of a SUF-depending sulfur component used by a CIA machinery might be unusual, examples for that are known. Especially in the case of Monocercomonoides sp. (Karnkowska et al. 2016), a protist synthesizing Fe–S clusters by a laterally acquired bacterial SUF system, which is, however, acting in the host-derived cytosol.

The evolution of organisms with complex plastids of red algal origin (“chromalveolates”, Cavalier-Smith 1999) is still a matter of debate and a monophyletic or even higher order origin have been proposed (Cavalier-Smith 1999; Burki et al. 2012; Gould et al. 2015, and references therein). However, it is accepted by most scientists that the plastid of chromalveolates is of red algal origin. In any case, the evolution of the chromalveolates cannot be deciphered by investigating the symbiont only. Therefore, the host genome might be informative, especially with regard to the presence of genes encoding for components of the core metabolism such as Fe–S cluster synthesis. In case of a CIA machinery, the scaffold for iron–sulfur protein assembly shows a specific protein composition. The central component for this scaffold is Nbp35, which interacts, according to the molecular analyses in yeast and human, with Cfd1 (e.g., Lill et al. 2015). Nevertheless, a recent report indicates that Nbp35 of Arabidopsis interacts with Dre2, another conserved component of the CIA machinery (Bastow et al. 2017). Cfd1 is encoded in Amoebozoa, ascomycetes, basidiomycetes, metazoans and others, but is generally absent in Archaeplastida and the SAR group (Tsaoosis et al. 2014). Thus, the presence or absence of Cfd1 might be an informative character at least for the evolution of chromalveolates. In this respect the presence of Cfd1 in cryptophytes in combination with several other recent studies might indicate a complex evolution of cryptophytes, which differs from that of other chromalveolates (figs. 2 and 3 and supplementary fig. S2, Supplementary Material online) (e.g., Bachvaroff et al. 2005; Bodily 2005; Teich et al. 2007; Burki et al. 2012; Petersen et al. 2014; Stiller et al. 2014). If this statement is correct, other CIA factors should show a non-chromalveolate grouping in phylogenies as well when cryptophytic factors are compared with homologs of the Archaeplastida and the SAR. This was indeed seen in respect to Nbp35. As shown in figure 2, representatives of Nbp35 group in one of the four Nbp35 clades, two of which are specific either for Opisthokonta or Archaeplastida. The third and fourth group (the SAR group I and SAR group II) include the diatom Nbp35 (SAR group I) whereas the PPC and host version of the chlorarachniophyte B. natans are members of the SAR group II clade. In contrast, the cryptophytic Nbp35 proteins group with the Opisthokonta (host Nbp35) and in the

![Diagram](image-url)
Archaeplastida branch close to red algae, as expected for the PPC version of Nbp35. Therefore, the cryptophytic host CIA machinery differs from the SAR group CIA systems not only in the presence of a Cfd1, but also in the inferred phylogenetic origin for Nbp35.

As mentioned, both Nbp35 of B. natans cluster in SAR group II. This might indicate that they have been subject to a duplication event. Complex plastids of chlorarachniophytes like B. natans are thought to have evolved from green algae ancestors (e.g., Curtis et al. 2012). Therefore, we rather would have expected one of these Nbp35s to cluster within Archaeplastida, comparable to the situation seen in G. theta. In case of a duplication event, the host version (SAR) might have been duplicated and have been equipped with a suitable BTS for PPC targeting. With this, the symbiont version of Nbp35 was not needed anymore and got lost.

We recently inspected the origin of several nucleus-encoded factors directed into the complex plastid of diatoms and other chromalveolates like SELMA (symbiont-specific ERAD-like machinery) and triosephosphate transporters (Hempel et al. 2009; Felsner et al. 2011; Moog et al. 2015). Our phylogenetic analyses revealed that these components, located in the PPCs and plastid membranes of chromalveolates, are of symbiont (red algal) origin. Interestingly, this is also the case with respect to the PPC-specific CIA factor Nbp35 in cryptophytes, indicating that although a host copy is present, the symbiont version of Nbp35 is of cell biological/biochemical importance and is maintained as long as they are needed. However, the situation is different in respect to predicted Nbp35 PPC version of B. natans, which is grouping in the SAR group and not together with green alga/land plants. A similar result was shown in the work of Hopkins et al. (2012), indicating that several PPC and stroma targeted proteins are of nongreen origin.

Conclusions

Taken together the here reported phylogenetic study of Nbp35 and the presence of a Cfd1 factor in G. theta suggest that cryptophytes are cellular mergers, which originated by the union of a red alga and a host cell, the latter with a different phylogenetic origin than in other members of the chromalveolates. More than a speculation is our finding that a PPC-specific CIA machinery correlates with the presence of a nucleomorph. Nevertheless, cryptophytes have a PPC-specific iron–sulfur burden, which necessitates a CIA machinery presumably fed by a sulfur component synthesized in the stroma.

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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