Most plastid proteins are encoded by their nuclear genomes and need to be targeted across multiple envelope membranes. In vascular plants, the translocons at the outer and inner envelope membranes of chloroplasts (TOC and TIC, respectively) facilitate transport across the two plastid membranes. In contrast, several algal groups harbor more complex plastids, the so-called secondary plastids, which are surrounded by three or four membranes, but the plastid protein import machinery (in particular, how proteins cross the membrane corresponding to the secondary endosymbiont plasma membrane) remains unexplored in many of these algae. To reconstruct the putative protein import machinery of a secondary plastid, we used the chlorarachniophyte alga Bigelowiella natans, whose plastid is bounded by four membranes and still possesses a relict nucleus of a green algal endosymbiont (the nucleomorph) in the intermembrane space. We identified nine homologs of plant-like TOC/TIC components in the recently sequenced B. natans nuclear genome, adding to the two that remain in the nucleomorph genome (B. natans TOC75 [BnTOC75] and BnTIC20). All of these proteins were predicted to be localized to the plastid and might function in the inner two membranes. We also show that the homologs of a protein, Der1, that is known to mediate transport across the second membrane in the several lineages with secondary plastids of red algal origin is not associated with plastid protein targeting in B. natans. How plastid proteins cross this membrane remains a mystery, but it is clear that the protein transport machinery of chlorarachniophyte plastids differs from that of red algal secondary plastids.
delivering preproteins from the ER into the plastid stroma across three or four envelope membranes. A few homologs of TOC and TIC components (TIC20, TIC22, TIC110, and/or TOC75) have been detected by comparative genome searches in several organisms bearing secondary plastids of red algal origin: the heterokont *Phaeodactylum tricornutum* (10), the cryptophyte *Guillardia theta* (26), and the apicomplexans *Plasmodium falciparum* (39) and *Toxoplasma gondii* (65). This suggests that the outermost membrane is crossed via the general ER-targeting/secretion pathway and that the third and fourth membranes are crossed using some kind of TOC/TIC complexes, but for most algae this still leaves one membrane unaccounted for: the second outermost membrane, which is homologous to the plasma membrane of the endosymbiont. In several lineages with red algal secondary plastids, ER-associated protein degradation (ERAD)-derived proteins (e.g., Der1-like proteins) have been identified as the preprotein translocon that mediates the crossing of this membrane (2). However, there are currently no data on the nature of this process for any secondary plastid of green algal origin. This is unfortunate, because this is arguably the only step where the parallel evolution of a similar pathway is not highly likely. The symbiont would import already-functional TOC and TIC systems, and because all secondary plastids reside in the endomembrane system, it is hardly unexpected that the host ER-targeting pathway was independently coopted for this targeting step. Crossing the second outermost membrane is different because neither the host nor the endosymbiont would have had a single, obvious, and ready-made system in place to carry out this step (which might explain why two lineages of algae, dinoflagellates and euglenids, have lost this membrane).

In this study, we characterize the Der1-like proteins, as well as all recognizable TOC and TIC components, of the chlorarachniophyte *Bigelowiella natans*. Chlorarachniophytes possess four-membrane plastids derived from a green algal endosymbiont, and these plastids retain a highly reduced nucleus, the so-called nucleomorph, between the inner and outer pair of membranes (50). Plastid preproteins of chlorarachniophytes, like those of other secondary algal groups, possess an N-terminal bipartite targeting sequence consisting of an SP and a TPL sequence (54). Our previous studies demonstrated that the preproteins are transported into the plastids via the ER and that a net positive charge of TPL sequences is essential for passing through the inner two plastid membranes (31, 33). This suggests the existence of translocons that selectively recognize the TPL sequence and facilitate preprotein transport across the two inner plastid membranes. The complete sequence of the nucleomorph genome encodes one component of each of the TOC and TIC complexes, TOC75 and TIC20, but no other targeting gene products (25). We have therefore comprehensively searched the recently completed *B. natans* nuclear genome to develop a picture of what the plastid translocons in this organism might look like. We show that neither Der1-like protein (based on the localization of green fluorescent protein [GFP] fusion proteins) appears to be specifically associated with plastid protein targeting. Moreover, we show that both TOC and TIC complexes are apparently reduced and that the TOC in particular has lost all putative receptor proteins so that only the channel protein remains. Overall, these results support the conclusion that protein targeting to chlorarachniophyte plastids differs in several ways from that of secondary plastids of a red algal origin, at least one of which (transport across the second outermost membrane) is likely fundamentally different. So although many aspects of the evolutionary integration of red and green algal plastid symbionts and their hosts have proceeded in parallel in different lineages, for other steps, different solutions have also been found, as expected for a system assembled by “tinkering.”

**MATERIALS AND METHODS**

**Identification of Der1, TOC, and TIC homologs in the Bigelowiella natans genome.** Homologs of the ERAD protein Der1 were identified by similarity with homologs from other eukaryotes using BLASTP searches against a library of predicted proteins from the *B. natans* genome sequencing project (sequenced at the DOE Joint Genome Institute). To identify the components of TOC and TIC homologs in *B. natans*, we used a collection of all proteins from both complexes from *P. sativum*, *A. thaliana*, *C. reinhardtii*, *O. lucimarinus*, and *O. tauri*, summarized in a previous study (38). These TOC and TIC sequences were used to search the predicted proteins from the *B. natans* genome using BLASTP. For a more sensitive manual survey to identify less-conserved potential homologs, we used the results of Smith-Waterman searches provided by JGI. Detected homologs to reciprocal BLASTP sequences were then used against *P. sativum* sequences in the NCBI server, and resulting E values are indicated (Table 1). Some predicted TOC and TIC homologs were absent from transcriptome surveys of *B. natans* (namely, *B. natans* TIC20 [BnTIC20], BnTIC21, BnTIC32, BnTIC40, and BnOMP85), and in these cases, we carried out 5′ and/or 3′ rapid amplification of cDNA ends (RACE) to confirm that they represent expressed genes. Total RNA was isolated from *B. natans* cells using TRIzol reagent (Invitrogen), and cDNA was synthesized using the FirstChoice RNA ligase-mediated (RLM)-RACE kit (Ambion) according to the manufacturer’s protocols. Each target cDNA was amplified by PCR using Econo Taq DNA polymerase (Lucigen) with specific primer sets (shown in Table S1 in the supplemental material). Resulting PCR products were cloned into the pSC-A-amp/km plasmid of a StrataClone PCR cloning kit and subsequently sequenced with an Applied Biosystems 3730 DNA analyzer and a BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems).

Predictions of N-terminal SP sequences were performed by the Hidden Markov Models (HMM) method of the SignalP 3.0 server (6), and TPL sequences were predicted by the ChloroP 1.1 server (20). The probability of each prediction is indicated in Table 1. The overall charge of each predicted TPL sequence was calculated using the Peptide Property Calculator (Innovagen). Transmembrane α-helices were predicted with the TMHMM server, v. 2.0 (46), and SOSUI, v. 1.11 (34). Predictions of α-helices and β-strands in BnTIC75-nucleomorph (Nm) and BnOMP85 were made with PROFsec of the PredictProtein server (55). Protein motifs and domains were identified using two databases: PFAM (23) and SMART (48). We collected TOC and TIC homologs from plants (*P. sativum*, *A. thaliana*, *O. sativa*, and *Physcomitrella patens*) and green algae (*C. reinhardtii*, *Chlorella variabilis*, *Micromonas pusilla*, *O. tauri*, *O. lucimarinus*, and/or *Volvox carteri*) using Phytozome v. 7.0 and JGI websites, and alignments of homologs were created by the ClustalW program in MEGA 4 (63). Sequence conservations were shown as logo plots generating by WebLogo v. 2.8.2 (19).

**Phylogenetic analyses.** Homologs of Der1, TIC20, and OMP85 (TOC75) were collected for apicomplexans, heterokonts, cryptophytes, red algae, green algae, and/or vascular plants using GenBank, JGI, or *C. megalae* genome project websites. Automatic alignment was performed by the L-INS-I method of the MAFFT package (41), and Gblocks (12) was used to automatically and reproducibly eliminate poorly aligned positions, with no gapped positions allowed, the minimum number of sequences for a conserved and a flank position set to 50% of the number of taxa plus one, the maximum of contiguous nonconserved positions set to 10, and the minimum length of a block set to 8; the resulting unambiguously aligned positions were manually inspected with SeaView (27). Maximum-likelihood phylogenetic analyses were performed with RAxML 7.2.8 (62), in combination with the rapid hill-climbing algo-
this interendomembrane sorting takes place in chlorarachniosome, the second outermost membrane. There is no direct evidence for how this is mediated, and the most mysterious, is the crossing of the outermost membrane. This is, according to all available evidence, mediated by the signal peptide in all sec-}

Using reciprocal BLAST and Smith-Waterman searches. We can validate for proteins potentially involved in preprotein translocons and/or ERAD-derived proteins of the...protein translocon or a general ERAD component, was sought. Specifically, the net charge calculation, since their TPL regions could not be predicted by ChloroP. The BLASTP E values against P. sativum TOC/TIC components are in the rightmost column.

### RESULTS AND DISCUSSION

To create a model of the plastid preprotein transport pathway in chlorarachniophytes, the B. natans genome sequence was sur-
volved in the crossing of the second outermost membrane, which is, according to all available evidence, mediated by the signal peptide in all secondary plastids and will not be discussed further. The second part of the pathway, and the most mysterious, is the crossing of the second outermost membrane. There is no direct evidence for how this interendomembrane sorting takes place in chlorarachniophytes, though two hypotheses have been proposed. In one, a duplicated ERAD complex controls this process, as is the case in at least some red secondary plastids (59), and in the other, a duplicated TOC complex carries it out (13). The third stage is the crossing of the inner two plastid membranes, and it is widely assumed that this is mediated by standard TOC and TIC complexes inherited from the primary algal endosymbiont, although few of the proteins making up this complex have been identified.

**ERAD-derived proteins of the B. natans genome.** Several algal groups (heterokonts, haptophytes, and cryptophytes) and a parasitic group (apicomplexans) harbor four-membrane-bounded plastids derived from a red algal endosymbiont, and it was from these that the first direct evidence for how preproteins cross the second membrane was found. Sommer et al. (59) described a nucleomorph-encoded Der1-like protein as the symbiont ERAD-like machinery (SELMA) and speculated it to be involved in the crossing of the second outermost plastid membrane in cryptophytes. Several studies have subsequently proposed that symbiont-specific Der1-like proteins commonly act as the second outermost membrane translocon of secondary plastids in heterokonts (30, 59), haptophytes (22), and apicomplexans (1, 60).

To determine whether Der1 is involved in crossing the second outermost membrane in chlorarachniophyte plastids, we searched for Der1 homologs in the B. natans genome. We detected two Der1-like proteins (BnDer1-49642 and BnDer1-92850) in the nuclear genome, and phylogenetic analyses suggested that these proteins were weakly related to nucleomorph-encoded Der1-like proteins of cryptophytes and symbiont-specific Der1-like proteins, respectively (Fig. 1A). This suggests that chlorarachniophytes might have independently coopted the same ERAD-derived system to target plastid preproteins as did red algal secondary plastids. However, the phylogenetic position of neither protein is statistically supported, so more direct evidence for the functional roles of these proteins, as a plastid translocon or a general ERAD component, was sought. Specifically, we tested the localizations of both B. natans Der1-like proteins using

### TABLE 1 Putative TOC and TIC components in the Bigelowiella natans genome*

| Component | P. sativum accession no. | B. natans accession no. | No. of aa | SP sequence (probability score) | TPL sequence (probability score) | TPL net charge (no. of aa) | E value |
|-----------|--------------------------|--------------------------|-----------|---------------------------------|---------------------------------|-----------------------------|--------|
| TOC34     | Q41009                   | —                        |           |                                 |                                 |                             |        |
| TOC64     | AAF62870 (JGI 89506)     | 784                      | Yes (0.998) | No (0.493)                     | 4 (50)                          | 7.00E−20                    |        |
| TOC75     | Q43715                  | ABA27321(Nm)             | 818       | No                              | No (0.427)                     | −2.9 (50)                   | 9.00E−24 |
| OMP85     | Q43715                  | JGI 79166               | 899       | Yes (0.786)                     | No (0.435)                     | 10 (50)                      | 0.002  |
| TOC159    | AF75761                 | —                        |           |                                 |                                 |                             |        |
| TIC20     | AAC64607 (Nm)           | 207                      | No        | No (0.458)                     | 9 (50)                          | 5.00E−5                     |        |
| TIC21     | JGI 115241              | 322                      | Yes (1.000) | Yes (0.549)                    | 3.1 (32)                        | 1.00E−5                     |        |
| TIC22     | JGI 66755               | 272                      | Yes (1.000) | Yes (0.518)                    | 7.1 (36)                        | 2.00E−4                     |        |
| TIC32     | JGI 115241              | 341                      | Yes (0.996) | Yes (0.530)                    | 8 (32)                          | 0.001                       |        |
| TIC40     | JGI 115241              | 330                      | Yes (0.948) | Yes (0.538)                    | 9.1 (53)                        | 0.15                         |        |
| TIC55     | JGI 66755               | 272                      | Yes (1.000) | Yes (0.518)                    | 9 (50)                          | 3.00E−32                    |        |
| TIC62     | JGI 66755               | 341                      | Yes (0.996) | Yes (0.530)                    | 9.1 (53)                        | 0.15                         |        |
| TIC110    | JGI 66754               | 322                      | Yes (1.000) | Yes (0.518)                    | 7.1 (36)                        | 2.00E−4                     |        |
| TIC110    | JGI 66754               | 322                      | Yes (1.000) | Yes (0.518)                    | 9 (50)                          | 3.00E−32                    |        |

*The GenBank accession or JGI protein identification number of each TOC/TIC component is recorded. “SP sequence” indicates the result of signal peptide prediction by the HMM method of SignalP, and the probability scores are shown in parentheses. “TPL sequence” is the result of transit peptide prediction by ChloroP. “TPL net charge” shows the net charge of the predicted TPL sequence, with the predicted length of the TPL sequence in parentheses (the N-terminal 50 aa of TOC64, TOC75, and TIC20 were used for the net charge calculation, since their TPL regions could not be predicted by ChloroP). The BLASTP E values against P. sativum TOC/TIC components are in the rightmost column.

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**Subcellular localization of B. natans Der1-like proteins.** To construct two GFP expression vectors, cDNA fragments of two Der1-like genes (BnDer1-49642 and BnDer1-92850 genes) were amplified by PCR with specific primers shown in Table S2 in the supplemental material, and then each fragment was inserted between HindIII and NcoI sites of the pLaRGfp vector (33). These plasmids were cloned in the DH5α strain of *Escherichia coli* and purified with a QiAprep Spin miniprep kit (Qiagen). To analyze the subcellular localization of GFP fusion proteins, we transformed a related chlorarachniophyte species, *Amorphochlora* amoebiformis, with these plasmids using a Biolistic PDS-1000/He particle delivery system (Bio-Rad) as described previously (32). Transiently transformed cells were observed under an Axioskop 2 fluorescence microscope (Carl Zeiss AG) with a 3CCD HD XL H1S video camera (Canon). GFP fluorescence and plastid autofluorescence were detected with filter 17 (excitation BP, 485/20 nm; emission LP, 546/12 nm; emission long-pass [LP], 590 nm), respectively.

**Nucleotide sequence accession numbers.** cDNA sequences from 5′- and/or 3′-RACE analyses are deposited in GenBank under accession no. JQ088188 to JQ088192.
a GFP fusion protein and the genetic transformation system of a related chlorarachniophyte, *Amorphochlora amoebiformis*. We created two plasmid constructs that express GFP fused with full-length BnDer1-49642 (217 amino acids [aa]) or BnDer1-92850 (251 aa). *A. amoebiformis* cells were transformed with each plasmid individually, and GFP localizations were examined. Both fusion proteins led to green fluorescence exclusively in the ER (Fig. 1B). This indicates that both *B. natans* Der1-like proteins function in the ER as general components of ERAD machinery and that neither of them is a translocon specifically at the second outermost membrane of the plastid. This result is not surprising, since organisms with red algal secondary plastids possess four homologs of Der1 proteins in their genomes: two of them are components of ERAD machinery and the other two are components of the SELMA, so there is a precedent for such algae using multiple ERAD-specific Der1 proteins. Although this does not reveal how proteins cross the second membrane in chlorarachniophytes, it does show that this lineage has evolved a system for transporting preproteins across the second plastid membrane that is dis-
distinct from the ERAD-derived system of red algal plastid-bearing organisms.

**TOC and TIC complexes in B. natans.** TOC and TIC protein sequences of vascular plants and green algae were used for an initial BLASTP analysis against all predicted protein sequences of the B. natans genome, and Smith-Waterman searches were used for more sensitive detections. The predicted TOC and TIC homologs of B. natans were subsequently analyzed by BLASTP analysis of P. sativum protein sequences, and the resulting E values are shown in Table 1. We also predicted targeting sequences and functional domains/motifs using bioinformatic tools and manual curations. Overall, we identified 11 members of TOC/TIC components from the B. natans genomes, and all of them were predicted to be localized in the plastid.

**TOC proteins.** The outer membrane translocon complex of vascular plants consists mainly of four TOC components. The B. natans nucleomorph genome encodes a copy of TOC75, and we found another homolog of OMP85 in the nuclear genome, but our searches did not clearly identify any additional component of the TOC complex.

TOC75 is a member of outer membrane protein 85 (OMP85) family and forms a protein-conducting channel. This protein has two typical structural domains; the N-terminal region contains polypeptide transport-associated (POTRA) domains, and the C-terminal bacterial surface antigen domain forms a transmembrane β-barrel (24). POTRA domains share the same strand-helix-helix-strand topology in their secondary structures (56), and the β-barrel domain contains 16 β-strands (21). A homolog of TOC75 in B. natans, BnTOC75-Nm, has previously been identified in the nucleomorph genome (25). This protein is predicted to possess an N-terminal POTRA domain and a bacterial surface antigen domain in its C-terminal region, according to PFAM searches (Fig. 2). Secondary-structure prediction based on PROFsec suggests that the putative POTRA domain consists of three β-strands, with the first and second strands separated by two α-helices, and that the C-terminal region contains 16 putative β-strands. However, it was difficult to predict regions of canonical POTRA1 and POTRA2 sequences and 16 β-strands from the alignment with FhaC, whose crystal structure has been reported (18) (Fig. 2). All together, these data suggest that BnTOC75-Nm probably forms a transmembrane β-barrel and functions as a protein-conducting channel. We also found a nucleus-encoded homolog of the OMP85 family, named BnOMP85, that was weakly predicted by reciprocal BLASTP searches (its E value against P. sativum TOC75 is 0.002 [Table 1]). Our phylogenetic analysis indicates that BnOMP85 is closely related with OMP85-like proteins of heterokonts and apicomplexans, but it is unclear whether these proteins are grouped with either TOC75 or OEP80 (Fig. S1). On the basis of the alignment with FhaC, this protein is predicted to include the POTRA2 domain and 16 β-strands in its
C-terminal bacterial surface antigen domain (Fig. 2), which implies that BnOMP85 also functions as a protein-conducting channel. In vascular plants, TOC75 preproteins carry a cleavable transit peptide followed by a polyglycine stretch at their N termini, and these proteins should be inserted into the outer membrane from the stroma side. It has been reported that both of these motifs play an essential role in the outer plastid membrane localization of TOC75 in *P. sativum* (5) and *A. thaliana* (36). Interestingly, nucleus-encoded BnOMP85 possesses an N-terminal bipartite plastid targeting sequence followed by a polyglycine stretch containing 8 glycine residues, which implies that BnOMP85 may be inserted into the third membrane (counting from the outside) in a way similar to that of plant TOC75 (Fig. 2, 3). In contrast, BnTOC75-Nm lacks both a canonical transit peptide and a polyglycine stretch. There is no experimental evidence for the localization of BnTOC75-Nm, but if it occupies a location homologous to that in plants and green algae, it would be found in the third membrane (Fig. 3), although how it is targeted there is unknown and seems to be different from that of plant TOC75.

TOC64 is an integral membrane protein composed of an N-terminal transmembrane domain, an inactive amidase domain, and C-terminal tetratricopeptide repeat (TPR) motifs (57). The *B. natans* nuclear genome encodes a single homolog of amidase with a canonical N-terminal bipartite targeting sequence followed by a polyglycine stretch containing 8 glycine residues, which implies that BnOMP85 may be inserted into the third membrane (counting from the outside) in a way similar to that of plant TOC75 (Fig. 2, 3). In contrast, BnTOC75-Nm lacks both a canonical transit peptide and a polyglycine stretch. There is no experimental evidence for the localization of BnTOC75-Nm, but if it occupies a location homologous to that in plants and green algae, it would be found in the third membrane (Fig. 3), although how it is targeted there is unknown and seems to be different from that of plant TOC75.

TOC64 is an integral membrane protein composed of an N-terminal transmembrane domain, an inactive amidase domain, and C-terminal tetratricopeptide repeat (TPR) motifs (57). The *B. natans* nuclear genome encodes a single homolog of amidase with a canonical N-terminal bipartite targeting sequence consisting of an SP and a weakly predicted TPL sequence (Fig. 3; Table 1). However, this protein lacks the TPR motifs, which are proposed to be a chaperone-docking site, as well as the N-terminal transmembrane domain. These suggest that the *B. natans* amidase homolog is probably targeted to the plastid but that it possibly lacks the function of an import receptor, so it is not certain that the TOC64 component is strictly required for protein transport into the chlorarachniophyte plastids. This view is supported by a few additional observations: the genomes of both a green alga (*C. reinhardtii*) and a red alga (*C. merolae*) also lack TOC64 (38), and it has been demonstrated that the *A. thaliana* TOC64 protein is not essential for efficient preprotein transport into plastids (4).

Both TOC34 and TOC159 are GTPase proteins that function as receptors of plastid preproteins and, together with the protein-conducting channel TOC75, constitute the core TOC components (3). Plastid preproteins are initially identified by TOC34 and TOC159 by the direct binding of transit peptides or an indirect binding to transit peptide associated with Hsp70 and 14-3-3 proteins. Bound proteins are then threaded into the TOC75 channel. In our survey, no homolog of either TOC34 or TOC159 was found in the *B. natans* genome. Although it is possible that the sensitivity of our survey was not sufficient to detect these proteins, overall, the evidence suggests that the *B. natans* genome lacks canonical plant-like receptors of plastid preproteins. We propose two possible implications for the absence of TOC34 and TOC159 receptors in *B. natans*. First, the TPL sequence of plastid preproteins might be directly identified by BnTOC75-Nm or BnOMP85 without receptors, since it has been reported that *P. sativum* TOC75 contains a transit peptide-binding site in its N-terminal region, including POTRA domains (21). Alternatively, *B. natans* might use novel preprotein receptors which are divergent from plant TOC components. Our previous studies have indicated that a net positive
charge of TPL sequences is essential for protein transport across the two inner plastid membranes (31, 33). This implies that novel receptors might recognize the TPL sequence net charge on the third outermost plastid membrane.

**TIC proteins.** In vascular plants, eight TIC components have been identified so far (45). In this study, we found 7 homologs of TIC components in the *B. natans* nuclear genome (TIC20, TIC21, TIC22, TIC32, TIC40, TIC55, and TIC62 were identified, and TIC110 was not identified). A homolog of TIC20 is also encoded in the nucleomorph genome (Table 1).

TIC110, TIC20, and TIC21 are proposed to be components of protein-conducting channels at the inner plastid membrane. TIC110 is the most abundant component of the TIC complex, acting as the main channel for protein transport (29). TIC20 is an alternative candidate for the inner membrane channel, which is an integral membrane protein with four α-helical transmembrane domains (14). TIC21 also contains four α-helices and has been postulated to be a putative channel (64). We detected single TIC20 and TIC21 homologs (BnTIC20 and BnTIC21) but no TIC110 in the *B. natans* nuclear genome (Table 1). A homolog of TIC20 is also encoded in the nucleomorph genome (Table 1).

TIC110, TIC20, and TIC21 are proposed to be components of protein-conducting channels at the inner plastid membrane. TIC110 is the most abundant component of the TIC complex, acting as the main channel for protein transport (29). TIC20 is an alternative candidate for the inner membrane channel, which is an integral membrane protein with four α-helical transmembrane domains (14). TIC21 also contains four α-helices and has been postulated to be a putative channel (64). We detected single TIC20 and TIC21 homologs (BnTIC20 and BnTIC21) but no TIC110 in the *B. natans* nuclear genome (Table 1). A homolog of TIC20 is also encoded in the nucleomorph genome (Table 1).

TIC110, TIC20, and TIC21 are proposed to be components of protein-conducting channels at the inner plastid membrane. TIC110 is the most abundant component of the TIC complex, acting as the main channel for protein transport (29). TIC20 is an alternative candidate for the inner membrane channel, which is an integral membrane protein with four α-helical transmembrane domains (14). TIC21 also contains four α-helices and has been postulated to be a putative channel (64). We detected single TIC20 and TIC21 homologs (BnTIC20 and BnTIC21) but no TIC110 in the *B. natans* nuclear genome (Table 1). A homolog of TIC20 is also encoded in the nucleomorph genome (Table 1).

TIC22 is a soluble intermembrane protein that is peripherally bound to the outer face of the inner plastid membrane (44). This protein has been postulated to serve as an adaptor between TOC and TIC complexes. In the *B. natans* genome, two homologs of TIC22 were weakly predicted by reciprocal BLASTP searches against *P. sativum* TIC22 (E values of 0.15 and 0.001 [Table 1]). Although the accuracy of BLASTP predictions is not high, these proteins share two conserved motifs with TIC22 homologs of plants and green algae (Fig. 4) and carry a canonical N-terminal bipartite targeting sequence consisting of an SP and a pronounced net positively charged TPL sequence (Table 1). Therefore, we postulate that these two proteins are TIC22 homologs of *B. natans* (BnTIC22A and BnTIC22B) and that they are most likely localized in the intermembrane space between the inner two plastid mem-

**FIG 4** Protein structures of BnTIC22, BnTIC40, BnTIC32, BnTIC55, and BnTIC62. Sequence conservations among TIC homologs of land plants (*Pisum sativum*, *Arabidopsis thaliana*, *Oryza sativa*, and *Physcomitrella patens*) and green algae (*Chlamydomonas reinhardtii*, *Chlorella variabilis*, *Micromonas pusilla*, *Ostreococcus tauri*, *O. lucimarinus*, and/or *Volvox carteri*) are shown as logo plots generating by WebLogo v. 2.8.2. *B. natans* TIC sequences are shown above the logo plots and conserved amino acids are indicated by asterisks.
branes. However, it remains unclear how these proteins are inserted into the intermembrane space.

TIC40 has been described as a cochaperone protein with an N-terminal α-helical transmembrane domain, followed by a TPR motif that interacts with TIC110 and a Sti1/Hip/Hop domain that mediates the interaction with Hsp93 (16, 17, 61). We identified a TIC40 homolog in *B. natans* (BnTIC40) that is predicted to possess these three characteristic domains as well as an N-terminal bipartite targeting sequence (Fig. 3; Table 1). Notably, the C-terminal sequence contains a TPR-like motif and a Sti1/Hip/Hop domain sharing a high degree of similarity with TIC40 proteins of plants and green algae (Fig. 4). BnTIC40 appears to function as a translocon at the innermost membrane but likely does not interact with TIC110 as in plants, since this protein is absent in *B. natans*.

Three TIC components, TIC32 (15, 35), TIC55 (11), and TIC62 (47), contain either NADP(H)-binding sites or Rieske-type iron-sulfur clusters, and these are hypothesized to be involved in preprotein transport regulation depending on the metabolic state of the plastid. We found one homolog of each of these TIC components, each predicted to carry an N-terminal targeting sequence, suggesting plastid localization (Fig. 3, Table 1). *B. natans* TIC55 (BnTIC55) contains a 2Fe-2S binding site and a mononuclear iron binding site in its N-terminal region, followed by a domain with similarities to pheophoribine a oxygenase (PAO), and two transmembrane helices were predicted to fall in the C-terminal region based on the alignment with TOC55 homologs of plants and green algae (the TMHMM server predicted a single transmembrane helix) (Fig. 4). BnTIC55 closely resembled TIC55 homologs in these features, but there is a striking difference; although TIC55 homologs possess a C-terminal conserved domain, including an R-Y-X$_2$-H-X$_3$-C-X$_2$-C motif of unknown function (28), two cysteine residues of this motif have been replaced by serine residues in BnTIC55 (Fig. 4). Both homologs of TIC32 and TIC62 in the *B. natans* genome (BnTIC32 and BnTIC62) were predicted to belong to the (extended) family of short-chain dehydrogenases (SDRs) containing conserved NADP(H)-binding sites (Fig. 3). BnTIC32 possesses the signature motif, T-G-X$_2$-G-X-G, for NADP(H) binding and catalytic tetrad residues consisting of asparagine, serine, tyrosine, and lysine (35) (Fig. 4). The C-terminal region of BnTIC32 is loosely defined as a calmodulin-binding site that is present in plant TIC32 homologs (15) (Fig. 4). BnTIC62 also contains the G-X$_2$-G-X$_2$-G motif within its predicted NADP(H)-binding domain but lacks a C-terminal ferredoxin-NADP reductase (FNR)-binding domain that has been reported for TIC62 of *P. sativum* and *A. thaliana* (47) (Fig. 4). These three TIC homologs, BnTIC55, BnTIC32, and BnTIC62, possess several domains with high resemblance to plant TIC components. However, a few domains having essential/unknown functions are weakly predicted or not detected in *B. natans* TIC homologs. We propose these three homologs to be putative TIC components, but it is unclear whether they have the same functions as plant TIC components.

**Reconstructing chlorarachniophyte plastid-targeting translocons in *B. natans***. Collectively, the *B. natans* nuclear and nucleomorph genomes encode two homologs of TOC75 and seven members of the TIC complex. These plant-like TOC and TIC components seem to be derived from the green algal endosymbiont via secondary endosymbiosis. It is noteworthy that only two components, TOC75 and TIC20, remain encoded in the nucleomorph genome and that these two proteins are also the putative protein-conducting channels. It is also noteworthy the way in which the *B. natans* TOC complex appears to have been reduced. The conserved pore is retained, but none of the receptors are evident. These proteins are present in the genome of *C. reinhardtii*, which is comparatively closely related to the green algal endosymbiont of chlorarachniophytes (38, 53), but were apparently lost in the *B. natans* lineage after the secondary endosymbiosis. Because primary plastids reside in the cytoplasm, their TOC complex must be able to distinguish plastid proteins from all other cytoplasmic and mitochondrial proteins. In contrast, secondary plastids reside within the endomembrane system and in chlorarachniophytes within a small residual volume of cytoplasm that is discontinuous with the rest of the host endomembrane system. Accordingly, the receptors at the third plastid membrane must distinguish plastid proteins only from the pool of nonplastid proteins that would reside in this compartment. Since this compartment is massively reduced compared to ordinary green algal cytoplasm, we hypothesize the receptors might require much less information to accurately recognize plastid proteins, which might in turn lead to a reduced demand for receptors.

An even more pressing problem is how plastid-targeted preproteins in chlorarachniophytes cross the second outermost membrane. We found no evidence to support the conclusion that ERAD-derived components, in particular, symbiont-specific Der1 proteins, are utilized to cross this membrane, as has been shown in all other algae that have retained it (2). While this may have been the most obvious possibility, we have shown that the localization patterns of *B. natans* Der1 proteins are consistent with a generalized ER function. However, the hypothesis that a duplicate TOC complex is located in the second membrane and mediates import is consistent with our findings, since two homologs of the OMP85 family (BnTOC75-Nm and BnOMP85) are encoded in the nuclear and the nucleomorph genome, respectively. It is possible that both putative channel proteins localize in the third membrane (counting from the outside). One paralogue of *A. thaliiana* TOC75, the so-called AtOEP80 protein, is directly inserted into the outer plastid membrane from the cytoplasm without an N-terminal transit peptide (37), and both AtTOC75 and AtOEP80 have the same topology in the plastid outer membrane despite the opposite directions of their insertions (ATTOC75 and ATOEP80 are delivered from the stroma side and the cytoplasm, respectively) (58). If BnTOC75-Nm is targeted in the same way, both it and BnOMP85 may localize to the third membrane. However, it is also possible that BnTOC75-Nm might be integrated into the second outermost membrane, because this protein lacks a canonical plastid membrane targeting sequence that BnOMP85 and plant TOC75 preproteins possess. In this case, BnOMP85 would fulfill the canonical role of TOC75 in the third membrane, while BnTOC75-Nm would serve to translocate proteins across the second membrane, as the SELMA system does in red algal secondary plastids. The detailed localization of these putative channels should be confirmed in the future, but currently there is no method available in chlorarachniophytes that would allow us to unambiguously distinguish between localizations in any of the four plastid membranes (which might be demonstrated by a self-assembling split GFP system, for example). Moreover, we currently have no means to transform the nucleomorph genome, so even if the location could be distinguished, it would not yet be possible to determine how BnTOC75-Nm is targeted. Neverthe-
less, these two putative channels are fascinating candidates to solve this interesting puzzle in protein targeting.

Further characterization of this important step in plastid targeting will be of great interest, not only from the perspective of chlorarachniophyte biology, but for helping to define the limits of the integration of endosymbionts, since this is arguably the only step where one might realistically expect to see a genuinely non-parallel development in green and red algal secondary plastids. The absence of a duplicated plastid-specific Der1 protein in chlorarachniophytes shows that they do use a nonhomologous system, and the onus is now on defining what that system is.

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