New Insights into the Unique Structure of the F$_0$F$_1$-ATP Synthase from the Chlamydomonad Algae Polytomella sp. and Chlamydomonas reinhardtii

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In this study, we investigate the structure of the mitochondrial F$_0$F$_1$-ATP synthase of the colorless alga Polytomella sp. with respect to the enzyme of its green close relative Chlamydomonas reinhardtii. It is demonstrated that several unique features of the ATP synthase in C. reinhardtii are also present in Polytomella sp. The α- and β-subunits of the ATP synthase from both algae are highly unusual in that they exhibit extensions at their N- and C-terminal ends, respectively. Several subunits of the Polytomella ATP synthase in the range of 9 to 66 kD have homologs in the green alga but do not have known equivalents as yet in mitochondrial ATP synthases of mammals, plants, or fungi. The largest of these so-called ASA (ATP Synthase-Associated) subunits, ASA1, is shown to be an extrinsic protein. Short heat treatment of isolated Polytomella mitochondria unexpectedly dissociated the otherwise highly stable ATP synthase dimer of 1,600 kD into subcomplexes of 800 and 400 kD, assigned as the ATP synthase monomer and F$_1$-ATPase, respectively. Whereas no ASA subunits were found in the F$_1$-ATPase, all but two were present in the monomer. ASA6 (12 kD) and ASA9 (9 kD), predicted to be membrane bound, were not detected in the monomer and are thus proposed to be involved in the formation or stabilization of the enzyme. A hypothetical configuration of the Chlamydomonad dimeric ATP synthase portraying its unique features is provided to spur further research on this topic.

F$_0$F$_1$-ATP synthase (EC 3.6.1.3) occurs ubiquitously on energy-transducing membranes, such as mitochondrial, thylakoid, and bacterial plasma membranes, and produces the majority of cellular ATP under aerobic conditions. The enzyme separates readily into two distinct parts: a membrane-embedded domain (F$_0$-ATP synthase) involved in proton translocation and a water-soluble domain (F$_1$-ATPase) that is the site of ATP synthesis or hydrolysis. The most investigated ATP synthase to date is that of Escherichia coli, consisting of eight different subunits with a stoichiometry of α$_3$β$_3$γδεαβ(c$^{10,12}$), which are all essential for its function (Foster and Fillingame, 1982; Schneider and Altendorf, 1985). Subunits-α, -β, -γ, -δ, and -ε constitute the F$_1$-ATPase, whereas subunits-a, -b, and -c constitute the F$_0$-ATP synthase. A central stalk that connects the two domains is formed by subunits-γ and -ε, while a second peripheral stalk that links the apex of the α$_3$β$_3$ hexagon to the F$_0$ domain is constituted by subunit-b and -δ.

The mitochondrial F$_0$F$_1$-ATP synthase is significantly more complex than the bacterial enzyme and consists of at least 15 different proteins. In addition to the eight essential subunits found in bacteria, the mitochondrial enzyme contains several so-called supernumerary subunits (<20 kD) that are required for the structure or regulation of the enzyme. For instance, the peripheral stalk of mitochondrial ATP synthase is composed of four subunits (b, d, F6, and oligomycin sensitivity conferral protein [OSCP]; OSCP is equivalent to subunit-δ in bacteria; Collinson et al., 1994, 1996) instead of two in bacteria (b and δ). Other supernumerary subunits are involved in the dimerization or oligomerization of the ATP synthase, a process that was shown to occur in the mitochondrial membrane (Allen et al., 1989; Paumard et al., 2002; Dudkina et al., 2005; Minauro-Sanmiguel et al., 2005). In yeast (Saccharomyces cerevisiae), dimerization involves the physical association of two neighboring F$_0$ domains via subunit 4 (subunit-b in mammals) and the associated proteins e and g (Spannagel et al., 1998; Paumard et al., 2002). These proteins as well as several other F$_0$ subunits are found in both mammals and yeast. In...
plants, fewer $F_0$ subunits have been identified so far, whereas a plant-specific subunit named $F_{A6L}$ was found (Heazlewood et al., 2003).

Compared to other known mitochondrial ATP synthases, those of Chlamydomonad algae show several unique structural features. First, the catalytic subunits-$\alpha$ and -$\beta$ in the green alga *Chlamydomonas reinhardtii* exhibit peculiar extensions at their N and C termini, respectively (Franzén and Falk, 1992; Nurani and Franzén, 1996). The $\alpha$-subunit in the nonphotosynthetic alga *Polytomella* sp. also exhibits an extended N terminus, the sequence of which is highly similar to that of its close relative, *C. reinhardtii* (Atteia et al., 1997). Second, both *C. reinhardtii* and *Polytomella* sp. mitochondrial ATP synthases, solubilized with dodecyl maltoside (1%–2%), run on blue native PAGE (BN-PAGE) exclusively as a dimer of approximately 1,600 kD (Atteia et al., 2003; van Lis et al., 2003, 2005). This is in clear contrast to the mitochondrial ATP synthases of beef, yeast, or plants, which, when solubilized with 1% dodecyl maltoside, occur on BN-PAGE as two major forms corresponding to the $F_0F_1$-ATP synthase monomer (600 kD) and the $F_1$-ATPase (400 kD; Jänsch et al., 1996; Arnold et al., 1998; Eubel et al., 2003).

Third, *C. reinhardtii* mitochondrial ATP synthase separated by BN-PAGE contains seven proteins in the range of 10 to 61 kD that have no clear equivalents in other mitochondrial ATP synthases (Funes et al., 2002; van Lis et al., 2003) and that were given the designation ASA (ATP Synthase-Associated; Cardol et al., 2005). Moreover, a recent study on the ATP synthase of *Polytomella* sp. revealed the presence of nine ASA proteins, whereas typical mitochondrial ATP synthase subunits-b, -$F_0$, and -d of the peripheral stalk and supernumery subunits-A6L, -e, -f, and -g are missing (Vázquez-Acevedo et al., 2006). Last, electron microscopy images of the dimeric mitochondrial ATP synthase of *Polytomella* sp. showed the presence of a very pronounced peripheral stalk (Dudkina et al., 2005), strongly disparate from the thin stalk found in the yeast ATP synthase (Dudkina et al., 2006).

Here, we further characterize the mitochondrial ATP synthase of the colorless alga *Polytomella* sp. in relation to the available data on this enzyme in *C. reinhardtii* and follow up on the electron microscopy and biochemical data obtained for the *Polytomella* ATP synthase (Atteia et al., 1997, 2003; Dudkina et al., 2005, 2006; Vázquez-Acevedo et al., 2006). As the composition of the ATP synthases from the two algae appear highly similar, data on the enzyme obtained from either source are assumed to be mostly interchangeable. This work gives new insights into the structure of the mitochondrial ATP synthase of the Chlamydomonad algae and opens the way to a testable hypothesis regarding the structural features and role of the identified atypical traits.

**RESULTS**

*Polytomella* sp. and *C. reinhardtii* $F_0F_1$-ATP Synthase Separated by Two-Dimensional BN/SDS-PAGE

BN gel bands containing the dimeric $F_0F_1$-ATP synthase from *Polytomella* sp. and *C. reinhardtii* (1,600 kD) were applied on Gly or Tricine SDS-PAGE. The *Polytomella* ATP synthase is resolved into 17 distinct polypeptides on glycine SDS-PAGE, ranging from 66 kD to 7 kD (Fig. 1). Recently, on Tricine SDS-PAGE, 17

![Figure 1. Polypeptide composition of the *Polytomella* (Ps) $F_0F_1$-ATP synthase dimer and comparison with that of *C. reinhardtii* (Cr). The subunits of the ATP synthase from BN gel were resolved on 2D Gly SDS-PAGE (15% acrylamide) or on Tricine SDS-PAGE (13% acrylamide) and Coomassie Blue stained. For *C. reinhardtii*, Tricine SDS-PAGE was used to allow reference to earlier studies using this system (Funes et al., 2002; van Lis et al., 2003). The identities of subunits-ASA8, -ASA9, and -e of *C. reinhardtii* on Tricine SDS-PAGE were not confirmed. The 12 largest *Polytomella* subunits on Gly SDS-PAGE were identified by N-terminal sequencing, whereas the rest (marked with *) were tentatively assigned based on the Tricine SDS-PAGE subunit profile (Vázquez-Acevedo et al., 2006). The *C. reinhardtii* subunits on Gly SDS-PAGE were identified by liquid chromatography-mass spectrometry/mass spectrometry analysis (A. Atteia, A. Adrait, S. Brugière, M. Tardiff, R. van Lis, L. Kuhn, O. Bastien, J. Garin, J. Joyard, and N. Rolland, unpublished data).
subunits, including nine ASA subunits, were found that were assigned after N-terminal sequencing (Vázquez-Acevedo et al., 2006). These N-terminal sequences corroborated those we obtained for the 12 largest proteins from the Polytomella ATP synthase (data not shown). The ASA subunit nomenclature used here is as previously reported (Cardol et al., 2005; Vázquez-Acevedo et al., 2006). The ASA designation with Ps or Cr prefix refers to the subunit in a particular alga and otherwise refers to the subunit in a general sense, having presumably a similar structure and function in both algae.

Notwithstanding some disparities, the ATP synthase subunit profiles on Gly SDS-PAGE are rather similar in the two algae. The α- and β-subunits migrate similarly in both algae (Fig. 1). Sequencing of the α- and β-subunit cDNAs of Polytomella sp. revealed the presence of atypical extensions highly similar to those of C. reinhardtii. The α-subunits in both algae contain an N-terminal extension of approximately 20 residues, whereas the β-subunits exhibit at their C termini a hydrophilic α-helical extension of about 64 residues (Fig. 2). The function of these extensions is not known. The γ-subunit of Polytomella sp. runs consistently higher on SDS-PAGE than that of C. reinhardtii (Fig. 1), although the predicted molecular masses of the γ-subunits from both algae are very similar (approximately 31 kD). Polytomella subunits-α, -β, and -γ all show highest sequence identity with their counterparts in C. reinhardtii. Subunit-6 of C. reinhardtii, which could only be detected by silver staining, migrates similarly to that of Polytomella sp. (18 kD; data not shown).

In C. reinhardtii, ASA1 migrates together with the β-subunit at 60 kD on SDS-PAGE (van Lis et al., 2003) but runs at 66 kD in Polytomella sp. (Fig. 1). Interestingly, it was inferred from the cDNA sequence of Polytomella ASA1 (PsASA1) that the protein shares 54% sequence identity with the C. reinhardtii ASA1 (CrASA1) but has a 52-residue insertion (L186-S238, precursor protein), which accounts for the 6-kD difference observed on SDS-PAGE.

In the low molecular mass range (<15 kD), the profiles are similar and consist of subunits-ASA5, -6, -8, -9, -e, and -c. C. reinhardtii ASA8 (9.9 kD) and ASA9 (12.1 kD) were newly identified on Gly SDS-PAGE using mass spectrometry (liquid chromatography-mass spectrometry/mass spectrometry) as part of an ongoing proteomics project that includes the C. reinhardtii ATP synthase separated on BN-PAGE (A. Atteia, A. Adrait, S. Bruguier, M. Tardiff, R. van Lis, L. Kuhn, O. Bastien, J. Garin, J. Joyard, and N. Rolland, unpublished data). CrASA8 was identified based on the similarity of its N-terminal sequence to that of

Figure 2. Subunits-α and -β in Chlamydomonas algae exhibit extensions. A, Multiple sequence alignment of the N-terminal part of mature α-subunits. Of the two predicted N-terminal α-helices, the second α-helix is amphipathic and likely interacts with the N terminus of the OSCP (Weber et al., 2004). The arrows indicate the position of unique Pro residues in the algal proteins: ●, conserved Glu residue substituted by a His residue in the algae; *, the presence of an unusual Cys residue in the algae. Sequences used are: BOVIN, Bos taurus (P19483); CHAVU, Chara vulgaris (Q7YAN8); CHLRE, C. reinhardtii (Q96550); MARPO, Marchantia polymorpha (P2685); PLASU, Platymonas subcordiformis (Q36517); POLSP, Polytomella sp. (CAI3486); PROWI, Prototheca wickerhamii (Q37628); RICPR, Rickettsia prowazekii (O50288); RHOSA, Rhodovalva salina (Q9G8W0); and YEAST (P00830). B, Multiple sequence alignment of the C-terminal part of different β-subunits. Sequences used are: CHLRE, C. reinhardtii (P38482); POLSP, Polytomella sp. (CAI3487); PLAF, Plasmidium falciparum (Q810V2); RICPR, R. prowazekii (O50290); ECOLI, E. coli (P0ABB4); YEAST (P00830); HUMAN, Homo sapiens (P04512); and ARATH, Arabidopsis (Q541W7). Note that the presence of Pro residues in predicted α-helices (as seen in the β extension) may in reality disrupt the helices.

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PsASA8. CrASA9 was inferred to correspond to PsASA9, because the complete sequences of all 17 C. reinhardtii ATP synthase subunits are known and the N-terminal sequences of all 17 Polytomella subunits, except PsASA9, were matched to their C. reinhardtii counterparts (Vázquez-Acevedo et al., 2006). The apparent molecular masses of PsASA9 and CrASA9 are approximately 9 and 10 kD, respectively.

Characteristics of the Chlamydomonad ASA Subunits

Using the complete amino acid sequences of all nine CrASA subunits that were retrieved from the C. reinhardtii genome sequence database (v3.0), no homologs were found after sequence similarity searches against nonredundant databases at the National Center for Biotechnology Information and others, including the genome sequences of the diatom *Thalassiosira pseudonana* and the green alga *Ostreococcus tauri* at the Joint Genome Institute (JGI). At present, the ASA subunits were found only in Chlorophycean algae (Vázquez-Acevedo et al., 2006). Conserved domain searches at the National Center for Biotechnology Information and different pattern and profile searches at the ExPASy proteomics server did not reveal any functional domains in the CrASA subunits that could hint to their role.

Based on the CrASA sequences, the set of uncharacterized ASA subunits covers a pI range of 5.66 to 9.27, a calculated molecular mass range of 60 kD (66 kD in *Polytomella* sp.) to 10 kD, and a grand average of hydrophobicity (GRAVY) score of −0.402 to 0.355 (Table I). ASA6 is predicted to exhibit two transmembrane (TM) segments, whereas ASA8 and ASA9 are largely hydrophilic proteins that, on the contrary, seem to contain one TM segment. Conversely, ASA2 is an overall hydrophilic protein that does not contain strongly predicted TM segments. ASA1, ASA4, and ASA6 possibly possess coiled-coil structures, which may be important for inter- or intrasubunit interactions. ASA5, ASA8, and ASA9 do not contain cleavable targeting sequences, which suggests their targeting to the ATP synthase complex directly via the intermembrane space instead of via the matrix (for review, see Herrmann and Neupert, 2003; Herrmann and Hell, 2005) or possibly via a differential translocation across the inner membrane, such as was described for the AAA protein BCS1 (Stan et al., 2003).

Both CrASA1 and PsASA1 are predicted to be soluble. This notion was supported by dissociation studies using mitochondrial membranes from *Polytomella* sp. Upon treatment of mitochondrial membranes with either heat (55°C, to dissociate the ATP synthase; see below), Na₂CO₃ (to release extrinsic proteins), or a combination of the two, ASA1 could be dissociated from the complex and released as a soluble protein, although heat alone results in relatively low levels of soluble ASA1 (Fig. 3A). The differential dissociation of ASA1 and the β-subunit could imply that ASA1 is anchored to the membrane independently from the F₁-ATPase. Solubility studies with the overexpressed ASA1 from both algae show that the protein is scarcely soluble at physiological pH (7–8). The protein is, however, soluble at high pH (with Na₂CO₃ or buffer only; shown in Fig. 3B for PsASA1), which supports its extrinsic nature. A hint that ASA1 is indeed not intrinsic comes from the fact that nonionic detergents such as Triton X-100 or dodecyl maltoside do not increase the solubility of overexpressed ASA1 (data not shown).

Heat Dissociation of the Strongly Dimeric ATP Synthase of *Polytomella* sp.

Unlike other known ATP synthases, those of *Polytomella* sp. and *C. reinhardtii* hardly dissociate into their F₁ and F₀ domains upon solubilization with

*Table 1. Physicochemical properties of the ASA subunits of the Chlamydomonad mitochondrial ATP synthase*

The presence of putative coiled-coil-forming sequences may indicate intersubunit interactions or homodimerization. N-term. (Res. No.), Residue number marking start of N terminus; MM, molecular mass calculated from sequence; GRAVY, higher scores indicate increasing hydrophobicity; TM prediction, indicates the number of predicted TM segments using SOSUI (S), TMpred (T); H, HMMTOP; Paircoil score, coiled-coil domain prediction; Ps, *Polytomella* sp.; Cr., *C. reinhardtii*.

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<th>Protein</th>
<th>N-Term. (Res. No.)</th>
<th>Protein Identification*</th>
<th>MM</th>
<th>pI</th>
<th>GRAVY</th>
<th>TM Prediction</th>
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*Gene model numbers of JGI Chlamy v3.0. bGenBank accession number.

dodecylmaltoside (Figs. 4 and 5A; Atteia et al., 2003; van Lis et al., 2003). Dissociation of the Polytomella ATP synthase dimer could, however, be observed after a short incubation of freshly isolated mitochondria at 55°C. The destabilization of the dimer was shown by BN-PAGE stained with Coomassie Blue and in-gel ATPase activity staining. The ATPase activity of untreated Polytomella mitochondria distributes into a main band of approximately 1,600 kD (Vd) and a faint dissociation product of approximately 800 kD (Vm) on BN gel (Fig. 4). After a 2-min heat treatment of Polytomella mitochondria, the intensity of Vm increased, while a new complex of approximately 400 kD (F1) appeared that showed ATPase activity. After 4 min exposure to 55°C, F1 was the major complex to exhibit ATPase activity (Fig. 4). Low amounts of ATP synthase monomer (Vm) are detected in untreated mitochondria, suggesting a slight dissociation that may be detergent induced. The effect of heat treatment on potato (Solanum tuberosum) mitochondria, used here as a control, was also followed (Fig. 4). The profile of potato protein complexes exhibiting ATPase activity shows the monomeric form of 580 kD and three different forms of the F1 domain at 350 to 450 kD that dissociate upon solubilization (Jänsch et al., 1996). Unlike for Polytomella sp., heat treatment did not alter significantly the presence of the potato ATP synthase (sub) complexes or the other oxidative phosphorylation (OXPHOS) complexes (Fig. 4). Also, for mitochondria from C. reinhardtii, no significant heat dissociation could be observed (data not shown). The difference in heat dissociation between Polytomella sp. and these other organisms may stem from differences in subunit structure and in membrane lipid composition.

Control and 2-min heat-treated mitochondria from Polytomella sp. were analyzed by two-dimensional (2D) BN/SDS-PAGE and silver staining using the Tricine system for its good resolution of smaller proteins (Schägger and von Jagow, 1987). The main changes induced by heat are the dissociation of the complexes that have large extramembrane entities, the ATP synthase and complex I, whereas complexes III and IV seemed relatively unaffected (Fig. 5, A and B). In the 2-min heat treatment 2D SDS-PAGE profile, the composition of Vm resembles that of Vd. Because Vm
includes all the typical mitochondrial ATP synthase subunits, it likely corresponds to the monomeric ATP synthase. The polypeptide profile of F₁ is typical of a F₁-ATPase with its five major subunits (Fig. 5, B and D). The behavior of ASA1 with respect to the β-subunit on 2D BN/SDS-PAGE after 2-min heat treatment was followed by immunoblotting. ASA1 was mainly detected in Vd and Vm, whereas low levels of ASA1 were found past Vm but not in F₂, indicating that some ASA1 may be associated in F₁ subcomplexes (Fig. 5C). However, clear evidence of this was not obtained from 2D SDS-PAGE analysis, likely because protein levels were too low. Most of subunit-β was present in Vd, Vm, and F₁, whereas a portion was found, unlike ASA1, in its free form at the bottom of the gel. As judged by the strong signal of the β-subunit in F₁, a sizable part of the ATP synthase had dissociated beyond the monomer after 2-min heat treatment. The ASA1 signal beyond Vm is proportionally much weaker than that of the β-subunit. It is therefore likely that an important fraction of ASA1 becomes insoluble when heat dissociation progresses beyond the monomeric form.

Although most subunits of Vd are present in Vm, it was found that in the latter form, two bands around 10 kD were missing. These subunits are thus hypothesized to be important for the dimerization of the ATP synthase or for the stabilization of the dimer. By comparing our Tricine SDS-PAGE subunit profile (Fig. 5D) to that reported previously (Vázquez-Acevedo et al., 2006), one subunit was identified as ASA6 (12 kD) and the other as ASA9 (9 kD), which is assumed to match ASA9 in C. reinhardtii (see above).

**DISCUSSION**

The F₁Fₒ-ATP synthases of *Polytomella* sp. and its photosynthetic counterpart *C. reinhardtii* share an atypical polypeptide composition. The unusual extensions that *C. reinhardtii* exhibit at the N and C termini of the catalytic subunits-α and -β, respectively (Franzén and Falk, 1992; Nurani and Franzen, 1996), are shown to be also present in *Polytomella* sp. The fact that the α- and β-subunit extensions of both algae are highly conserved suggests functional constraints. Based on sequence homology with the IF₁ inhibitor protein and the inhibitory peptide of the *E. coli* e-subunit, a role of the β extension as reversible inhibitor of ATP hydrolysis was hypothesized (Atteia et al., 1997). Based on new sequence and secondary structure data for IF₁ and subunit-e, this could, however, not be further confirmed. The possibility exists that the β extension is required to interact with the ASA proteins.

In the F₁Fₒ-ATP synthase, the N termini of the α-subunit and the OSCP interact (Carbajo et al., 2005). It was proposed that four conserved residues at the N terminus of the OSCP and its equivalent in chloroplast and bacteria, subunit-δ, are important for the interaction with the α-subunit (Y10, A11, A13, R84, *E. coli* numbering; Hong and Pedersen, 2003). Surprisingly, these residues are not present in the OSCP of the chlorophytes, which suggests a different interaction with the α-subunit. Twin Pro at positions 41 and 42 may enable a specific reorientation to accommodate this atypical interaction, in which the α extension and/or ASA proteins may play a role.

The occurrence of the ASA proteins and the lack of typical subunits of the peripheral stalk in the mitochondrial ATP synthase, an enzyme that has been extensively studied, are puzzling. Because these atypical subunits lack significant sequence homology to known proteins, questions arise as to their function and localization within the complex and whether they are genuine subunits. In several studies on the ATP synthase of *C. reinhardtii* and *Polytomella* sp. using whole mitochondria, mitochondrial membranes, or the isolated complex, the ASA proteins were consistently found in the ATP synthase subunit profile on 2D BN/SDS-PAGE in a similar stoichiometry (Atteia et al., 2003; van Lis et al., 2003, 2005; Vázquez-Acevedo et al., 2006). In addition, in *Polytomella*, the respective amounts of ATP synthase and respiratory complexes vary with the pH of the growth medium, but this does not affect the presence of the ASA proteins (Atteia et al., 2003). Taken together, these data suggest that the ASA proteins are genuine subunits of the algal ATP synthases.

Different indirect clues exist as to the localization and function of at least some of the ASA subunits in the ATP synthase complex. Of the subunits that constitute the peripheral stalk in the bovine ATP synthase (OSCP, b, d, and F6), only the OSCP is present in the algae; it therefore follows that ASA subunits fulfill a structural role in forming part of the peripheral stalk. A typical mitochondrial subunit-b (approximately 20 kD) contains two TM segments at its N terminus, while the rest of the protein is hydrophilic (Arnold et al., 1998; Burger et al., 2003; Heazlewood et al., 2003). CrASAT (13 kD) possibly contains an N-terminal TM segment with the remainder of the proteins being hydrophilic; the protein may be anchored in the membrane and protruding into the matrix, similar to subunit-b. Electron microscopy images of dimeric ATP synthase from *Polytomella* sp. show a very pronounced peripheral stalk or possibly two stalks (Dudkina et al., 2005). In comparison, the yeast ATP synthase possesses no pronounced peripheral stalks but a considerably more bulky F₆ domain. It seems thus that the *Polytomella* membrane domain is too small to host the larger ASA subunits, which are more likely part of the observed pronounced peripheral stalk(s). For the *Polytomella* enzyme, a large globular domain is visible at the apex of the F₆ domain that may be attributed to ASA1. The fact that we established ASA1 to be a soluble protein adds to the notion that the protein most likely has a considerable portion of its surface exposed to the solvent, as seems to be the case with the observed globular domain.

The dissociation of the *Polytomella* ATP synthase dimer (Vd) by heat treatment allowed to have insights into the composition of the monomer (Vm). Vm (800 kD)
contains seven ASA subunits (1, 2, 3, 4, 7, 5, and 8), which causes its apparent molecular mass to be at least 200 kD greater than the ATP synthase monomer of yeast, beef, and Arabidopsis (*Arabidopsis thaliana*). Two subunits found in Vd, ASA6 and ASA9, were lacking in Vm and are therefore proposed to be involved in the dimerization. Dimer-specific PsASA9 is assumed to match CrASA9, identified by mass spectrometry (A. Atteia, A. Adrait, S. Brugi`ere, M. Tardiff, R. van Lis, L. Kuhn, O. Bastien, J. Garin, J. Joyard, and N. Rolland, unpublished data). Both ASA6 and ASA9 are predicted to be membrane bound. In yeast, GXXXG motifs inside the single TM segments of dimer-specific subunits-g and -e are essential for ATP synthase dimerization, whereas the coiled-coil structure of subunit-e stabilizes the dimer (Arselin et al., 2003; Everard-Gigot et al., 2005). In ASA6, a possible coiled coil is predicted that could function in the dimerization. In the predicted TM segments of ASA6 and ASA9, no GXXXG motifs are present. In addition, no other specific elements were identified that could point to their role in dimer stability, which should thus be further assessed in *Polytomella* sp. using biochemical methods.

Heat treatment (60°C) was also used by Vázquez-Acevedo et al. (2006) to dissociate the purified *Polytomella* ATP synthase. Although heat seems to have a similar effect on the ATP synthase whether it is in the mitochondrial membrane or purified, discrepancies exist between the outcomes of the two approaches. First, the above-mentioned authors stated that Vd was identical to Vm (although this was not clearly shown), whereas we show Vm to be devoid of ASA6 and ASA9. The presence of dodecyl maltoside and glycerol in the preparations of purified ATP synthase can affect subunit interactions. Glycerol is known to enhance hydrophobic interactions, whereas dodecyl maltoside may favor TM segment association when used close to the critical micelle concentration of 0.2 mM (Fisher et al., 2003), possibly causing a different dissociation behavior of ASA6 and ASA9. Second, the authors observed an ASA1/3/5/8/a/c10 subcomplex of 200 kD that we have, however, never seen during several years of heat dissociation experiments; immunodetection of ASA1 at or near the expected position of the subcomplex on 2D BN/SDS-PAGE in 2-min heat-treated mitochondria is negligible (this work). The ASA1/3/5/8/a/c10 subcomplex released upon heat treatment in the presence of detergent is apparently soluble, but when membranes are heat treated, the subcomplex likely aggregates, because it cannot be solubilized from the membranes and observed on BN-PAGE.

**Figure 5.** 2D BN/SDS-PAGE analysis of *Polytomella* mitochondria incubated at 55°C. A to C, A whole lane from a BN gel was loaded on a Tricine denaturing gel (13% acrylamide). A, No heat treatment. B, A 2-min heat treatment. Arrows indicate the subunits of the F1-ATPase. C, Immunoblotting of proteins on 2D gels after 2-min heat treatment. Pieces from a BN gel containing the different ATP synthase forms that were present after 2 min of heat treatment were cut out and loaded on Tricine-SDS-PAGE. Dimer-specific subunits ASA6 and ASA9 are boxed. Roman numbers I to V, Vd, Vm, as in Figure 4; ?, not identified and only detected by silver staining; *, contaminating bands resulting from overlapping protein components in the 2D protein profile. Subunit assignment on Tricine SDS-PAGE was confirmed from Vázquez-Acevedo et al. (2006).
Figure 6. Working model of dimeric mitochondrial ATP synthase in Chlamydomonad algae, based on our current knowledge of the enzyme from both C. reinhardtii and Polytomella sp. The monomers are shown rotated 180° one from another (around a vertical axis), as proposed previously for the yeast enzyme (Paumard et al., 2002). Arrows indicate the α and β extensions. The α extension is shown interacting with the OSCP and the β extension with ASA subunits of the peripheral stalk. ASA1 is placed in the matrix because it is an extrinsic protein. ASA2 is a hydrophilic protein and probably associated to the membrane. ASA3 is somewhat hydrophilic but without specific features. ASA4 is a possible homodimer-forming subunit based on its prediction for coiled-coil formation (Table I). ASA5 may be anchored in the membrane and protruding into the matrix. ASA6 and ASA9 are proposed to be dimer specific and membrane bound, whereas ASA7 and ASA8 may be associated to or traversing the membrane.

Based on our analysis of the characteristics of the ATP synthase subunits, which are assumed to be similar in both algae, an attempt has been made to provide structural details to the working model proposed by Dudkina et al. (2005, 2006) involving the presence of two peripheral stalks (Fig. 6). Vázquez-Acevedo et al. (2006) also propose a model of the Polytomella ATP synthase that differs significantly in that it places the ASA1/3/5/8/a/c10 subcomplex entirely in the membrane and ASA subunits 2, 4, and 7 forming a single thin peripheral stalk. We propose that the ASA1/3/5/8/a/c10 subcomplex constitutes the (double) peripheral stalk mostly intact, with the other ASA subunits elsewhere in the complex (Fig. 6, legend). This may help to imagine why the subcomplex stalk is destabilized in the absence of detergent: the large matrix-exposed and membrane-bound moieties in the stalk could easily aggregate when the interaction with F1-ATPase is disrupted by heat treatment. Complexes without large extramembrane portions such as complex III and IV seem not notably affected by heat.

Although the ATP synthases of C. reinhardtii and Polytomella sp. are expected to be similar, the fact that the C. reinhardtii ATP synthase dimer is not readily dissociated by heat illustrates that differences exist. Also, PsASA1 and CrASA1 are highly similar in the algae, but PsASA1 contains a 52-residue insertion (this work). It is therefore desirable to obtain the complete sequences of the PsASA subunits. There are other examples of differences in the respiratory chain of the two algae, which are thought to reflect their evolutionary divergence from a photosynthetic ancestor (Round, 1980; van Lis et al., 2005).

Although questions remain as to what is the advantage of a highly stable dimer, the atypical Chlamydomonad mitochondrial ATP synthase is expected to bring new incentives to the matured field of ATP synthase research.

**MATERIALS AND METHODS**

**Isolation of Mitochondria**

Mitochondria were isolated from Polytomella sp. cells (198.80, E.G. Pringsheim) grown on acetate (van Lis et al., 2005) and from Chlamydomonas reinhardtii cells grown on Tris-acetate phosphate medium as previously described (Eriksson et al., 1995). Potato (Solanum tuberosum) tuber mitochondria were isolated as reported (von Stedingk et al., 1997), omitting the Percoll gradient step. Protein concentration was determined using the bicinchoninic acid kit (Pierce). Mitochondrial membranes from Polytomella sp. were prepared as described (Atteia et al., 2003).

**BN-PAGE Analysis and ATPase Staining**

Isolated mitochondria were washed in 0.25 m sorbitol, 15 m Bis-Tris, pH 7.0, solubilized with 2% (w/v) dodecyl maltoside (4 g detergent/g protein), and supplemented with 0.5% (w/v) Coomassie Blue after ultracentrifugation for 20 min at 60,000g. Subsequently, BN-PAGE was done as described (Schägger and von Jagow, 1991; Atteia et al., 2003). For in-gel ATPase activity, BN gels were rinsed twice in HEPES-KOH 50 mM, pH 8.0, and then incubated in this buffer containing 30 mM CaCl2 and 10 mM ATP at room temperature. Calcium phosphate precipitates appear within 20 min and continue to intensify up to 1 h.

**2D Gel Electrophoresis, Immunoblotting, and N-Terminal Protein Sequencing**

2D analysis of BN gel lanes containing mitochondrial proteins of Polytomella sp. and C. reinhardtii was done using 15% Gly SDS-polyacrylamide gels (Laemmli, 1970) or 13% Tricine SDS-polyacrylamide gels (Schägger and von Jagow, 1987). For N-terminal protein sequencing, the gels were blotted onto polyvinylidene difluoride membrane (Millipore) and stained with Porceen Red. The proteins were subjected to N-terminal sequencing by automated Edman degradation using a LFF3000 Beckman sequencer. For immunoblotting, the proteins from Tricine 2D gels were transferred onto nitrocellulose and decorated with the antibodies indicated in Figure 5. The antibodies against the β-subunit and COX2a were previously described (van Lis et al., 2005; Atteia et al., 2006).

**Dissociation Studies**

Isolated mitochondria were resuspended in their own breaking buffer at a concentration of 25 mg protein/mL. Aliquots of 3 mg of mitochondria in breaking buffer, containing protease inhibitors phenylmethylsulfonyl fluoride (0.1 mM), benzamidine (0.5 mM), and hexanoic acid (1 mM), were treated at 55°C for the indicated times and placed immediately on ice. Subsequently, the samples were processed for BN-PAGE as described above. For ASA1 dissociation studies, mitochondrial membranes from Polytomella sp. were resuspended at a concentration of 1.7 mg/mL, either in PM buffer (5 mM potassium phosphate, pH 7.0, 200 mM mannitol) or in 100 mM Na2CO3, pH 11.4, with the above-mentioned protease inhibitors added. Membranes, part of
which were treated at 55°C for 2 min, were left on ice for 30 min with occasional vortexing. The membranes were then centrifuged at 100,000 g for 30 min, after which the supernatant was kept, whereas the membranes were collected after washing once in PM buffer. The protein samples were separated using Gly SDS-PAGE (12% acrylamide), transferred onto nitrocellulose, and analyzed by Ponceau Red staining and subsequent immunoblotting.

Screening of a Polytomella cDNA Library for ATP Synthase Subunits

Screening of a Polytomella k-ZAPII cDNA library (Atteia et al., 2005) was carried out by plaque hybridization following standard protocols, as using the cDNAs coding for the corresponding subunits of C. reinhardtii. The isolated cDNAs were sequenced and deposited in the DDBJ/EMBL/GenBank databases.

Overexpression, Antibody Production, and Solubility Assays of ASA1

Using the cDNA of ASA1 as template, a PCR product containing the coding sequence for the entire mature protein from C. reinhardtii was obtained with the forward primer 5′-GGAATTCATATGGATGACCGGCGCTG-3′ and reverse primer 5′-ATGCTGCAGCGAGCGCCGCGCCG-3′, containing, respectively, the Ndel and XhoI restriction sites (underlined) for subsequent cloning. The PCR product was cloned into the expression vector pET24a (Novagen). A PCR product containing the coding sequence for the entire mature protein from Polytomella sp. was obtained with forward primer 5′-GATCCATGGTTACCTTGCCCCCCTCCGC-3′ and reverse primer 5′-CTAAAGCATTCTAGTGGACCGCCCGCC-3′, containing, respectively, the Ncol and HindIII restriction sites (underlined) for subsequent cloning. The PCR product was first cloned into the vector pQE60 (Qiagen), from which it was excised using restriction enzymes Ncol and HindIII (partial restriction, as HindIII cuts once in the PsaASA1 cDNA). This fragment now contained a C-terminal His tag, which was then cloned into expression vector pACYC-Duet (Novagen). Overexpression of CrASA1 and PsASA1 was done using the Escherichia coli strain BL21 (DE3) (Stratagene) at 37°C for 4 h, induced by 1 mM isopropyl β-D-1-thiogalactopyranoside. For antibody production, CrASA1 was purified using nickel-nitrilotriacetic acid agarose resin (Sigma) according to standard protocols but in the presence of 2 mM urea to maintain complete solubility of the protein. Antibodies against CrASA1 were produced in rabbit at Charles River Laboratories. For solubility studies, E. coli cell pellets overexpressing ASA1 were resuspended in lysis buffer containing 300 mM NaCl, 1 mg/mL lysozyme, and 1 mM phenylmethylsulfonyl fluoride in addition to either 20 mM Tris, pH 8.0, 100 mM NaClO4, pH 11.4, or 50 mM 3-(cyclhexylamino)propanesulfonic acid (CAPS), pH 11.0, incubated for 30 min on ice and lysed using sonication. Additives such as nonionic detergents were used with Tris buffer at pH 8.0. After centrifugation, the supernatant was kept, whereas the membranes were collected after washing once in PM buffer and analyzed by Ponceau Red staining and subsequent immunoblotting.

Sequence Analysis

Molecular mass, pl, GRAVY scores, and amino acid composition were determined using the ProtParam program. For the prediction of secondary structures, the SOPMA program was used (Combet et al., 2000), and the Paircoil program was used for the prediction of coiled coil regions (Berger et al., 1995). For TM segment prediction, the SOSUI (Hirokawa et al., 1998), TMPred (Hofmann and Stofel, 1993), and HMMTOP (Tusnady and Simon, 2001) programs were applied. Multiple sequence alignments were performed with ClustalW (Thompson et al., 1997) and refined manually. These software packages are all available from the ExPaSy proteomics Web site. For the analysis of α-helices, use was made of the helical wheel projection and amphipathic moment plot options of the program Winpep v.3.01 (Hennig, 1999). Representations of multiple alignments were done using the BOX-SHADE program. The genome sequences of C. reinhardtii (v 3.0), Thalassiosira pseudonana, and Ostreococcus tauri are available at the Joint Genome Institute.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers CAI3486 (α-subunit), CAI3487 (β-subunit), CAF6302 (γ-subunit), and CAD90138 (ASA1).

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