Chaperones of F_1-ATPase

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Mitochondrial F_1-ATPase contains a hexamer of alternating α and β subunits. The assembly of this structure requires two specialized chaperones, Atp11p and Atp12p, that bind transiently to β and α. In the absence of Atp11p and Atp12p, the hexamer is not formed, and α and β precipitate as large insoluble aggregates. An early model for the mechanism of chaperone-mediated F_1 assembly (Wang, Z. G., Sheluho, D., Gatti, D. L., and Ackerman, S. H. (2000) EMBO J. 19, 1486–1493) hypothesized that the chaperones themselves look very much like the α and β subunits, and proposed an exchange of Atp11p for α and of Atp12p for β; the driving force for the exchange was expected to be a higher affinity of α and β for each other than for the respective chaperone partners. One important feature of this model was the prediction that as long as Atp11p is bound to β and Atp12p is bound to α, the two F_1 subunits cannot interact at either the catalytic site or the noncatalytic site interface. Here we present the structures of Atp11p from Candida glabrata and Atp12p from Paracoccus denitrificans, and we show that some features of the Wang model are correct, namely that binding of the chaperones to α and β prevents further interactions between these F_1 subunits. However, Atp11p and Atp12p do not resemble α or β, and it is instead the F_1 γ subunit that initiates the release of the chaperones from α and β and their further assembly into the mature complex.

Mitochondrial F_1-ATPase consists of three α and three β subunits occupying alternate positions in a hexamer that surrounds a rod-like element containing one each of γ, δ, and ε subunits (1–3). Three nucleotide-binding catalytic sites (CS) and three noncatalytic sites (NCS) alternate at the six α/β interfaces. Early work with respiratory-deficient strains of Saccharomyces cerevisiae (4) revealed that two additional mitochondrial proteins, Atp11p and Atp12p, which are not integral subunits of the enzyme, are nonetheless necessary for the assembly of F_1-ATPase. Besides their failure to assemble F_1, a particularly interesting feature of atp11 and atp12 mutants is that they accumulate α and β subunits as high molecular weight aggregates (4) that can be recognized as densely stained inclusion bodies in the mitochondrial matrix (5). Subsequent studies in yeast have shown that Atp12p binds to F_1 α (6) and that Atp11p binds to β (7); these interactions include binding determinants in the nucleotide binding domains (NBD) of the two F_1 subunits. On this basis, it is now recognized that Atp11p and Atp12p are members of two new families of molecular chaperones, pfam06644 and pfam07542 (8), which are required for the assembly of mitochondrial ATP synthase in all eukaryotes. In fact, the first nuclear genetic lesion associated to a defect of mitochondrial ATP synthase in humans was identified in the locus ATPA12F for Atp12p and was responsible for the death of a 14-month-old infant (9). Atp12p is also present in the α subdivision of Proteobacteria, consistent with the proposed origin of mitochondria from this ancestral line (10).

The nature of the interactions between the F_1 subunits and Atp11p and Atp12p has remained elusive because of the lack of structural information for these chaperones. As α and β aggregate in the absence of Atp11p and Atp12p, it is usually assumed that the F_1 subunits are themselves poorly soluble, and that the two chaperones maintain them in a dispersed state until they are incorporated in the mature enzyme. Based on the analysis of the distribution of hydrophilic and hydrophobic areas on the surface of the α and β subunits of F_1, and on the interaction energies between these subunits at the interfaces that provide the CS and NCS sites, Wang et al. (6) have proposed a model of F_1 assembly in which Atp11p binds at the region of the β subunit that contributes to the CS site, and Atp12p binds at the region of the α subunit that contributes to the NCS site. One consequence of this particular binding of Atp11p and Atp12p to the F_1 subunits is that as long as Atp11p is bound to β and Atp12p is bound to α, the two F_1 subunits cannot interact at either the CS or the NCS interface. Since no other modulators of chaperone release are known, the Wang model requires an exchange of Atp11p for α and of Atp12p for β. Implied in this model is that the chaperones must themselves look very much like the α and β subunits, and that the driving force for the exchange must simply be a higher affinity of α and β for each other than for the respective chaperone partners. Here we present the structures of Atp11p from Candida glabrata and...
Atp11p from Paracoccus denitrificans, and we show that some features of the Wang model are correct, namely that binding of the chaperones to $H_9251$ and $H_9252$ prevents further interactions between these $F_1$ subunits. However, Atp11p and Atp12p do not resemble $H_9251$ or $H_9252$, and it is instead the $F_1$ subunit that initiates the release of the chaperones from $H_9251$ and $H_9252$ and their further assembly into mature complex.

**EXPERIMENTAL PROCEDURES**

Protein Purification and Structure Determination—Mature (leaderless) C. glabrata Atp11p was produced in Escherichia coli Rosetta (DE3) cells at 37 °C. The 5′-end of the recombinant gene (Table 1) encodes the sequence Met-Gly-Gly-Ser-(His)$_6$-Ser-Ser-Gly in front of residues 24–309 of Atp11p, such that His tag-Atp11p is produced without the 23-amino acid targeting peptide that is removed when the yeast protein is imported into mitochondria. The protein was purified via Talon Cobalt affinity column (Clontech). A selenomethionine derivative was produced in modified LeMaster media containing selenomethionine in place of Met. Both native and selenomethionine C. glabrata Atp11p crystallize in 30% PEG 400, 100 mM sodium acetate, 100 mM MES, pH 6.5, over 2–4 weeks at room temperature by sitting, hanging, or sandwich drops. The Atp11p structure was determined at 1.8 Å by single wavelength anomalous dispersion phasing with SOLVE/RESOLVE (11) at the selenium absorption peak with data collected at the APS beamline 22ID (SER-CAT). The resolution was extended to 1.5 Å with new data from the same beamline, and the structure was refined with SHELX (12) (Table 2).

Full-length P. denitrificans Atp12p was produced in E. coli from a plasmid that encodes an upstream His$_6$ tag sequence and a thrombin recognition site (Table 1) and purified via Talon affinity chromatography. A selenomethionine derivative of this protein was produced as described for C. glabrata Atp11p. Crystals grow from around 5–20% PEG 4000, 50–250 mM sodium acetate, 100 mM HEPES, pH 7.5, at 4 °C. The Atp12p structure was determined at 1.9 Å by single wavelength anomalous dispersion phasing with SOLVE/RESOLVE (11) at the selenium absorption peak with data collected at the APS beamline 22ID (SER-CAT). A different crystal form was obtained from 28 to 30% PEG 4000, 50 mM MgCl$_2$, 100 mM Tris-Cl, pH 8.5, in sitting drops at 4 °C. A data set at the resolution of 1.0 Å was collected at the APS beamline 21ID-F (LS-CAT). The structure was determined at 1.8 Å by Molecular Replacement with CCP4/Molrep (13), and the resolution was extended to 1.0 Å with SHELX (Table 2). Crystals of D202K Atp12p grow around 20% PEG 4000, 50–250 mM sodium acetate, 100 mM HEPES, pH 7.5, at 4 °C with 20 mM spermidine as additive. A data set at 1.8 Å was collected at the APS beamline 21ID-G (LS-CAT), and the structure refined with CNS version 1.2 (14) (Table 2).

Electrostatic Potentials—Electrostatic potentials at the solvent-accessible surface of proteins were calculated with PDB2PQR/APBS (15) using a nonlinear Poisson-Boltzmann continuum model with protein dielectric constant $\varepsilon_p = 2$, solvent dielectric constant $\varepsilon_s = 78$. Protein charges were assigned on the basis of the PARSE force field (16), assuming a pH of 8.0 for the mitochondrial matrix (17, 18), and by calculating the derivative was produced in modified LeMaster media containing selenomethionine in place of Met. Both native and selenomethionine C. glabrata Atp11p crystalize in 30% PEG 4000, 100 mM sodium acetate, 100 mM MES, pH 6.5, over 2–4 weeks at room temperature by sitting, hanging, or sandwich drops. The Atp11p structure was determined at 1.8 Å by single wavelength anomalous dispersion phasing with SOLVE/RESOLVE (11) at the selenium absorption peak with data collected at the APS beamline 22ID (SER-CAT). The resolution was extended to 1.5 Å with new data from the same beamline, and the structure was refined with SHELX (12) (Table 2).
TABLE 1
Yeast strains and plasmids

| Yeast         | Genotype               |
|---------------|------------------------|
| W303-1A*      | MAT$a$ ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 |
| $\Delta$atp11* | MAT$a$ ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 atp12::His3 |
| $\Delta$atp12* | MAT$a$ ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 atp11::His3 |
| $\Delta$γ*    | MAT$a$ ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 atp12::LEU2 |
|               | MAT$a$ ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 atp3::His3 |

| Plasmid       | Description                                      |
|---------------|--------------------------------------------------|
| pET28-cgap11  | E. coli expression plasmid pET28a. Encodes C. glabrata Atp11p without the mitochondrial leader peptide and with a His tag at the N terminus |
| pET28-pdatp12  | E. coli expression plasmid pET28a. Encodes P. denitrificans Atp12p with a His tag and thrombin cleavage site at the N terminus |
| pET28-pdatp12(D202K) | Identical to pET28-pdatp12 with the exception of a D202K mutation that was created by PCR mutagenesis using the QuikChange protocol |
| YEp-cgap11    | Yeast/E. coli shuttle plasmid YEp352 (2 $\mu$) with URA3 marker gene. Encodes full-length C. glabrata Atp11p |
| YEp-pdatp12    | Yeast/E. coli shuttle plasmid YEp352 (2 $\mu$) with URA3 marker gene. Encodes P. denitrificans Atp12p fused to the C terminus of the S. cerevisiae Atp11p mitochondrial leader peptide |
| pRS314-pdatp12 | Yeast/E. coli shuttle plasmid pRS314 (CEN) with URA3 marker gene. Encodes P. denitrificans Atp12p peptide |
| pαFLAG        | Yeast/E. coli shuttle plasmid YEp352 (2 $\mu$) with URA3 marker gene. Encodes S. cerevisiae F1 $\alpha$ subunit with FLAG epitope fused to the C terminus |
| pβHIS6*       | Yeast/E. coli shuttle plasmid pRS314 (CEN) with TRP1 marker gene. Encodes a chimeric protein composed of the S. cerevisiae OSCP mitochondrial leader peptide (targeting signal) followed by the S. cerevisiae F1 $\beta$ subunit that bears a His tag at the N terminus |

*From R. Rothstein, Department of Human Genetics, Columbia University.
*From Ref. 37.
*From Ref. 38.
*From Ref. 39.
*From Ref. 40.

pK$\alpha$ values of all ionizable residues with the program PROPKA (19). Calculations were carried out on a uniform grid of −0.5 Å spacing.

**Coprecipitation Experiments**—These experiments utilized the yeast mutant, $\Delta$γ/$\gamma$-pαFLAG+$\gamma$HIS6, which harbors a disrupted allele ($atp3::HIS3$) for F1 $\gamma$ in the chromosome and carries two episomal plasmids, pαFLAG (URA3-linked) and pβHIS6 (TRP1-linked), for the simultaneous production of, respectively, F1 $\alpha$ that is tagged at the C terminus with the FLAG epitope (DYKDDDDK) and F1 $\beta$ that is tagged at the N terminus with 6 histidine residues (see Table 1). Plasmid pβHIS6 was a gift from Dr. David Mueller (Rosalind Franklin University of Medicine and Science, Department of Biochemistry and Molecular Biology, The Chicago Medical School, North Chicago, IL). To make pαFLAG, the QuikChange method was used to insert the DNA coding for the FLAG epitope immediately in front of the TAA stop codon of yeast ATP1. The plasmid template for mutagenesis (pG500/ST2) carries the entire ATP1 coding sequence for yeast F1 $\alpha$ cloned in the BamHI and EcoRI sites of YEp352 (20). The sequences of the complementary primers that were used for mutagenesis are as follows: 5′-GCTACCTGAATCTTTTGC-3′ (sense strand) and 5′-ttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttttt
with TALON metal affinity resin, 75 μl of soluble mitochondrial extract was combined with 30 μl of resin (prepared as a 50% slurry) in a final volume of 500 μl of buffer that contained 50 mM sodium phosphate, pH 7.0, 300 mM NaCl, 0.5% Triton X-100, 4 mM ATP, and 5% (w/v) sucrose. Experimental incubations with anti-FLAG M2 affinity gel were identical except that the pH was increased to 7.4 and NaCl was reduced to 150 mM. Samples were incubated end-over-end at 4 °C overnight and subsequently centrifuged for 5 min at 13,000 × g to sediment the affinity beads. The initial (post-bead) supernatant was collected from the post-bead supernatant and first wash fractions by precipitating the samples (500 μl) with a 1:4 mixture chloroform:methanol. Chloroform:methanol and affinity bead precipitates were suspended in 45 μl of SDS gel loading buffer, boiled for 5 min, centrifuged, and loaded (10 μl) on a 12% SDS-polyacrylamide gel.

Miscellaneous Methods—Yeast transformations employed the lithium acetate procedure (22). The detection in Western blots of F1, α, F1 β, Atp11p, and Atp12p and assays for molecular weight standards (hemoglobin and lipoamide dehydrogenase) used in sucrose gradients were performed as described previously (6, 7).

RESULTS

Crystals of C. glabrata Atp11p and P. denitrificans Atp12p were obtained after screening several homologous proteins, including those from Homo sapiens, Mus musculus, S. cerevisiae, Candida albicans, Kluyveromyces lactis, Rhodobacter capsulatus, and Arabidopsis thaliana. C. glabrata Atp11p and P. denitrificans Atp12p are 79.9 and 44.2% similar and 56.3% and 23.4% identical to the corresponding proteins in S. cerevisiae (Fig. 1). When produced from episomal plasmids C. glabrata Atp11p and P. denitrificans Atp12p rescue the respiratory defect of S. cerevisiae atp11 or atp12 deletion strains (Table 1; Fig. 2), suggesting that the mechanism by which the chaperones facilitate the assembly of F1 is maintained across evolutionary lines.

Structure of Atp11p—The crystal structure of C. glabrata Atp11p was initially determined at the resolution of 1.8 Å by single wavelength anomalous dispersion phasing of diffraction data from a single crystal of selenomethionine derivatized recombinant Atp11p. The resolution was subsequently extended to 1.5 Å with diffraction data from a crystal of native Atp11p (Fig. 3A; Table 2). Although the full-length protein produced in E. coli can be recovered from the crystals, the first 93 residues (out of 298, PDB numbering) are not visible in the crystal structure. However, there is an empty cavity in the density map that is large enough to harbor the disordered domain. The remaining residues 94–298 correspond to the part of S. cerevisiae Atp11p that was shown by deletion studies to retain full chaperone activity (23). The elements of sequence identity between the C. glabrata and the S. cerevisiae protein (Fig. 1) are distributed evenly throughout this region (Fig. 3B). A loop (residues 163–176) that is poorly conserved or absent in Atp11p from different sources is also disordered. The overall fold of Atp11p resembles an α/β "taco" (an incomplete antiparallel...
Chaperones of $F_1$-ATPase

TABLE 2

X-ray data collection and refinement statistics

Each data set was collected from a single crystal at 100 K and processed with HKL2000 (41) or XDS (42, 43). $R_{meas} = \Sigma |F_{obs} - F_{calc}|/\Sigma |F_{obs}|$, $R_{ave}$ was calculated on 10% (CNS) or 5% (SHELX) of the data omitted from refinement. In all the structures values in parentheses refer to the highest resolution shell.

| Atp11p | Atp12p | Atp12p | D202K Atp12p |
|--------|--------|--------|--------------|
| Data collection/reduction | SER-CAT/HKL2000 | SER-CAT/HKL2000 | LS-CAT/XDS | LS-CAT/HKL2000 |
| Space group | I4 | P1 | P2, 2, 2 | P1 |
| Cell dimensions | $a = b = 63.187 \text{ Å}$ | $a = 42.85 \text{ Å}$ | $a = 43.55 \text{ Å}$ | $a = 43.160 \text{ Å}$ |
| | $c = 153.577 \text{ Å}$ | $b = 50.652 \text{ Å}$ | $b = 51.047 \text{ Å}$ | $b = c = 67.117 \text{ Å}$ |
| Resolution | 50-1.5 Å (1.5-1.5 Å) | 50-1.9 Å (2.02-1.9 Å) | 50-1.0 Å (1.06-1.0 Å) | 50-1.8 Å (1.83-1.8 Å) |
| Unique reflections | 44,482 (2,450) | 38,168 (4,842) | 127,208 (12,549) | 48,230 (2,304) |
| (Redundancy) | 10.3 (3.6) | 3.8 (3.2) | 14.5 (14.3) | 3.8 (3.6) |
| Completeness | 92.8% (51.2%) | 97.3% (92.4%) | 99.8% (99.9%) | 97.5% (95.6%) |
| $\langle f_{o} - f_{c} \rangle$ | 30.5 (3.1) | 23.3 (1.7) | 25.4 (4.0) | 15.3 (2.3) |
| $R_{merge}$ | 5.5% (34.6%) | 6.0% (53.2%) | 6.2% (74.5%) | 8.2% (55.5%) |
| Refinement | SHELEX | CNS | SHELEX | CNS |
| PDB entry | 2P4F | 2P4A | 2P4I | 2P4D |
| $R_{merge}$ | 14.1% (26.8%) | 18.7% (29.6%) | 11.9% (21.8%) | 22.5% (29.0%) |
| $R_{free}$ | 17.5% (NA$^*$) | 23.0% (36.9%) | 14.2% (NA) | 27.9 (34.3) |
| Amino acids | 204 | 468 | 235 | 468 |
| Waters | 170 | 349 | 527 | 534 |
| ($B$) | 30.2 Å$^2$ | 41.2 Å$^2$ | 16.7 Å$^2$ | 28.8 Å$^2$ |
| $B$ from Wilson plot | 18.8 Å$^2$ | 22.6 Å$^2$ | 5.9 Å$^2$ | 20.7 Å$^2$ |
| Coordinate error | 0.08 Å | 0.22 Å | 0.05 Å | 0.24 Å |
| r.m.s.d. bond length | 0.013 Å | 0.009 Å | 0.014 Å | 0.009 Å |
| r.m.s.d. angles | 2.4 | 1.4 | 2.4 | 1.4 |
| r.m.s.d. dihedral | 22.3 | 20.9 | 22.1 | 20.7 |
| r.m.s.d. improper | 1.27 | 0.99 | 1.96 | 1.01 |
| Ramachandran | Preferred | 97.33% | 96.76% | 97.50% | 95.69% |
| | Allowed | 2.14% | 2.16% | 1.88% | 3.02% |
| | Outliers | 0.53% | 1.08% | 0.62% | 1.29% |

*Free $R$ value in the high resolution shell is not available in SHELEX. NA means not applicable.

β-barrel whose concave face is filled by two α-helices) flanked by two helical domains. Only one chain of Atp11p is present in the asymmetric unit, and a survey of all symmetry mates with PISA (24) does not reveal a dimer or a higher order assembly that might be stable in solution. This conclusion is consistent with earlier sedimentation analysis of purified recombinant yeast Atp11p and of native yeast Atp11p in mitochondrial extracts, which concluded that Atp11p is a monomer (25). Structural homology searches against the entire PDB with the VAST (26) and MATRAS (27) engines do not reveal relatives for the entire protein. Searches with the individual domains reveal poor structural matches of the first helical domain (residues 94–130) with the 3-helix bundle headpiece of villin (PDB code 1WY3), of the α/β taco (residues 131–262) with the C-terminal domain of phosphoglucomutase (residues 423–547 in PDB code 3PMG), and of the C-terminal helical domain (residues 263–298) with the two C-terminal helices of E. coli HSP90 (PDB code 1SF8). At this point it does not appear that any of these weak structural homologies might be of mechanistic importance.

Structure of Atp12p—The structure of P. denitrificans Atp12p was determined from two crystal forms at the resolution of 1.9 Å (sg P1, crystal form I) and 1.0 Å (sg P2, 2, 2, crystal form II) (Table 2). In the asymmetric unit of crystal form I, there are two chains of Atp12p related by a local 2-fold axis, but they are loosely packed and unlikely to represent a functional dimer. A single chain occupies the asymmetric unit of crystal form II, with no neighbors in the unit cell in a position suitable to form a dimer with a large buried surface. Differences between the two crystal forms of Atp12p are distributed throughout the structure (C-α r.m.s.d. 3–238 = 0.782 Å; r.m.s.d. 3–83 = 0.685 Å; r.m.s.d. 84–238 = 0.653 Å) but are also associated to a small rotation of the N- and C-domains with respect to each other (see below). Analysis of both structures with PISA also indicates that P. denitrificans Atp12p is a monomer, in agreement with earlier sedimentation, cross-linking, and two-hybrid studies of yeast and human Atp12p (28, 29). The shape of Atp12p resembles a boxing glove, with the wrist and hand sections provided by two domains encompassing residues 3–83 (N-domain) and 84–238 (C-domain) (Fig. 3C). The hand domain of Atp12p displays a concave face, the “palm,” and a convex face, the “dorso.” Looking at the palm face of the Atp12p, there is a sharp difference in surface charge between the C-domain, which is uniformly negative, and the N-domain, which is uniformly positive (Fig. 3D). In contrast, the dorso surface is almost uniformly negatively charged (Fig. 3E).

A protein from Agrobacterium tumefaciens (PDB code 2R6l) very similar to P. denitrificans Atp12p was identified by the Midwest Center for Structural Genomics but was not characterized. Based on our study, 2R6l can now be assigned as the Atp12p chaperone of that bacterium. A structural homology search of Atp12p against the entire PDB bank identified at low threshold 2 additional neighbors for the N-domain and 67 additional neighbors for the C-domain; superposition of the best scoring neighbors onto each domain of Atp12p did not reveal a level of similarity sufficient to draw mechanistic information from the search. However, it is of interest that the core of the C-domain consists of a central helix surrounded by six other...
helices (Fig. 4A). This topology is reminiscent of the fold of the membrane domains of colicins, pore-forming toxins, and Bcl-2 apoptotic proteins, all of which are thought to become variably associated with biological membranes upon some conformational change (30–32). Despite this similarity, there is no evidence at this time that in vivo Atp12p might be associated with the mitochondrial inner membrane or the bacterial cell membrane.

Deletion of the sequence of S. cerevisiae Atp12p corresponding to the first 37 residues of the P. denitrificans sequence (Fig. 3C, light blue shading) produces a protein that supports ~80% of the wild type level of F₁ assembly in vivo but is barely detectable by Western blot in isolated mitochondria (29). This finding suggests that the N-domain is primarily necessary for stability. A W94R mutation was identified in a recessive allele of the ATPAF2 locus that caused the death of a 14-month-old homozygous patient (9). Trp-94 (corresponding to Trp-57 in P. denitrificans) and to a Phe or Trp in all known sequences of Atp12p is localized in the N-domain (Fig. 3C, yellow surface), and thus it is conceivable that the human syndrome originates from enhanced degradation of Atp12p rather than inactivation.

Deletion of the last 42 residues of the sequence of S. cerevisiae Atp12p, corresponding to the last two helices of the P. denitrificans protein (shown in magenta in Fig. 3C), completely eliminates the chaperone activity and prevents F₁ assembly (29). Interestingly, in the 1.0 Å structure of Atp12p (Table 2), the last helix is clearly visible in two positions corresponding to an ~3.0 Å shift of the helix axis (Fig. 4B). It is also worth noting that a negatively charged residue (Asp-202, shown as a green surface in Fig. 3C) of P. denitrificans Atp12p is situated immediately after a sharp kink in the penultimate helix of the protein, almost at the beginning of the C-terminal helical region whose deletion completely suppresses the chaperone activity. Substitution with Lys of Glu-289 in the C-domain of S. cerevisiae Atp12p (corresponding to Asp-202 in P. denitrificans) completely eliminates F₁ assembly in vivo without decreasing the amount of Atp12p detected in isolated mitochondria (29). Other substitutions at this position, E289Q and E289A, are tolerated to the extent that there is enough ATP synthase assembled to support the growth of the Δatp12 transformants that produce the mutant proteins. However, in both cases the level of F₁F₀ in mitochondria is significantly decreased. Instead, the phenotype of strains that produce the yeast E289D variant is indistinguishable from wild type, which is in accord with the fact that some members of the Atp12p family, like P. denitrificans Atp12p, have an aspartic acid at this position. The effect of this mutation has also been studied in the context of purified human Atp12p (E240K in this case), and it was found to interfere with its chaperone activity in assays with the model substrate citrate synthase (28). We have determined the crystal structure of the D202K mutant form of P. denitrificans Atp12p at the resolution of 1.8 Å (Table 2); the overall protein architecture is unchanged in D202K Atp12p (Cₐ r.m.s.d. 5–238 Å). The most notable difference with the wild type is the long lysine side chain at position 202 (Fig. 4, C and D), which, by replacing the negative charge of aspartic acid with a positive charge, strikingly disrupts the uniformly negatively charged surface of the C-domain in the palm-facing view (Fig. 3F). The D202K mutation also affects the last two helices, which in different structures of the wild type and in this mutant structure are in slightly different positions (Fig. 4B). These position shifts appear also...
Chaperones of F₁-ATPase

among the nine lowest frequency normal modes of Atp12p (see supplemental material) and therefore reflect intrinsic vibrational properties of the protein. Altogether, the effects of the C-terminal deletion and of the E289K mutation in Atp12p suggest that the region of Atp12p encompassing the last two helices is directly involved in the chaperone activity.

Intermediates of F₁ Assembly—Earlier hypotheses on the role of Atp11p and Atp12p were based on the assumption that the regions of F₁ α and β subunits most likely to promote aggregation are hydrophobic patches sequestered at the subunit interfaces in the αβ₁ structure, and that Atp11p and Atp12p mimic, respectively, the α and β subunit, temporarily masking these hydrophobic regions (6). However, side-by-side analysis of the structures of both F₁ and the two chaperones shows no similarity of shape between Atp11p and α and Atp12p and β. This observation is inconsistent with an assembly mechanism in which the chaperone is released (e.g. Atp11p from β) in exchange for the incoming partner subunit (e.g. α) (6), simply by virtue of a phenomenon of molecular mimicry. The likelihood that the exchange step of F₁ assembly may require a specific “trigger” event was studied in yeast cells in which the nuclear gene encoding the F₁ γ subunit was disrupted (Table 1), as we had observed that these cells also fail to assemble F₁. Despite α and β being present in a normal amount, there is little or no hexameric F₁ in soluble mitochondrial extracts from Δγ cells (Fig. 5, upper panel). Instead, most of α and β accumulate as soluble protein complexes of ∼100 kDa (Fig. 5). It appears that these complexes are stable intermediates along the path to F₁ assembly, and that the γ subunit is necessary for their final progression into mature F₁. When the complexes are analyzed by means of a high resolution sedimentation gradient, α and β do not comigrate; furthermore, the centers of the distributions of Atp12p and Atp11p in the gradient fractions correspond well to the centers of the distributions of α and β, respectively (Fig. 5, lower panel). As α and β have molecular masses of ∼55 and ∼52 kDa, and Atp12p and Atp11p have molecular masses of ∼34 and ∼31 kDa, the comigrations of α with Atp12p (at a position corresponding to a higher molecular weight) and of β with Atp11p (at a position corresponding to a lower molecular weight) suggest that there are two distinct populations of complexes represented, respectively, by α:Atp12p (∼89 kDa) and β:Atp11p (∼83 kDa) heterodimers.

Experiments in which affinity resins were used to precipitate tagged α or β subunits from soluble mitochondrial extracts of the Δγ mutant also provided evidence that α:Atp12p and β:Atp11p complexes, but not α:β hetero-oligomers, are stable intermediates in the assembly pathway (Fig. 6). For this work, Δγ yeast was doubly transformed with two autonomously replicating plasmids, bearing different genetic markers, for the coproduction of FLAG-tagged α (α-FLAG) and His-tagged β (β-HIS₉) in the mutant cells. Soluble mitochondrial extracts were prepared from doubly transformed Δγ and mixed end-over-end with either TALON metal affinity resin (HIS₉ affinity beads) or with anti-FLAG M2 affinity gel (FLAG affinity beads) overnight at 4 °C. The affinity beads (Fig. 6, lane B) were collected by centrifugation, washed, and analyzed in Western blots, along with samples of the post-bead supernatants (lane S)
DISCUSSION

Atp11p and Atp12p are present in the mitochondrial matrix in small amounts, comparable with those of unassembled α and β subunits. In the absence of one of these subunits, as F1 cannot be assembled, the concentration of the other subunit exceeds that of the cognate chaperone and the subunit aggregates. In S. cerevisiae strains harboring a disruption of the nuclear gene encoding either the α or the β subunit, insoluble aggregates are formed even in the presence of normal amounts of functioning Atp11p and Atp12p (4). During F1 assembly, the formation of Atp11p:β and Atp12p:α intermediates may not promote directly the formation of the physiological α:β dimers, but rather act as decoys preventing the formation of nonphysiological α:α, β:β, and α:β complexes. Although this action would be critical to prevent aggregation, it also requires an initial trigger to form at least one physiological α:β dimer devoid of chaperones. After this initial event, it is conceivable that successive bindings of an Atp11p-β or Atp12p-α complex to an α:β dimer might produce a conformational change that decreases the affinity of the chaperones for their target proteins. Analysis of the F1 assembly intermediates in yeast cells, in which the nuclear gene encoding the F1 γ subunit was disrupted (Figs. 5 and 6), suggests that the γ subunit is the trigger factor necessary for the initial chaperone release. In the absence of structures of the Atp11p:β and Atp12p:α intermediates, it is not possible to know exactly how the γ subunit interacts with these complexes. However, a side-by-side visual inspection of Atp12p and γ reveals that the most peripheral segment of the long coiled-coil of the γ subunit is remarkably similar to the C-terminal helical fragment of Atp12p (Fig. 7A), which was shown to be essential for activity. If the C-terminal helices of Atp12p are superimposed to the coiled-coil tail of the γ subunit, the interaction surface of Atp11p (Fig. 7B), providing a reasonable guess of how the two proteins interact. The finding of a possible molecular mimicry between the C-terminal region of Atp12p and the coiled-coil tail of γ raises the possibility that F1 assembly may start with the γ subunit displacing Atp12p from the Atp12p:α heterodimer.

It had been assumed that the main reason for the existence of Atp11p and Atp12p is to protect hydrophobic regions of the surface of F1 α and β that otherwise would drive the aggregation of these subunits. However, there are no dramatic differences in the extent and distribution of hydrophobic patches between the regions of F1 α and β subunits exposed to solvent, and those buried at the subunit interface (supplemental Fig. 1). Further-
Chaperones of F₁-ATPase

more, hydrophobic areas account for less than 50% of the total contact surface between these subunits. Thus, other forces, besides hydrophobic interactions, play an important role in F₁ assembly. When electrostatic potentials are calculated for the entire F₁, the oligomer surface appears almost uniformly negative, with small positive regions (data not shown). However, when electrostatic potentials are computed separately for each individual subunit of F₁, the regions of the α subunits that contact the β subunits appear mostly positively charged, and the corresponding regions of the β subunits appear mostly negatively charged (supplemental Fig. 1). These observations suggest that electrostatic attraction plays a major role in the interaction of F₁ α and β with their cognate chaperones and in the formation of the correct interfaces in F₁. In humans, one fatal syndrome has been attributed to a defect in an F₁ chaperone (9), and intermediate phenotypes with partially impaired assembly of F₁ are likely to exist, in which a metabolic defect will appear only under stress. Other unrelated pathologic conditions are also characterized by the formation of nonphysiological aggregates. For example, it has been suggested that coulombic interactions play a key role in the formation of amyloid Aβ fibrils, which is the central event in the development of Alzheimer disease (33), and at least two distinct chaperones have been implicated in preventing the formation of these fibrils (34, 35).

Thus, the mechanism by which Atp11p and Atp12p prevent the chaotic aggregation of F₁ may be representative of a more general mechanism shared with other dedicated chaperones that also act primarily by masking electrostatic interactions.

REFERENCES

1. Abrahams, J. P., Leslie, A. G., Lutter, R., and Walker, J. E. (1994) Nature 370, 621–628
2. Blanchet, M. A., Hullinen, J., Pedersen, P. L., and Amzel, L. M. (1998) Proc. Natl. Acad. Sci. U. S. A. 95, 11065–11070
3. Kabaleswaran, V., Puri, N., Walker, J. E., Leslie, A. G., and Mueller, D. M. (2006) EMBO J. 25, 5433–5442
4. Ackerman, S. H., and Tzagoloff, A. (1990) Proc. Natl. Acad. Sci. U. S. A. 87, 4986–4990
5. Lefebvre-Legendre, L., Salin, B., Schaeffer, J., Bréthes, D., Dautant, A., Ackerman, S. H., and di Rago, J. P. (2005) J. Biol. Chem. 280, 18386–18392
6. Wang, Z. G., Sheluho, D., Gatti, D. L., and Ackerman, S. H. (2000) EMBO J. 19, 1486–1493
7. Wang, Z. G., and Ackerman, S. H. (2000) J. Biol. Chem. 275, 5767–5772
8. Marchler-Bauer, A., Anderson, J. B., DeWeese-Scott, C., Federova, N. D., Geer, L. Y., He, S., Hurwitz, D. I., Jackson, J. D., Jacobs, A. R., Lanzczyki, C. J., Liebert, C. A., Liu, C., Madej, T., Marchler, G. H., Mazumder, R., Nikolskaya, A. N., Panchenko, A. R., Rao, B. S., Shoemaker, B. A., Simon, V., Song, I. S., Thiessen, P. A., Vasudevan, S., Wang, Y., Yamashita, R. A., Yin, J. I., and Bryant, S. H. (2003) Nucleic Acids Res. 31, 383–387
9. De Meirleir, L., Seneca, S., Lissens, W., De Clercq, I., Eyskens, E., Gerlo, E., Smet, J., and Van Coster, R. (2004) J. Med. Genet. 41, 120–124
10. Lang, B. F., Gray, M. W., and Burger, G. (1999) Annu Rev. Genet. 33, 351–397
11. Terwilliger, T. C., and Berendzen, J. (1999) Acta Crystallogr. D Biol. Crystallogr. 55, 849–861
12. Sheldrick, G. M. (2008) Acta Crystallogr. A 64, 112–122
13. CCP4 (1994) Acta Crystallogr. D Biol. Crystallogr. 50, 760–763
14. Brünger, A. T., Adams, P. D., Clore, G. M., DeLano, W. L., Gros, P., Grosse-Kunstleve, R. W., Jiang, J. S., Kazusawa, J., Nitzges, M., Pannu, N. S., Read, R. J., Rice, L. M., Simonson, T., and Warren, G. L. (1998) Acta Crystallogr. D Biol. Crystallogr. 54, 905–921
15. Baker, N. A., Sept, D., Joseph, S., Holst, M. J., and McCammon, J. A. (2001) Proc. Natl. Acad. Sci. 98, 10037–10041
16. Sitkov, D., Sharp, K. A., and Honig, B. (1994) J. Phys. Chem. 98, 1978–1988
17. Metoki, K., and Hommes, F. A. (1984) J. Inherited Metab. Dis. 7, 9–11
18. Lloipis, J., McCaffery, J. M., Miyawaki, A., Farquhar, M. G., and Tsien, R. Y. (1998) Proc. Natl. Acad. Sci. U. S. A. 95, 6803–6808
19. Li, H., Robertson, A. D., and Jensen, J. H. (2005) Proteins 61, 704–721
20. Hill, J. E., Myers, A. M., Koerner, T. J., and Tzagoloff, A. (1986) Yeast 2, 163–167
21. Ackerman, S. H., and Tzagoloff, A. (2007) Methods Mol. Biol. 372, 363–377
22. Schiesl, R. H., and Gietz, R. D. (1989) Curr. Genet. 16, 339–346
23. Wang, Z. G., and Ackerman, S. H. (1996) J. Biol. Chem. 271, 4887–4894
24. Kasssil, E., and Henrick, K. (2007) J. Mol. Biol. 372, 774–797
25. White, M., and Ackerman, S. H. (1995) Arch. Biochem. Biophys. 319, 299–304
26. Gibrat, J. F., Madej, T., and Bryant, S. H. (1996) Curr. Opin. Struct. Biol. 6, 377–385
27. Kawabata, T. (2003) Nucleic Acids Res. 31, 3367–3369
28. Hinton, A., Gatti, D. L., and Ackerman, S. H. (2004) J. Biol. Chem. 279, 9016–9022
29. Wang, Z. G., and Ackerman, S. H. (1998) J. Biol. Chem. 273, 2993–3002
30. Cascales, E., Buchanan, S. K., Duché, D., Kleanthous, C., Llobés, R., Postle, K., Riley, M., Slatin, S., and Cavard, D. (2007) Microbiol Mol. Biol. Rev. 71, 158–229
31. Alouf, J. E. (2003) Folia Microbiol. 48, 5–16
32. Chao, D. T., and Korsmeyer, S. J. (1998) Annu. Rev. Immunol. 16, 395–419
33. Lazo, N. D., Grant, M. A., Condron, M. C., Rigby, A. C., and Teplow, D. B. (2007) Protein Expr. Purif. 48, 3367–3369
34. Wang, Z. G., and Ackerman, S. H. (1998) J. Biol. Chem. 273, 2993–3002
35. Kallhoff, V., Peethumonsin, E., and Zheng, H. (2007) Mol. Neurodegener. 2, 6
36. Ackerman, S. H., Martin, J., and Tzagoloff, A. (1992) J. Biol. Chem. 267, 7386–7394
37. Ackerman, S. H., Martin, J., and Tzagoloff, A. (1992) J. Biol. Chem. 266, 7517–7523
38. Paul, M. F., Ackerman, S., Yue, J., Arselin, G., Velours, J., Tzagoloff, A., and Ackerman, S. (1994) J. Biol. Chem. 269, 26158–26164
39. Mueller, D. M., Puri, N., Kabaleswaran, V., Terry, C., Leslie, A. G., and Walker, J. E. (2004) Protein Expr. Purif. 37, 479–485
40. Otwinowski, Z., and Minor, W. (1997) in Methods in Enzymology (Carter, C. W., and Sweet, R. M., eds) pp. 307–326, Academic Press, New York
41. Kabsch, W. (1988) J. Appl. Crystallogr. 21, 67–72
42. Kabsch, W. (1993) J. Appl. Crystallogr. 26, 795–800