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The Stator Complex of the A₁A₀-ATP Synthase—Structural Characterization of the E and H Subunits

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Archaeal ATP synthase (A-ATPase) is the functional homolog to the ATP synthase found in bacteria, mitochondria and chloroplasts, but the enzyme is structurally more related to the proton-pumping vacuolar ATPase found in the endomembrane system of eukaryotes. We have cloned, overexpressed and characterized the stator-forming subunits E and H of the A-ATPase from the thermoacidophilic Archaeon, Thermoplasma acidophilum. Size exclusion chromatography, CD, matrix-assisted laser desorption ionization time-of-flight mass spectrometry and NMR spectroscopic experiments indicate that both polypeptides have a tendency to form dimers and higher oligomers in solution. However, when expressed together or reconstituted, the two individual polypeptides interact with high affinity to form a stable heterodimer. Analyses by gel filtration chromatography and analytical ultracentrifugation show the heterodimer to have an elongated shape, and the preparation to be monodisperse. Thermal denaturation analyses by CD and differential scanning calorimetry revealed the more cooperative unfolding transitions of the heterodimer in comparison to those of the individual polypeptides. The data are consistent with the EH heterodimer forming the peripheral stalk(s) in the A-ATPase in a fashion analogous to that of the related vacuolar ATPase.

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Keywords: A-ATPase; ATP synthase; rotary molecular motor; stator complex; protein structure

Introduction

The archaeal ATP synthase (A₁A₀-ATP synthase; A-ATPase) is the functional homolog to the ATP synthase found in bacteria, mitochondria and chloroplasts (F₁F₀-ATP synthase; F-ATPase) but is structurally more related to the vacuolar proton-pumping ATPase (V₁V₀-ATPase; V-ATPase) found in the endomembrane system of eukaryotic organisms.¹⁻⁴ The A-ATPase is organized in two domains, a water-soluble A₁ and a membrane-bound A₀, in analogy to the related F₁/V₁ and F₀/V₀ domains of the F- and V-ATPases, respectively. The subunit composition of the A-ATPase is A₃B₃C₁DₓEᵧ for A₁ and CₓKᵧ for A₀.† The A₁ domain is formed by

† In the V-ATPase, subunit H of the A₁ is called G and subunits C, I and K of the A₀ are called d, a and c, respectively. The V-ATPase subunits C and H have no homologs in the A-ATPase.
the catalytic A and noncatalytic B subunits, which are arranged in a pseudo hexagonal barrel surrounding a hydrophobic cavity within which is located part of the D subunit. The A0 domain is made of multiple copies of the membrane-bound K subunits (proteolipids), which are arranged in a ring; an I subunit, which contains a C-terminal membrane-bound and a cytoplasmic N-terminal domain; and subunit C. The A1 and A0 domains are joined by a central stalk that is made of subunits D, F and C. The central stalk connects to the membrane-bound ring of the K subunits via the C subunit, which binds in the cytoplasmic opening of the K subunit ring. F, V- and (presumably) A-ATPases are rotary molecular motor enzymes5,6 (reviewed in Ref. 7); sequential nucleotide binding and catalysis taking place on the water-soluble domains (F1, V1 or A1) is coupled to ion translocation across the membrane-bound domains (F0, V0 or A0) via rotation of the central stalk and proteolipid ring. Ion translocation takes place at the interface of the membrane-bound domain of the I subunit and the ring of K subunits, which each have a lipid-exposed carboxyl residue (Asp or Glu). As in the F- and V-ATPases, peripheral stators or stalks function to hold the rotary molecular motors together during turnover. While there is general agreement as to the subunit structure of the motors together during turnover. While there is general agreement as to the subunit structure of the E and H subunits and the EH complex.

Results

Purification of the E and H subunits and the EH complex

Subunits E and H of the T. acidophilum A-ATPase were expressed as N-terminal fusions with Escherichia coli maltose binding protein (MBP). Fusion proteins were isolated by amylose resin affinity chromatography, protease cleaved, and purified by size exclusion chromatography. The EH complex was prepared by reconstituting purified E and H subunits at a 1:1:1 molar ratio followed by gel filtration to remove excess of H subunit. Figure 1 summarizes the purification of the E and H subunits as well as the EH complex. As can be seen from the SDS-PAGE analysis (Fig. 1a), the individual subunits and the complex are all highly purified. Figure 1b shows the elution profiles of the proteins from the size exclusion chromatography steps. Both individual subunits E and H eluted around 40–42 ml, corresponding to a relative molecular size of 75,000–65,000 (assuming globular proteins). Interestingly, the EH complex eluted later, at a volume of around 45 ml, corresponding to a size of around 60,000. The significantly lower than predicted elution volumes indicate that both individual subunits behave much larger than their predicted molecular weights. This behavior might be due to the formation of oligomeric aggregates or the presence of an unusually elongated shape, or a combination of both effects. From the fact that the EH complex elutes later than the individual subunits, the first possibility seems more likely (see also below).

MALDI-TOF mass spectrometry

In order to confirm the purity, identity and oligomeric state of the individually expressed polypeptides, we performed mass spectrometry (MS) with the purified subunits and the subunit complex. The spectra for subunits H and E and the EH complex are shown in Fig. 2a, b and c, respectively. As can be seen, the most intense peaks match expected m/z values for singly charged species for the individual subunits, irrespective of whether individual subunits or the subunit complex was analyzed. The measured molecular weights are very close to the predicted values (measured 21,861 versus predicted 21,886 for E and 13,698 versus 13,688 for H, respectively), confirming the identity of the polypeptides. Furthermore, next to the monomer peaks of E and H, peaks for subunit dimers, trimers and higher oligomers (for H) can be observed, consistent with the size exclusion chromatography results (see above). The matrix chosen (ferulic acid) has been shown to allow detection of native protein–protein
interactions using laser-induced desorption and ionization MS,\textsuperscript{17} indicating that at least some of the observed oligomeric species were already present in the aqueous solutions of the subunits used for sample preparation. MALDI-TOF of the EH complex resulted in a peak for the complex (observed mass 35,558; expected mass 35,555) next to peaks for the individual subunits and low-abundance oligomeric adducts. Dissociation of some of the complex into individual subunits is not surprising, given the high-energy input during the laser irradiation for sample desorption and ionization.

Analytical ultracentrifugation

To analyze the shape and molecular weight of the individual subunits and the EH complex in solution, proteins were subjected to analytical ultracentrifugation (AUC) experiments. Both sedimentation velocity and equilibrium centrifugation runs were performed. Initial sedimentation velocity runs of the individual E and H subunits gave poorly defined boundaries, consistent with a polydisperse behavior of both subunits (data not shown); AUC with the individual subunits was therefore not further pursued. Ultracentrifugation of the EH subunit complex, on the other hand, showed a very different behavior. Figure 3a shows the results of a representative sedimentation velocity experiment conducted with a protein concentration of 0.9 mg/ml and a temperature of 20 °C. The well-defined boundary and the flat plateau regions indicate that the sample is monodisperse, lacking higher aggregates. Fitting of the data to the modified Fujita–MacCosham function using the program SVEDBERG (see Experimental Procedures) revealed the single species to have an $s_{20,w}$ of 2.14 S. A virtually identical result ($s_{20,w}=2.15$ S) was obtained from another of the cells in this run, loaded with the EH heterodimer at a concentration of 1.7 mg/ml, indicating the lack of significant concentration-dependent aggregation. Given the expected mass of the complex (35,555 Da), the frictional ratio was calculated to be 1.71, indicating an extended structure. Analysis of sedimentation velocity runs carried out at 40 °C led to similar but slightly lower $s_{20,w}$ values of 2.07 and 2.08 S at the lower and higher concentrations, respectively, indicating that any structural changes over this temperature range were minor.

Sedimentation equilibrium analyses were conducted in triplicate at rotor speeds of 20,000 and 25,000 rpm at 20 °C, and at 40 °C after equilibration at 25,000 rpm. Figure 3b shows a sample data set collected at 20,000 rpm; analysis of these data gave a good fit for a single species of molecular weight 34,680 with a standard error of 500. The randomness of the residuals (inset) indicates the appropriateness of the single-species model. Analysis of the triplicate data sets under three conditions gave average values (± standard deviations) as follows: 20 °C at 20,000 rpm, 34,200±380; 20 °C at 25,000 rpm, 33,870±560; and 40 °C at 25,000 rpm, 32,840±520. The slight decrease in average molecular weight at 40 °C could indicate a slight reduction in affinity between subunits at that temperature, but it should be remembered that high-speed sedimentation tends to exaggerate the influence of trace low molecular weight species, and the magnitude of the change is borderline in significance. Overall, the sedimentation data indicate that the EH complex behaves as an extended, nonaggregating, high-affinity heterodimer.
Secondary-structure analysis by CD spectroscopy

The secondary-structure content of the stalk subunits was estimated by CD spectroscopy experiments. The results are shown in Fig. 4a. As can be seen, spectra of both individual E and H subunits as well as the EH subunit complex show an evidence of α-helical structure based on the distinct minima at 222 and 208 nm. However, the magnitude of the molar ellipticity of the EH complex at the two minima around 221 and 209 nm (~25,052 and ~25,681 deg cm$^2$ dmol$^{-1}$, respectively) is significantly greater compared to the individual subunits (approximately ~14,000 and ~12,500 for E and H, respectively), indicating that EH complex formation leads to a dramatic increase in α-helical secondary structure and/or its stabilization. Using the program K2d (see Experimental Procedures), the amount of α helix, β strand and random coil was estimated to be 37%, 15% and 58% for E, 27%, 15% and 58% for H and 81%, 0% and 19% for the EH complex, respectively. Interestingly, the EH complex showed a ratio of the magnitudes of the CD minima near 222 and 208 nm ($\theta_{222}/\theta_{208}$) of 0.99, indicative of the presence of coiled-coil secondary structure. In the isolated E and H subunits, the $\theta_{222}/\theta_{208}$ ratios are 0.92 and 0.86, respectively. The significantly
lower $\theta_{222}/\theta_{208}$ ratios for the isolated subunits suggest that individual subunit oligomerization probably does not involve formation of stable $\alpha$-helical coiled-coil secondary structure.

Thermal stability of the individual subunits and the EH complex

To estimate the thermal stability of the $T$. *acidophilum* A-ATPase stalk subunits, the CD signal at 222 nm was recorded as a function of temperature. Figure 4b shows thermal denaturation curves for the individual subunits and the heterodimeric complex. Subunit H showed a monotonic decrease in CD signal with increasing temperature. The absence of any cooperative melting behavior is consistent with some portions of subunit H forming solvent-exposed $\alpha$ helices. Thermal unfolding of subunit E showed a weak cooperative transition around 25 °C followed by a monotonic decrease in CD signal up to 70 °C. The melting behavior of the EH subunit complex was more complicated in that it seemed to involve more than one cooperative transition. Thermal denaturation of the individual subunits and the EH complex was therefore analyzed in more detail by DSC (Fig. 4c and d). Figure 4c shows the change in specific heat capacity for the individual subunits and the EH complex as a function of temperature. Again, for subunit H, no distinct transition can be observed, consistent with the lack of any cooperativity during thermal unfolding of the subunit. Subunit E shows virtually no change in specific heat capacity up to 75 °C, followed by a strong exothermic transition that would imply aggregation/precipitation of the subunit around this temperature. Samples recovered at the end of the experiment confirmed that precipitation had occurred. The temperature-dependent change in specific heat capacity for the EH complex, on the other hand, was positive throughout the entire temperature range with multiple transitions above 60 °C. Figure 4d shows fitting of the temperature dependence of the excess specific heat capacity for the EH complex after baseline subtraction. As can be seen, the best fit to the data indicates three transitions occurring at around 63, 83 and 92 °C, respectively. Nevertheless, the data indicate that the EH complex is relatively stable up to $\sim 60$ °C, which is the temperature where $T$. *acidophilum* shows optimum growth behavior.19

Binding affinity of the E and H subunits by isothermal titration calorimetry

In order to estimate the affinity of the two polypeptides, isothermal titration calorimetry (ITC) experiments were performed. The results of a representative experiment are shown in Fig. 5. In the experiment, subunit H at a concentration of 125 $\mu$M was injected 29 times in 6 $\mu$l aliquots into 1.4 ml of subunit E at a concentration of 10.6 $\mu$M, with one preinjection of 3 $\mu$l (Fig. 5a). The amounts of heat release for the titration were fitted to a model with one set of binding sites, resulting in a dissociation constant of 40±20 nM, a value close to the technical limit of the instrument (Fig. 5b). The value for the stoichiometry from the fit was given as approximately 0.85:1 (H:E). A possible explanation for the deviation from the expected 1:1 ratio might be due to the observation that the individual E and H subunits exist as oligomeric species (see above), and therefore, some of the heat generated during binding will be consumed to break up the subunit E and H oligomers. Indeed, when injecting subunit H into buffer alone, a significant positive heat change was observed (Fig. 5a, inset). That also means that the fitted association constant is probably underestimated and that the true affinity between E and H subunits is even greater. Trying to determine the EH affinity by diluting small amounts of preformed EH complex into the calorimeter cell did not result in measurable heat absorption, again consistent with the high affinity of the subunit interaction (data not shown).
Solution NMR spectroscopy of subunit E

As shown in Fig. 4b, subunit E undergoes a cooperative melting transition at around 25 °C. To determine whether this transition might be accompanied by a dissociation of the subunit E oligomers, $^{15}$N heteronuclear single quantum coherence (HSQC) spectra of uniformly labeled subunit E were recorded below (10 °C), at (25 °C) and above (37 °C) the transition temperature. NMR spectra were recorded and processed under identical conditions. As can be seen in Fig. 6, there is virtually no signal in the spectrum recorded at 10 °C (Fig. 6a), consistent with subunit E existing as high molecular weight oligomers. Here, the lack of signal might be due to intermediate chemical exchange and/or slow tumbling of the subunit E oligomers. At 37 °C (Fig. 6b), on the other hand, subunit E produced a $^{15}$N HSQC spectrum typical for a protein containing solvent-exposed \( \alpha \) helix (poorly resolved peaks in the center around 8 ppm) and some \( \beta \) sheet (peaks downfield of 9 ppm). An intermediate situation is found in the spectrum recorded around the transition temperature of 25 °C (Fig. 6c). Overall, these data are consistent with subunit E forming more stable secondary structure at low temperature by interacting with itself to produce dimers and/or higher oligomers. At temperatures above 25 °C, some of these oligomeric species will dissociate to produce monomeric subunit E, which is tumbling at
a fast enough rate to produce observable peaks in the HSQC NMR spectra. At the same time, however, some of the α-helical secondary structure present at 10 °C is probably destabilized, leading to the poorly resolved NMR signals in the center of the spectrum.

**Discussion**

In the present study, we have characterized the peripheral stalk subunits E and H of the A-ATPase form *T. acidophilum*, a moderately thermophilic acidophilic Archaeon growing optimally at 59 °C and a pH of 2. Gel filtration, AUC, MS and NMR spectroscopy experiments all indicate that the individual subunits in absence of their binding partner tend to interact with themselves to form dimers and higher oligomers. Furthermore, we provide evidence that when the two subunits are allowed to interact with equimolar stoichiometry, they form a stable, heterodimeric subunit EH complex with a $K_d$.
in the nanomolar range. The resulting subunit EH complex, as shown by AUC experiments, presented itself as a well-behaved, monodisperse species under the conditions tested. In addition, CD spectroscopy experiments showed that upon complex formation, there was a dramatic increase in α-helical secondary structure, most likely due to α-helical coiled-coil formation between the two subunits as indicated by the increased ratio of \( \theta_{222}/\theta_{208} \). While the estimated α helix content in the isolated subunits was significantly lower than expected based on secondary-structure prediction (see Fig. 7a), for the EH heterodimer it was 81%, very close to the predicted value of 78%. The relatively low degree of α helicity estimated for the isolated subunits (especially H) is expected, given the generally accepted model for formation of coiled coils, which involves two largely unfolded monomers coming together to form a folded (hetero)dimer.\(^{20,21}\) Additional evidence that the resulting EH heterodimer is the physiologically relevant species comes from thermal denaturation experiments, which show that the EH complex exhibits cooperative unfolding behavior, a hallmark of native protein structure. On the other hand, the cooperative loss of CD signal observed for the isolated E subunit probably does not indicate the unfolding of a native structure as it occurs at a temperature of \( \sim 25^\circ C \), which is significantly below the optimum growth temperature for \( T.\ acidophilum \) (59 \( ^\circ C \)).\(^{19}\) Instead, the NMR data shown in Fig. 6 suggest that this transition is the result of the reduction of oligomeric species. Taken together, the data strongly suggest that the heterodimer of subunits E and H functions as a structural element in the intact A-ATP synthase complex.

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**Fig. 7.** Building the stator of the A-, F- and V-ATPase. (a) Secondary-structure analysis of the peripheral stalk-forming subunits of the A-, V- and F-ATPase. Secondary-structure prediction was carried out using the PsiPred server [http://bioinf.cs.ucl.ac.uk/psipred] Vma4p and Vma10p, E and G subunits of the yeast vacuolar ATPase. Bov_E and Bov_G1, E subunit and G1 subunit isoform of the bovine clathrin-coated vesicle vacuolar ATPase. Ec_δ and Ec_b, δ and b subunits of the \( E.\ coli \) F,F0-ATP synthase. The structure of the \( E.\ coli \) F-ATPase δ subunit N-terminal domain (residues 1–106; indicated in cyan) has been determined by NMR spectroscopy.\(^{27}\) Structural models of the (b) bacterial F- and the (c) archaeal A-ATPase. Subscripts N and C next to the subunit name (EN, EC) stand for N- and C-terminal domain, respectively. For reasons of simplicity, the coiled-coil nature of the interaction between subunit H and the N-terminal domain of subunit E (E_N) has been omitted from the model. The collar observed in EM images of the intact A-ATPase\(^{11}\) is formed in our model by the hydrophilic domain of subunit I plus contributions from the N-terminal domains of the EH heterodimer that interact with the I subunit hydrophilic domain. For details, see the text.
What might be the function of this heterodimer in the A-ATPase? From earlier studies, it had been suggested that homodimers of subunits E, H, and the related G subunit of the V-ATPase are key structural elements of the A- and V-ATPase peripheral stalks, but these conclusions were all based on experimental data obtained for isolated, individual subunits. For example, isolated subunit H from *M. jannaschii* A-ATPase was shown to exist as a dimer in solution, a structural model of which was determined by small-angle X-ray scattering. The low-resolution model showed an elongated, elbow-like structure and it was speculated in that study that one of the peripheral stalks is constituted by the subunit H dimer alone. In the case of subunit E, dimer formation has been observed in a crystal structure obtained for the C-terminal domain of the subunit from the *Pyrococcus horikoshii* A-ATPase. The structure contains residues 81–198 and is composed of four β strands and six α helices, consistent with the secondary-structure prediction shown in Fig. 7a. The dimer interface is generated via domain swapping in that the C-terminal α helix of one monomer interacts with the N-terminal α helix of the other monomer and vice versa.

However, the evidence provided here, namely, that E and H form a stable complex with high affinity, casts doubt on the physiological significance of the homodimers. Dimerization of E subunit C-terminal domain may result from the missing N-terminal domain, as the results from the current study indicate that the dimerization seen in the crystal structure does not occur in the intact EH subunit complex in solution under the conditions tested. Similarly, while we have observed dimer (and higher oligomer) formation for isolated subunit H, addition of subunit E is able to break up this interaction of subunit H with itself, indicating preferential formation of the EH heterodimer.

Of course, for the EH complex to be able to function as peripheral stalk in the A-ATPase, its structure would have to be highly elongated, reaching from the bottom to the very top of the complex, a distance of more than 100 Å. The biophysical characterization of the EH complex provided here is consistent with this structural requirement. First, the complex is highly α helical, including a region of coiled coil (see above). Secondly, the complex elutes at a much higher molecular weight on gel filtration, and thirdly, a frictional ratio of 1.71 was obtained from AUC experiments. All these observations are consistent with an elongated shape for the subunit EH complex.

Further evidence that the EH complex plays the role of peripheral stalk in the A-ATPase comes from a comparison of the E and H subunit sequences to those of stator-forming subunits of the related F- and V-ATPase. While there is significant amino acid sequence homology between F-, A- and V-ATPase for the catalytic subunits and the proteolipids, the rotor- and stator-forming subunits of the related ATPases show little sequence conservation. The situation becomes clearer, however, when conducting the comparison on the level of (predicted) secondary structure. Figure 7a shows a schematic comparison of the secondary-structure prediction of subunits E and H of the A-ATPase with those of subunits E and G of V-ATPase and δ and b of bacterial F-ATPase.

In the bacterial F-ATPase, it is now well established that the peripheral stalk is formed by the δ and b subunits. In the *E. coli* enzyme, the peripheral stalk is anchored to the membrane via the N-terminal transmembrane α helices of the b subunits. The C-terminal domain of one of the b subunits interacts with the C-terminal domain of the δ subunit, while the N-terminal domain of the δ subunit is bound to the F1 via the N-terminus of one of the three α subunits. There is evidence that the two b subunits dimerize by forming a right-handed coiled coil for much of their length. In the V-ATPase, which is structurally more similar to the A-ATPase than to the F-ATPase, the number and nature of the peripheral stalk(s) is still not fully understood, but there is evidence from chemical cross-linking that the stator is formed by subunits E and G, that subunits E and G interact to form an elongated, stable heterodimer and that there are at least two copies of the EG heterodimer in the V1-ATPase complex from yeast.

The situation in the A-ATPase is probably similar to that in the V-ATPase except that the number of EH heterodimers has yet to be addressed in a rigorous manner, even though electron microscopy had suggested the presence of at least two peripheral stalks. While the secondary-structure analysis shown in Fig. 7a points to a structural relationship between subunits b, G and H of the F-, V- and A-ATPase, the comparison of subunits δ and E reveals a clear difference between the F-ATPase stator subunit on one hand and the related V- and A-ATPase subunits on the other. The situation can be clarified, however, when looking at the structure of the δ subunit. It consists of a compact, N-terminal six-helix bundle (residues 1–106 in *E. coli*), indicated by the * in Fig. 7a) and a C-terminal domain, for which there is no structure available yet. The N-terminal domain of δ has been shown to occupy the dimple formed by the N-terminal domains of the α and β subunits of the F1-ATPase, while the C-terminal domain of δ has been shown to interact with the C-terminal domain of the b subunits. This is illustrated in Fig. 7b. Interestingly, even though there is poor conservation on the level of the primary sequences between the peripheral stalk subunits of the A-, V- and F-ATPase, secondary-structure analysis reveals the similarity of the structural elements of the stators in the members of the three ion-translocating ATPase families.

Our current working model of the stator composition of the A-ATPase is the following (illustrated in Fig. 7c): each stator in the A-ATPase is formed by a heterodimer of the E and H subunits. The two subunits interact by forming a coiled coil (as evident from CD analysis; see Fig. 4a) involving the α-helical
N-terminal domain of the E subunit (E\textsubscript{N}) and an as of yet unspecified region in the H subunit. Evidence supporting the existence of an unusual right-handed coiled-coil structure in the \(b\) dimer of eubacterial ATP synthase has been presented. While our studies indicate formation of a heterodimeric coiled coil by the E and H subunits, we have no evidence that would allow us to position the region of the coiled coil or to distinguish between left-handed and right-handed models. The C-terminal domain of the E subunit, on the other hand, is responsible for interaction with the ATPase domain.

What might have led to this domain interchange within the stator-forming subunits between F- and A/V-ATPase? A possible answer to that question might be found when looking at the number of peripheral stalks present in the two ATPase families. The single N-terminal domain of F-ATPase \(\delta\) subunit occupies the top of the \(F_1\) and therefore leaves room for only one stator. For a reason yet unknown, multiple stators are found in \(A\) - and V-ATPase. Consequently, when going from F- to A/V-ATPase, the N-terminal domain of F-ATPase \(\delta\) had to be eliminated in order to accommodate multiple stalks. At the same time, the interaction between the stator and the ATPase domain had to be moved from the \(\delta\) subunit N-terminal domain to the E subunit C-terminal domain. The nature of the interaction between the peripheral stalk and catalytic domain in the A- and V-ATPase is not known at this point, but might be similar to what is seen in the bacterial flagellar motor and type III secretion systems. For example, in the bacterial flagellum, a protein with weak homology to subunit E, named FlIH, has been found, and it has been shown that this protein can bind via its C-terminal domain to FlJ, a flagellar component with homology to the F-ATPase catalytic \(\beta\) subunit. On the other hand, a close proximity of subunits A and H near the top of the \(A_1\) domain has been observed based on chemical cross-linking experiments, suggesting that subunit H might also play a role in the interaction between peripheral stator and catalytic domain. Experiments to identify the interaction of peripheral stator and ATPase domain in the A-ATPase from \textit{T. acidophilum} are ongoing in our laboratories.

### Experimental Procedures

#### Materials

The cDNA for open reading frames Ta\textsubscript{0001} and Ta\textsubscript{0007a} (subunits E and H of the A-ATPase from \textit{T. acidophilum}) was kindly provided by Drs. Dennis Revis and Robert Richards, California Lutheran University, Thousand Oaks, CA. The QuikChange kit was from Stratagene Inc. [\(^{15}\text{N}\)]Ammonium chloride was from Spectra Stable Isotopes. PrecisionProtease was from GE Healthcare. Plasmid pMAL-C2E (New England Biolabs) containing a PrecissionProtease sequence downstream of the C-terminus of MBP was a kind gift from Dr. Gino Cingolani, SUNY Upstate Medical University. All other reagents were of analytical grade.

### Protein expression and purification

The open reading frames for the E and H subunits were PCR amplified using primer pairs GCTCAGTGATTGTCCTCAGAAAGTGCTGAAG, CGAGTCAGCTTTCAATCCTCTATCCAGAATTAAAGC and GCTCAGGTACCGAGCTGTGATGCTGAT, CGAGTCAGCTTTTACCTCTCTATGATACCATCTCATATTA for Ta\textsubscript{E} and Ta\textsubscript{H}, respectively. 5'-KpnI and 3'-HindIII sites (underlined) were used for insertion into the modified pMal-c2e plasmid. Removal of the MBP fusion from the target protein resulted in a five-amino-acid N-terminal linker (GPKVP) on the target protein. All biophysical characterization experiments presented here were conducted with E subunit in which the endogenous cysteine at position 60 was changed to alanine via the PCR-based QuickChange protocol. Earlier experiments, which were conducted with wild-type E subunit, indicated essentially identical characteristics except that after prolonged storage, the cysteine-containing subunit had the tendency to precipitate due to intermolecular disulfide bond formation. In the following text, E subunit therefore stands for E, C64A. All PCR and site-directed mutagenesis was verified by DNA sequencing. Plasmid DNA was introduced into Rosetta pLacI \textit{E. coli} cells (Novagen) containing the pRare plasmid. A 10-ml overnight culture was used to inoculate 1 liter of rich broth medium supplemented with 100 \(\mu\text{g} / \text{ml} \) ampicillin and 34 \(\mu\text{g} / \text{ml} \) chloramphenicol to maintain the expression and pRare plasmids, respectively. Cells were grown at 37 \(^\circ\text{C} \) in rich broth containing 100 \(\mu\text{g} / \text{ml} \) ampicillin and 34 \(\mu\text{g} / \text{ml} \) chloramphenicol to an OD\textsubscript{600} of \(-0.8\) at which time expression was induced by addition of IPTG to a final concentration of 0.5 mM. After \(\sim 4\) h at 37 \(^\circ\text{C}\), cells were harvested by centrifugation, resuspended in 20 ml of column buffer (CB; 20 mM Tris-HCl, pH 7.4, 200 mM NaCl, 1 mM EDTA) and stored at \(-20\) \(^\circ\text{C} \) until further use. Cells were incubated with lysozyme and cell lysis was accomplished by sonication in the presence of DNAase I and 1 mM PMSF. Cell debris was removed by centrifugation at \(-15,000g\) for 30 min at 4 \(^\circ\text{C} \) after which time the supernatant was diluted fivefold with CB and passed over a 15-ml amylose column (NEB) at \(-1\) \(\text{ml} / \text{min} \). The column was washed with 12 column volumes of CB at \(-3\) \(\text{ml} / \text{min} \) and the fusion protein was eluted with CB + 10 mM maltose in ten 3 ml fractions at \(-1\) \(\text{ml} / \text{min} \). Fusion-protein-containing fractions were pooled and incubated with PrecisionProtease overnight at 4 \(^\circ\text{C} \) according to the manufacturer’s instructions. Most of the MBP was removed by ammonium sulfate precipitation (40–60% cut for E and 55% for H) and the reisolated proteins were further purified by size exclusion chromatography (Superdex 75, GE Healthcare; 16/30 column) in 25 mM sodium phosphate, pH 7, 1 mM EDTA and 0.02% sodium azide. Fractions were analyzed by SDS-PAGE and E- and H-subunit-containing fractions were pooled and concentrated or dialyzed against the appropriate buffer. Isotope ([\(^{15}\text{N}\)]-labeled E subunit was produced in an analogous fashion, except that bacterial cultures were grown in M9 minimal medium containing [\(^{15}\text{N}\)]ammonium chloride as sole nitrogen source. Complex formation was carried out by combining purified E subunit with \(-10\%\) excess of subunit H. The mixture was incubated on ice for 20 min and concentrated by ultrafiltration (10k MWCO, Vivaspin); excess H subunit was removed by gel filtration using Superdex 75 as described above. The gel filtration column was calibrated using blue dextran (molecular weight \(-2 \times 10^9\)), bovine serum albumin (67,000), MBP (44,000), chymotrypsin (25,000) and aprotinin (6500). Initially, EH
complex was also prepared by coexpression of subunits H and E (subunit E was expressed with an N-terminal histidine tag) from a bicistronic plasmid. However, the purity of the complex prepared this way was not as good compared to the above-described method (data not shown).

**CD spectroscopy**

Concentrated proteins were exchanged into 25 mM sodium phosphate, pH 7, by Sephadex G25 centrifugation columns. CD spectra were recorded on an Aviv 202 spectrometer with a 0.2- or 0.1-cm path-length cuvette. Scans were recorded from 250 to 195 nm with a step size of 1 nm at 25 °C. Buffer scans were subtracted from the raw data, which were then converted to molar ellipticity using concentration values determined from the absorbance at 277 nm of Gd HCl denatured samples. The values were 0.30, 0.36 and 0.37 mg/ml for subunits E, H and EH complex, respectively. Temperature melts of the proteins were observed at 222 nm over the temperature range of 10 to 70 °C (90 °C for the EH complex) using a 1 °C step size and a rate of temperature change of 5 °C/min. CD data from 240 to 200 nm were used to estimate the secondary-structure composition with the K2d¶ program. Secondary-structure prediction was carried out using the PSIPred server.§

**Mass spectrometry**

Purified subunits and subunit complex, each diluted to concentrations of ~5, 10 and 50 μM, were spotted onto a Bruker 384 target with an equal amount of saturated ferulic acid in 30:70 acetonitrile/trifluoroacetic acid (0.1%). Spectra were collected on an Autoflex TOF (Bruker) spectrometer using an m/z range of 10,000 to 80,000. Four hundred fifty spectra were summed for each protein and the program Mmass.

**Analytical ultracentrifugation**

Sedimentation velocity and equilibrium experiments were conducted on a Beckman Optima XL-A Analytical Ultracentrifuge (Biomolecular Interactions and Conformations Facility, University of Western Ontario, Canada). Two-channel and six-channel cells with Epon-charcoal centerpiecees were used for the velocity and equilibrium runs, respectively, in an An60Ti rotor. Samples were dialyzed against 25 mM sodium phosphate, pH 7, 100 mM NaCl, 1 mM EDTA, 0.02% sodium azide buffer and diluted to the desired concentration. Dialysis buffers were saved for use in the reference channels. Sedimentation equilibrium and velocity experiments were carried out at 20 and at 40 °C. The buffer density and partial specific volumes of the target proteins were calculated as described in Ref. 35. For sedimentation velocity, absorbance was monitored at 273 nm and data were collected in 0.002-cm radial steps, averaged over 10 replicates. Data were collected at 20 °C after equilibration at rotor speeds of 20,000 and 25,000 rpm, then the temperature was increased to 40 °C and data were collected after equilibration at 25,000 rpm. Data were analyzed using a single ideal species model that was built into Prism 4 (Graphpad) as described previously.¶

**Isothermal titration and differential scanning calorimetry**

DSC and ITC analysis were carried out on VP-DSC and VP-ITC instruments (MicroCal). For DSC, reconstituted EH complex was prepared as described for the AUC experiments. DSC thermograms were generated for samples of the EH complex as well as individual E and H subunits at concentrations of 48, 33 and 65 μM, respectively, over a range of 10 to 115 °C, at a rate of 60 °C/h. Raw data were processed by nonlinear least squares regression analysis in Origin 5.0 (MicroCal), by first subtracting a buffer–buffer reference scan and then fitting to a non-two-state transition model. For isothermal titration, E and H subunits were dialyzed against 10 mM sodium phosphate buffer, pH 7. Protein concentrations were determined by measuring the absorbance at 277 nm in 6 M Gd·HCl. Subunit H at a concentration of 125 μM was injected in one 3-μl and 29 6-μl aliquots into a 1.4-m1 cell containing subunit E (10.6 μM). A subunit H-only injection experiment was subtracted from the final data used for fitting. ITC data were analyzed in Origin 7.0 (MicroCal).

**NMR spectroscopy**

Uniformly 15N labeled E subunit was concentrated in 25 mM sodium phosphate buffer, pH 7, 1 mM EDTA and 0.02% sodium azide and D2O was added to a final concentration of 7%. 15N HSQC spectra were recorded at 10, 25 and 37 °C on a Bruker Avance 600-MHz spectrometer equipped with a HCN triple-axis gradient probe. NMR data were processed with Felix2004. Proton resonances were referenced to 2,2-dimethyl-2-silapentane-5-sulfonate and the 15N zero point frequency was calculated as described.†

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¶ http://www.embl.de/andrade/k2d.html
§ http://bioinf.cs.ucl.ac.uk/psipred
∥ http://mmass.biographics.cz/ was used for data analysis.
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Protein Eng. 6, 383–390.

