

Title: Structural snapshots of V/A-ATPase reveal a new paradigm for rotary catalysis.

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Abstract

V/A-ATPase is a motor protein that shares a common rotary catalytic mechanism with F₀F₁ ATP synthase. When powered by ATP hydrolysis, the V₁ moiety rotates the central rotor against the A₃B₃ hexamer, composed of three catalytic AB dimers adopting different conformations (AB_{open}, AB_{semi}, and AB_{closed}). Here we have determined the atomic models of 18 catalytic intermediates of the V₁ moiety of V/A-ATPase under different reaction conditions by single particle Cryo-EM, which revealed that the rotor does not rotate immediately after binding of ATP to the V₁. Instead, three events proceed simultaneously with the 120° rotation of the shaft: hydrolysis of ATP in AB_{semi}, zipper movement in AB_{open} by the binding ATP, and unzipper movement in AB_{closed} with release of both ADP and Pi. This indicates the unidirectional rotation of V/A-ATPase by a ratchet-like mechanism owing to ATP hydrolysis in AB_{semi}, rather than the power stroke model proposed previously for F₁-ATPase.

Main

The proton translocation ATPase/synthase family includes F-type enzymes found in eubacteria, mitochondria, and chloroplasts, and the V/A type enzymes found in archaea and some eubacteria¹⁻⁵ (Fig. 1a). These ATPases produce the majority of cytosolic ATP from ADP and *Pi* using energy derived from the transmembrane proton motive force generated by cellular respiration⁶. These ATPases share a common molecular architecture, consisting of a hydrophilic V_1/F_1 moiety responsible for ATP hydrolysis or synthesis, and a hydrophobic V_o/F_o domain housing a proton translocation channel⁷⁻⁹. The chemical reaction (ATP hydrolysis/synthesis) in V_1/F_1 is tightly associated with proton movement through V_o/F_o using a rotary catalytic mechanism, where both reactions are coupled by rotation of the central rotor complex relative to the surrounding stator apparatus, which includes the ATPase active hexamer^{6,10,11} (Fig. 1b).

The V/A-ATPase from the thermophilic bacterium, *Thermus thermophilus* (*Tth*) is one of the best-characterized ATP synthases^{3,12}. The overall architecture and subunit composition of V/A-ATPase is more similar to that of the eukaryotic V-ATPase, rather than F-type ATPase. However, the *Tth* V/A-ATPase has a simpler subunit structure than eukaryotic V-ATPase and shares the ATP synthase function of F-type ATPase¹³ (Fig. 1a). The V_1 moiety

of *Tth* V/A-ATPase ($A_3B_3D_1F_1$) is an ATP driven rotary motor where the central DF shaft rotates inside the hexameric A_3B_3 containing three catalytic sites, each composed of an AB dimer. The V_o domain ($E_2G_2d_{1a_{1c_{12}}}$) is composed of stator parts including the a subunit and two EG peripheral stalks and the $d_{1c_{12}}$ rotor complex, which consists of a central rotor complex with the DF subunits of V_1 ¹⁴⁻¹⁶. When ATP hydrolysis by A_3B_3 powers the DF shaft, the reverse rotation of the central rotor complex drives proton translocation in the membrane-embedded V_o domain (Fig. 1b).

According to the binding change mechanism of ATP synthesis⁶, the three catalytic sites in ATP synthases are in different conformations but interconvert sequentially between three different conformations as catalysis proceeds. Indeed, our previous structure demonstrated that the A_3B_3 hexamer in the V/A-ATPase adopts an asymmetrical structure composed of three different AB dimers, termed open (AB_{open}), semi-closed (AB_{semi}), and closed (AB_{closed})^{16,17}. The asymmetrical structure of the V_1 implies that sequential interconversion between these three different AB dimers is synchronized with rotation of the central DF shaft. Experimental studies using specific rotational probes attached to DF, revealed that ATP-driven rotation of the central shaft was unidirectionally clockwise when viewed from the V_1 side¹⁰. At low ATP concentrations where ATP binding is rate-limiting, rotation proceeds

in steps of 120° , commensurate with the three catalytic sites of AB dimers¹⁸. When using 40 nm gold beads with almost negligible viscous resistance, V_1 also pauses every 120° even at a ATP concentration around K_m without a sign of substeps¹⁹. These single-molecule experiments on V_1 suggest that the both catalytic events, ATP hydrolysis and product (ADP and P_i) release occur at an individual ATP binding position, and imply the presence of chemo-mechanically stable catalytic intermediates (Fig. 1c and Supplementary Fig. 1).

However, single-molecule observation experiments only allow us to see the motion of the shaft to which the observation probe is bound, and do not tell us what events are occurring at each catalytic site. To elucidate the entire rotational mechanism of V/A-ATPase, we must determine the structures of catalytic intermediates of the rotary ATPase during rotation. There are many reaction intermediates of the enzyme during turnover, and this structural heterogeneity makes successful crystallization of a specific state very challenging.

Technological breakthroughs in single particle Cryo-EM, such as the development of direct electron detectors, and advances in image processing and automation^{20,21}, have triggered a revolution in structural biology, making this the technique of choice for large and dynamic complexes unsuitable for crystallization. In addition, by freezing Cryo-EM grids at different time points or under different reaction conditions, it is possible to trap

intermediate states and thus build up a picture of the chemo-mechanical cycle of biological macromolecular complexes step by step. To date, there are few examples of studies that have successfully captured such details of a catalytic cycle at atomic resolution using Cryo-EM^{22,23}.

Here we report several key, and thus far uncharacterized, intermediate states of V/A-ATPase, obtained under different reaction conditions. Comparison of these structures provides insight into a novel cooperativity between the three catalytic sites and demonstrates a rotary catalytic mechanism powered by ATP hydrolysis.

Results

Sample preparation for Cryo-EM structural analysis

We previously determined the Cryo-EM structures of the wild-type V/A-ATPase containing an ADP in the catalytic site of AB_{closed}^{16,17}. The V/A-ATPase bound to the inhibitory ADP exhibits no ATPase activity until the ADP is removed^{13,16,24}. Partial ADP removal from AB_{closed} is possible by dialysis against an EDTA-phosphate buffer, but it is difficult to obtain a homogenous nucleotide-free V/A-ATPase after such a treatment, due to the high binding affinity of the ADP to AB_{closed} (Supplementary Table 1). To obtain a homogeneous ATPase

active enzyme, mutant V/A-ATPase (A/S232A, T235S) with reduced nucleotide-binding affinity was purified from *T. thermophilus* membranes¹⁰. The mutated V/A-ATPase exhibits higher K_m values for nucleotide in both the ATP hydrolysis and synthesis reactions than the wild-type enzyme, but the enzymatic and rotational properties are almost the same as those of the wild-type enzyme²⁴. The mutated V/A-ATPase is fully activated for ATPase activity after EDTA/phosphate dialysis; no ADP or ATP was found in the enzyme by quantitative analysis of nucleotides (Supplementary Fig. 2 and Table 1). We incorporated the nucleotide-free V/A-ATPase into nanodiscs comprising the MSP1E3D1 scaffold protein and DMPC. The resulting V/A-ATPase obeys simple Michaelis-Menten kinetics and exhibits ATPase activity of 22 s^{-1} and the K_m of $394\text{ }\mu\text{M}$ ATP (Supplementary Fig. 2b).

The nucleotide-free V/A-ATPase (Nucfree) was used for cryo-grid preparation under different ATPase reaction conditions (Supplementary Fig. 1c). Results of the structural analysis of the protein under each set of reaction conditions are summarized in Supplementary Figs. 3a-d.

The structures of V/A-ATPase without nucleotide (V_{nucfree})

The flow charts showing image acquisition and reconstitution of the 3D structure of V/A-ATPase without nucleotide are summarized in Supplementary Fig. 3a. We obtained

structures of three rotational states of V/A-ATPase without nucleotide; state1 at 3.1 Å, state2 at 4.7 Å, and state 3 at 6.3 Å resolution, with the DF shaft positions differing by 120° in each case (Fig. 2). Using signal subtraction of the V_o domain, we achieved resolution of 3.0 and 4.1 Å for the V₁ moiety including half of the EG stalk in state1 and 2, allowing us to build atomic models of the V₁ domain of these states.

The three AB dimers in the V₁ moiety adopted open (AB_{open}), semi-closed (AB_{semi}), and closed (AB_{closed}) states, respectively (Fig. 2b and c). The tip of the C-terminal helix bundle (CHB) of A_{open} is in contact with the C-terminal helix of the D subunit, and the wide part of the CHB of B_{open} is in contact with the N-terminal helix of the D subunit, respectively (Supplementary Fig. 4 a-c). The AB_{semi} and AB_{closed} also interact with the coiled-coil of subunit D in specific regions of the CHB, respectively (Supplementary Fig. 4 d-g).

The differences in the structures of the three AB dimers when superimposed on the β barrel domains of both A and B subunits are the result of movement of the N-terminal bulge domain, the nucleotide binding domain (NB) of the A subunit and the CHBs of both the A and B subunits (Fig. 2d). When comparing the structure of AB_{open} and AB_{semi}, both the NB and CHB of the A_{semi} are in closer proximity to B_{semi} than B_{open}, resulting in a closed structure of AB_{semi} (Fig. 2c and d). The structure of B_{open} is very similar to B_{semi}, as shown in Fig.

2d. In the AB_{closed}, both the CHB and NB domains of A_{closed} are in closer proximity to B_{closed}, and the CHB of B_{closed} moves to A_{closed}, resulting in the more closed structure of AB_{closed} compared to AB_{semi} (Fig. 2c and d).

In the AB_{closed} and AB_{semi} dimers, densities for the catalytic side chains are well resolved, but no density corresponding to nucleotide was observed (Fig. 3a). Hereafter we refer to the structure as the V_{nucfree}. The structure of V_{nucfree} is very similar to the previously reported ADP inhibited structure^{16,17}. For state1, the *rmsd* value for the C_α chains of A₃B₃DF of the V_{nucfree} and ADP inhibited structures is 1.98 Å (Supplementary Fig. 5). In addition, the V_{nucfree} is also similar to the structures under saturated-ATP condition determined in this study, with the positions of the catalytic side chains almost identical in both cases (Supplementary Fig. 6). This indicates that the V₁ moiety adopts the same conformation, including the arrangement of the DF shaft in the A₃B₃ and the geometry of the catalytic side chains, irrespective of the presence or absence of bound ATP.

In the density maps obtained for state1 of V_{nucfree}, the CHB of the AB dimers were slightly blurred, likely due to structural heterogeneity. To classify the probable substates of state1, we performed focused 3D classification using a mask covering AB_{open} and B_{semi} (Supplementary Fig. 3a). We identified an atomic resolution structure of the original state1

from 39,902 particles at 3.1 Å resolution and another substate from 24,101 particles at 3.1 Å resolution. We termed the substates reconstructed from these major particle classes as state1-1 and state1-2, respectively. The atomic model initially constructed as state1 is identical to the atomic model of state1-1. The structures of the sub-states are very similar, with most differences due to the movement of the CHB of the A and B subunits. Therefore, we quantified the difference in the structure of the CHB observed when the structures were superimposed on the N-barrel domain (Supplementary Table 2 and 3). Substates were also obtained under other reaction conditions (see below) and the *rmsd* values shown in Supplementary Table 2 and 3 are used to discuss which subunits are responsible for the differences in structure of the substates obtained under different reaction conditions.

Structures obtained at a saturating ATP concentration

Cryo-grids were prepared using a reaction mixture of Nucleotide-free V/A-ATPase, containing the regenerating system and ATP at a saturating concentration of 6 mM. The reaction mixture was incubated for 120 sec at 25 °C, and then loaded onto a holey grid, followed by flash freezing.

We determined three rotational states followed by focused refinement using a V₁EG mask for each state (Supplementary Fig. 3b). In the density maps obtained for each state, the

amino acid residues of the nucleotide binding sites in both AB_{closed} and AB_{semi} were well resolved, but the CHB domains of the AB dimers were blurred due to structural heterogeneity, as with the V_{nucfree} . For state1, we identified an atomic resolution structure of state1-1 from 40,831 particles at 3.1 Å resolution and state1-2 from 28,801 particles at 3.2 Å resolution by further 3D classification without alignment (Supplementary Fig. 3b). The same classification analysis was performed for state2 and state3, yielding state 2-1 (3.0 Å resolution) and state 2-2 (3.4 Å resolution), and state 3-1 (3.0 Å resolution) and state 3-2 (3.4 Å resolution) respectively. In these structures, nucleotide densities have been identified in the three catalytic sites. Hereafter, we refer to the structures obtained at ATP saturating condition as $V_{3\text{nuc}}$.

The structure of AB_{open} of $V_{3\text{nuc}}$ state1-1 is almost identical to that of V_{nucfree} state1-1 (Supplementary Fig. 6). This is confirmed by the fact that the *rmsd* values in the CHB of A_{open} and B_{open} for $V_{3\text{nuc}}$ state1-1 and V_{nucfree} are less than 1 Å (Supplementary Table 2 and 3). The AB_{open} of state1-2 adopts a slightly more closed conformation compared to that of state1-1, which results from a movement of CHB of B_{open} towards the β-barrel domain (Fig. 4c). Nevertheless, the AB_{open} of $V_{3\text{nuc}}$ with bound ATP retains the interaction with the DF shaft, indicating that ATP binding to the AB_{open} does not move the DF shaft.

The AB_{semi} in state1-2 has a more closed structure than that in state1-1 mainly due to the movement of CHB in B_{semi} (Fig. 4b, Supplementary Table 3). In summary, V_{3nuc} state1-2 has a more closed structure than state1-1 due to the movement of the CHB of both A_{semi} and B_{semi}, but the slightly closed conformation of state1-2 is independent of ATP binding to the AB_{open}.

Structures of the catalytic sites at AB dimers of V_{3nuc}

In both state1-1 and state1-2 structures obtained under ATP saturating conditions, a bound ATP molecule is clearly observed in the catalytic site of AB_{open} (Fig. 3b and Supplementary Fig. 7c). The catalytic sites in the AB_{closed} and AB_{semi} in both the state1-1 and state1-2 also contained density corresponding to an ATP molecule, and in these cases the associated magnesium ions were visible (Fig. 3b and Supplementary Fig. 7a-b). In the V_{3nuc} structure, we did not find density corresponding to nucleotides between the D and A subunits as reported in a previous paper²⁵ (Supplementary Fig. 8).

In the catalytic site of AB_{semi} of V_{3nuc}, the density of each nucleotide phosphate atom was easily identifiable (Supplementary Fig. 7b), indicating that the ATP molecule occupies the catalytic site in AB_{semi}. The protein structure is sufficiently clear to also provide a detailed picture of the configuration of the catalytic side chains (Figs. 5a-c). The γ -phosphate of ATP and the magnesium ion are coordinated by the A/K234 and A/S235 residues on the P-loop,

which contains the conserved nucleotide-binding motif^{7,26}. The aromatic ring of A/F230, not conserved in F type ATPase, is oriented away from the triphosphate moiety, allowing access of the guanidium group to the arginine finger (Supplementary Fig. 9). Considering clear EM density for the γ -phosphate of the ATP bound in AB_{semi}, hydrolysis of ATP is unlikely to proceed in AB_{semi}.

The nucleotide-binding site of the AB_{closed} is shown in Fig. 5b. The geometry of ATP binding in AB_{closed} is very similar to that found in the AB_{semi}, however, the carbonyl group of A/E257 is closer to the γ -phosphate by about 1 Å than AB_{semi}. In state1-2 of V_{3nuc}, the γ -phosphate of ATP bound to the AB_{closed} appears to be separated from β -phosphate when a relatively high density threshold is used (Supplementary Fig. 10). These findings strongly suggest that ATP bound to AB_{closed} is either already hydrolyzed or in the process of being hydrolyzed. The state of ATP in AB_{closed} is discussed further in Supplementary text.

In the nucleotide-binding site of the AB_{open} of V_{3nuc}, the adenosine moiety of ATP is occluded as in the AB_{semi} and AB_{closed}, with A/F415, A/Y500, and A/V236 forming the adenine binding pocket, however, the hydrogen bonding of the ribose moiety to the side chain of B/N363 is lost due to movement of the CHB of A_{open} (Fig. 5c). Unlike in the AB_{closed} and AB_{semi}, the phenyl group of A/230F in the AB_{open} is closer to the tri-phosphate group of

ATP due to torsion of the main chain, resulting in formation of a hydrophobic barrier between the catalytic side chains and the triphosphate moiety of ATP (Supplementary Fig. S9). Compared with AB_{semi}, the side chains of catalytic residues A/E257, A/R258, and B/R360 of AB_{open} are much further away from the γ -phosphate of ATP (10.0, 10.0 and 7.1 Å, respectively). Consequently, the configuration of the catalytic residues in the nucleotide-binding site of AB_{open} is not appropriate for hydrolysis of the bound ATP. Instead, the bound ATP has the potential to zipper the AB interface via interaction with the surrounding catalytic residues, which ultimately results in the transition of AB_{open} to a more closed form via a typical zipper conformational change.

Structures of V/A-ATPase waiting for ATP to bind

To determine the ATP-waiting structure of V/A-ATPase, we prepared a cryo-grid with a reaction mixture containing 4 μ M enzyme, 50 μ M ATP, and the ATP regeneration system, pre-incubated for 300 sec. We reconstructed three rotational states (state1, 2.7 Å, state2, 3.3 Å, and state3 3.6 Å resolution) from the single-particle images of the holo-complex and three rotational states of V₁EG (state1, 2.8 Å, state2, 3.1 Å, and state3, 2.8 Å) by focused masked refinement. For state1, two substates (state1-1, 2.9 Å and state1-2, 3.0 Å) were separated by focused classification using AB_{open} and AB_{semi} masks (Supplementary Fig. 3c).

In both AB_{semi} and AB_{closed} of state1-1, apparent density of ATP-magnesium was observed, but the density of the γ -phosphate at the AB_{closed} is weaker than that at the AB_{semi} (Fig. 3c and Supplementary Figs. 7a and b). In contrast, density was not observed in the nucleotide-binding site of AB_{open}. For state1-2, as in state1-1, nucleotides are present in both AB_{semi} and AB_{closed}, while AB_{open} is empty. Hereafter we refer to the structure as V_{2nuc}.

The overall structure and geometry of the catalytic residues of state1-1 of V_{2nuc} are largely identical to state1-1 of V_{nucfree} and V_{3nuc} (Supplementary Fig. 11). This structural similarity between V_{2nuc}, V_{nucfree}, and V_{3nuc} is confirmed by the low *rmsd* values when comparing the CHB of the A and B subunits of these structures (Supplementary Table 2 and 3). The similarity of the structures of these substates indicates that the structural polymorphism of V₁ moiety is independent of the binding of ATP to AB dimers.

Structures obtained at a saturating concentration of ATP γ S (V_{prehyd})

The V₁-ATPase from *T. thermophilus* is capable of hydrolyzing ATP γ S, however, the turnover rate of ATP γ S is much lower than that of ATP due to the decrease in hydrolysis rate¹⁸ (Supplementary Fig. 2d). Thus, pre-hydrolysis structures of V/A-ATPase can be obtained at 4 mM ATP γ S. The cryo-grid was prepared by blotting of the reaction mixture comprising Nucleotide-free V/A-ATPase and 4 mM of ATP γ S in the absence of the

regenerating system in order to exclude any effect of regenerated ATP produced from hydrolyzed ATP γ S. We reconstructed three rotational states from the acquired EM images using the CRYOARM300 (JEOL). After the focused masked refinement of the V₁EG domain, we obtained atomic resolution structures of each state (state1, 2.7 Å, state2, 3.4 Å, and state3, 3.6 Å), respectively (Supplementary Fig. 3d). We refer to these structures as V_{prehyd}. For state1, two sub-states, state1-1 and state1-2, were obtained at 2.7 and 2.9 Å resolution respectively, by focused classification using a mask with AB_{open} and AB_{semi} (Supplementary Fig. 3d). For the AB_{open} of V_{prehyd}, bound ATP γ S is clearly observed in the catalytic site, which has an almost identical structure to that of the ATP bound state of V_{3nuc} (Fig. 3b).

The nucleotide-binding sites of the AB_{closed} and AB_{semi} of V_{prehyd} are almost identical to those of V_{3nuc}, respectively, as shown in Fig. 5. The γ -phosphate group of the bound ATP γ S molecule in the AB_{semi} is well resolved as seen in for ATP in V_{3nuc} and V_{2nuc} (Figs. 3d and Supplementary Fig. 7b). In contrast, the density of γ -phosphate of ATP γ S at the AB_{closed} is faint (Supplementary Fig. 7a), suggesting that the ATP γ S in the AB_{closed} has already been hydrolyzed and the bound nucleotide in the AB_{closed} is ADP. This indicates that the AB_{semi} is in the pre-hydrolysis conformation, waiting for ATP hydrolysis.

Discussion

V/A-ATPase has a very similar structure irrespective of the nucleotide binding status.

We have obtained catalytic intermediates of V_1 moiety, V_{nucfree} , $V_{3\text{nuc}}$, $V_{2\text{nuc}}$, and V_{prehyd} , with different nucleotide occupancy. Despite the different nucleotide occupancy of these structures, their overall conformations are very similar. For instance, the *rmsd* of the C_α chains of A_3B_3 in V_{nucfree} and $V_{3\text{nuc}}$ state1-1 is 1.98Å. In addition, the relative position of the central DF shaft within the asymmetric A_3B_3 is almost the same in the V_{nucfree} and $V_{3\text{nuc}}$ structures. These findings demonstrate that the configuration between the DF shaft and individual AB dimers is independent of the state of nucleotide occupancy of each AB dimer. In other words, the structure of the V_1 moiety adopts three rotational states, 1, 2, and 3, during continuous ATP hydrolysis, with the conformational changes in the A_3B_3 hexamer driven by ATP hydrolysis, being discrete rather than continuous.

Chemo-mechanical cycle of the V/A-ATPase powered by ATP hydrolysis

The $V_{3\text{nuc}}$ structure, obtained under ATP saturation conditions shows all three catalytic sites occupied by ATP or the products of hydrolysis (ADP + P_i). Since the hydrolyzed P_i is clearly visible in the AB_{closed} of $V_{3\text{nuc}}$ (Supplementary Fig. 10), it is assumed that $V_{3\text{nuc}}$ is the structure before dissociation of P_i from the catalytic site in AB_{closed} . We also obtained the

V_{2nuc} structure in which ATP and product(s) are bound to AB_{semi} and AB_{closed} , respectively, but AB_{open} is empty. The V_{2nuc} is therefore assumed to be the structure of the protein awaiting ATP binding to AB_{open} . When using ATP γ S as a substrate, which has a very slow hydrolysis rate, the high-resolution atomic structure of the V_1 moiety allowed visualization of ATP γ S molecules bound to the catalytic sites of AB_{open} and AB_{semi} , as well as identification of the hydrolyzed ATP γ S at the AB_{closed} . The V_{prehyd} reveals both that the AB_{closed} adopts the post-hydrolysis state where the product of phosphate (P_i) is dissociated, and that the AB_{semi} is awaiting ATP hydrolysis.

The structures provide important insights into the chemo-mechanical cycle of V/A-ATPase. The V/A-ATPase undergoes a unidirectional conformational change from state1 to state2 to state3 when powered by ATP. Thus, V_{3nuc} of state1, in which three catalytic sites are already occupied by nucleotides, should change to state2 of V_{2nuc} , following ATP hydrolysis at AB_{semi} , and the subsequent or simultaneously dissociation of ADP and P_i by the discrete structural transition of AB_{closed} to AB_{open} (Fig. 6). This demonstrate that rotation of the rotary ATPase proceeds via the tri-site model with the protein progressing through a two nucleotide bound state and a three nucleotide bound state, settling the long-standing

debate on whether the bi-site model or tri-site model is appropriate for rotary ATPases^{5,6,27-}

³⁰.

Based on previous single molecular observation experiments for both F_1 - and V_1 -ATPase, ATP binding onto the enzyme directly triggers the first 120° rotation step of the DF shaft^{18,19,30} (Supplementary Fig. 1a). In light of our findings presented here, this scheme needs to be redrawn; the rotor does not immediately travel 120° as a result of ATP binding to enzyme.

The next catalytic event after ATP binding is ATP hydrolysis in AB_{semi} . Each conformational change, from AB_{open} to AB_{semi} , AB_{semi} to AB_{closed} , and AB_{closed} to AB_{open} occurs simultaneously, with the rotation of the shaft, and with the hydrolysis of ATP in the AB_{semi} and release of products (ADP and P_i) from the AB_{closed} (Fig. 6 and Supplementary Movie 1). This is in marked contrast to the classical rotary model, where catalytic events occur in sequence at the three catalytic sites, until now the broadly accepted mechanism of action of the F_1 -ATPase^{6,27,31,32}.

In the $V_{2\text{nuc}}$ and $V_{3\text{nuc}}$, two sub-states, state1-1 and state1-2 were identified. These substates were also identified in V_{nucfree} , thus the conformational dynamics of the V_1 moiety are independent of ATP binding. In other words, state1-1 and state1-2 are in thermally

equilibrium state, irrespective of nucleotide occupancy in each catalytic site. Both AB_{semi} and AB_{open} in state1-2 adopt more closed structures than those in state1-1, suggesting that state1-2 of $V_{3\text{nuc}}$ is likely an intermediate structure just prior to the 120° rotation step of the DF shaft. Compared to state1-2 of $V_{3\text{nuc}}$, state1-2 of V_{prehyd} exhibits slightly more closed structures of AB_{open} and AB_{semi} (Supplementary Fig. 13), likely to be associated with the progress of the catalytic reaction in AB_{closed} , i.e., the dissociation of the phosphate. In this respect, state1-2 of V_{prehyd} may be another reaction intermediate structure in which the P_i in the AB_{closed} is released prior to the 120° step (Supplementary Fig. 14).

Rotation Mechanism of the V/A-ATPase

Based on the catalytic intermediates of the V_1 moiety of V/A-ATPases obtained under four different reaction conditions, we propose a model for the ATP-driven rotation mechanism of V/A-ATPases.

When ATP binds to V_{nucfree} , which is in a stable initial state (ground state), the enzyme transits into the steady state for ATP hydrolysis (Fig. 7, *upper row*). The $V_{3\text{nuc}}$ structure is formed by binding of ATP to the AB_{open} of $V_{2\text{nuc}}$ but the binding of ATP itself does not cause structural transitions between AB dimers associated with the 120° step of the DF shaft.

In the V_{3nuc} structure, where nucleotides are bound to all three AB dimers, three distinct but associated catalytic events occur at the three AB dimers simultaneously and these events are coupled to the first 120° rotation step of the DF shaft. One of the driving events for this transition is the conformational change from the ATP-bound AB_{open} to the more closed AB_{semi} , which can be explained by a zipper motion of AB_{open} occurring upon ATP binding. From our structures, a comparison between AB_{open} and AB_{semi} implies that new hydrogen bonds form between the triphosphate moiety of ATP and the surrounding side chain groups of B/R360, A/258R, A/257E, and A/K234. (Fig. 5).

The V_{prehyd} structures indicate that the ATP bound to AB_{semi} is awaiting hydrolysis. The conformational change from AB_{semi} to AB_{closed} should occur spontaneously because it involves ATP hydrolysis, an exergonic reaction. In contrast, ADP bound to AB_{closed} hampers the unzipper motion in AB_{closed} , thereby preventing the overall structural transition of the V_1 moiety. This is supported by the fact that V/A-ATPase adopts the ADP inhibited state in which ADP is entrapped in AB_{closed} (Fig. 7, upper line). The enzyme in the ADP-inhibited state does not show ATP hydrolysis activity even at saturated ATP concentration^{13, 16}.

In summary, the ATP-driven unidirectional rotation of V/A-ATPase proceeds by a discrete structural transition between the three rotational states, i.e., the potential barrier to

the structural transition of AB_{closed} to AB_{open} , accompanied by release of ADP and P_i , is overcome by both a zipper motion of AB_{open} by the bound ATP and ATP hydrolysis in AB_{semi} . Since the ATP hydrolysis reaction is a heat dissipation process, the structural transition of AB_{semi} to AB_{closed} associated with the ATP hydrolysis occurs spontaneously and irreversibly, resulting in a unidirectionality of the 120° steps of rotor. In other words, our model explains the unidirectional rotation by a ratchet-like mechanism driven by ATP hydrolysis, rather than the power stroke model proposed previously for F_1 -ATPase^{5,33}.

V/A -ATPase and F_0F_1 are molecular machines based on the same construction principle, and thus are likely to share the same rotary mechanism. Importantly, for the thermophilic F_1 , the first 120° rotation step also includes $\sim 80^\circ$ and $\sim 40^\circ$ substeps, suggesting the existence of at least one additional catalytic intermediate of F_1 ^{5,27,34}. A recent structural study of thermophilic F_1 -ATPase indicated a possible intermediate structure responsible for the substeps³². In the V/A -ATPase, any intermediate structure containing phosphate after ADP or P_i release is likely to be an unstable state, and therefore studies on the ATP driven rotation of V/A -ATPase have failed to reveal the presence of any substep¹⁹.

Online Methods

Preparation of *Tth* V/A-ATPase for Biochemical assay and Cryo-EM imaging.

The *Tth* V/A-ATPase containing His3 tags on the C-terminus of each c subunit and the TSSA mutation (S232A and T235S) on the A subunit was isolated from *Thermus thermophilus* membranes as previously described²⁴ with the following modifications. The enzyme, solubilized from the membranes with 10 % Triton X-100 was purified by Ni²⁺-NTA affinity with 0.03 % dodecyl- β -D-maltoside (DDM). For bound nucleotide removal, the eluted fractions containing *Tth* V/A-ATPase were dialyzed against 200 mM Sodium phosphate, pH 8.0, 10 mM EDTA, and 0.03% DDM over night at 25°C with three buffer changes, followed by dialysis against 20 mM Tris-Cl, pH 8.0, 1 mM EDTA, and 0.03% DDM (TE buffer) prior to anion exchange chromatography using a 6 ml Resource Q column (GE healthcare). The *Tth* V/A-ATPase was eluted by a linear NaCl gradient using a TE buffer (0-500 mM NaCl, 0.03% DDM). The eluted fractions containing *holo-Tth* V/A-ATPase were concentrated to ~ 10 mg/ml using Amicon 100 k molecular weight cut-off filters (Millipore). For nanodisc incorporation, the 1,2-Dimyristoyl-sn-glycero-3-phosphorylcholine (DMPC, Avanti) was used to form lipid bilayers in reconstruction as previously described¹⁶. Purified *Tth* V/A-ATPase solubilized in 0.03 % n-Dodecyl- β -D-maltoside (DDM) was mixed with

the lipid stock and membrane scaffold protein MSP1E3D1 (Sigma) at a specific molar ratio V_0V_1 : MSP: DMPC lipid = 1: 4: 520 and incubated on ice for 0.5 h. Then, 200 μ L of Bio Beads SM-2 equilibrated with a wash buffer (20 mM Tris-HCl, pH8.0, 150 mM NaCl) was added to the 500 μ L mixture. After 2 hours incubation at 4°C with gentle stirring, an additional 300 μ L of Bio Beads was added and the mixture incubated overnight at 4°C to form the nanodiscs. The supernatant of the mixture containing nanodisc-*Tth* V/A-ATPase (nd-V/A-ATPase) was loaded onto the Superdex 200 Increase 10/300 column equilibrated with wash buffer. The peak fractions were collected, analyzed by SDS-PAGE and concentrated to ~4 mg/mL. The prepared nd-V/A-ATPase was immediately used for biochemical assay or cryo-grid preparation, since nd-V/A-ATPase aggregates within a few days.

Biochemical assay

The quantitative analysis of bound nucleotides of *Tth* V/A-ATPase was carried out using anion exchange high performance liquid chromatography¹³. Bound nucleotides were released from the enzyme by addition of 5 μ L of 60% perchloric acid to 50 μ L of the enzyme solution. Thereafter, the mixture was incubated on ice for 10 min. Then, 5 μ L of 5 M K_2CO_3 solution was added and the mixture incubated on ice for 10 min. The resulting

pellet was removed by centrifugation at 4°C. The supernatant was applied to a Cosmopak-200 column equilibrated with 0.1 M sodium-phosphate buffer (pH 7.0). The column was eluted isocratically with the same buffer at a flow rate of 0.8 ml/min. The nucleotide was monitored at 258 nm. The peak area was determined by automatic integration.

ATPase activity was measured at 25°C with an enzyme-coupled ATP-regenerating system, as described previously¹³. The reaction mixture contained 50 mM Tris-HCl (pH 8.0), 100 mM KCl, different concentrations of ATP-Mg, 2.5 mM phosphoenolpyruvate (PEP), 50 µg/ml pyruvate kinase (PK), 50 µg/ml lactate dehydrogenase, and 0.2 mM NADH in a final volume of 2 ml. The reaction was started by addition of 20 pmol *nd* V/A-ATPase to 2 ml of the assay mixture, and the rate of ATP hydrolysis monitored as the rate of oxidation of NADH determined by the absorbance decrease at 340 nm.

Cryo-EM imaging of *Tth* V/A-ATPase.

Sample vitrification was performed using a semi-automated vitrification device (Vitrobot, FEI). For *nd*-V/A-ATPase that underwent nucleotide removal, hereafter referred as nucfree *nd*-V/A-ATPase, 2.4 µl of sample solution at a concentration of 3 mg/ml (2 µM) was applied to glow discharged Quantifoil R1.2/1.3 molybdenum grid discharged by Ion Bombarder (Vacuum Device) for 1 min. The grid was then automatically blotted once from

both sides with filter paper for 6 s blot time. The grid was then plunged into a liquid ethane with no delay time.

The reaction basal buffer (RB buffer) containing 50 mM Tris-Cl, pH 8.0, 100 mM KCl, and 2 mM MgCl₂ was used for different reaction conditions. For saturated ATP or ATP waiting condition, 4 μM of nucfree *nd-V/A*-ATPase was mixed with same volume of x2 RB buffer containing 10 mM PEP, 200 μg/ml of PK, 12 mM or 100 μM of ATP-Mg. Then the mixtures were incubated for 120 sec or 300 sec at 25 °C, followed by blotting and vitrification, respectively. For the ATPγS saturated condition, 4 μM of nucfree *nd-V/A*-ATPase was mixed with the same volume of x2 RB buffer containing 8 mM ATPγS-Mg, then incubated for 300 sec at 25 °C, followed by the blotting and vitrification.

With the exception of the saturated ATPγS condition, cryo-EM imaging was performed with a Titan Krios (FEI/Thermo Fisher) operating at 300kV acceleration voltage and equipped with a direct K3 (Gatan) electron detector in electron counting mode (CDS). Data collection was carried out using SerialEM software³⁵ at a calibrated magnification of 0.88 Å pixel⁻¹ (x 81,000) and total dose of 50.0 e⁻ Å⁻² (or 1.0 e⁻ Å⁻² per frame) (where e⁻ specifies electrons) with total 5 s exposure time. The defocus range was -0.8 to -2.0 μm. The data were collected as 50 movie frames.

For the saturated ATP γ S condition, Cryo-EM movie collection was performed with a CRYOARM 300 (JEOL) operating at 300 keV accelerating voltage and equipped with a K3 (Gatan) direct electron detector, in electron counting mode (CDS) using the data collection software serialEM. The pixel size was 1.1 Å/pix (x60,000) and a total dose of 50.0 e⁻ Å⁻² (1.0 e⁻ Å⁻² per frame) with a total 3.0 s exposure time (50 frames) with a defocus range of -1.0 to -3.5 μm.

Image processing.

Image processing steps for each reaction condition are summarized in Figure S3 A-D. Image analysis software, Relion 3.1 and Cryosparc 3.2, were used^{36,37}. CTFFIND 4.1 and MotionCor2 were used for CTF estimation and movie correction in Relion^{38,39}. Topaz software was used for machine-learning based particle picking⁴⁰. We started with 15,317 movies for the nucleotide free enzyme (nucfree *nd*-V/A-ATPase), 13,164 movies for saturated ATP condition, 15,711 movies for saturated ATP condition, and 17,522 movies for ATP γ S condition. The software used in the steps is indicated in the figure. Autopicking based on template matching or based on Topaz machine-learning resulted in 4,354,341 particles for the nucfree *nd*-V/A-ATPase, 2,300,834 particles for the ATP saturated condition, 1,671,397 particles for the ATP waiting condition, and 4,677,284 particles for the ATP γ S waiting

condition. Particles were extracted at 5x the physical pixel size from the movie-corrected micrographs and selected using 2D or 3D classification (nucfree nd-V/A-ATPase; 132,904 particles, saturated ATP; 188,673 particles, ATP waiting; 186,928 particles, ATP γ S; 197,960 particles). The selected particles were extracted at full pixel size and subjected to 3D auto-refinement refollowed by CTF refinement by Bayesian polishing. Another round of 3D auto-refine, CTF refinement, and a final round of masked auto-refinement gave *holo*-V/A-ATPase maps at between 2.7 ~ 6.3 Å resolution. The membrane domain was visible but not particularly clear compared to the hydrophilic V₁ domain in a *holo*-enzyme map. This seemed to be due to the structural flexibility between the membrane domain and V₁ in the *holo*-enzyme. Focused refinement with signal subtraction targeting the V₁EG region improved the map quality of the V₁EG region (Figure S3A-D). The refinements provided the density maps for V₁EG under each condition at 2.8 - 4.1 Å resolution. After the focused refinement, masked classification on A_{open} and B_{semi} subunits was carried out to classify the conformational differences. Resolution was based on the gold standard Fourier shell correlation = 0.142 criterion.

Model building and refinement.

To generate the atomic model for the V₁EG region of V/A-ATPase, the individual subunits of the V₁EG model from the previous structure of V/A-ATPase (PDBID: 6QUM) were fitted into the density map as rigid bodies²⁵ with particular focus on the N terminal region of EG stalk (E; 1-77 aa., G; 2-33 aa.). The rough initial model was refined against the map with Phenix suite phenix.real_space_refine program⁴¹. The initial model was extensively manually corrected residue by residue in COOT⁴¹ in terms of side-chain conformations. Peripheral stalks were removed due to low resolution in this region. The corrected model was again refined by the phenix.real_space_refine program with secondary structure and Ramachandran restraints, then the resulting model was manually checked by COOT. This iterative process was performed for several rounds to correct remaining errors until the model was in good agreement with geometry, as reflected by the MolProbity score of 1.08~1.74 and EMRinger score of 1.59 ~3.94^{42,43}. For model validation against over-fitting, the built models were used for calculation of FSC curves against both half maps, and compared with the FSC of the final model against the final density map used for model building by phenix.refine program. The statistics of the obtained maps and the atomic model were summarized in Table S4. RMSD values between the atomic models were calculated using UCSF chimera⁴⁴. All the figures were rendered using UCSF chimeraX⁴⁵.

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Author contributions

KY, JK, A.Nakanishi and A.Nakano designed, performed and analyzed the experiments. JK, A.Nakanishi, KY, A.Nakano, AF, and SS analyzed the data and contributed to the preparation of the samples. TK and KM provided technical support and conceptual advice.

1 KY designed and supervised the experiments and wrote the manuscript. All authors
2 discussed the results and commented on the manuscript.

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4 **Declaration of interests**

5 The authors declare no conflicts of interest associated with this manuscript.

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Figure legends

Fig. 1: Schematic representation of rotary ATPases and the conventional rotary

mechanism. a, Illustration of subunit composition of different types of rotary ATPases;

prokaryotic V/A-ATPase (*left*), eukaryotic V-ATPase (*middle*), prokaryotic F-ATPase

(*right*). The stators are represented in various colors and the rotors are represented in grey.

b, A schematic model of the rotary catalytic mechanism of the V/A-ATPase. When

powered by ATP, the central rotor composed of $D_1F_1d_1c_{10}$ (grey) rotates against a

surrounding stator composed of $A_3B_3E_2G_2a_1$ (white), coupled with proton translocation

across the membrane. **c**, The conventional catalytic cycle of V/A-ATPase. At low ATP

concentration, the ATP binding dwell time is increased. $ATP\gamma S$ also prolongs the ATP

hydrolysis dwell.

Fig. 2: Cryo-EM density map and the atomic model for nucleotide free V/A-ATPase.

a, Cryo-EM density map of whole V/A-ATPase of state1 in the absence of nucleotide

($V_{nucfree}$ structure). **b**, Cryo-EM density map of V_1EG of state1 without nucleotide (*left*).

Cross sections of the nucleotide binding sites (*right upper*) and A_3B_3 C-terminal region

(*right lower*) viewed from top. **c**, Comparison of the AB dimer structures in $V_{nucfree}$. AB

dimers are shown as space filling models and superimposed on the β barrel domain (A subunit 1-70 a.a.). *Left*; AB_{open} (solid) vs AB_{semi} (semi-transparent), *middle*; AB_{open} (solid) vs AB_{closed} (semi-transparent), and AB_{semi} (solid) vs AB_{closed} (semi-transparent). **d**, Comparison between each AB subunit in V_{nucfree}. The subunits are shown as wire representation. A and B subunits are superimposed on the β barrel domain.

Fig. 3: Structures of nucleotide binding sites obtained in each condition. *upper panels*: V_{nucfree} (**a**), V_{prehyd} (**b**), V_{3nuc} (**c**), and V_{2nuc} (**d**) viewed from the cytosolic side. Scale bar is 20 Å. Magnified views of the three nucleotide binding sites (AB_{open}, AB_{semi}, and AB_{closed}) in each structure are shown in the rows below. Cryo-EMmaps are represented as semi-translucent. Bound nucleotides and Mg ions are shown in ball-and-stick and sphere representation, respectively. Scale bar is 4 Å

Fig. 4: Comparison between state1-1 and 1-2 in each subunit of V_{3nuc}. The subunits of state1-1 and 1-2 were superimposed on the β barrel domain (A: 1-70 a.a., B: 1-70 a.a.). Ribbon models are colored by the *rmsd* values calculated for the atoms of the main chain; gray (small changes) to red (large changes). Magnified views of the CHBs are represented

in the lower panels as wire models. The models of state1-1 are represented in gray, and state1-2 are represented in the different colors. **a**, subunits in AB_{closed}, **b**, subunits in AB_{semi}, **c**, subunits in AB_{open}.

Fig. 5: Coordination of nucleotides in the binding sites of V_{3nuc} and V_{prehyd}. *Left*

panels; Comparison of the three nucleotide binding sites (AB_{open} (**a**), AB_{semi} (**b**), and AB_{closed} (**c**)) of state1-1 of V_{3nuc} shown with colored (green, blue and pink) atoms and bonds, and main chain, and state1-1 of V_{prehyd} shown with grey atoms, bonds and main chain. *Right panels*; Schematic representations of the coordination of the ATP group in the three binding sites of V_{3nuc} and V_{prehyd} in parentheses. The distances between the atoms are shown in dotted lines. All distances are in Å.

Fig. 6: Chemo-Mechanical cycle of V/A-ATPase driven by ATP hydrolysis. The

structures of V/A-ATPase viewed from the cytosolic side are shown as ribbon models. The coiled coil of the DF subunits is shown in grey. The bound ATP molecules are highlighted in sphere representations. State1-1 and 1-2 of V_{2nuc} are in equilibrium and are fluctuating. These structures transit to state 1-1 and 1-2 of V_{3nuc} by ATP binding to AB_{open},

without a 120° rotation step of the DF rotor. V_{3nuc} in state1-1 and state1-2 are also in equilibrium. ATP hydrolysis at AB_{semi} and zipper motion at AB_{open} occur simultaneously. This triggers the transition of V_{3nuc} in state1-2 to V_{2nuc} state2-2 together with the 120° rotation step and simultaneous release of ADP and P_i . State2 of V_{2nuc} returns to state1 via state3 of V_{2nuc} by the same process. Asterisks indicate the structures which were not identified in this study.

Fig. 7: The rotary mechanism of V/A-ATPase powered by ATP hydrolysis. The schematic models of AB_{open} , AB_{semi} , and AB_{closed} are shown in green, pink, and blue, respectively. The coiled coil region of the D subunit in contact with A_3B_3 is shown in grey. In the ADP inhibited state, the entrapped ADP in AB_{closed} hampers structural transition of AB_{open} to AB_{semi} by binding of ATP to AB_{open} . The $V_{nucfree}$ in ground state is activated by the binding of ATP to the catalytic sites. In V_{2nuc} awaiting ATP binding, binding of ATP to AB_{open} does not induce the 120° rotation step. In V_{3nuc} , both zipper motion of AB_{open} and ATP hydrolysis in AB_{semi} induce unzipper motion of AB_{closed} accompanying with the release of ADP and P_i . The catalytic events in the three AB

- 1 dimers occurs simultaneously with the 120° step of the DF shaft, resulting in structural
- 2 transition of state1 of V_{3nuc} to state2 of V_{2nuc} .
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Figures

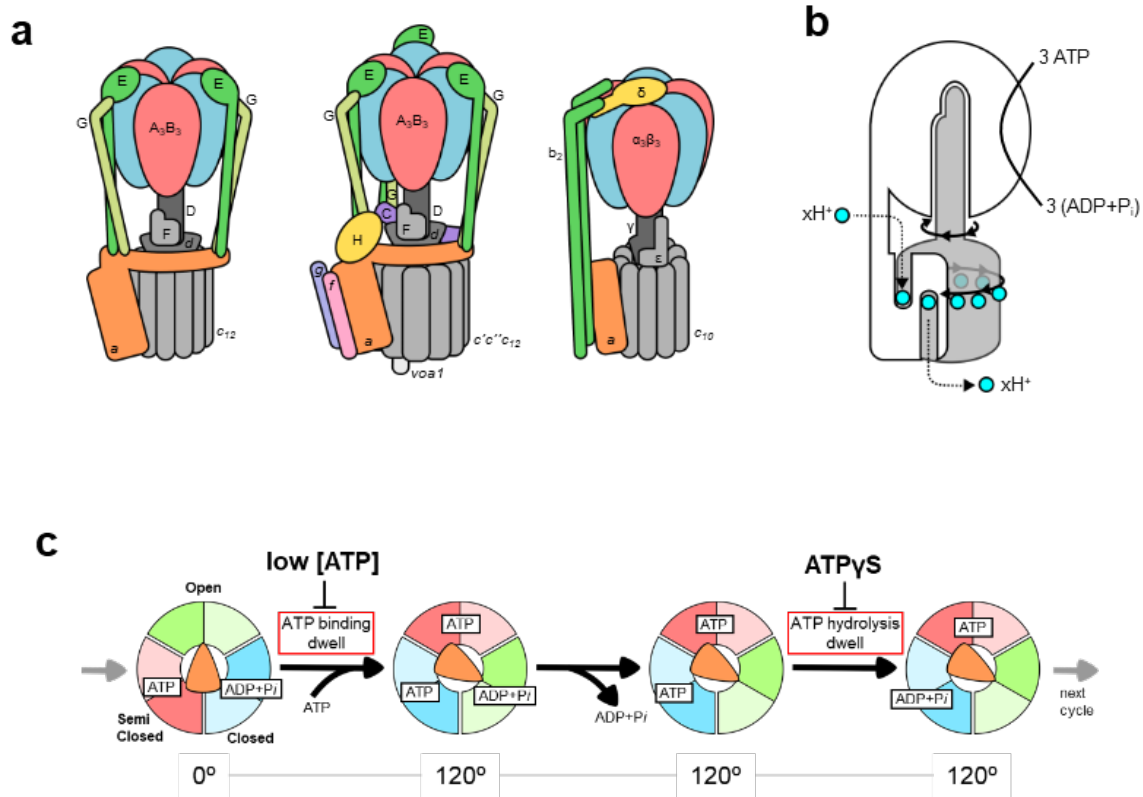


Fig. 1: Schematic representation of rotary ATPases and the conventional rotary mechanism.

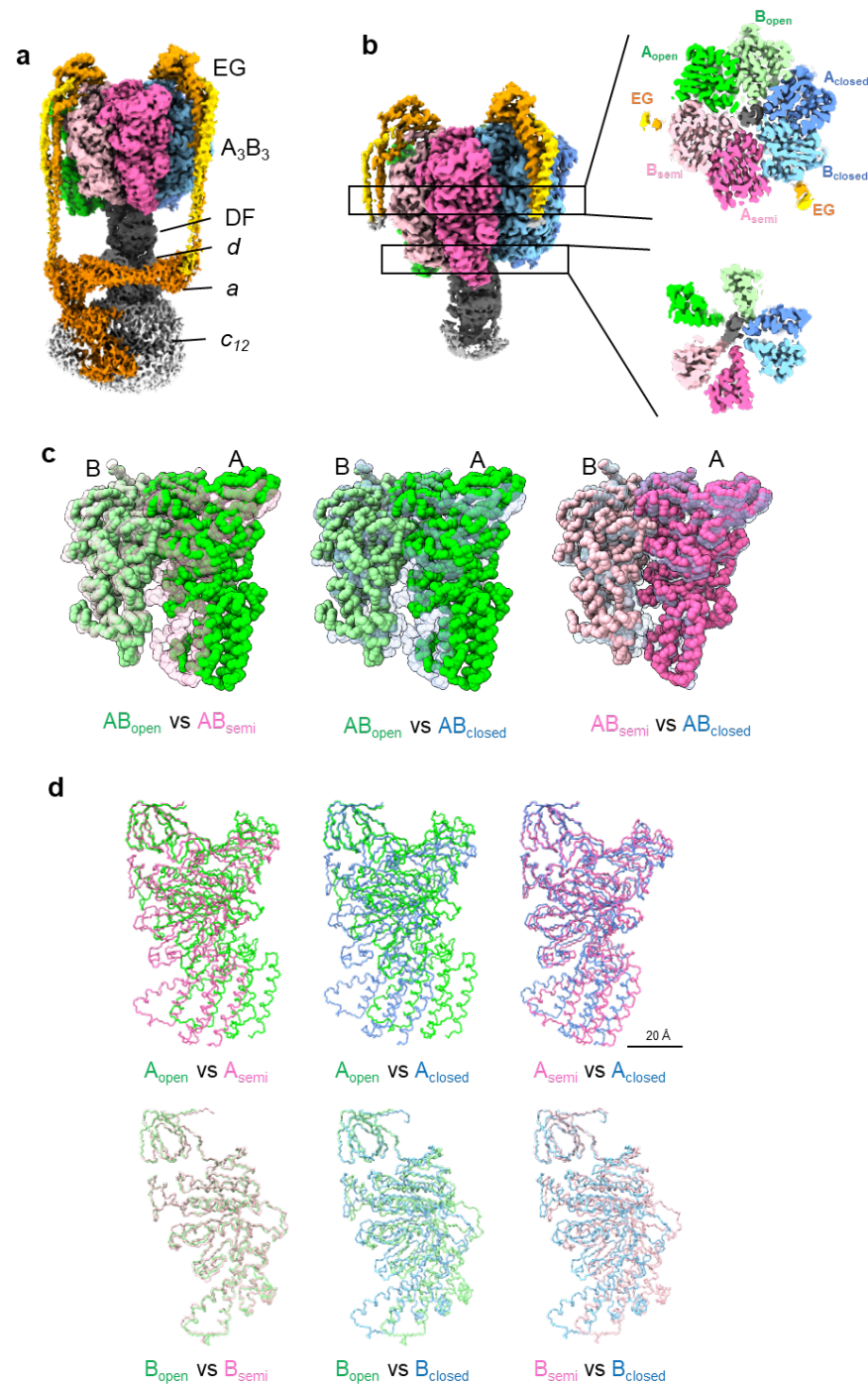


Fig. 2: Cryo-EM density map and the atomic model for nucleotide free V/A-ATPase.

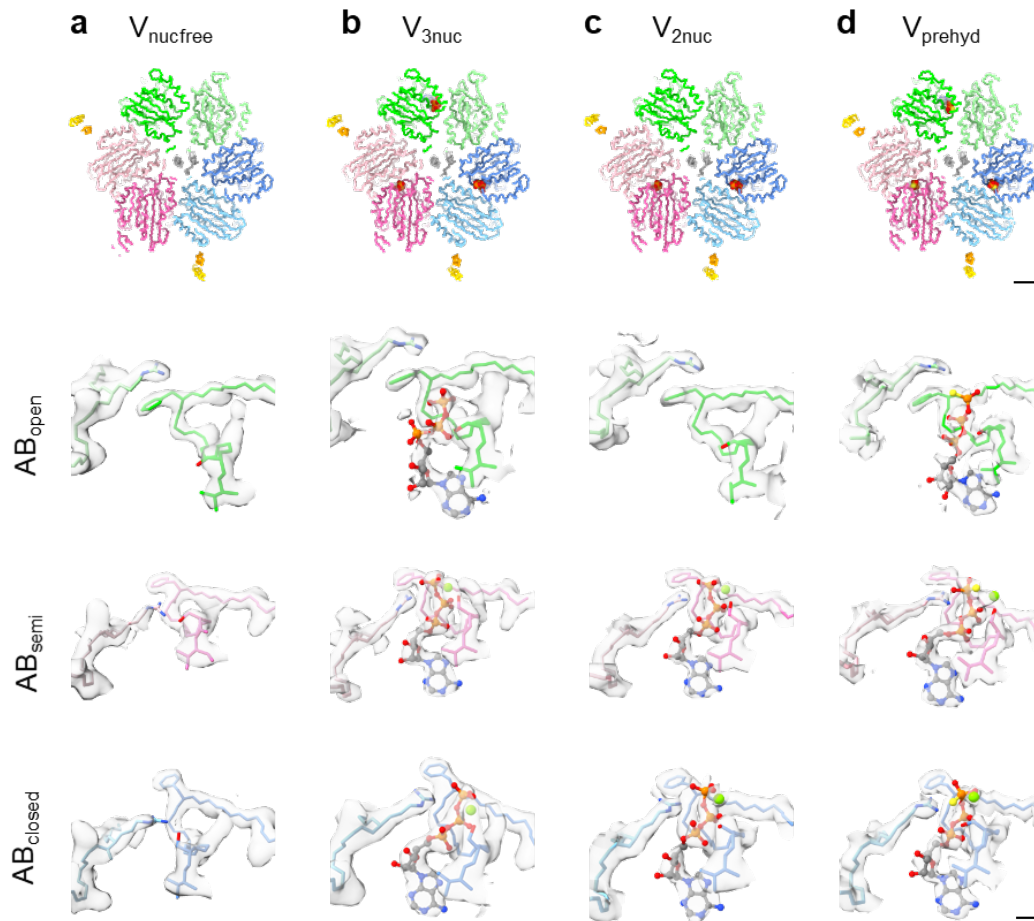


Fig. 3: Structures of nucleotide binding sites obtained in each condition.

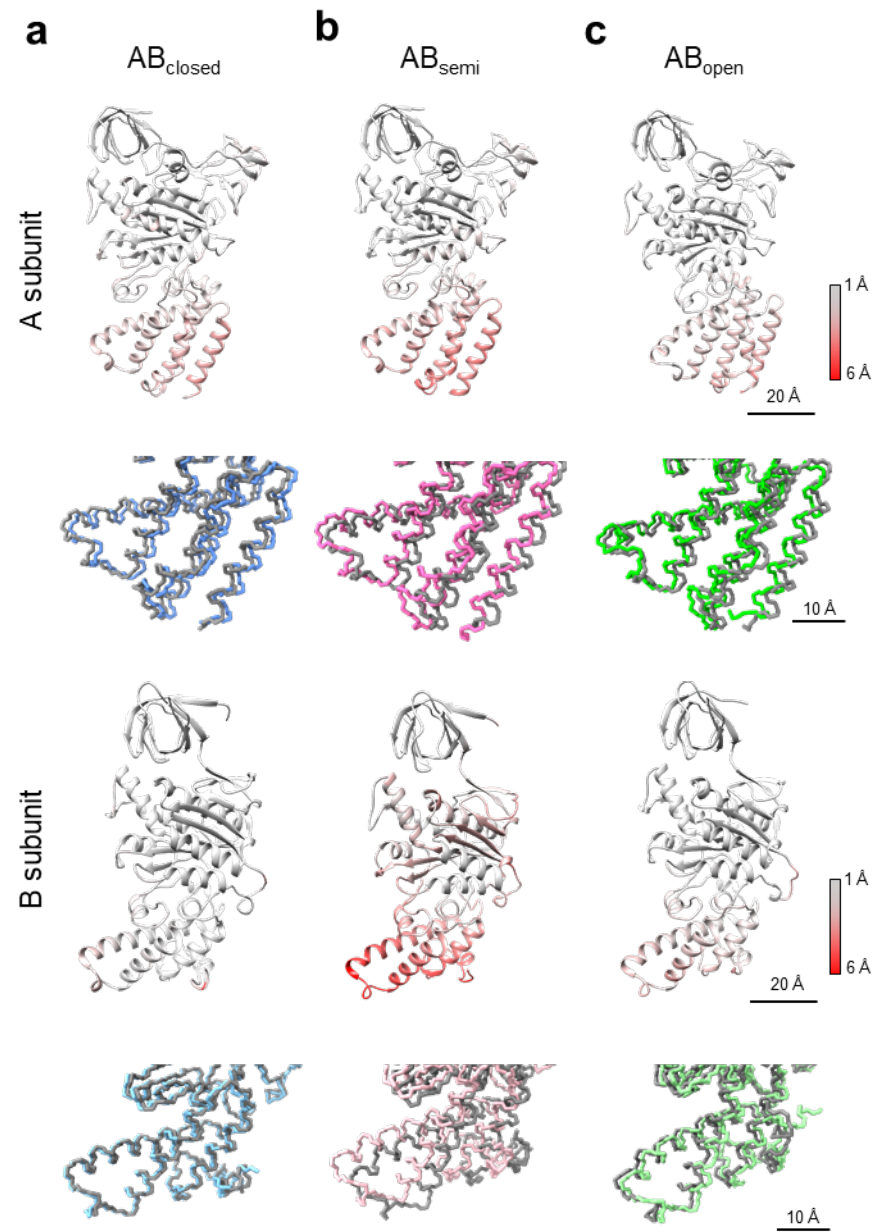
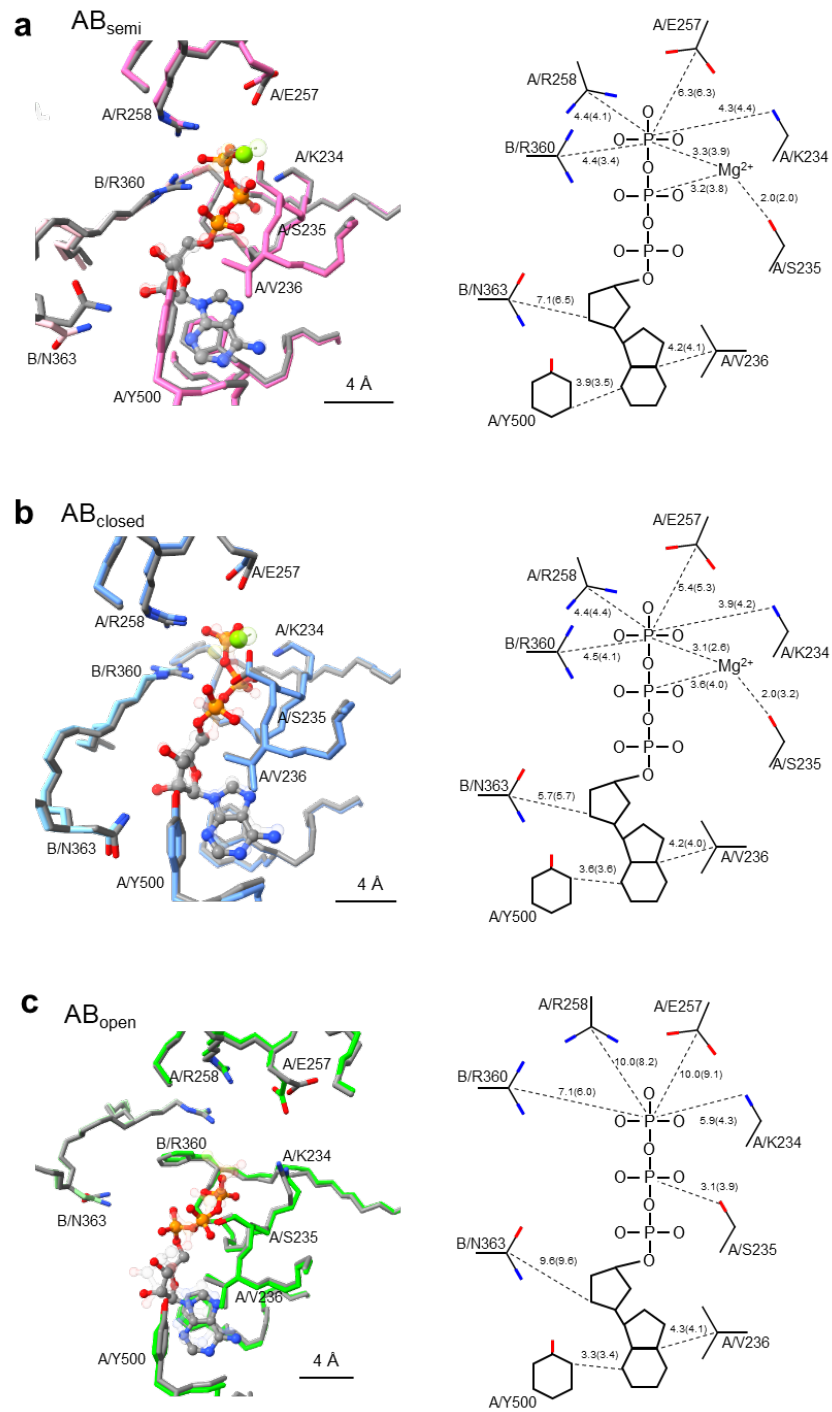
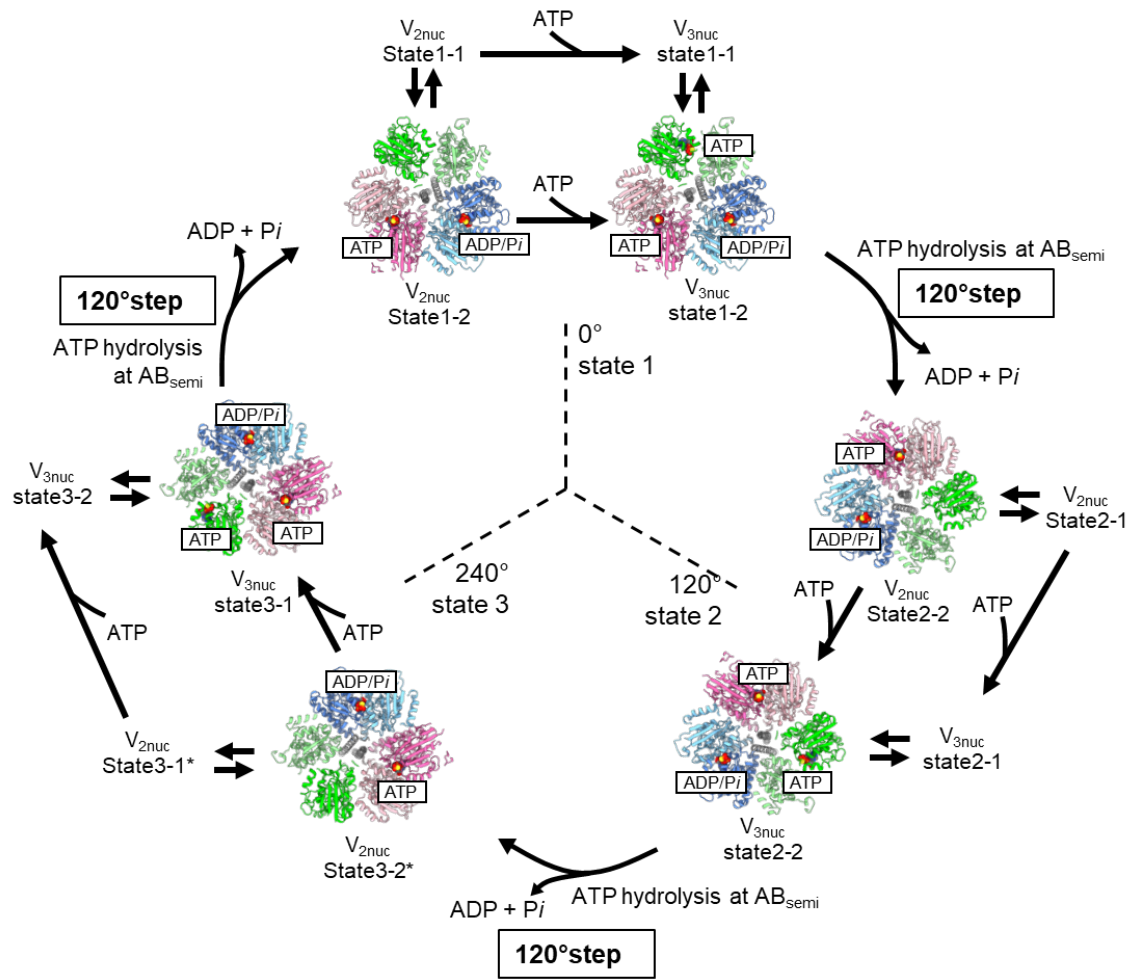


Fig. 4: Comparison between state1-1 and 1-2 in each subunit of V_{3nuc} .



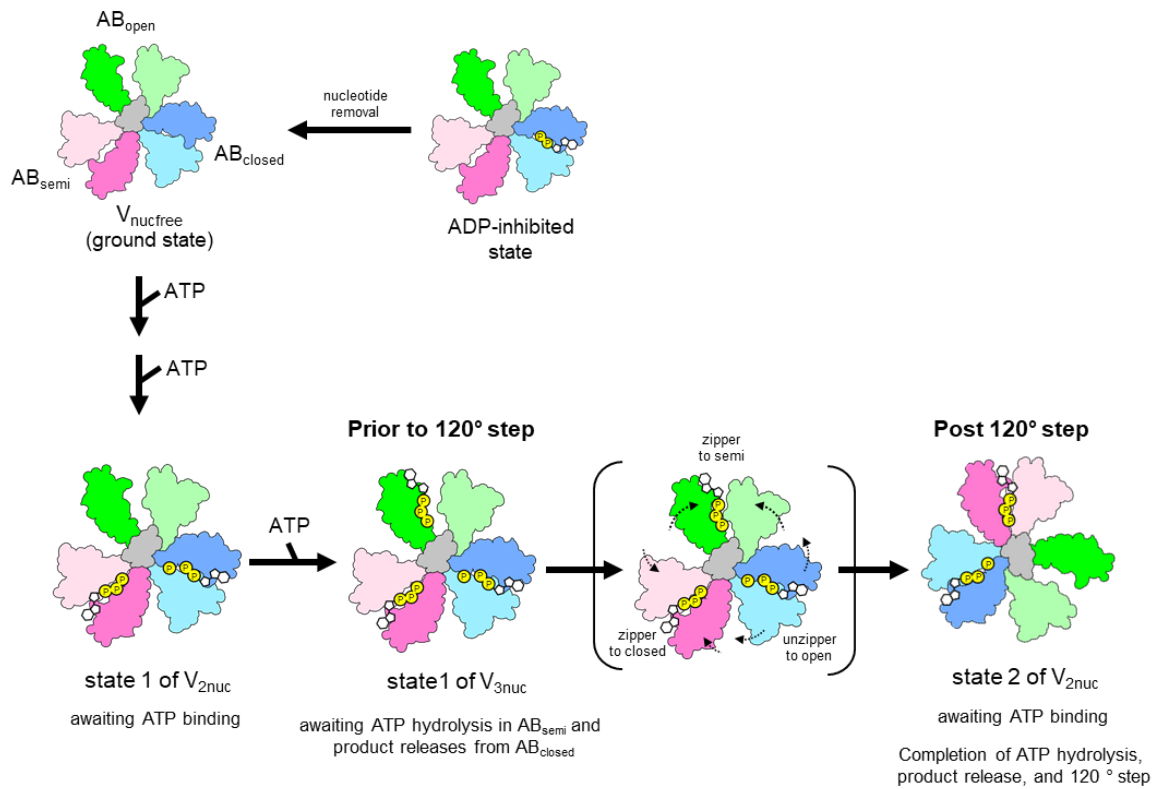
2 **Fig. 5: Coordination of nucleotides in the binding sites of V_{3nuc} and V_{prehyd}.**

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2 **Fig. 6: Chemo-Mechanical cycle of V/A-ATPase driven by ATP hydrolysis.**

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2 **Fig. 7: The rotary mechanism of V/A-ATPase powered by ATP hydrolysis.**

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