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Eukaryotic yeast V_1 -ATPase rotary mechanism insights revealed by high-resolution single-molecule studies

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Vacuolar ATP-dependent proton pumps (V-ATPases) belong to a super-family of rotary ATPases and ATP synthases. The V_1 complex consumes ATP to drive rotation of a central rotor that pumps protons across membranes via the V_o complex. Eukaryotic V-ATPases are regulated by reversible disassembly of subunit C, V_1 without C, and V_o . ATP hydrolysis is thought to generate an unknown rotary state that initiates regulated disassembly. Dissociated V_1 is inhibited by subunit H that traps it in a specific rotational position. Here, we report the first single-molecule studies with high resolution of time and rotational position of *Saccharomyces cerevisiae* V_1 -ATPase lacking subunits H and C ($V_1\Delta HC$), which resolves previously elusive dwells and angular velocity changes. Rotation occurred in 120° power strokes separated by dwells comparable to catalytic dwells observed in other rotary ATPases. However, unique $V_1\Delta HC$ rotational features included: 1) faltering power stroke rotation during the first 60°; 2) a dwell often occurring ~45° after the catalytic dwell, which did not increase in duration at limiting MgATP; 3) a second dwell, ~2-fold longer occurring 112° that increased in duration and occurrence at limiting MgATP; 4) limiting MgATP-dependent decreases in power stroke angular velocity where dwells were not observed. The results presented here are consistent with MgATP binding to the empty catalytic site at 112° and MgADP released at ~45°, and provide important new insight concerning the molecular basis for the differences in rotary positions of substrate binding and product release between V-type and F-type ATPases.

KEYWORDS

eukaryotic V_1V_o ATPase, V_1 -ATPase, single-molecule studies, rotary molecular motor, yeast vacuolar ATPase

Introduction

Vacuolar H^+ -ATPase (V-ATPase) is an ATP-dependent proton pump that regulates the pH of organelles in eukaryotic cells including Golgi, endosomes, lysosomes, and vacuoles (Kane, 2006). The plasma membranes of certain mammalian cells specialized for proton secretion also contain V-ATPases to aid in proton export from the cell (Breton and Brown, 2013; Collins and Forgac, 2020). V-ATPases are critical for a plethora of cellular processes, including protein processing and secretion, endocytosis and vesicle trafficking, zymogen activation, and autophagy (Kane, 2006; Forgac, 2007). V-ATPases are especially important

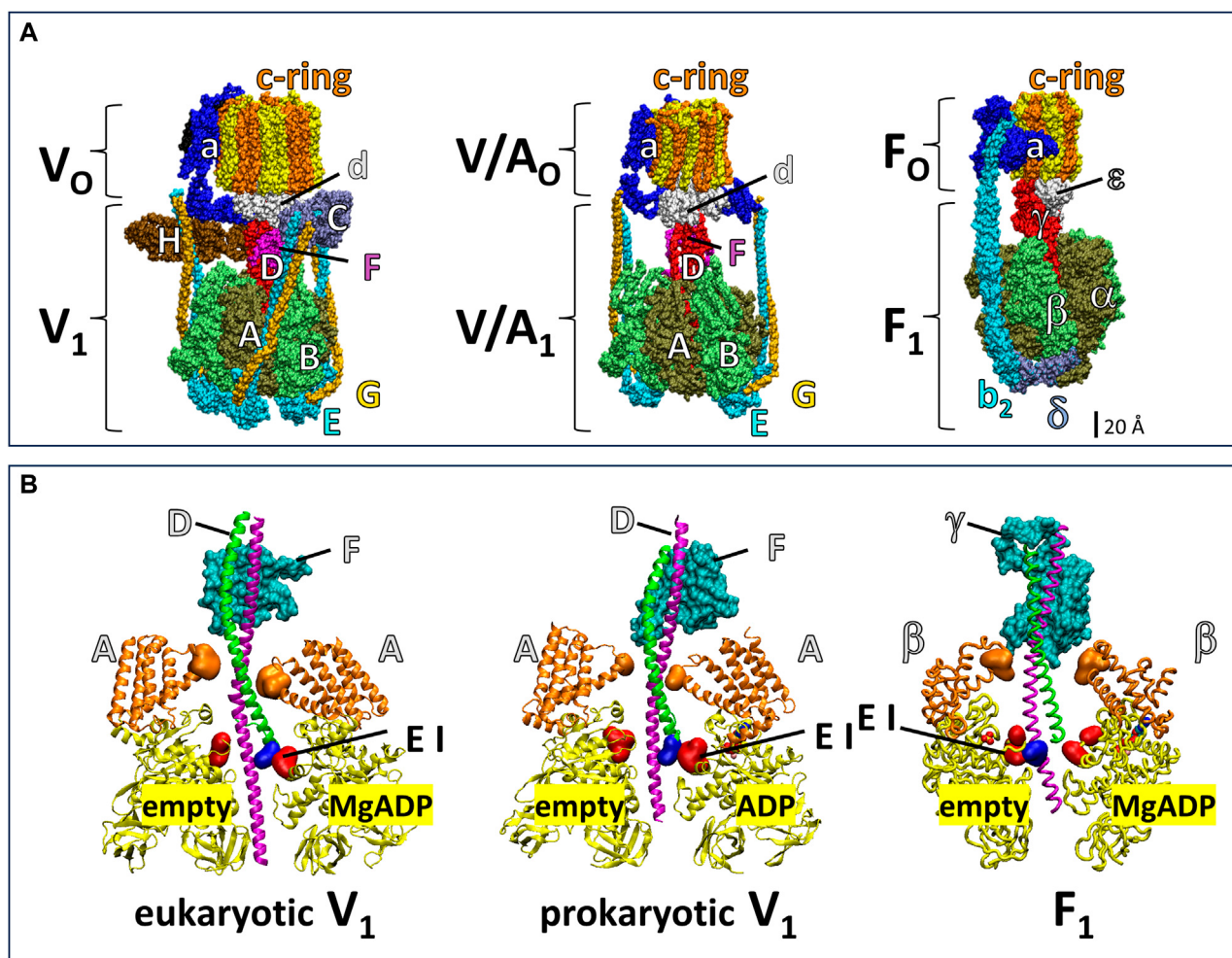


FIGURE 1

Subunit composition of eukaryotic V_1V_0 ATPase from *Saccharomyces cerevisiae* (PDB-ID 6O7W), V_1A_0 ATP synthase from *Thermus thermophilus* (PDB-ID 5GAR), and F_1F_0 ATP synthase from *E. coli* (PDB-ID 6OQR). **(A)** The rotors of V_1 and V_1A_0 are composed of subunit D and subunit F. The former has an α -helical coiled-coil that extends through the core of the $(AB)_3$ -ring, while the latter globular protein helps dock subunit D to the V_0 or V_1A_0 rotor c-ring. Analogous to subunits D and F, subunit γ serves as the F_1 rotor, which contains a coiled-coil domain that extends through the F_1 $(\alpha\beta)_3$ -ring and a globular domain that docks to the c-ring of F_0 . The eukaryotic V_1V_0 ATPase has unique regulatory subunits H and C and three peripheral stalks (EG)₃ connecting V_1 and V_0 . V_1A_0 ATP synthase has two peripheral stalks (EG)₂, and F_1F_0 ATP synthase has one (b₂) peripheral stalk. **(B)** Cross section of eukaryotic V_1 from *Saccharomyces cerevisiae* (PDB-ID 7TMO), prokaryotic V_1 from *E. hirae* (PDB-ID 5KNB); and F_1 from *E. coli* (PDB-ID 3OAA). The helical coiled-coil of the rotors is comprised of the N-terminal helix (green) and C-terminal helix (pink) of V_1 subunit F and F_1 subunit γ and the globular foot of the rotor (cyan) is V_1 subunit D and the globular domain of F_1 subunit γ . Catalytic A subunits (V_1) and β subunits (F_1) have a helical domain (orange) and a Catch-Loop (red). The electrostatic interaction (EI) occurs between the rotor (blue) and the catch loop of the subunit-A about to release MgADP in V_1 , and between the rotor and the empty subunit β in F_1 .

in human disease (Hinton et al., 2009; Alper, 2010; Breton and Brown, 2013; Hayek et al., 2014; Kartner and Manolson, 2014; Cotter et al., 2015; Cotter et al., 2016; Licon-Munoz et al., 2018).

The V-ATPase belongs to the super family of rotary ATPases (Muench et al., 2011; Schep et al., 2016; Sobti et al., 2020; Vasanthakumar et al., 2019) that also include the F-type, A-type and V/A-type (Figure 1). The eukaryotic V-ATPase is composed of the integral membrane V_0 complex that provides the pathway for proton translocation, which is docked to the peripheral V_1 complex (V_1V_0) (Benlekbir et al., 2012; Wang et al., 2020). The *Saccharomyces cerevisiae* V_0 complex consists of subunits a, d, e, f in addition to ten proteolipids (subunits c, c', c''), which form a ring structure (c-ring) (Mazhab-Jafari et al., 2016; Roh et al., 2018). The V_1 complex is comprised of eight different subunits

$A_3B_3CDE_3FG_3H$ (Zhang et al., 2003; Zhang et al., 2008; Benlekbir et al., 2012). Alternating subunits A and B form a catalytic hexameric ring consisting of three AB heterodimers, each with a catalytic site to hydrolyze ATP (Arai et al., 2013). The V_1 rotor (subunits D and F) is in the center of the $(AB)_3$ -ring and docks to the c-ring via V_0 subunit d for catalytic coupling. Three peripheral stalks (E/G heterodimers) connect the V_1 -ring of three AB heterodimers to the V_0 -proton transfer domain. The peripheral stalks provide the docking site for the V_1 regulatory H and C subunits in the V_1V_0 holocomplex (Sagermann et al., 2001; Drory et al., 2004; Diepholz et al., 2008; Oot et al., 2012). Together, the EG₁₋₃ heterodimers, subunit H, subunit C, and N-terminal domain of subunit a provide structural support between the ATPase and proton channel (Balakrishna et al., 2015).

The F_1F_0 and A_1A_0 complexes assemble into similar heterodimer rings that contain three catalytic sites surrounding a central rotor that is attached to a c-ring, which uses subunit a to facilitate proton translocation (Figure 1). However, the F-type and A-type motors have one and two peripheral stalks, respectively, and lack the eukaryotic V-type regulatory subunits H and C (Muench et al., 2011). The bacterial V/A-type ATP synthase, found in *Thermus thermophilus*, and bacterial V-type ATPase Na^+ pump found in *Enterococcus hirae* contain two peripheral stalks and also lack the subunits H and C that are essential for regulation of eukaryotic V-types (Muench et al., 2011; Schep et al., 2016; Vasanthakumar et al., 2019; Sobti et al., 2020). While F-, A-, and V/A-type ATP synthases can synthesize and hydrolyze ATP, eukaryotic V-type ATPases are ATP hydrolysis-dependent proton pumps *in vivo* (Kane, 2006; Forgac, 2007), even though *Arabidopsis thaliana* V_1V_0 has been shown to synthesize ATP at exceptionally low ATP synthase rates *in vitro* (Hirata et al., 2000).

Eukaryotic V_1 catalyzes ATP hydrolysis at the nucleotide-binding domains within the interface of the subunits A and B dimers (Arai et al., 2013; Wang et al., 2020), and most residues that facilitate ATP hydrolysis reside on V_1 subunit A. Due to the different conformations of the three A subunits relative to subunit D of the rotor, these residues catalyze ATP hydrolysis via an alternating site mechanism (Kayalar et al., 1977; Boyer, 2002) where each catalytic site is in a different conformation (open/empty state, tight/substrate bound state, and loose/product release state) at any given time. ATP hydrolysis drives V-ATPase rotation of subunits DFd and the c-ring (Hirata et al., 2003). Bacterial *T. thermophilus* V/A₁ was observed to rotate in an equivalent manner (Yokoyama et al., 2003). During rotational catalysis, carboxyl groups on the c-ring take up protons from the V_1 side of the membrane via the input channel in V_0 subunit a, which upon completing a full rotation, are deposited on the other side of the membrane via the subunit a output channel (Kawasaki-Nishi et al., 2001; Forgac, 2007; Oot et al., 2017). This enables V-ATPase-dependent proton pumping to maintain a non-equilibrium pH gradient across the cytoplasm and organellar membranes.

Eukaryotic V_1V_0 is regulated by a unique mechanism among the rotary ATPase superfamily, which results in the reversible dissociation of the V_1 -ATPase complex from the V_0 proton-transfer complex (Kane, 1995; Sumner et al., 1995; Parra and Kane, 1998; Parra et al., 2014; Hayek et al., 2019). Reversible dissociation helps to maintain cellular pH homeostasis in coordination with the metabolic state of the cell (Parra and Kane, 1998; Hayek et al., 2019). Nutrient stress conditions such as limiting glucose induce ATP hydrolysis dependent dissociation of the V_1 subunit C from V_1V_0 , which in turn prompts dissociation of V_1 from V_0 and halts proton pumping until glucose is restored and the V_1V_0 holocomplex reassembles. Upon disassembly, a conformational change of V_1 subunit H inhibits the futile hydrolysis of cytosolic ATP by the dissociated V_1 that is uncoupled from proton transport (Parra et al., 2000; Oot et al., 2016; Oot et al., 2017). In this conformation, subunit H bridges the V_1 rotor and a stator to halt rotation and trap an inhibitory Mg-ADP in one catalytic site of the autoinhibited V_1 . Glucose-dependent V-ATPase disassembly requires ATP hydrolysis (Parra and Kane, 1998), while reassembly involves the RAVE V-ATPase exclusive

assembly factor (Jaskolka et al., 2021). The protein Oxr1p appears to aid in V-ATPase disassembly (Khan et al., 2022).

However, the mechanism of how rotational catalysis mediates disassembly and reassembly remains elusive. Structural studies have identified three states of *S. cerevisiae* V_1V_0 that differ by 120° rotational positions of the rotor relative to the asymmetric stator (Zhao et al., 2015). The rotary positions of these states are thought to correspond to the catalytic dwell positions of V_1V_0 when ATP hydrolysis occurs at one of the three catalytic sites. Only one of these rotary states appears to be optimal to initiate disassembly (Mazhab-Jafari et al., 2016; Oot et al., 2016). These structures captured snapshots of individual protein conformations, but single-molecule rotation studies are required to capture the nuances of the rotary mechanism.

Although progress has been made regarding rotational studies of bacterial V/A₁-ATPase from *T. thermophilus* and V_1 -ATPase from *E. hirae* (Furuie et al., 2011; Iida et al., 2019), much less is known about the rotational mechanism of eukaryotic V_1 rotation. Single molecules of *S. cerevisiae* V_1V_0 were observed to undergo ATPase-dependent counterclockwise rotation as viewed from the membrane (Hirata et al., 2003), which was in the same direction as has been observed for F_1 , A_1 , and V/A₁ motors (Imamura et al., 2003; Spetzler et al., 2009; Furuie et al., 2011; Sielaff et al., 2016). In the *S. cerevisiae* V_1V_0 experiments, the c-ring was attached to the slide and rotation was monitored by an actin filament attached to subunit G (Hirata et al., 2003). However, the drag imposed by the actin filament and the limitation of measurement time resolution obscured other rotational details of this motor including changes in its angular velocity or whether dwells interrupt rotation. To date, only the F_1 -ATPase, A_1 -ATPase and bacterial V_1 -ATPase rotary motors have been characterized by single-molecule rotation studies under conditions that resolve dwells and angular velocity changes (Spetzler et al., 2009; Ishmukhametov et al., 2010; Sielaff et al., 2016; Ragunathan et al., 2017; Iida et al., 2019). The rotary positions of ATP binding and product release have been found to differ among the F_1 , A_1 , and bacterial V_1 motors (Furuie et al., 2011; Martin et al., 2014; Suzuki et al., 2014; Sielaff et al., 2016; Martin et al., 2018; Iida et al., 2019; Kobayashi et al., 2020; Zarco-Zavala et al., 2020), although the mechanistic basis for these differences is a major unresolved question.

Arrhenius analysis of F_1 rotation indicated that the first 60° of the 120° power stroke resulted from release of elastic energy, which was postulated to result from interactions between the rotor coiled-coil domain and the surrounding catalytic sites (Martin et al., 2018). The electrostatic interaction between highly conserved residues of the C-terminal helix of the rotor and catch loop residues (Greene and Frasch, 2003) of the empty catalytic site is thought to contribute significantly to the elastic energy that powers the first 60° of rotation. Notably, this electrostatic interaction in eukaryotic and prokaryotic V_1 -ATPases occurs between the N-terminus of the subunit F rotor and the catch loop of the catalytic site conformation that releases bound MgADP (Figure 1B).

Here, we characterized rotational dynamics of the eukaryotic *S. cerevisiae* V_1 -ATPase that lacks regulatory subunit H and subunit C using single-molecule studies with high resolution of time and rotational position. Rotation was observed in 120° power strokes separated by dwells comparable to the power strokes and catalytic dwells observed in F_1 , A_1 , and bacterial V_1 and V/A₁ ATPases.

However, the power strokes were interrupted by dwells at rotary positions that occurred most frequently 45° and 112° after the end of the catalytic dwell. The results support a mechanism in which the 45° dwell results from dissociation of ADP while the 112° dwell late in the power stroke is the result of ATP binding to the empty site.

Materials and methods

Construction of a His-tagged $V_1\Delta$ Hc and site-directed mutagenesis

To generate *S. cerevisiae* that express the V_1 -ATPase for single molecule measurements, the regulatory subunits C and H were deleted, two cysteine substitution mutations (Y73C and T123C) were made on subunit D for biotinylation, and a 6xhis tag was added to c-terminus of subunit G for purification.

First, the genes encoding subunits C and H (*VMA5* and *VMA13*, respectively) in the *S. cerevisiae* strain SF838–5Aa genomic DNA were replaced with *NAT* and *KanMX* selectable marker genes through homologous recombination, respectively. Additionally, the subunit G gene (*VMA10*) was replaced with the *URA3* gene so the wild type subunit G would not compete for assembly in V_1 because only the mutant containing the 6xhis tag was expressed. Mutant colonies were selected by inoculating the cells on SC + nourseothricin + kanamycin-uracil plates.

To generate V_1 with a 6xhis tag on subunit G, yeast genomic DNA was isolated from SF838–5Aa cells. The subunit G gene (*VMA10*) was PCR amplified using primers that contained the His tag sequence, the PCR product was restriction digested, and the DNA fragment was subcloned the expression vector. Then, 5Aa *VMA5Δ::NAT*, *VMA10Δ::URA3*, *VMA13Δ::KanMX* cells were transformed with the recombinant plasmid. Mutant colonies were selected by inoculating the cells on SC-uracil-leucine plates.

To generate the cysteine substitution mutation for the biotinylation, the subunit D gene (*VMA8*) was PCR amplified from the genomic DNA, the PCR product was restriction digested, and the DNA fragment was subcloned into the pRS313 expression vector. The two cysteine substitution mutations (Y73C and T123C) were introduced through site-directed mutagenesis. This plasmid was used as a template to PCR amplify the mutant *VMA8* gene along with the *HIS3* gene from the vector using primers containing the 5' and 3' flanking regions of the *VMA8* gene. Then, the double cysteine mutant *VMA8-HIS3* PCR product was integrated into the genome of 5Aa *VMA5Δ::NAT*, *VMA10Δ::URA3*, *VMA13Δ::KanMX* cells containing pRS315-*VMA10-6His* through homologous recombination. Mutant colonies were selected by inoculating the cells on SC-uracil-leucine-histidine plates. Mutations were confirmed after each step with agarose gel electrophoresis and DNA sequencing.

$V_1\Delta$ Hc purification

Saccharomyces cerevisiae 5Aa *VMA5Δ::NAT*, *VMA8-Y73C-T123C-HIS3*, *VMA10Δ::URA3*, *VMA13Δ::KanMX* - pRS315-*VMA10-6His* cells were grown in six 1L SC-histidine-leucine-uracil growth media at 30°C while shaking until the OD₆₀₀ was on average 1.0 OD/

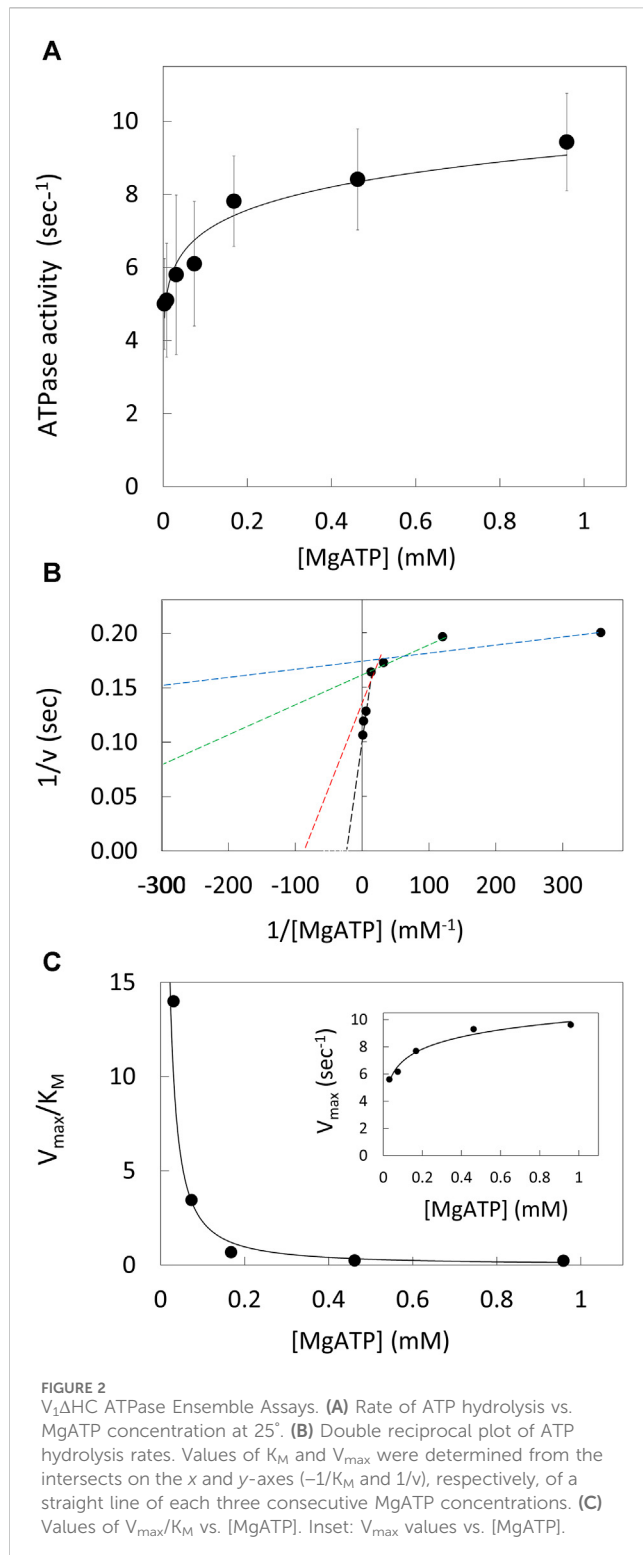
mL. The cells were spun down at 5,000 rpm at room temperature for 5 min. The cell pellets were resuspended in 200 mL of spheroplast pretreatment buffer (100 mM Tris/HCl pH 9.4, 10 mM DTT) and spun down in the same condition. The cells were resuspended in 2% glucose solution and spun down. The cells were resuspended in spheroplast buffer (10 mM Tris/HCl pH 7.5, 1.2 M sorbitol, 40% glucose) to the final concentration of 15 OD/mL. Then, 1.5 U/μL zymolyase was added to the cell suspension to the final concentration of 1U/10 OD of cells. The cells were incubated at 30°C while shaking at 80 rpm for 60 min. After incubation, the spheroplasts were spun down at 3,000 rpm in 4°C for 5 min. The pellets were resuspended in spheroplast wash buffer and spun down in the same condition and the washing step was repeated two more times (spheroplast wash buffer: 6.8 mg/mL Yeast Nitrogen Base, 50 mM sodium phosphate dibasic, 50 mM succinic acid/NaOH pH 5.0, 2% glucose, 1.2 M sorbitol, 0.02 mg/mL histidine, 0.12 mg/mL leucine, 0.02 mg/mL adenine, 0.06 mg/mL lysine, 0.02 mg/mL arginine, 0.02 mg/mL tryptophan, 0.03 mg/mL tyrosine, 0.2 mg/mL threonine, 0.02 mg/mL methionine, 0.05 mg/mL phenylalanine, 0.02 mg/mL uracil). The following steps were done at 4°C. The cell pellet was homogenized in 20 mL of lysis buffer (PBS +1% (w/v) C₁₂E₉ 1mM PMSF, 5 μg/mL aprotinin, 2 μg/mL chymostatin, 1 μg/mL pepstatin A, 1 μg/mL leupeptin) and incubated in ice for 10 min. The samples were centrifuged at 30 k rpm (109,000 xg). After centrifugation, 10X binding buffer (0.5 M Tris/HCl pH 8.0, 1 M KCl, 400 mM imidazole, 50 mM MgCl₂) was added to the supernatant to make the final concentration 1X. Finally, V_1 -ATPase was purified from the mixture by Ni-NTA chromatography, 1 mg of biotin maleimide was added to the column elution and incubated at 4°C for 15 min while shaking, and the sample was run through Sephadex G50 column equilibrated with storage buffer (50 mM Tris/HCl pH 8.0, 20 mM KCl, 2 mM ATP, 1 mM MgCl₂, 15% glycerol). The purified, biotinylated V_1 -ATPase samples were aliquoted into 20 μL, quickly frozen, and stored at –80°C until use.

ATP hydrolysis assay

The rate of ATP hydrolysis by the purified $V_1\Delta$ Hc was measured with an ATP-regenerating NADH-coupled assay (Lotscher et al., 1984). The measurement was made with the final concentration of 25 mM Tris/HCl (pH 8.0), 60 mM KCl, 2.5 mM phosphoenolpyruvate, 0.3 mM NADH, 17.5 Units pyruvate kinase (rabbit muscle, Sigma Aldrich), 25 Units L-lactate dehydrogenase (rabbit muscle, Sigma Aldrich), at varying concentrations of ATP including 1, 0.5, 0.2, 0.1, 0.05, 0.02, 0.01 mM, twice the MgCl₂ concentration for each corresponding ATP concentration, and 3.22×10^{-5} mM of purified $V_1\Delta$ Hc (0.0174 mg/mL) in a final volume of 2.5 mL. The rate was determined in three replicates as the change in absorbance at 340 nm using a Cary 100 spectrophotometer with Peltier temperature control at 25°C. MgATP concentration was determined by the Maxchelator program MgATP calculator v1.3 using constants from NIST database #46 v8 (UC Davis Health).

Single molecule gold nanorod rotation assay

The rotation of individual $V_1\Delta$ Hc molecules were observed with a single-molecule rotation assay using gold nanorods under a dark



field microscope (Spetzler et al., 2009; Martin et al., 2014). Purified $V_1\Delta HC$ molecules were immobilized on a microscope cover slip by the His-tag on the G-subunits, unbound molecules were washed off the slide with wash buffer (30 mM Tris/HCl pH 8.0, 10 mM KCl). The surface area of the cover slip that remained exposed around the bound $V_1\Delta HC$ molecules was then coated with BSA-C, which prevented the gold nanorods from binding nonspecifically to the

surface. The 80×40 nm gold nanorod (A12-50-600 purchased from Nanopartz) coated with Neutravidin was bound to the biotinylated subunit D, excess gold nanorods were washed off with the wash buffer, and rotation buffer (1 mM $MgCl_2$, 2 mM ATP, 30 mM Tris/HCl pH 8.0, 10 mM KCl) was added to the cover slip (Spetzler et al., 2009). The rotations of individual molecules were observed by measuring the fluctuation of polarized red light scattered off the AuNR using a single-photon detector. In each molecule observed, the orientation of the polarizing filter was adjusted to align with the minimum light intensity position that corresponded to one of the three catalytic dwells. The sinusoidal fluctuation of the polarized red-light intensity was measured as the gold nanorod rotated from 0° to 90° relative to the catalytic dwell position. Measurements were taken in the form of 5 s datasets at a frame rate of 100 kHz. The standard error measurements of histograms of the intensity of red light scattered from a single nonrotating nanorod fixed to a slide as a function of the rotational position of the polarizer varies between about 0.02 and 0.12° as the scattered light intensity varied between minimum and maximum values (Ishmukhametov et al., 2010).

Results

A strain of *S. cerevisiae* lacking the genes for the V_1 regulatory subunit H and subunit C was used to express the V_1 -ATPase complex (hereafter $V_1\Delta HC$). $V_1\Delta HC$ resembles the V_1 naturally found in the cytosol, which lacks subunit C. However, unlike the cytosolic V_1 , $V_1\Delta HC$ is not inhibited by subunit H. For single-molecule rotation studies, subunit G of $V_1\Delta HC$ was genetically modified to add a 6xhis tag to the C-terminus, and subunit D mutations Y73C and T123C were made to enable covalent modification by biotin maleimide. The $V_1\Delta HC$ construct exhibited ensemble ATP hydrolysis commensurate with V-ATPases and single molecule rotational ATPase activity consistent with the general mechanism of the rotary ATPases. However, $V_1\Delta HC$ rotation differed in notable ways.

Ensemble ATPase assays

The ATPase activity of purified $V_1\Delta HC$ versus the MgATP concentration (Figure 2A) was measured using an ensemble coupled assay with pyruvate kinase and lactic dehydrogenase at $25^\circ C$. The apparent V_{max} was observed at $990 \mu M$ MgATP. The double-reciprocal plot of ATPase activity versus [MgATP] was not linear (Figure 2B). As such, $V_1\Delta HC$ did not exhibit simple Michaelis-Menten kinetics, which would have a linear dependence in a double reciprocal plot defined by Eq. 1,

$$1/v = (K_M/V_{max})(1/[MgATP]) + 1/V_{max} \quad (1)$$

where v is the observed velocity at a given MgATP concentration, V_{max} is the apparent maximum velocity and K_M is the Michaelis constant. The kinetic values were determined from Eq. 1 for each of three consecutive ATP concentrations in the double-reciprocal plot. The V_{max} increased with MgATP concentration to a maximum turnover number of $9.6 s^{-1}$ (Figure 2C, inset). The time required to hydrolyze an ATP at $990 \mu M$ MgATP (saturating) as well as at $490 \mu M$ and $5.7 \mu M$ MgATP, which were rate-limiting, is shown in

TABLE 1 Durations of $V_1\Delta$ HC power strokes, dwells and ATP consumption derived from single-molecule measurements compared to those derived from the ensemble ATPase assay *versus* MgATP concentration.

[MgATP] (μ M)	Single-molecule rotation measurements					Ensemble ATPase assays	
	Number of power strokes	Number of $V_1\Delta$ HC ^a	Average power stroke duration (μ s)	Average catalytic dwell duration (ms)	Average ms/ATP ^b	ms/ATP ^c	% $V_1\Delta$ HC active ^d
990	10,274	48	625	7.2	7.8	105.2	7.4
490	1,024	8	745	12.3	13.0	118.9	11
5.7	1,625	14	910	13.4	14.3	200.0	7.2

^aNumber of $V_1\Delta$ HC, molecules examined.
^bAverage ATP, consumption time (sum of power stroke and catalytic dwell durations).
^cAverage ensemble ATP, consumption time (1/apparent V_{max}).
^dPercent of $V_1\Delta$ HC, molecules actively consuming ATP, in ensemble assay (single-molecule ms/ATP/ensemble ms/ATP).

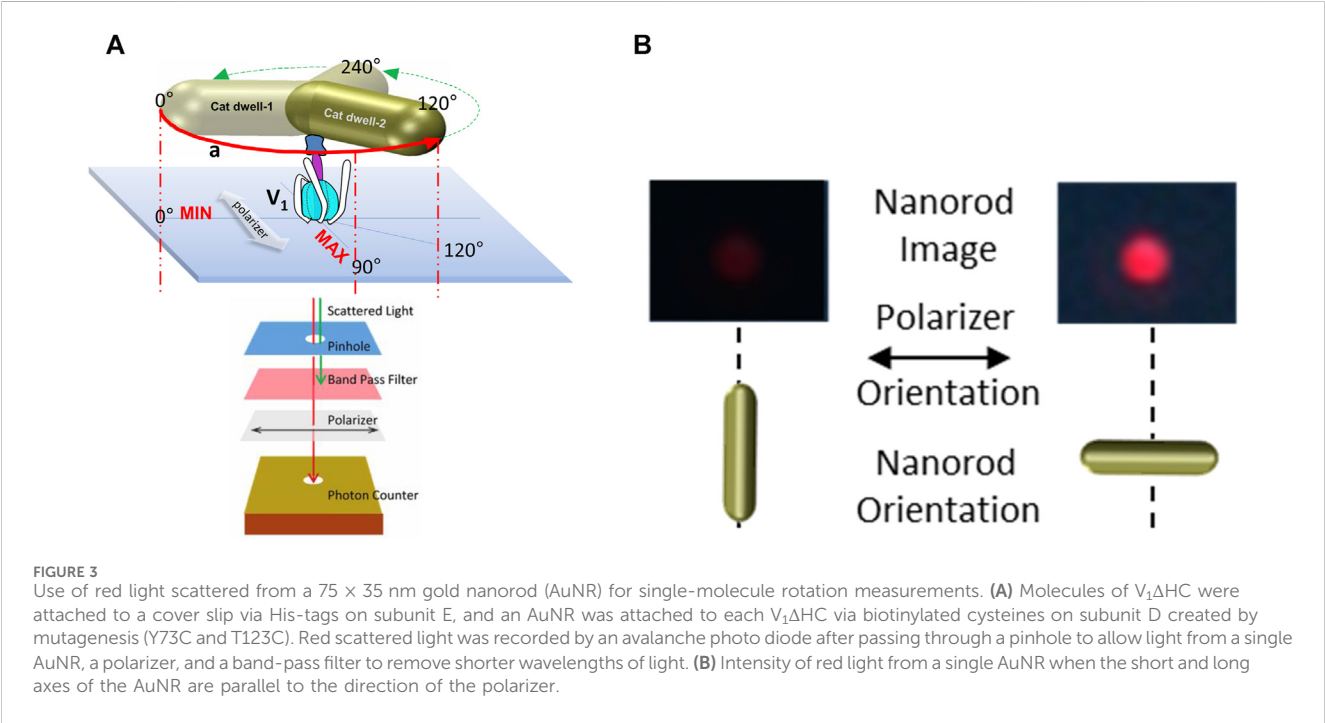
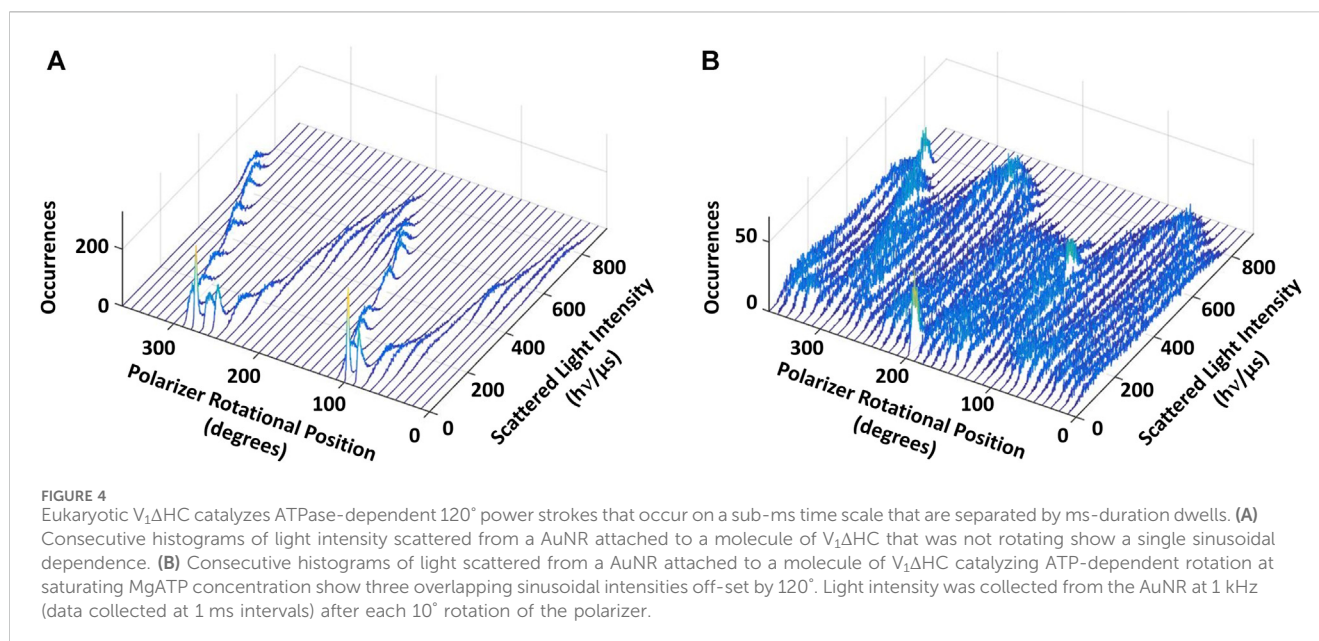


Table 1. The V_{max}/K_M values decreased 30-fold *versus* MgATP in a non-linear manner (Figure 2C), which indicates that the affinity of the empty catalytic site to bind MgATP decreases as $V_1\Delta$ HC approaches saturating MgATP concentrations. At saturating MgATP concentrations, the K_M was 41.7 μ M. Changes in both V_{max} (~2-fold) and K_M (100-fold) contributed to the decrease in V_{max}/K_M *versus* MgATP.

Single molecule rotation assays

To measure ATP hydrolysis-dependent rotation of subunit D, purified $V_1\Delta$ HC molecules were immobilized on a microscope cover slip by the His-tags on the three G subunits, and a 35 × 75 nm NeutrAvidin-coated gold nanorod (AuNR) was attached to the biotinylated Y73C and T123C mutations of subunit D

(Figure 3A). Rotation of single $V_1\Delta$ HC molecules was observed in the presence of MgATP by the changes in intensity of polarized red light that scatters from the AuNR. While concentrations of the MgATP complex used in ensemble ATPase measurements and single-molecule experiments were the same, the Mg:ATP ratio was 2:1 and 1:2, respectively. The 1:2 ratio used in the single-molecule experiments was intentional to minimize MgADP inhibition. These measurements were also carried out in the absence of an ATP regenerating system because the small number of V_1 molecules on the cover slip did not consume significant amounts of ATP during the assay. After the scattered red light passed through a high wavelength band-pass filter to remove light of wavelengths shorter than 600 nm, and a polarizing filter that could be rotated to specific orientations, the light intensity changes were quantified by an avalanche photodiode with a 50 ns time resolution. In this manner the intensity of scattered



red light was at a minimum and maximum when the polarizer was oriented perpendicular and parallel to the long axis of the AuNR (Figure 3B).

Figure 4 shows the results of a polarizer rotation measurement when the AuNR is attached to subunit D of a single $V_1\Delta$ Hc molecule. In this experiment, the light intensity scattered from the polarizer was rotated 10° in successive 5 s intervals for a total of 360°. During this time, the intensity of light scattered from the AuNR was acquired by the single-photon counter at 1 kHz (equivalent to 1,000 fps). Since red light scatters specifically from the long axis of the AuNR, the scattered light intensity when subunit D was not rotating in this experiment (Figure 4A, control) varied in a sinusoidal manner relative to the rotational axis of the polarizer (Spetzler et al., 2006; Ishmukhametov et al., 2010; Frasch et al., 2022).

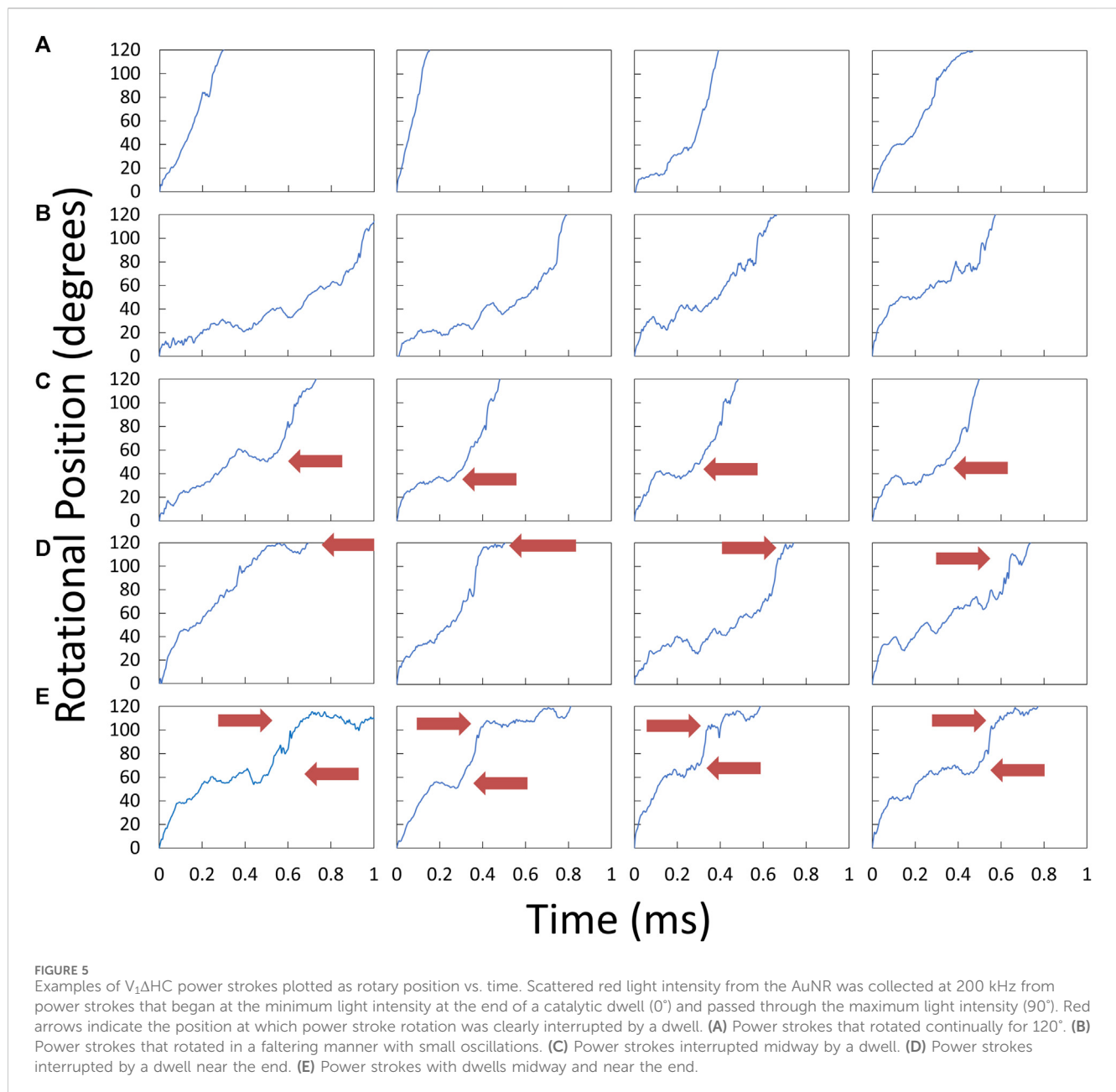
In the presence of 990 μM (saturating) MgATP (Figure 4B), three sinusoidal intensity curves were observed in the polarizer rotation measurement, which were off-set from each other by 120°. The 1 kHz data acquisition rate provides a time resolution of 1 ms such that the light intensities reported the rotational position of the AuNR primarily when it was in the same position for more than 1 ms. This indicates that, in the presence of saturating MgATP, subunit D stopped at three rotary positions separated by 120° that each lasted >1 ms. To do so, subunit D rotated between these dwells with a power stroke that occurred too fast for the 1 kHz data acquisition rate to record as much scattered light as was observed during the dwells. These dwells are hereafter referred to as catalytic dwells, since they are comparable to the duration and rotary positions of catalytic dwells observed between the 120° ATPase-driven power strokes of F_1 , A_1 , V/A_1 and bacterial V_1 ATPases (Spetzler et al., 2006; Furuike et al., 2011; Minagawa et al., 2013; Sielaff et al., 2016; Zarco-Zavala et al., 2020).

To resolve the intermediate positions of the 120° rotational events between the dwells observed in Figure 4, the intensity of light scattered from the AuNR was sampled at 200 kHz (5 μs per

data point). Prior to the 5 s data acquisition of each $V_1\Delta$ Hc molecule, the rotational position of the polarizer was set so that the scattered light intensity was at a minimum at one of the three catalytic dwell positions. As a result, the light intensity of the subsequent power stroke increased from a minimum through a maximum as the AuNR rotated from 0° to 90° relative to that catalytic dwell, then decreased until it reached the next catalytic dwell upon rotating 120°. An arcsine^{1/2} function of light intensity (Sielaff et al., 2016; Martin et al., 2018) was used to calculate rotational position *versus* time. Changes in rotary position *versus* time (angular velocity) for each power stroke were then averaged and binned to every 3° of rotation, where 0° and 120° refer to catalytic dwell positions.

Examples of $V_1\Delta$ Hc power strokes that were used to determine the angular velocity profiles *versus* rotary position are shown in Figure 5. Many $V_1\Delta$ Hc power strokes rotated continuously (Figure 5A), which are comparable to *E. coli* F_1 power strokes at saturating MgATP (Martin et al., 2018). However, many other $V_1\Delta$ Hc power strokes rotated in a faltering manner (Figure 5B) with frequent small oscillations. Some contained a clearly defined dwell during the first 60° of the power strokes (Figure 5C) even in the presence of saturating MgATP, while other power strokes contained a dwell near the end of the power stroke (Figure 5D). Although the power strokes shown were observed in the presence of 5.7 μM MgATP, all four types were present at all three of the MgATP concentrations examined.

The distribution of rotary positions during the power stroke when $V_1\Delta$ Hc dwells occurred (Figure 6) shows that dwells were most commonly observed ~45° and 112° after the catalytic dwell. The distribution of the former dwell was significantly broader than the latter, and the peak of the distribution observed at 990, 490, and 5.7 μM MgATP appeared to occur at 45°, 50°, and 40°, respectively (hereafter designated the 45° dwell). This is the approximate rotary position at which F_1 -ATPases gives rise to an “ATP-binding” dwell when MgATP is rate-limiting (Yasuda et al., 2001; Bilyard et al., 2013; Martin et al., 2014; Sobti et al., 2021).

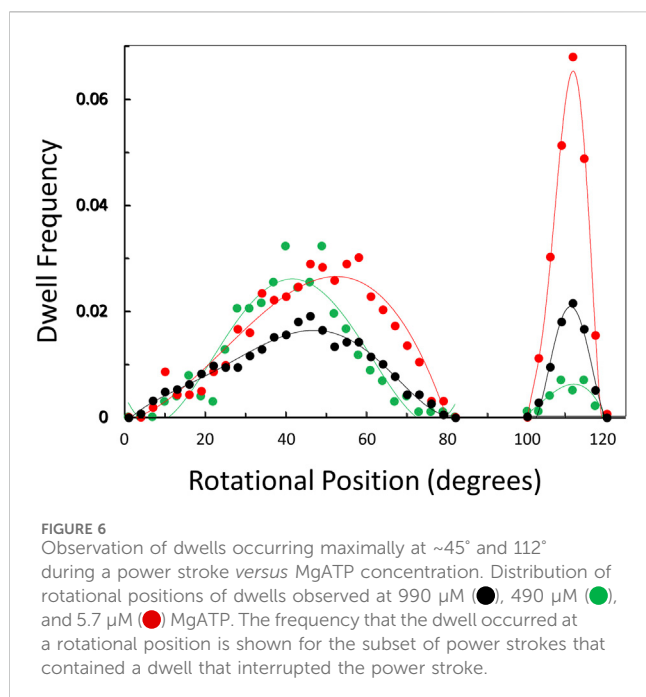


The percentage of power strokes examined that contained a 45° dwell increased from 24% at saturating MgATP by 6% and by a further 8% at the rate-limiting MgATP concentrations of 490 and $5.7 \mu M$ MgATP, respectively (Table 2). The $184 \pm 2 \mu s$ average duration of these dwells did not change significantly at the rate-limiting MgATP concentrations. However, the occurrence of the 112° dwell increased by ~ 3 -fold at $5.7 \mu M$ MgATP with respect to that of the 8.5% occurrence observed at saturating MgATP. The duration of these dwells increased by 21% ($391 \mu s$) at $5.7 \mu M$ MgATP.

The average angular velocity of the $V_1\Delta HC$ power stroke vs. rotary position in the presence of saturating ($990 \mu M$) MgATP was calculated from the data of 10,274 power strokes examined from 48 $V_1\Delta HC$ molecules (Figure 7A). The initial average velocity as the catalytic dwell ended was $\sim 350^\circ \cdot ms^{-1}$. The $V_1\Delta HC$ decelerated to

$\sim 200^\circ \cdot ms^{-1}$ at $\sim 13^\circ$ (d1-deceleration), then slowly decelerated further from $\sim 23^\circ$ to $\sim 180^\circ \cdot m^{-1}$ at 60° (d2-deceleration). The rate subsequently accelerated to $\sim 470^\circ \cdot ms^{-1}$ at 85° (a1-acceleration), and then rapidly accelerated to briefly reach a rate of $\sim 1,200^\circ \cdot ms^{-1}$ at $\sim 90^\circ$ (a2-acceleration) before decelerating at $\sim 93^\circ$ (d3-deceleration). The rate returned to $300^\circ \cdot ms^{-1}$ at $\sim 100^\circ$ then decelerated to $100^\circ \cdot m^{-1}$ (d4-deceleration) as it approached the next catalytic dwell at 120° . The $V_1\Delta HC$ angular velocity profile was closely similar to that of the *E. coli* F_1 -ATPase (Martin et al., 2018) with the exception that the angular velocity of the latter was significantly slower during the first 70° , and was 21% slower during the spike in velocity at 90° (Figure 7A).

Changes in the angular velocity profile of the $V_1\Delta HC$ power stroke vs. rotary position were observed when measured at $490 \mu M$ MgATP and at $5.7 \mu M$ mgATP (Figures 7B, C), which gave rise to



ATPase rates that were 86% and 65% of that observed at $990\ \mu\text{M}$ MgATP, respectively. In the presence of $490\ \mu\text{M}$ MgATP (Figure 7B), the rates after the d2-deceleration, after the a1-acceleration, and after the d3-deceleration were $\sim 100^\circ\cdot\text{ms}^{-1}$ at $\sim 45^\circ$, $\sim 200^\circ\cdot\text{ms}^{-1}$ at 85° , and $\sim 180^\circ\cdot\text{ms}^{-1}$ at 100° . These rates corresponded to rate decreases of 2-fold, 2.4-fold, and 1.7-fold, respectively, from those observed at saturating MgATP. When measured at $5.7\ \mu\text{M}$ MgATP (Figure 7C), the average $V_1\Delta\text{Hc}$ angular velocity profile showed additional decreases in velocity from that observed $490\ \mu\text{M}$ MgATP between rotary positions 60° and 120° . Notably, the $5.7\ \mu\text{M}$ MgATP velocities at 85° , 90° , and 100° were $230^\circ\cdot\text{ms}^{-1}$, $440^\circ\cdot\text{ms}^{-1}$, and $110^\circ\cdot\text{ms}^{-1}$, respectively, which represents decreases of 2.0-fold, 2.7-fold, and 2.7-fold from that observed at saturating MgATP.

The average time required for power strokes to rotate between catalytic dwells were calculated from the angular velocity profiles at each MgATP concentration (Table 1), since each profile reports the time required for each three degrees of rotation. At saturating MgATP ($990\ \mu\text{M}$ MgATP), the average power stroke duration was $625\ \mu\text{s}$ while the average power stroke durations at the rate-limiting MgATP concentrations of $490\ \mu\text{M}$, and $5.7\ \mu\text{M}$ MgATP were 745 and $910\ \mu\text{s}$, respectively. The power stroke durations at

these limiting MgATP concentrations were 1.19-fold and 1.46-fold longer than that measured at saturating MgATP. These increases were comparable to the 1.16-fold and 1.53-fold longer times required to consume an ATP as determined from the ensemble ATPase measurements in the presence of $490\ \mu\text{M}$ and $5.7\ \mu\text{M}$ MgATP, respectively (Table 1). These results support the conclusion that the additional time required for MgATP to bind to the empty catalytic site when MgATP is rate-limiting is evident as a decrease in angular velocity during the power stroke.

The angular velocity profiles (Figure 8) were determined from the average of several thousand power strokes. Consequently, the decreases in average angular velocities observed at limiting MgATP may be due to slower rotation or may occur if a proportion of the power strokes briefly stop rotating, which would be observed as a dwell. The extent that limiting MgATP caused decreases in the angular velocity *versus* rotary position was determined by taking the difference angular velocity profiles at limiting MgATP from that at saturating MgATP (Figures 8A, B). When compared to the distribution of dwells (black squares) there is a clear correlation between the incidence of dwells and the decreases in angular velocity of the power stroke, but the frequency did not change as MgATP became increasingly limited. This suggests that the 45° dwell does not result from MgATP binding. However, the largest decreases in power stroke angular velocity when MgATP is rate-limiting was observed between 80° and 100° when dwells did not occur. Consequently, these results indicate that the major contribution to the decrease in ATPase rate when MgATP is limiting occurs between 80° and 120° .

The consumption of each ATP requires a consecutive catalytic dwell and power stroke. The power strokes analyzed here are one of three required for a complete rotation of subunit D. Based on the number of power strokes analyzed for each $V_1\Delta\text{Hc}$ molecule during the 5 s data acquisition period, and the average power stroke durations obtained directly from the angular velocity profiles, the average duration of catalytic dwells was determined by subtraction of the time consumed by the power stroke from the total data acquisition time (Table 1). The average catalytic dwell durations in the presence of $990\ \mu\text{M}$, $490\ \mu\text{M}$, and $5.7\ \mu\text{M}$ MgATP were calculated to be $7.2\ \text{ms}$, $12.3\ \text{ms}$, and $13.4\ \text{ms}$, respectively. The average durations of the power strokes and catalytic dwells were consistent with the polarizer rotation results that have a minimum time resolution of $1\ \text{ms}$ (Figure 4).

The average time required to consume an ATP molecule was also calculated from the sum of the average durations of the power stroke and the catalytic dwells (Table 1). In the presence of 990 , 490 , and $5.7\ \mu\text{M}$ MgATP, the average times to consume an ATP were

TABLE 2 Occurrence and duration of dwells that interrupt the $V_1\Delta\text{Hc}$ power stroke *versus* MgATP concentration.

[MgATP] (μM)	45° dwell		112° dwell	
	% occurrence ^a	Average duration (μs)	% occurrence ^a	Average duration (μs)
990	24.0	184 ± 2	8.5	322 ± 5
490	30.2	183 ± 6	3.4	265 ± 25
5.7	38.9	194 ± 5	23.6	391 ± 10

^aPercent occurrence of dwell *versus* total power strokes analyzed at each MgATP concentration.

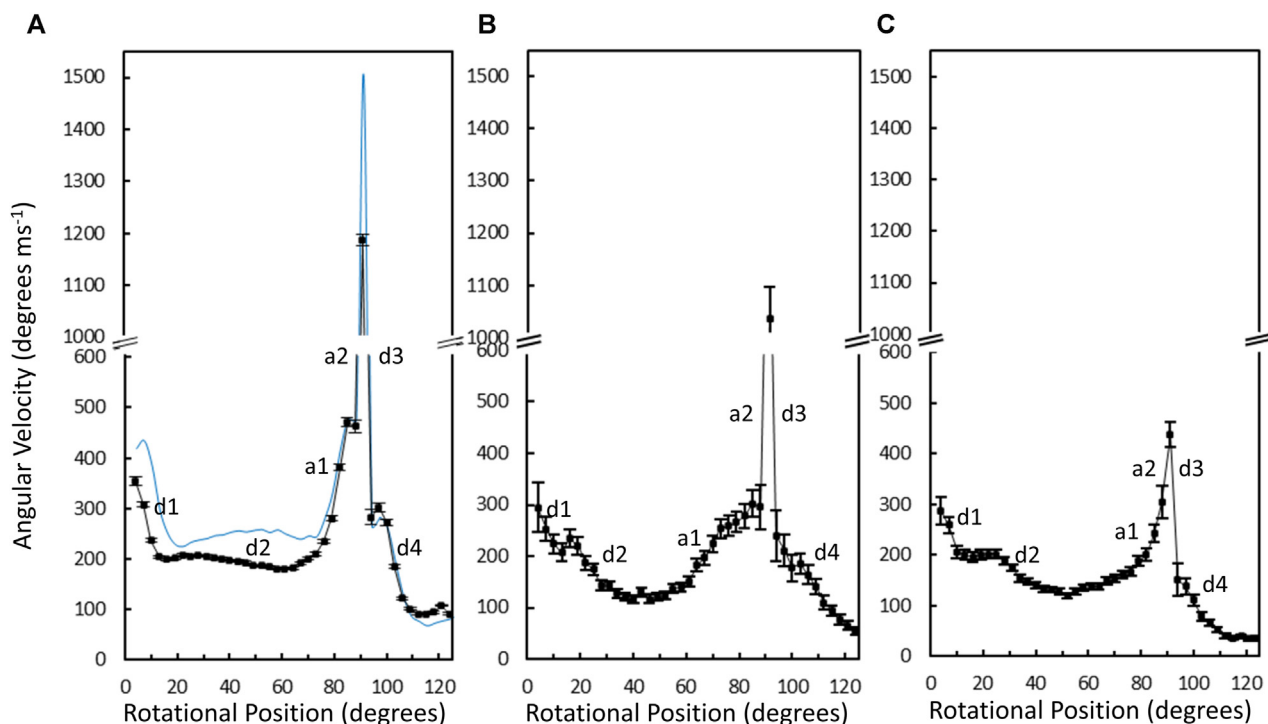


FIGURE 7
Average angular velocity vs. rotational position of *V*₁ΔHC (black) versus *E. coli* F₁ (blue) power strokes at 990 μM Mg-ATP (A), 490 μM MgATP (B), and 5.7 μM MgATP (C). Rotational positions of decelerations (d) and accelerations (a) are numbered in the order that they occur during the power stroke.

calculated from the single-molecule data to be 7.8, 13.0, and 14.3 ms, respectively. It is noteworthy that the times required to consume ATP as measured by the ensemble ATPase assay (Figure 2) at 990, 490, and 5.7 μM MgATP were 105.2, 118.9, and 200 ms (Table 1). These times were considerably longer than those determined by other single-molecule studies, where each molecule was known to be undergoing ATPase-dependent rotation. The ensemble assays reported the average of many *V*₁ΔHC molecules without knowing how many molecules are actively consuming ATP. By comparing the ensemble and single-molecule results, we estimate that 7%–11% of the *V*₁ΔHC molecules were actively consuming ATP at any moment in the ensemble assay (Table 1).

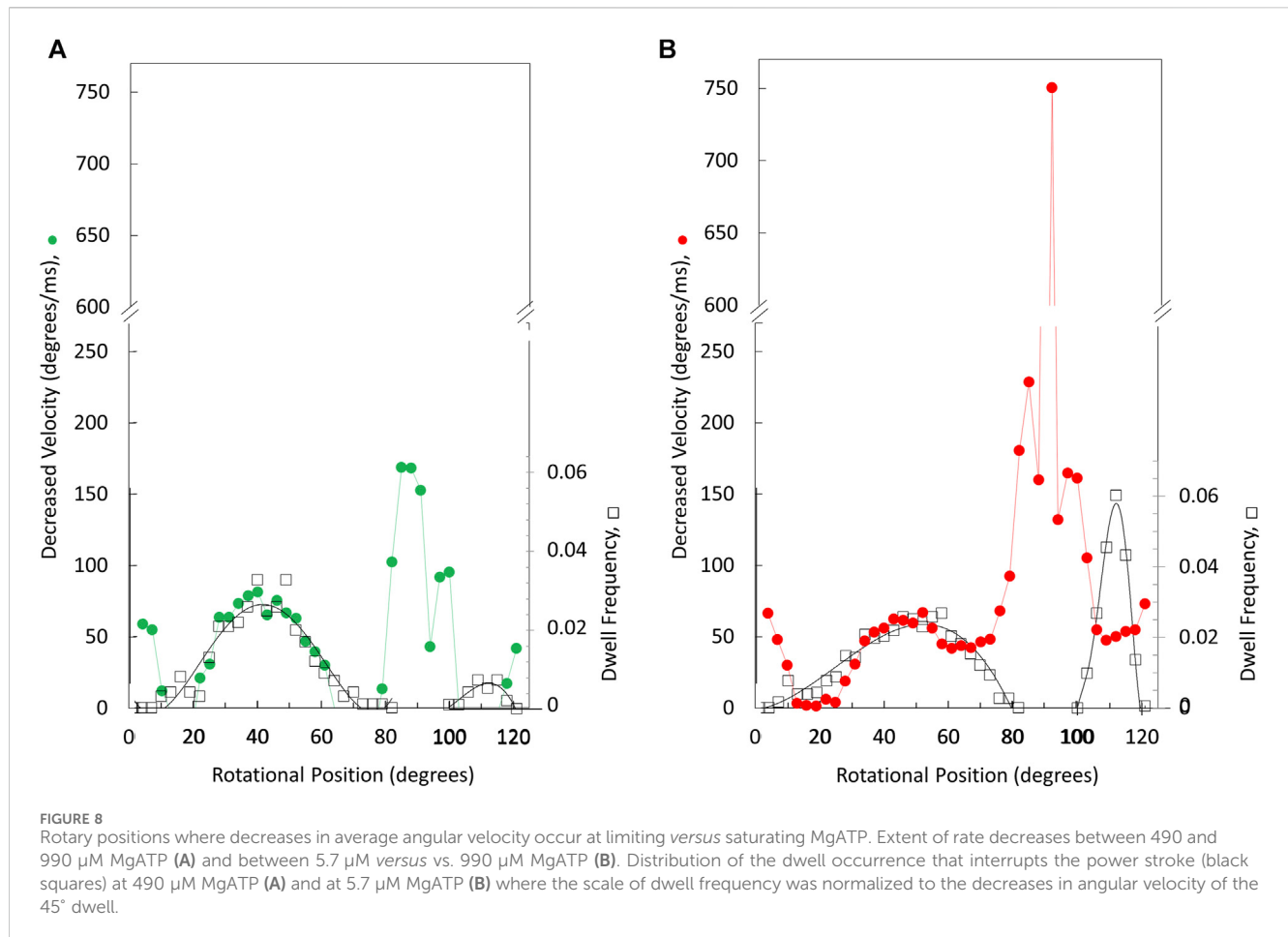
Discussion

The single-molecule results of eukaryotic *V*₁ΔHC ATPase-dependent rotation presented here are consistent with a mechanism in which subunits D and F rotate in 120° power strokes separated by catalytic dwells when ATP hydrolysis occurs. This is supported by the presence of 120° power strokes that last for 0.63 ms–0.91 ms separated by longer 7.2–13.4 ms duration dwells. These power strokes have the same velocity profile as those of the F₁-ATPases and A₁-ATPases examined except for the magnitudes of the velocities (Sielaff et al., 2016; Ragunathan et al., 2017; Martin et al., 2018). These longer dwells are also consistent with catalytic dwells observed in F₁-ATPases. These mechanistic features are shared by all members of the super family of rotary ATPases examined to date (Figure 9) even though these rotary ATPases

have been found to vary in the rotational positions where ATP binds, as well as where ADP and Pi are released. The results presented here support the mechanism of the *V*₁ΔHC shown in Figure 9A, and provide important new insight concerning the molecular basis for the differences in rotary positions of substrate binding and product release among these rotary motors.

The *V*₁ΔHC rotation differed in notable ways from the more thoroughly studied F₁-ATPase. First, rotation during the *V*₁ΔHC power strokes often faltered with small back and forth oscillations during the first 60° of rotation subsequent to the catalytic dwell. Second, the *V*₁ΔHC power strokes often contained a dwell occurring 40°–50° and/or a dwell at 112° after the catalytic dwell (8° before the subsequent catalytic dwell). Third, as the MgATP concentration became increasingly limited, the duration of the power stroke lengthened not only as the result of an increase in the occurrence and/or duration of 45° and 112° dwells, but also because the angular velocity decreased between 80° and 100° when dwells did not occur (Figure 8).

Interactions between the rotor and the catalytic sites provide clues to the differences in *V*₁ΔHC rotation (observed here) from that of F₁ ATPases. In both ATPases, the rotors are surrounded by contacts at the tip of the C-terminal helical domains (CHDs) of each subunit in the (AB)₃-ring and by the (αβ)₃-ring, where the catalytic sites are primarily located on the A-subunits and β-subunits. Both *V*₁ and F₁ have a strong electrostatic interaction between highly conserved catch-loop residues from one of the three catalytic sites with their respective rotors. These residues are A/D422, A/S424, and A/D425 and βD301, βT304, and βD305 in *S. cerevisiae* *V*₁, and *E. coli* F₁, respectively. However, the location of the rotor residues that

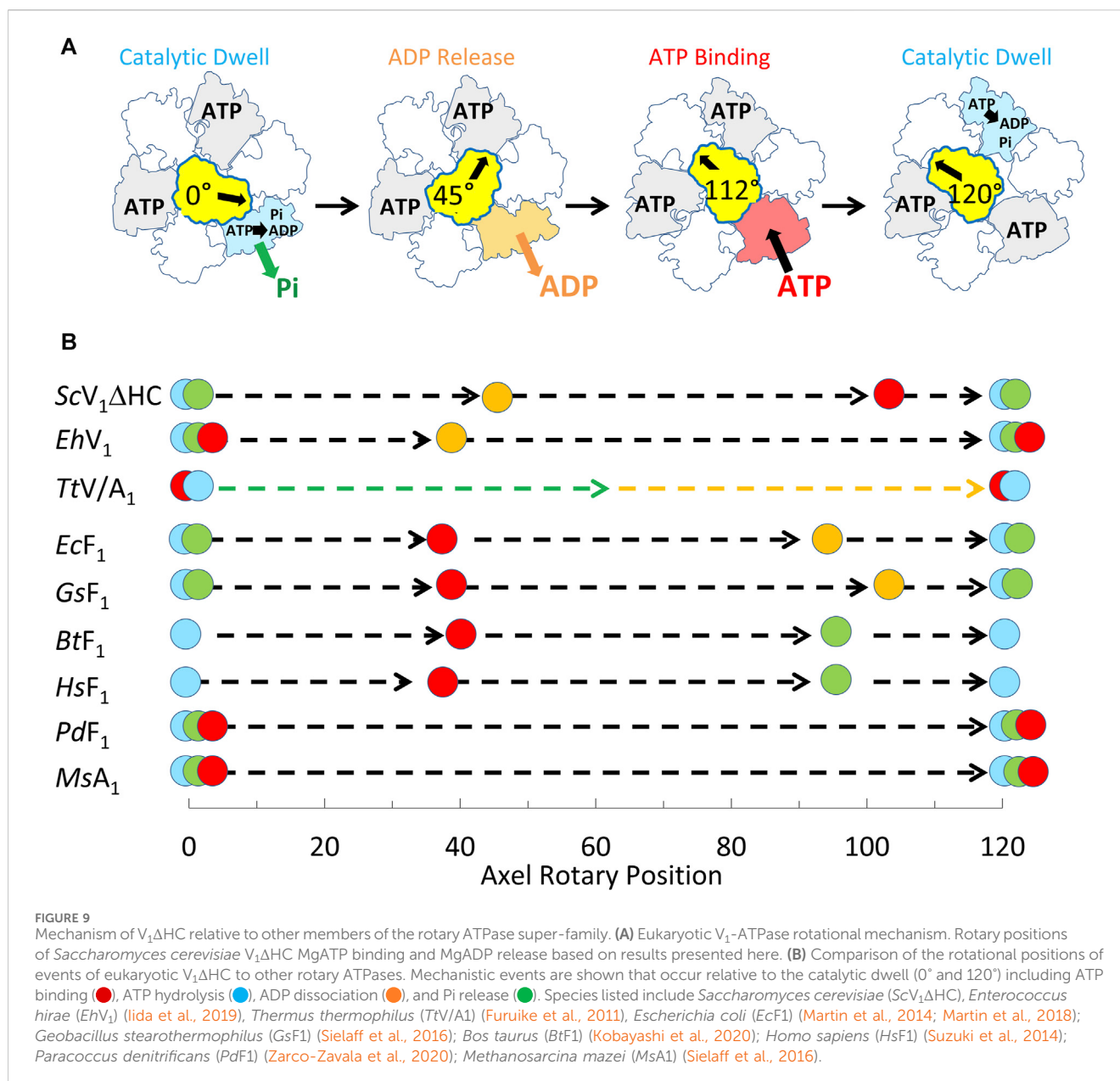


form the electrostatic interaction differs significantly. In *S. cerevisiae* V_1 -ATPase, these residues (D/R12) are on the shorter N-terminal helix of the subunit D coiled-coil (Vasanthakumar et al., 2022) while these F_1 residues (γ Q269 and γ R268 in *E. coli*) are on the longer C-terminal helix of the coiled-coil. As a result, V_1 subunit D forms electrostatic interactions with the catch-loop of the subunit A catalytic site that is about to release ADP (loose conformation), while F_1 subunit γ interacts electrostatically with the catch-loop of the empty β subunit (open conformation). Mutations of any of the residues that comprise this electrostatic interaction in *E. coli* F_1 and *S. cerevisiae* V_1 -ATPase result in dramatic losses of catalytic activity (Greene and Frasch, 2003; Boltz and Frasch, 2006; Arsenieva et al., 2010).

Elastic coupling powered by ATP hydrolysis provides an explanation for faltering rotation (throughout first 60°) during *S. cerevisiae* $V_1\Delta\text{HHC}$ power strokes, which result from small oscillations and the occurrence of 45° dwells even at saturating MgATP. In F_1 , the restraints on the rotor imposed by the surrounding CTHs and by the catch loop interactions impose elastic strain by twisting the coiled-coil of the rotor. Unwinding this coiled-coil spring then powers the first 60° of rotation (Martin et al., 2018; Frasch et al., 2022). If the first 60° of V_1 rotation is also powered by unwinding the coiled-coil of the rotor, the spring constant is likely to be decreased due to the subunit D short helix location and its electrostatic interaction to the ADP release catalytic site.

The V_1 subunit D electrostatic link to the catalytic subunit interaction that releases ADP also suggests that the 45° dwells result from the dissociation of ADP. *E. hirae* V -ATPase has equivalent subunit D residues forming the electrostatic interaction with the ADP-release conformation (Suzuki et al., 2016). Of the rotary ATPases studied to date (Figure 9B), the *E. hirae* V -ATPase appears most closely related to eukaryotic V -ATPases. *E. hirae* V_1V_O is an ATPase-dependent Na^+ pump that is incapable of ATP synthesis. Single-molecule studies of purified *E. hirae* V_1 show the presence of a dwell that occurs 40° after the catalytic dwell, which results from ADP dissociation, while ATP-binding occurs at the catalytic dwell (Iida et al., 2019). It is noteworthy that ADP-dependent rotational backsteps are observed during *E. hirae* V_1 power strokes, which are much larger than the small oscillations of the $V_1\Delta\text{HHC}$ power strokes observed here. Recent structures of the *T. thermophilus* A/ V_1 , which also have the subunit D electrostatic interaction at the catalytic site that releases ADP indicate that ADP and P_i dissociate during the 120° power stroke, although at positions that are undefined to date (Figure 9B).

The $V_1\Delta\text{HHC}$ results presented here are consistent with MgATP binding to the empty catalytic site at 112°. The duration and occurrence of the 112° dwell increased when MgATP became rate-limiting, and large decreases in angular velocity were observed during the final 60° of rotation. Supporting further this conclusion, the duration of the 45° dwells did not increase with decreasing MgATP, even though the occurrence of these dwells increased to some extent. *E. hirae* V_1



single-molecule studies suggested that ATP binding occurs during catalytic dwells (Iida et al., 2019). It is noteworthy that the *S. cerevisiae* $V_1\Delta HC$ catalytic dwell duration did not change significantly when the major decrease in ATP hydrolysis occurred (from $490\ \mu M$ to $5.7\ \mu M$ MgATP). However, dwell duration did increase by 1.86-fold relative to $990\ \mu M$ MgATP (Table 1) suggesting that some $V_1\Delta HC$ molecules may bind MgATP during the catalytic dwell 8° later when MgATP is saturating.

The 112° dwell is unique in several ways. Its frequency of occurrence is MgATP dependent. The dwell occurrence is significantly more frequent when MgATP concentration is limiting at $5.7\ \mu M$ (23.6% occurrence) than saturating at $990\ \mu M$ (8.5% occurrence) (Table 2). In addition, the dwell is distinctly preceded by a major angular velocity reduction (Figure 8) that is remarkably steep at limiting MgATP. A nucleotide binding dwell 112° after the catalytic dwell has not been observed in other rotary

ATPases to our knowledge. The eukaryotic V-ATPase is uniquely regulated by reversible disassembly of V_1 and V_O , which is an important regulatory mechanism that requires V_1V_O ATP hydrolysis (Parra and Kane, 1998) that traps the dissociated V_1 complex in a specific rotational state (Oot et al., 2016; Vasanthakumar et al., 2022). More work is required to determine whether this distinct 112° dwell is a functional adaptation of the eukaryotic rotary V-ATPases, and/or has been observed here as the result of increased resolution of our single-molecule assay.

Scope statement

V-ATPases (V_1V_O -ATPases) are conserved rotary molecular motors that regulate cellular pH and play crucial roles in a large repertoire of physiological processes and human illnesses.

Developing therapies that target V_1V_O -ATPase with precision requires understanding dynamics of V-ATPase rotation at high resolution at a molecular level. Here we report single-molecule rotation studies of the yeast V_1 -ATPase complex with high resolution of time and rotational position. Single molecules of V_1 -dependent rotation occurred in 120° power strokes similar to those of other rotary ATPases. However, these 120° rotational steps were interrupted by dwells at 45° and 112° when the product (ADP) was released and a new substrate (ATP) bound, respectively. This nucleotide binding sub-step at 112°, which may be unique to eukaryotic V_1 -ATPases, was distinctly preceded by a major reduction in angular velocity. This is important because current V-ATPase inhibitors immobilize the V_1V_O assembled state, although V-ATPases are regulated by reversibly disassembling V_1 and V_O *in vivo*. These results will help to design drugs that target a specific rotational sub-step to prevent reassembly and trap a disassembled and naturally inhibited state, leading the way in the development of a new generation of treatments that reversibly control V-ATPase function.

Data availability statement

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

Author contributions

SY: Writing—original draft, Writing—review and editing, Formal Analysis, Investigation, Methodology, Validation, Visualization. ZB: Writing—review and editing, Data curation, Formal analysis, Software, Validation, Visualization. KP: Writing—review and editing, Conceptualization, Formal Analysis, Funding acquisition, Methodology, Project administration, Resources, Supervision. WF: Writing—original draft, Writing—review and editing,

Conceptualization, Formal Analysis, Funding acquisition, Methodology, Project administration, Resources, Supervision, Validation, Visualization.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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