Rotary catalysis of bovine mitochondrial F₁-ATPase studied by single-molecule experiments

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The reaction scheme of rotary catalysis and the torque generation mechanism of bovine mitochondrial F₁ (bMF₁) were studied in single-molecule experiments. Under ATP-saturated concentrations, high-speed imaging of a single 40-nm gold bead attached to the γ subunit of bMF₁ showed 2 types of intervening pauses during the rotation that were discriminated by short dwell and long dwell. Using ATPγS as a slowly hydrolyzing ATP derivative as well as using a functional mutant βE188D with slowed ATP hydrolysis, the 2 pausing events were distinctively identified. Buffer-exchange experiments with a nonhydrolyzable analog (AMP-PNP) revealed that the long dwell corresponds to the catalytic dwell, that is, the waiting state for hydrolysis, while it remains elusive which catalytic state short pause represents. The angular position of catalytic dwell was determined to be at +80° from the ATP-binding angle, mostly consistent with other F₁s. The position of short dwell was found at 50 to 60° from catalytic dwell, that is, +10 to 20° from the ATP-binding angle. This is a distinct difference from human mitochondrial F₁, which also shows intervening dwell that probably corresponds to the short dwell of bMF₁ at +65° from the binding pause. Furthermore, we conducted “stand-and-release” experiments with magnetic tweezers to reveal how the binding affinity and hydrolysis equilibrium are modulated by the γ rotation. Similar to thermophilic F₁, bMF₁ showed a strong exponential increase in ATP affinity, while the hydrolysis equilibrium did not change significantly. This indicates that the ATP binding process generates larger torque than the hydrolysis process.

F₅αF₆ATP synthase (or ATP synthase) is one of the most ubiquitous enzymes found in the mitochondrial inner membrane, chloroplast thylakoid membrane, and bacterial plasma membranes (1, 2). F₅αF₆-ATP synthase catalyzes the ATP synthesis reaction coupled with H⁺ (or sodium in some bacteria) translocation, which is driven by proton motive force (pmf) across membranes. This enzyme is structurally and functionally separated into 2 components, F₁ and F₀, both of which are rotary molecular motors. F₁ is the protruding portion from the membrane and possesses catalytic reaction centers for ATP synthesis. F₀ is the membrane-embedded portion and conducts H⁺ translocation across the membrane. In the whole ATP synthase complex, F₁ and F₀ interconvert the free energy of ATP hydrolysis and pmf, via the mechanical rotation of the rotor complex. When the free energy of ATP hydrolysis per turn of the rotor complex exceeds the pmf per turn, F₅αF₆-ATP synthase catalyzes the reverse reaction and hydrolyzes ATP, pumping H⁺ to generate pmf.

F₅αF₆-ATPase (F₅αF₆), when isolated from F₅αF₆, hydrolyzes ATP to ADP and inorganic phosphate (P₀). Upon catalysis, F₁ rotates the rotor complex against the surrounding stator ring, on which the catalytic reaction centers are located. The subunit composition of F₁ is αβ₃γδε in both bacterial and mammalian types. However, the δ and the ε subunits are not equivalent in bacterial and mammalian types. The minimum complex as a rotary motor is the αβ₃γ subcomplex.

The atomic structures of F₁ have been intensively studied by X-ray crystallography since the first report on bovine mitochondrial F₁, bMF₁, in 1994 (3). The first crystal structure revealed most of the basic structural features of F₁, which were repeatedly confirmed in later structural studies on bMF₁ and other F₁s (4–6). F₁ is composed of the α/β₃ stator ring and the central rotor complex of the γε in bacterial types and the γδε in mammalian types. In the α/β₃ stator ring, the α and the β subunits are arranged alternately. The catalytic sites reside on one side of the α/β interface, while the other side of the α/β interface binds to ATP; however, it is catalytically impotent and thereby termed the noncatalytic site. The catalytic residues are mostly located on the β subunit, except for the catalytically critical arginine residue termed the “arginine finger” on the α subunit (3, 7–9). Among the 3 β subunits, 2 β subunits bind to nucleotides: one β binds to the ATP analog AMP-PNP and the other binds to ADP [later revealed to also bind to azide (5)]. These β subunits, termed β₁ and β₂, respectively, adopt so-called closed conformation, in which the C-terminal helical domain rotates inwardly to the rotor γ subunit. The third subunit, pₚempt, has no bound nucleotide and adopts an open conformation, swinging the C-terminal domain outwardly. From the structural features, it has been proposed that ATP binding triggers the open-to-closed conformational transition of the β subunit, which is a major power-stroking motion. The conformational transition of the β subunit was later visualized using the single-fluorescence polarization technique (10), Förster resonance energy transfer (11), and high-speed atomic force microscopy (12).

Significance

The gold-standard model for structural analysis of F₅αF₆-ATPase has been bovine mitochondrial F₁ (bMF₁), but its rotational dynamics remain elusive. This study analyzes rotational characteristics of bMF₁. bMF₁ showed 3 distinct dwells in rotation, “binding dwell,” “catalytic dwell,” and “short dwell,” in each 120° step of rotation. While the positions of binding and catalytic dwell are similar to those of human mitochondrial F₁ (hMF₁), bMF₁ shows short dwell at a distinctively different position from the corresponding dwell of hMF₁, implying variation in the timing of the putative reaction at short dwell, phosphate release or ADP release. Single-molecule manipulation experiments revealed that the affinity change of ATP is a major torque-generating step.

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Later crystallographic studies showed that \( \beta_{DP} \) can bind to AMP-PNP (4) or a transition-state analog (13), while \( \beta_{TP} \) predominantly binds to AMP-PNP unless AMP-PNP is omitted from the crystallization medium. Therefore, \( \beta_{DP} \) is thought to represent the catalytically active conformational state for cleavage of bound ATP. This contention is supported by several studies (7, 14, 15). In some crystal structures, \( \beta_{empty} \) has a small anionic ligand: phosphate (6), thiophosphate (16), with the implication that \( \beta_{empty} \) represents the pre-/post-\( P_i \)-release state, as supported in a theoretical study (17).

Since its first visualization, the rotary dynamics of F\(_1\) have been well studied by single-molecule rotation assay (18). In a typical rotation assay, the \( \alpha\beta\gamma \) ring is immobilized on the glass surface, and a probe for rotational imaging is attached onto the protruding component of the \( \gamma \) subunit. For high-speed imaging of rotation practically not affected with viscous friction against a rotating probe, nanoprobes are used such as 40-nm gold colloid (19, 20) or nanorods (21). For the torque measurement or rotation manipulation, 100- to 500-nm-diameter polystyrene beads or magnetic beads are used. Single-molecule rotation analysis has elucidated the basic features of \( F_1 \) rotation and the chemomechanical coupling scheme of rotation.

Among \( F_1 \)'s characterized by the rotation assay thus far, thermophilic Bacillus PS3 (TF\(_1\)) is the best characterized due to its high conformational stability and clear stepping behaviors. The rotation analysis of TF\(_1\) established the reference reaction scheme, although some variations for different \( F_1 \)'s have been observed as described later. The unitary step size of the rotation is 120°, each coupled with a single turnover of ATP hydrolysis, reflecting the pseudo 3-fold symmetry of the structure. The 120° step rotation is divided into 2 substeps of 80° and 40° (19), each intervened by ATP-waiting dwell (binding dwell) or catalysis-waiting dwell (catalytic dwell), respectively (22). During binding dwell, another β releases ADP (23, 24), and P\(_i\) release is suggested to occur during catalytic dwell (24, 25). Considering that each β exerts a single turnover of ATP hydrolysis upon a single turn of the rotor, and the reaction phase is different by 120° among 3 βs, the reaction scheme is proposed as shown in Fig. 1D (25), although another scheme has also been proposed (26).

Recent statistical analysis (27) revealed that TF\(_1\) makes a small rotation upon catalysis during catalytic dwell that is too small to be detected in conventional image analysis, suggesting that the catalytic dwell is split into hydrolysis and P\(_i\)-release dwells. The split of the catalytic dwell was also proposed in studies on the rotation of yeast mitochondrial F\(_1\) (28) and human mitochondrial F\(_1\) (hMF\(_1\)) (29). The work on hMF\(_1\) showed that the 120° rotation was resolved into 3 substeps: 65°, 25°, and 30°. Each step was initiated by ATP binding, presumably P\(_i\) release and hydrolysis. Therefore, the dwells before the 65°, 25°, and 30° substeps are referred to as the binding dwell, P\(_i\)-release dwell, and catalytic dwell, respectively. The reaction scheme of hMF\(_1\) was proposed as shown in Fig. 1B. Due to the close sequence homology of hMF\(_1\) and bMF\(_1\) (99% in the \( \alpha \) and \( \beta \) subunits and ~95% in the \( \gamma \) subunit (16)), it is expected that the reported rotation behavior of hMF\(_1\) is similar to bMF\(_1\). From the viewpoint of the structure–function relationship of \( F_1 \), the correlation between dwells and conformational states found in crystal structures is important to determine. Assays with inhibitors suggest that the P\(_i\)-release dwell corresponds to the state found in the majority of bMF\(_1\) crystal structures, including the first crystal structure (3), ground-state structure (4), and thiophosphate-bound structure (16). However, there are still differences in amino acid sequences between hMF\(_1\) and bMF\(_1\), and the investigation of the exact correlation between rotary dynamics and atomic structure of F\(_1\) requires a rotation assay with F\(_1\) from the same species used in the crystal structure analysis. Although a preliminary study on bMF\(_1\) was reported (30), basic characteristics of bMF\(_1\) have not been analyzed.

A single-molecule rotation assay of F\(_1\) enabled not only detailed kinetic analysis of stepping rotation but also manipulation experiments when combined with a magnetic tweezers system. The manipulation experiment of TF\(_1\) was first conducted for the direct demonstration of ATP synthesis upon the reverse rotation of the \( \gamma \) subunit (31, 32). After that, it has become a major focus how F\(_1\) modulates the rate and equilibrium constants of elementary reaction steps: binding, hydrolysis, and product releases. To assess this issue, a stall-and-release experiment was conducted to determine the rate constant and equilibrium constant of ATP binding or hydrolysis of ATP bound on the catalytic site as a function of rotary angle that formed a basis for following theoretical studies (17, 33–38). Significantly larger angle dependence of ATP binding than hydrolysis revealed that TF\(_1\) generates larger torque in the ATP binding step than in the hydrolysis step. However, the stall-and-release experiments have been conducted only for TF\(_1\) (39) and the generality of these findings remains unclear.

In this study, we investigated the \( \gamma \) rotation of bMF\(_1\) and found several differences in rotation dynamics between bMF\(_1\) and hMF\(_1\), from which we propose the reaction scheme for bMF\(_1\) shown in Fig. 1C. Based on the reaction scheme, we also analyzed the angle dependence of ATP affinity change as well as the modulation of the equilibrium constant of ATP hydrolysis by

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**Fig. 1.** Chemomechanical coupling rotation schemes of TF\(_1\) (A), hMF\(_1\) (B), and bMF\(_1\) (C). Each circle and arrow represents the catalytic state of the β subunit and the angular positions of the \( \gamma \) subunit, respectively. 0° is defined as the position of \( \gamma \) subunit where a β subunit (orange) binds to ATP. The asterisks following “ATP” represent the catalytically active state to undergo hydrolysis of bound ATP.
conducting a stall-and-release experiment. The single-molecule manipulation analysis revealed the general features of angle dependence of binding and catalysis are well-conserved across the species, suggesting that the torque generation mechanism is common among F₈s, although the stepping behaviors have some variations.

Results

Construct of bMF₁ for Rotation Assay. Recombinant bovine mitochondrial F₁, composed of α, β, γ, δ, and ε subunits was coexpressed with assembly factors, AF1 and AF2, and purified according to a previous report (30) with slight modifications (Materials and Methods). Two cysteines were introduced in the protruding part of the γ subunit at A99 and S191. They were specifically biotinylated to residues (His-tag) were introduced at the N terminus of the subunits. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis showed the genuine subunit composition was retained after dilution, showing the complex of bMF₁ is stable at the nanomolar condition where the single-molecule rotation assay is to be conducted (SI Appendix, Fig. S1).

ATP-Driven Rotation. To observe genuine rotation of bMF₁, 40-nm gold colloid attached to the γ subunit of bMF₁ was observed under various [ATP] values at 23 ± 2 °C (Fig. 2C) with a dark-field microscope (20). Images were recorded at 125 to 45,000 frames per second (fps) (22 to 8,000 μs per frame), depending on [ATP]. Red data points in Fig. 2B show the Michaelis–Menten curve of the rotation rate, in which the maximum rotation rate (V_max) and Michaelis constant (K_m) were 707 revolutions per second (rps) and 77 μM, respectively. The maximum rotation rate of 707 rps was comparable with that of human mitochondrial F₁ (hMF₁), 741 rps, and remarkably faster than those of bacterial F₁s: 129 rps for thermophilic Bacillus PS3 (TF₁) (19) and 449 rps for E. coli F₁ (EF₁) (40). Considering the coupling ratio of 3 ATPs per turn, the maximum rotation rate corresponded to the ATP hydrolysis rate of 2,121 per s. We also measured the ATP hydrolysis rate of bMF₁ in solution with ATP-regeneration system (blue data points in Fig. 2B) and determined V_{ATP} and K_{ATP} as 1,037 ± 2° C (Fig. 2B) (19) and hMF₁ (2.7 ± 10^7 M⁻¹ s⁻¹) (29). At high [ATP] over K_m, several bMF₁ molecules showed 6 pauses as found in the rotation of hMF₁ (29), although many of the molecules did not show 6 clear pauses. Subpauses were detected in the angle histograms by eye in 4 of 23 molecules. Fig. 2D showed x-y projections of a trajectory and the corresponding histogram of angular position observed at 3 mM ATP. The time course also shows multiple pauses within one revolution. Three of the 6 pauses should correspond to catalytic dwell as found in TF₁ and hMF₁. The estimated time constant of ATP binding at 3 mM should be less than 10 μs, too short to be detected. Thus, the intervening pause is not binding dwell. These suggest that bMF₁ makes an intervening pause in addition to catalytic dwell. Note that the response time of the 40-nm gold nanoparticle was ∼0.1 ms, and thereby submillisecond events are principally blurred and difficult to resolve. Therefore, the dwell-time analysis at high [ATP] was impractical.

ATPγS-Driven Rotation. To resolve the rotation and dwells more clearly, we observed rotation in the presence of ATPγS, which is a slowly hydrolyzable ATP analog. The previous rotation assays showed that ATPγS slows the ATP hydrolysis on TF₁ (22) and also presumably release of thiophosphate on hMF₁ (29). Rotation rates of bMF₁ were determined at various [ATPγS] to draw the Michaelis–Menten curve (SI Appendix, Fig. S3A). V_max and K_m were determined to be 20.3 rps and 2.2 μM, respectively. As expected, V_max was largely suppressed at about 35 times slower than that of the ATP-driven rotation. The binding constant of ATPγS, K_{ATPγS}, was estimated from 3 x V_max/K_m to be 3.0 ± 10^7 M⁻¹ s⁻¹, which was almost identical to K_ATP.

At high [ATPγS] over K_m, bMF₁ showed distinct pauses separated by 120° steps, corresponding to 3 dense clusters in the x-y angles. 

Fig. 2. ATP-driven rotation of bMF₁. (A) A schematic image of the single-molecule rotation assay of bMF₁. The α6βγ-ring is immobilized on a glass surface, and a detection probe is attached to the γ subunit via biotin–streptavidin interaction. (B) [ATP] versus the rate of rotation (red) or ATPase/3 (blue). The mean value and the SD for each data point are shown with circles and error bars, respectively (n = 20 to 25 for measurement of rotation rate, n = 3 for measurement of ATPase). Solid lines represent Michaelis–Menten fittings; V_max = 707 ± 5 rps, K_m = 77 ± 2 μM for rotation rate; V_ATPase = 346 ± 11 s⁻¹, k_ATPase = 218 ± 26 μM for ATPase/3 (fitted parameter ± fitting error). (C and D) x-y plot, angular histogram, and time course of rotation found at 300 nM ATP (C) and at 3 mM ATP (D). The recording rate was 500 and 45,000 fps, respectively.
plot of rotation (Fig. 3 A, Inset). A closer look at the time courses showed additional short pauses during 120° steps (Fig. 3A), showing that bMF1 makes 2 types of dwells, hereafter referred to as long dwell (blue in Fig. 3A) and short dwell (orange in Fig. 3A) in a 120° rotation. Short dwell was not always observed in each 120° step due to the short lifetime. To identify the dwells and steps of rotation objectively, we employed a nonparametric change-point (CP) analysis to detect angular changes in the rotary traces based on the permutation test (27). Unlike other parametric-level detection methods in single-molecule time series analysis, such as hidden Markov modeling (43), our nonparametric CP analysis does not require any a priori assumption of the noise model and the number of pauses/steps in the rotary trace. CP analyses have also been shown to outperform the commonly used method of binning and thresholding the time series that may introduce artifacts to the waiting time kinetics in low signal-to-noise cases (44).

Details of our CP detection method are given in SI Appendix, Supplementary Information Text and Fig. S4.

The presence of short dwells was confirmed in the CP analysis. Fig. 3B shows angular histograms of a representative molecule. Fig. 3 B, Upper shows a conventional angular histogram prepared from all data points of a time course trajectory. Fig. 3 B, Lower shows the histogram constructed from the CP intervals, denoted as a histogram of CP intervals, in which the angular positions between 2 successive CPs are represented by the median angle of the interval. Each CP interval gives a single count in the histogram regardless of the dwell length, contrary to the conventional angle histogram where a longer dwell provides more counts. It is evident from Fig. 3B that the histogram of CP intervals clearly showed 3 clusters corresponding to the short dwells (orange) between distinctly high peaks of long dwells (blue). The angular position of short dwell was 60° from a long dwell at the left side (clockwise side) (SI Appendix, Fig. S5A). The dwell time was analyzed for long and short dwells, both of which showed single exponential decay functions, giving time constants of 14 to 15 ms for long dwell and 0.7 to 0.9 ms for short dwell (Fig. 3C and SI Appendix, Fig. S6). It should be noted that in addition to long and short dwells, distinctively long pauses over 1 s were also observed occasionally (SI Appendix, Fig. S7 A–C). We attributed the occasional long pause to ADP inhibition, considering that all characterized F1βs in the rotation assay showed long dwells during rotation by ADP inhibition on the order of seconds (40, 41, 45, 46).

The position of the ADP-inhibition dwell coincided with the position of long dwell (SI Appendix, Fig. S7C), suggesting that long dwells correspond to catalytic dwell where F1 executes the ATP cleavage reaction (22).

Rotation was observed at [ATPγS]s below K_{cat}, which was expected to resolve the rotation into binding dwell and long dwell. The recording rate was set at 1,000 fps to preserve image data storage of the high-speed camera. This allowed long-time observations. As expected, 2 types of dwells were found during 120° rotations in both the angle histogram and time course (Fig. 3D and E). In the region below K_{cat}, the binding pause limited the overall rotation rate, showing longer dwells than long dwells. In the time course and angle histogram, the longer pauses were identified as binding dwells (pink). Relatively shorter pauses were assigned as long dwells. The angle distance between long dwell and binding dwell was 77° (Fig. 3E, Lower). This is mostly consistent with hMF1 (29) and TF1 (22). As expected, when [ATPγS] was decreased, the duration time of binding dwell was lengthened. Fig. 3F and SI Appendix, Fig. S6 show the dwell-time histogram for binding dwell and long dwell. The time constant
of long dwell was constant at 11 to 13 ms, consistent with the abovementioned value (14 ms). The length of binding dwell depended on [ATP] as expected, giving the rate constants of ATP$^+$ binding ($k_{\text{on}}^{\text{ATP}}$), $2.9 \times 10^7$ M$^{-1}$s$^{-1}$, well consistent with $k_{\text{on}}^{\text{ATP}}$ (3.0 $\times 10^7$ M$^{-1}$s$^{-1}$) determined from the Michaelis–Menten analysis. In the rotation assay at low [ATP$^+$], a distinctly long pause attributable to ADP inhibition was again observed at the angle of long dwell (SI Appendix, Fig. S7 D–F), suggesting that long dwell corresponds to catalytic dwell. Short dwells were not detected throughout the rotation assay with low [ATP$^+$], probably because short dwells of ATP$^+$ rotation are too short to be detected with the recoding frame rate (1 ms per frame).

**Rotation of bMF$_1$(βE188D).** To confirm that long dwell is +80° from binding angle, we tested a mutant F$_1$(βE188D) in the rotation assay. This glutamic acid is highly conserved in primary sequences among all F$_1$s. In crystal structures, this glutamic acid interacts with the γ phosphate via a coordinated water molecule. A quantum mechanics/molecular mechanics study revealed that this glutamic acid accelerated the ATP cleavage reaction, promoting the rate-limiting proton relay (7). When βE190 of TF$_1$ (equivalent to βE188 of bMF$_1$) was replaced with aspartic acid (D), the rate constant of ATP cleavage step was greatly slowed over 320-fold (22).

**SI Appendix, Fig. S3B** shows the rotation rates of bMF$_1$(βE188D) at various [ATP$^+$]. The data points were well fitted with the Michaelis–Menten curve with $V_{\text{max}}$ and $K_m$ of 1.2 rps and 1.2 μM, respectively. As expected, $V_{\text{max}}$ was largely suppressed, which was about 600 times lower than that of wild-type bMF$_1$. The ATP binding constant, $k_{\text{on}}^{\text{ATP}}$, was estimated to be $3.2 \times 10^6$ M$^{-1}$s$^{-1}$, which was 10 times lower than $k_{\text{on}}^{\text{ATP}}$ of the wild type.

The stepping behaviors of bMF$_1$(βE188D) were well consistent with those found in the ATP$^+$-driven rotation of the wild-type bMF$_1$. At high [ATP$^+$], we again observed long and short dwells during 120° steps (Fig. 4 A–C). The dwell position histogram based on CP analysis showed that short dwell was located at 48° between long dwells (Fig. 4 B, Lower and SI Appendix, Fig. S5B). Histograms of durations of long and short dwells showed a single exponential decay function with time constants of 220 to 280 ms and 6 to 12 ms, respectively (Fig. 4 F and SI Appendix, Fig. S8). At low [ATP$^+$], bMF$_1$(βE188D) showed that long dwell occurred at +84° from binding dwell (Fig. 4 D–F). Dwell-time histograms determined the time constants of long dwell to be 180 to 230 ms and binding dwell to be 251 ms for 1 μM ATP, 75 ms for 3 μM ATP, and 27 ms for 10 μM ATP. The rate constant of ATP binding was determined to be $4.3 \times 10^6$ M$^{-1}$s$^{-1}$, which is mostly consistent with that estimated from the abovementioned Michaelis–Menten analysis. Thus, rotation assay of ATP$^+$ and bMF$_1$(βE188D) confirmed that bMF$_1$ makes long dwell at +80° from binding dwell and short dwell at +50° to 60° from long dwell, that is, +10° to 20° from binding dwell.

**Stall by AMP-PNP.** The rotation assays with ATP$^+$ or bMF$_1$(βE188D) showed that long dwell occurred at +80° from binding angle. In addition, the coincidence of long-dwell angle with the angle of the ADP-inhibited state suggested that long dwell represented catalytic dwell where F$_1$ executes the cleavage reaction. To further confirm these findings, we investigated the pause positions of bMF$_1$ by blocking rotation with AMP-PNP, a nonhydrolyzable ATP analog to stall rotation at the angle of cleavage.

The rotation of the γ subunit of bMF$_1$ was visualized with magnetic beads as a rotation probe because AMP-PNP–inhibited bMF$_1$ could be reactivatable with magnetic tweezers, which...
allowed repeated experiments for the molecules. Rotation was observed at 100 nM ATP, where clear pauses at the ATP binding dwell were observed at 3 positions (Fig. 5A). Recording rate was 30 fps. After confirming the 3 pauses as binding dwell in a turn, the solution of 100 nM ATP plus 500 nM AMP-PNP was gently introduced into a flow cell to minimize interference of rotation by buffer flow. Typically, molecules stopped rotation within 3 min after buffer exchange. Once lapsed into AMP-PNP inhibition, bMF1 molecules never resumed rotation unless forcibly rotated over +360° with magnetic tweezers. It should be noted that ADP inhibition is rarely observed at 100 nM ATP (SI Appendix, Fig. S9A). The mean duration time of ADP inhibition observed at 2 nM ATP was ~25 s, which is evidently shorter than the duration time of AMP-PNP inhibition, which is over 4 min (SI Appendix, Fig. S9B). In this experiment, after confirming that the pause lasted over 4 min, we defined the pause as an AMP-PNP stall. The pause angle of AMP-PNP inhibition was evidently different from the angles for binding dwell. The angular distance of the AMP-PNP stall from the nearest binding dwell on the left side was +76° (Fig. 5B), which is consistent with the position of long dwell. Thus, the angular position of ATP hydrolysis was confirmed at +80° from the angle of binding dwell, which is the same position as long dwell.

**Angle-Dependent Modulation of Reaction Rates and Equilibriums.** Identification of rotation angles for ATP binding and ATP hydrolysis is fundamental to elucidate how F1 interconverts chemical energy of ATP hydrolysis into mechanical rotation. One of the most distinctive features that discriminate F1 from other molecular motor proteins is that F1 largely modulates chemical equilibria of catalytic reaction steps depending on rotary angle to achieve ATP synthesis upon reversed rotation (31, 32). In a previous study (39), we established a "stall-and-release" experiment with magnetic tweezers, which allows for measurements of the rate constant and equilibrium constant of ATP binding or ATP hydrolysis as a function of rotary angle. This experiment revealed quantitative aspects of the "binding-change mechanism." It was shown that TF1 exponentially increased affinity to ATP by 235-fold upon rotation by 60°, while it increased the equilibrium constant of ATP hydrolysis/synthesis only by 3-fold. From these results, the contribution of affinity change for torque generation was estimated to be 21 to 54 pN nm, while that of hydrolysis was only 4 to 17 pN nm (2).

To investigate the angle-dependent modulation of affinity change and hydrolysis equilibrium of bMF1, we conducted a "stall-and-release" experiment. The experimental procedure was as follows. Rotation was observed under conditions where the target reaction, ATP binding or hydrolysis, was the rate-limiting step in the overall rotation rate. For ATP-binding, [ATP] was lowered to 100 nM, in which the mean waiting time for ATP binding was 0.9 s, while other reaction steps should be completed within 1 ms. For ATP-hydrolysis measurement, the intrinsic time constant for ATP hydrolysis, less than 0.5 ms, is too short for manipulation. Therefore, we observed rotation of bMF1 (βE188D) in the presence of ATPγS, in which the catalytic dwell was prolonged to 4.0 s. When F1 paused to wait for the target reaction to occur, we stalled the rotation of bMF1 at the targeted angle with magnetic tweezers. After the set time period lapsed, bMF1 was released from the magnetic tweezers. Principally, bMF1 showed 2 behaviors: returning to the original waiting angle or stepping to the next waiting angle. Returning indicated that F1 had not executed the waiting reaction during the stall. We refer to that case as "OFF." Stepping indicated that F1 had already executed the reaction and torque had been generated on the magnetic beads. That is referred to as an "ON" case. By determining the probability of ON cases (P_{ON}), we measured the probability of reaction as a function of rotary angle.

**Fig. 5**. Pause positions stalled by AMP-PNP. (A) Experimental procedure. After observing binding dwell at 100 nM [ATP] (Left), 500 nM AMP-PNP and 100 nM ATP was introduced into the reaction mixture (Right). Blue data points represent the positions when rotation was blocked with AMP-PNP. (B) The angular distance (Δθ) of AMP-PNP inhibition from the nearest binding dwell (pink) (n = 27). Values are mean ± SD.
The temperatures of ATP binding and ATP hydrolysis in bMF1. (A) Time course of \( P_{\text{on}} \) at 100 nM ATP in bMF1(WT). Each data point was obtained from 21 to 67 trials using 5 to 13 molecules. (B) Time course of \( P_{\text{on}} \) at 1 mM ATP/\( P_{\text{S}} \) in bMF1(βE188D). Each data point was obtained from 28 to 64 trials using 3 to 8 molecules. (C–E) Angle dependence of ATP binding and ATP hydrolysis. 0° represents the initial position of ATP binding or hydrolysis before manipulation. The directions for "forward" and "reverse" reactions are defined as that for ATP hydrolysis (counterclockwise) and ATP synthesis (clockwise), respectively. Pink and blue represent ATP binding/ATP release and ATP hydrolysis/ATP synthesis. (kATP)

The temperature dependence of the rotation rate of bMF1 is essentially the same as those of TF1. The \( Q_{10} \) factor of the rotation rate of bMF1 (1.3 to 1.9) is almost the same as the \( Q_{10} \) factors of ATP hydrolysis (1.9) and Pi release (1.6) of TF1 (47). As a result, the activation free energy, calculated from \( \Delta G = \Delta H - T \Delta S \), was 56 to 72 kJ/mol, also well consistent with the values obtained previously for TF1 (47), and EF1 (48, 49). These results indicate that the transition states of the catalytic dwell of bMF1 are the same as those for TF1.

**Discussion**

**Catalytic Event in Long Dwell and Binding Dwell.** This study investigated the fundamental features of bMF1 rotation under 3 conditions: in the absence of ATP, in the presence of ATPγS, and, by using a mutant F1, bMF1(βE188D). The latter 2 conditions were employed to slow down the cleavage step for resolving rotation into clear substeps. In all conditions, we observed long and short dwells under substrate-saturated conditions and long dwell and binding dwell in the region below \( K_m \). Although the short dwell in the ATP-driven rotation was too short for analysis, the rotation assays with ATPγS or with bMF1(βE188D) showed coincident angle assignments for short and long dwells: When the angular position for binding dwell was defined as 0°, long dwell was at +80° and short dwell at +10 to 20°. ATPγS and the βE188D mutation are known to specifically slow down the hydrolysis step, although several studies suggested that the release step of thiophosphate or phosphate was also kinetically bottlenecked reaction determining the overall reaction rate from 17 °C to 35.5 °C. This suggests that the reaction scheme found at room temperature is valid at a wide range of temperatures.
solved down to some degree (7, 22). Therefore, it is reasonable to identify long pause as the hydrolysis waiting state. The combination of ATPγS and the βγE188D mutation supports this assertion: In the presence of ATPγS, bMF1(βγE188D) showed long dwell of 4.0 s. This is extended from the original long pause (0.5 ms) by a factor of 8,000, which is very close to the expected value (12,000) from multiplication of individual extension factors: 30 by ATPγS and 400 by βγE188D. The agreement well supports that both ATPγS and βγE188D slowed down the same reaction step at which the bound ATP is hydrolyzed. Another support is the observation of ADP-inhibited pause: The rotating bMF1 transiently stopped at the angle of long dwell for a few seconds. To date, all F1s characterized in the rotation assay have shown the transient pause by ADP inhibition at catalytic angle. This was confirmed in the inhibition experiment with AMP-PNP, which halts F1 rotation at the hydrolysis step. Unlike the case of ADP inhibition, once bMF1 stopped rotation it never resumed rotation even after forcible rotation with magnetic tweezers. The pause angle of bMF1 inhibited by AMP-PNP was found at the angle for long dwell. All experimental results show that long dwell represents the hydrolysis waiting state of bMF1. This is consistent with the findings of TF1 and hMF1, in which the catalytic pause is at +80 to 90° from the binding angle.

The remaining uncertainty is the short pause. Considering the high similarity of amino acid sequences between bMF1 and hMF1, it was expected that the short pause would correspond to the intermitting pause found in the rotation assay of hMF1 (referred to as the “1s dwell” or the “Pdwell” in a previous study (29))). Actually, the short pause of bMF1 and the 1s dwell of hMF1 were both found between ATP-binding angle and catalytic angle. This suggests that short pause of bMF1 is also the pause for phosphate release, as considered for the 1s dwell of hMF1. However, the addition of phosphate or thiophosphate at concentrations from 10 μM to 100 mM in the assay solution did not show a clear impact on the rotation behavior of bMF1 in current conditions (SI Appendix, Fig. S10). Thus, the chemical state of short pause of bMF1 remains to be elucidated.

**Rotation Scheme of bMF1.** Fig. 1C shows the proposed reaction scheme of bMF1. Considering the findings of the present study as well as the crystal structures of bMF1 in which 2 of 3 catalytic sites are always occupied with bound nucleotides, ATP hydrolysis is assigned to be at 200° when the angle for ATP binding on the catalytic site is defined as 0°. This is also along the reaction schemes of TF1 and hMF1 (Fig. 1A and B). One prominent difference in the reaction scheme among species is the number of substeps: TF1 shows 2 distinctive substeps. A clear difference between mammalian F1s is the position of the pause between binding dwell and catalytic dwell: +65° from binding dwell in hMF1 and +10 to 20° in bMF1. As a result, substep size is also different: 65°, 25°, and 30° substeps for hMF1, while bMF1 makes 10 to 20°, 50 to 60°, and 40° substeps. Note while experimentally this had not been determined, careful data analysis based on a data-mining method found that TF1 also makes small substeps during catalytic dwell (27).

There are also some distinctive differences in the kinetics of rotation between bMF1 and hMF1, although overall kinetic parameters such as $V_{	ext{max}}$ and $K_m$ are mostly the same. In the rotation of bMF1, the duration time of catalytic dwell was always longer than short dwell, although the reverse is true in the rotation of hMF1. The 1s dwell was longer than catalytic dwell in hMF1. The source of these differences found in substeps and kinetics between bMF1 and hMF1 is unknown. The amino acid sequences are overall quite similar between bMF1 and hMF1. The α and β subunits share mostly identical sequences (99%), whereas the γ subunit shows relatively lower homology, 93% (16). Therefore, the most probable explanation is that the structural difference of the γ subunit causes differences in the kinetics and stepping behavior.

**Correlation with Crystal Structures.** The present work reveals that bMF1 has at least 3 distinctive conformational states: binding dwell state, short dwell state, and catalytic dwell state. Obviously, the catalytic dwell state principally corresponds to the bMF1 crystal structures. It has been suggested that the current resolved crystal structures of bMF1 represent the catalytically active state, as supported by several studies, including the crystal structure with transient state analog, beryllium fluoride, and theoretical analysis (13). However, variations among crystal structures have been reported to date. They differ in bound nucleotides, inhibitors, inorganic ligands, and conformational states of subunits. Particularly, there is variety in rotational orientation of the γ subunit in crystal structures. Although it should depend on methods for structural alignment and analysis, the maximum difference in the angular orientation of the γ subunit has been reported to be over 30°. Particularly, when bγ binds to thio phosphate the γ subunit is positioned at −30° (16) from that found in crystal structures with mitochondrial inhibitor proteins (50, 51). This feature is almost consistent with the rotation scheme proposed for hMF1 where phosphate release triggers rotation from the phosphate-releasing state at +65° from the binding site to the hydrolysis waiting state at +90°. However, the actual bMF1 does not show a dwell at around −30° from the catalytic dwell position. Although it is possible to assign short pause as the phosphate-releasing state, the angular distance between short and catalytic dwells, 50 to 60°, is too large. Thus, it is still unclear which crystal structure(s) exactly correspond to the catalytic dwell state.

**Angle Dependence of Catalytic Power of bMF1.** One of the most remarkable features of F1-ATPase that differentiates it from other molecular motors is that F1-ATPase can reverse the catalytic reaction to synthesize ATP from ADP and phosphate when the rotation is reversed. This means that all catalytic reaction steps should be modulated with the rotation angle. To investigate this characteristic feature in detail, we developed a “stall-and-release” experiment. In the previous study on TF1, we found that the affinity to ATP was exponentially increased with forward rotation, while the equilibrium constant of hydrolysis was only slightly increased (39). The estimated free energy release upon
from 2 cover glasses (18 x 18 mm² and 24 x 32 mm²; Matsunami Glass) using double-sided tape as a spacer. The bMF₁, of ~1 nM in the basal buffer was infused into the flow cell and incubated for 5 to 10 min. After that, unbound bMF₁ molecules were washed out with the basal buffer containing 10 mM BSA. Then, 40-nm gold nanoparticles or magnetic beads were infused and incubated for 5 to 10 min. Unbound beads were washed out with the basal buffer containing indicated concentrations of substrate. The basal buffer for bMF₁ assay contained 50 mM Hepes-KOH (pH 7.5), 50 mM KCl, and various concentrations of MgCl₂. When ATP was used, an ATP-regenerating system (1 mM phosphoenolpyruvate and 50 μg/mL pyruvate kinase) was added to the reaction mixture.

In the rotation assays with the 40-nm gold colloid, the rotating colloid particle that was attached to the γ subunit of bMF₁ was observed using a dark-field microscope with a 60x objective lens at the recording rate of 125 to 45,000 fps (FASTCAM-1024PCI; Photron). The localization precision was 1 to 2 nm with signal-to-noise ratio ranging from 60 to 100 (20). For observation of the rotary catalytic subunits, a phase-contrast microscope (Olympus) with a 100x objective lens at 30 fps (FC300M; Takeux) was used. The rotation assay was performed at 23 ± 2 °C, room temperature, except for 17 °C, 30 °C, and 35.5 °C in the temperature-dependence experiment (Fig. 7). For assay at 17 °C, the microscope room was cooled with an equipped air conditioner, and temperature was monitored with a thermometer attached on the flow cell on the microscopic stage. For assay at 30 °C or 35.5 °C, an objective lens heater (MATS-7SR, Tokai Hit Corp.) was used. Actual temperature of the sample was monitored the same as in the assay at 17 °C.

Data Analysis. To suppress the effect of focus drifts on analysis, we have corrected data using nonspecific binding molecules on a glass surface before analysis. To avoid undesired fluctuation, the median filter (±2 frames) was applied to the time course in Figs. 3A and 4A. To estimate time constants, the histograms of the dwell time were fitted by a single exponential decay function. Pause angles were determined by fitting the angle distribution with a Gaussian function in Figs. 3E, 4E, and 5B. For the visualization and estimation of short pause between long pauses, CP analysis was applied to the time traces shown in Figs. 3A and 4A, as described in SI Appendix, Supplementary Information Text.

Manipulation with Magnetic Tweezers. Magnetic tweezers were equipped on the microscope stage by stage actuators, controlled by stage actuators with a Gaussian function in Figs. 3A and 6A. and 6B, kinetic parameters were determined using single-exponential function according to the reversible reaction scheme. The SD of Power in Fig. 6A and B is given as $\sqrt{\text{Power (100 -- Power)}/N}$, where N is the number of trials for each experiment.

Data Availability. Data are available in the Dryad Digital Repository (https://doi.org/10.5061/dryad.pg4f4qrjk).

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