An Alternative Reaction Pathway of F_1 -ATPase Suggested by Rotation without 80°/40° Substeps of a Sluggish Mutant at Low ATP

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ABSTRACT F_1 -ATPase, a water-soluble portion of F_2F_1 -ATP synthase, is a rotary motor driven by ATP hydrolysis. The central y-subunit rotates in the $\alpha_3\beta_3$ cylinder by repeating four stages of rotation: ATP-binding dwell, rapid 80° substep rotation, catalytic dwell, and rapid 40° substep rotation. In the catalytic dwell, at least two catalytic reactions occur—cleavage of the enzyme-bound ATP and presumably release of the hydrolyzed product(s) from the enzyme—but we found that a slow ATP cleavage mutant of F_1 -ATPase from thermophilic Bacillus PS3 rotates at low ATP concentration without substeps and the catalytic dwell. Analysis indicates that in this alternative reaction pathway the two catalytic reactions occur during the preceding long ATP-binding dwell. Thus, F_1 -ATPase can operate through (at least) two competing reaction pathways, not necessarily through a simple consecutive reaction.

INTRODUCTION

 F_1 is a molecular rotary motor driven by ATP hydrolysis (1). In vivo, F_1 is connected with its partner F_0 , a membraneembedded portion, and constitutes a holoenzyme F_0F_1 -ATP synthase. F_0F_1 -ATP synthase synthesizes ATP from ADP and inorganic phosphate (Pi) using proton-motive force across membranes (2,3). The F₁ portion $(\alpha_3\beta_3\gamma\delta\varepsilon)$ has catalytic sites responsible for ATP synthesis reaction, and the F_0 portion (ab₂c₁₀) has a path for proton flow. A downhill proton flow through F_0 drives the rotation of F_1 in a direction reverse to hydrolysis, resulting in production of ATP. When isolated, F_1 only catalyzes the ATP hydrolysis reaction and is often called F_1 -ATPase. The crystal structure of F_1 shows that the rod-shaped γ -subunit is surrounded by a cylinder made of three α - and three β -subunits arranged alternately (4). The catalytic sites are located mainly on β -subunits, and some residues from adjacent α -subunits also contribute.

Rotation of the γ -subunit against the $\alpha_3\beta_3$ cylinder by ATP hydrolysis was first proposed by Boyer (5), was supported by results of biochemical (6) and spectroscopic experiments (7), and was finally visualized by single-molecule observation under an optical microscope (8). The mechanism of rotary catalysis by F_1 is known to have four principal features: First, the γ -subunit makes a 120° step when F₁ hydrolyzes one ATP molecule (9) . Second, this 120° step is further divided into four stages: ATP-binding dwell, 80° substep rotation, catalytic dwell, and 40° substep rotation (10). F_1 rotates by repeating these four stages. Third, binding

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of ATP occurs during the ATP-binding dwell, and at least two catalytic events occur during the catalytic dwell (10). One of these has been shown to be cleavage of ATP at a catalytic site (11), and the other is currently assumed to be release of hydrolyzed product(s) from F_1 (10). Fourth, in a single 120° rotation, ATP binding and ATP cleavage occur in two different β -subunits (12). Three β -subunits probably cooperate with each other to complete one 120° rotation; each of three β -subunits carries out one of the elementary reactions in ATP hydrolysis including ATP binding, ATP cleavage, and release of the hydrolyzed product(s).

In the above scenario, three β -subunits carry out their individual roles, faithfully obeying the order of sequential catalytic events in a single reaction pathway. Indeed, a single mechanism seems able to account for rotation of F_1 at any [ATP], because time-averaged rotation rates of F_1 show simple Michaelis-Menten dependence on [ATP] from the nanomolar to the millimolar range (10,13) and the duration of the catalytic dwell is constant $(\sim 2 \text{ ms})$ at [ATP] from the micromolar to the millimolar range (10). However, in the course of studying the mutant F_1 from thermophilic *Bacillus* PS3 that can hydrolyze ATP only at a very low rate, we noticed that this was not always the case. This mutant F_1 has a very long catalytic dwell during which ATP cleavage (time constant (τ_0) = 320 ms) and another event (τ_0 = 20 ms), presumably the product release, occur (11). When we observed rotations of this mutant at low [ATP] where the ATP-binding dwell was more than several seconds, substeps and catalytic dwell disappeared, and the mutant F_1 rotation included a repetition of the ATP-binding dwell and a rapid 120° rotation. ATP cleavage and product release can probably occur in the period when F_1 is still in the ATP-binding dwell. Therefore, against the previous assumption that the order of sequential catalytic events of F_1 is always

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maintained in a single reaction pathway, F_1 can take at least two catalytic pathways in ATP hydrolysis.

MATERIALS AND METHODS

Materials

ATP, phosphoenolpyruvate, and bovine serum albumin (BSA) were purchased from Sigma (St. Louis, MO). Biotin-PEAC₅-maleimide and Maleimide-C₃-NTA were from Dojindo (Kumamoto, Japan). Pyruvate kinase, streptavidin, and gold colloid (diameter $= 80$ nm) were from Roche Diagnostics, Wako (Osaka, Japan) and British Biocell International (Cardif, UK), respectively. We used $\alpha_3\beta_3\gamma$ subcomplex derived from thermophilic *Bacillus* strain PS3. Here, we call this subcomplex F_1 . The mutant F_1 $(F_1(\beta-E190D))$ containing a replacement of Glu-190 with Asp in the β -subunit was purified as reported previously (11). Streptavidin-coated gold colloid was prepared as previously described (14).

Rotation assay

For the rotation assay, two cysteines of the γ -subunit were biotinylated (11). A flow chamber was made of a Ni²⁺-NTA-coated coverglass (16 \times 16 mm²; Matsunami Glass, Osaka, Japan) and a slide glass $(76 \times 26 \text{ mm}^2)$ with a spacer of 50- μ m thickness. Biotinylated F₁(β -E190D) (0.5–2 pM) in solution A (50 mM MOPS-KOH, 50 mM KCl) containing 10 mg/ml BSA was infused into the flow chamber, and after 10 min, $F_1(\beta$ -E190D) unbound to the glass surface was washed away by 100 μ l of solution A. Then 2 pM streptavidin-conjugated gold colloid in solution A containing 10 mg/ml BSA was infused. After a 10-min incubation, unbound gold colloids were washed away by 100 μ l of solution A containing 2 mM MgCl₂, ATP-regenerating system (50 μ g/ml pyruvate kinase and 2.5 mM phosphoenolpyruvate), and the indicated amount of ATP. The rotating gold colloid was observed by dark-field microscopy (IX-70; Olympus, Tokyo, Japan) with a condenser (numerical aperture, $1.2-1.4$) and $\times 100$ objective lens (numerical aperture, 0.5–1.35). The images of the rotating gold colloid were recorded directly to the hard disc of a computer as an eight-bit Audio Video Interleave file with a fast-framing camera (Hi-Dcam II; NAC Image Technology, Tokyo, Japan) at 500–1000 frames per second. Custom software (created by R. Yasuda, Duke University) was used for analyses of the bead movements and dwells.

RESULTS

Rotation at saturating [ATP]

 $F_1(\beta$ -E190D) hydrolyzes ATP slowly because of its sluggish ATP cleavage. It exhibits a characteristic rotation where the duration of the catalytic dwell is 140 times longer than that of the wild-type (11). Using this mutant, we can therefore observe the catalytic dwell more easily at the recording rate of 500–1000 frames per second. At 2 mM ATP, a V_{max} condition where ATP binding should take place in 150 μ s (= $(k_{on} \times 2 \text{ mM})^{-1}$), we observed the rotation with discrete 120° steps pausing at the position of the catalytic dwell. (Some mutant F_1 molecules showed the unusually long pauses $(>100 \text{ ms})$ at the angle of the ATP-binding dwell. Since the average duration of these long pauses varied from molecule to molecule, we considered this pause not to be an intrinsic property of F_1 and did not use these molecules for analysis.) The histogram of the dwelling time corresponded to a single exponential curve (Fig. 1 A). We attribute the time

FIGURE 1 Catalytic dwell of an $F_1(\beta - E190D)$ molecule at 2 mM (A) and 500 nM (B) ATP. In the first bin, the number of catalytic dwells \leq 3 ms is indicated by gray. Inset in B shows the ATP-binding dwell of the same molecule as in B. Red lines show a fit with $y = C \times \exp(-t/\tau_0)$. C and τ_0 are a proportional constant and a time constant, respectively. (C) Typical time courses of the stepwise rotation of an $F_1(\beta - E190D)$ molecule at 500 nM ATP at 1000 frames per second. Horizontal lines are 120° apart. The ATP-binding dwells are shown in gray and the catalytic dwells in orange. The 120 rotations without catalytic dwells are indicated by arrowheads.

constant obtained, 305 ± 11 ms (mean \pm SE), mainly to the slow cleavage of the bound ATP in $F_1(\beta$ -E190D) (11). To simplify, we set the bin width of the histogram to 80 ms in this case, and the other time constant of 20 ms, which we have presumed to represent the product release, was not resolved because of the 80-ms bin size (11). At [ATP]s above 20 μ M, the histograms showed a similar single exponential decay (data not shown).

Rotation at [ATP] below K_m

As [ATP]s decreased under K_m (1.4 μ M), however, the histograms started to deviate from a single exponential decay. At 500 nM ATP, for example, the first bin $(< 80 \text{ ms})$ contained many more events than expected from a single exponential decay that were fitted well with the data after the second bin (Fig. 1 B). A closer inspection of the rotation trajectory revealed that 120° step rotations without catalytic dwell occurred at significant frequency (Fig. 1 C, indicated by arrowheads). Out of 317 120° rotations, 51 showed no obvious (\leq 3 ms) catalytic dwells (gray portion of the *bar* in Fig. 1 B). This result is in sharp contrast with the rotation at 2 mM ATP, where only 4 out of 324 catalytic dwells were $<$ 3 ms. Therefore, rotation of F₁(β -E190D) appears to proceed through two parallel reaction pathways, one with and one without catalytic dwell. We named the former the conventional pathway and the latter an alternative pathway. The time constant of the dwell-time histogram in Fig. 2 B is 338 ± 20 ms, a value close to that of rotation at 2 mM ATP, and corresponds to the time constant for cleavage of ATP of the conventional pathway.

Unlike those of the catalytic dwell, histograms of the ATP-binding dwell at low [ATP]s were fitted well by a single exponential curve (Fig. $1 B$, *inset*) and were not affected by the duration of the preceding catalytic dwell (see below). The rate constant k_{on} for ATP binding was determined to be $(3.3 \pm 0.3) \times 10^6$ M⁻¹ s⁻¹ (data for seven molecules). This value is one order of magnitude smaller than that of the wild-type and is consistent with the previous estimates (11). This result suggests that the ATP-binding dwell is terminated by the binding of ATP as reported previously in both of the two pathways (10).

Dependence on the ATP-binding dwell

The alternative pathway was more frequently observed as [ATP] decreased. As [ATP] decreased, ATP came to the catalytic site less frequently, and the ATP-binding dwell became longer. We therefore assume that duration of the ATPbinding dwells determines the population of $F_1(\beta$ -E190D) that takes the alternative pathway in the following 120° rotation. To determine whether this was in fact the case, we analyzed the correlation between the duration of the ATPbinding dwell and occurrence frequency of the alternative pathway (Fig. 2). We classified the ATP-binding dwells (2311 dwells, 7 molecules, 500 nM–2 μ M ATP) into 10 groups according to their durations and made histograms of the catalytic dwells that followed the individual ATPbinding dwells. The catalytic dwells longer than 5 s (6 out of 2311 dwells) were neglected because they were probably attributable to the ADP-inhibited state, in which F_1 fails to release ADP, lapsing into pauses of rotation over several tens of seconds (15). When the ATP-binding dwell was short, \leq 50 ms (Fig. 2 A), the histogram showed a single exponential decay ($\tau = 337 \pm 19$ ms) as observed at 2 mM ATP. As the ATP-binding dwell became longer, the first bin of the histogram became larger than the values extrapolated from a single exponential (Fig. 2, A–F). When the ATP-binding dwell was over 3 s, most catalytic dwells (36 out of 42) were included in the first bin (Fig. $2 F$). We defined catalytic dwells \leq 3 ms (gray portion of the first bin) as belonging to the alternative pathway and plotted their proportion relative to the total counts of catalytic dwells as a function of the ATP-binding dwell duration (Fig. 2 G). As shown, the proportion of the alternative pathway increases as the duration of the preceding ATP-binding dwell becomes longer. This increase can be simulated with a single exponential curve

FIGURE 2 Correlation between the catalytic dwell and the ATP-binding dwell. Durations of the ATP-binding dwells are (A) , 0–0.05 s; (B) , 0.2–0.3 s; (C), 0.3–0.5 s; (D), 0.5–1 s; (E), 1–2 s; and (F), >3 s. In the first bin, the number of catalytic dwells \leq 3 ms is indicated in gray. (G). The proportion of the alternative pathway as a function of the duration of the ATP-binding dwell. Sample numbers for the points are, left to right, 295, 235, 368, 246, 358, 418, 268, 63, 32, and 10. The red line shows a fit with a single exponential, $y = 100 \times (1 - \exp(-t/\tau_0))$. We assumed that occurrence of the conventional pathway is 100% at ATP-binding dwell $= 0$ s.

with a time constant of 3.0 \pm 0.3 s. The enzyme therefore tends to choose the alternative pathway rather than the conventional pathway when the enzyme must wait for ATP for longer than 3 s.

DISCUSSION

Can wild-type F_1 also take the alternative pathway?

Our results clearly show that the mutant $F_1(\beta$ -E190D) can rotate according to either of two reaction pathways. In the newly discovered alternative pathway of the mutant $F_1(\beta$ -E190D), rotation does not pause at the 80° position but proceeds with a single 120° step rotation. The rotation with the alternative pathway is currently observed only for the sluggish $F_1(B-E190D)$ mutant at low [ATP]. The immediate question is therefore whether this alternative pathway is allowed only for the sluggish mutant or is a general feature for the wild-type F_1 . In addition, the apparent contradiction between our results and those of two previous reports on rotation of the wild-type F_1 should be explained (10,13). Sakaki et al. recently reported that time-averaged rotation rates of F_1 show simple Michaelis-Menten dependence on [ATP] from the nanomolar to the millimolar range, suggesting that a single mechanism governs rotation over this [ATP] range (13). However, at low [ATP], the binding of ATP is rate-limiting in turnover of catalysis both for conventional and alternative pathways, and the interim between 120° rotations is governed by the ATP-binding dwell with little difference between the two pathways. Therefore apparent simple Michaelis-Menten kinetics does not exclude

the possibility of the alternative pathway at low [ATP]. Yasuda et al. showed that the duration of the catalytic dwell is not affected by the duration of the ATP-binding dwell and is constant at [ATP] from 2 μ M to 6 mM (10). In our case, [ATP] was below μ M, and k_{on} for ATP of the mutant was ~1 order of magnitude smaller than that of the wild-type. As a consequence, the ATP-binding dwell we observed, for example, 6.1×10^2 ms at 500 nM ATP, was longer by 40fold than that of the wild-type at 2 μ M ATP (17 ms). Therefore, almost all rotations of the wild-type should proceed through the conventional pathway in the previous experiments at 2 μ M ATP. Detection of the alternative pathway for the wild-type F_1 at lower [ATP], for example 20 nM, might be practically difficult because the ATP-binding dwell would be 1000-fold longer than the catalytic dwell and the two dwells would be difficult to distinguish by ordinary measurement methods. Altogether, we think it likely that the alternative pathway exists in the wild-type.

What is the alternative pathway?

The alternative pathway is characterized by the rotation without the catalytic dwell at the 80° position (Fig. 1). It is unlikely that the two events, ATP cleavage and product release, occur at the 80° position too rapidly for the time resolution of our measurement because this behavior would not explain why the alternative pathway is observed only when the preceding ATP-binding dwell is long. More likely is that ATP hydrolysis can take place during the ATPbinding dwell even though very rarely. The time constant for the transition from the conventional pathway to the

FIGURE 3 Two models of the reaction pathways of the ATP-hydrolysis reaction in $F_1(\beta$ -E190D). $F_1(\beta$ -E190D) are represented as three β -subunits (blue) and a y-subunit (orange oval with an arrow). The rate constants obtained from this work and the previous works (11) are shown. The rate constant 50 s⁻¹ is assumed to be unchanged between the conventional and alternative pathway. The models represent the pathways of $F_1(\beta$ -E190D) but the wild-type F_1 probably has similar alternative pathways as well (see the text). (Model 1) ADP is released during the rotary motion upon ATP-binding. (Upper path) The conventional pathway. (Lower path) The alternative pathway. (Model 2) ADP release and ATP binding occur at a different stage in the reaction pathway. (Upper path) The conventional pathway. The 40° substep rotation is coupled with release of the products. (Lower path) The alternative pathway. The products are released without accompanying rotation. Note that occupancy of the catalytic sites by ATP (or ADP) during catalysis in this alternative pathway is one or two (bisite catalysis).

alternative pathway is 3.0 s (Fig. 2 G). This value corresponds to the time constant for the cleavage of ATP when the mutant is at the ATP-binding dwell. We assume that product release also occurs during the ATP-binding dwell because the 20-ms event, which is presumably the product release, also disappeared from the catalytic dwell in the alternative pathway.

As discussed in Sakaki et al. (13), we currently assume that the rotation in the conventional pathway proceeds as follows (Fig. 3, model 1, *upper path*). ATP binds to the β_1 subunit of F_1 in the ATP-binding dwell and drives the 80 $^{\circ}$ substep rotation of the γ -subunit, which accompanies ADPrelease from the β_3 subunit. During the catalytic dwell, cleavage of the previously bound ATP to ADP and Pi occurs at the β_2 subunit. After the cleavage, the β_3 subunit releases Pi, an action coupled with the 40° substep rotation. In the alternative pathway (Fig. 3, model 1, lower path), hydrolysis of ATP at the β_2 subunit and release of Pi from the β_3 subunit occur during the ATP-binding dwell, and ADP is released from the β_3 subunit during the 120 \degree rotation that is triggered by the ATP-binding to the β_1 subunit. Another conventional pathway in which release of ADP is coupled with the 40° substep rotation is also possible (Fig. 3, model 2, upper path). In this pathway, three catalytic sites are occupied by nucleotides during the catalytic dwell (trisite catalysis). The sequence of events in the alternative pathway corresponding to this conventional pathway is as follows (Fig. 3, model 2, lower path): ATP cleavage at the β_2 subunit, product release from the β_3 subunit, ATP-binding to the β_1 subunit, and 120° rotation. In this pathway, one or two catalytic sites are occupied by ATP or ADP during catalysis (bisite catalysis). If model 2 is really functional, F_1 works with the trisite mode under usual conditions but does so with the bisite mode when [ATP] is extremely low.

F_1 reaction is not a simple consecutive reaction

This work revealed that F_1 can operate through (at least) two pathways. We have long assumed the faithfulness of F_1 in keeping the order of sequential catalytic events tracking a single reaction pathway. Actually, we stated in the previous work (16) ''During a long period in which one slow β -subunit waits for ATP, other β -subunits are also just waiting without letting the next catalytic event start." But the 80 $^{\circ}$ dwell with short lifetime $(\sim 2 \text{ ms})$ was not detectable with the experimental setup in Ariga et al. (16) and this statement should be corrected. Our finding implies that catalytic events of all three catalytic sites are coordinated not as a single, linear consecutive reaction but as diverging-converging reaction pathways. Recently, two pathways were revealed in myosin V as well (17). Two heads of myosin V work cooperatively, and existence of two pathways may not be rare among motor proteins in which the catalytic subunits work cooperatively. Two reaction pathways are competing with each other and slowing down one of them increases the weight of other pathways, which may originally make small contributions to the overall flux. The weight of each reaction pathway may also change with substrate concentrations. Characterization of multiple reaction pathways and their physiological relevance need to be further clarified.

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