

Activation of pausing F_1 motor by external force

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A rotary motor F_1 , a catalytic part of ATP synthase, makes a 120° step rotation driven by hydrolysis of one ATP, which consists of 80° and 40° substeps initiated by ATP binding and probably by ADP and/or P_i dissociation, respectively. During active rotations, F_1 spontaneously fails in ADP release and pauses after a 80° substep, which is called the ADP-inhibited form. In the present work, we found that, when pushed $>+40^\circ$ with magnetic tweezers, the pausing F_1 resumes its active rotation after releasing inhibitory ADP. The rate constant of the mechanical activation exponentially increased with the pushed angle, implying that F_1 weakens the affinity of its catalytic site for ADP as the angle goes forward. This finding explains not only its unidirectional nature of rotation, but also its physiological function in ATP synthesis; it would readily bind ADP from solution when rotated backward by an F_o motor in the ATP synthase. Furthermore, the mechanical work for the forced rotation was efficiently converted into work for expelling ADP from the catalytic site, supporting the tight coupling between the rotation and catalytic event.

ADP inhibition | ATP synthase | F_1 -ATPase | magnetic tweezers | single-molecule observation

The molecular rotary motor F_1 -ATPase generates rotary torque driven by chemical energy liberated from ATP hydrolysis (1–4). In cells, F_1 constitutes the ATP synthase binding with its partner motor, F_o , which is embedded in a biomembrane and rotates its rotor ring driven by a proton flux down to the electrochemical potential across the membrane. F_1 and F_o are connected to each other through their rotor and stator parts, whereas their rotary directions are opposite. Thus, they push each other intramolecularly. Under physiological conditions where the proton-motive force is larger than the free energy obtained from ATP hydrolysis, F_o can generate a larger torque than F_1 and enforces F_1 to rotate in the reverse direction. Consequently, F_1 reverses its catalytic reaction to synthesize ATP from P_i and ADP. In contrast, when the free energy of ATP hydrolysis is larger, F_1 hydrolyzes ATP, inversely rotating F_o to pump protons in the opposite direction. Thus, F_1 is a reversible molecular machine for converting energy between the chemical potential of ATP and mechanical work of rotation. It has been proved that mechanically reversing F_1 molecules leads ATP synthesis (5). We recently revealed that the coupling efficiency of the mechanical ATP synthesis is very high with single-molecule manipulation and microfabrication techniques (6).

F_1 has a subunit composition of $\alpha_3\beta_3\gamma\epsilon$. Its minimum complex as ATPase is the $\alpha_3\beta_3\gamma$ subcomplex, which is referred to hereafter as F_1 . Walker's group (7–9) solved F_1 's crystal structures to reveal that three α - and β -subunits are alternately arranged in a hexagonal cylinder in which the rotor subunit, γ , penetrates at the center. The catalytic site is located mainly on the β -subunit. Three β -subunits hydrolyze ATP in a cooperative manner to make a unidirectional rotation of the γ -subunit (10, 11). The kinetic model for its ATP hydrolysis coupled with the rotation was originally proposed by Boyer (12). The γ -rotation in F_1 hydrolyzing ATP was strongly supported by biochemical cross-linking (13) and a spectroscopic technique (14), and finally visualized at a single-molecule level (15). The direct observation of the rotation revealed F_1 makes a 120° stepping rotation upon each ATP hydrolysis (16). High-speed imaging of the F_1 rotation with no viscous load resolved the 120°

step further, into 80° and 40° substeps (17). The kinetic analysis in these studies has made allowance for the idea that the 80° substep arises from ATP binding, and the 40° substep is triggered by ADP and/or P_i dissociation. Recently, it was demonstrated that ATP hydrolysis occurs between the 80° and 40° substeps by using a mutant F_1 that hydrolyzes ATP at an extremely slow rate (18).

In our previous work, we observed that the F_1 lapses into an inactive state during rotation and showed it to correspond to the ADP-inhibited form (19), which had been characterized by many biochemical studies (20, 21). The ADP-inhibited form is the state in which the F_1 fails to dissociate ADP to stop catalysis and rotation. The ADP-inhibited F_1 pauses after the 80° substep (19) in agreement with the idea that the 40° substep arises from the release of the product(s). The inactivation is a reversible process; the F_1 in pause spontaneously becomes active to rotate again. Recent single-molecule works showed that some other enzymes are also in equilibrium between catalytic active and inactive or less active forms (22, 23). If we identify the structural feature of an inactive enzyme and mechanically manipulate its conformation back to the catalytic active state, in principle, it is possible to activate the enzyme. However, there have been few experimental attempts to do so, one of which is a report by the Bustamante group (23) showing that the probability of the transcriptional pause of RNA polymerase is affected by external force directions. In the ATP synthase, F_o turns the F_1 's catalysis direction by rotating the γ -subunit; therefore, F_1 is expected to modulate kinetic parameters upon the forced rotation of the γ -subunit. In the present study, we found that the ADP-inhibited F_1 can be mechanically activated by pushing the rotor γ -subunit in the forward direction. An elaborate analysis of this phenomenon revealed that the mechanical stress on the γ -subunit opens the catalytic site to dissociate inhibitory ADP, and that the rate constant of ADP dissociation increases with the angle of the γ -subunit. These findings have important implications for the model of the F_1 's unidirectional rotation and its ATP synthesis in the ATP synthase.

Methods

Materials. A mutant $\alpha_3\beta_3\gamma$ subcomplex (α -C193S, β -His-10 at N terminus, γ -S107C/I210C) from a thermophilic *Bacillus* PS3 (referred to as F_1 unless otherwise specified) was expressed and purified as described (24). ATP, ADP, and phosphoenolpyruvate were purchased from Sigma. Pyruvate kinase and streptavidin-coated magnetic beads (0.73 μ m) were purchased from Roche Diagnostics and Seradyn (Indianapolis), respectively.

Magnetic Tweezers. A schematic diagram of magnetic tweezers is shown in Fig. 1*a*. The tweezers comprise four electromagnets, each constructed of a soft iron core (10 \times 10 \times 40 mm) and a copper wire with 100 turns around the core. Each pair of tweezers was electrically connected in a series and separated by an interval of 15 mm. The two electromagnet pairs were crossed at each center space and positioned 10 mm above the microscope stage. The microscope

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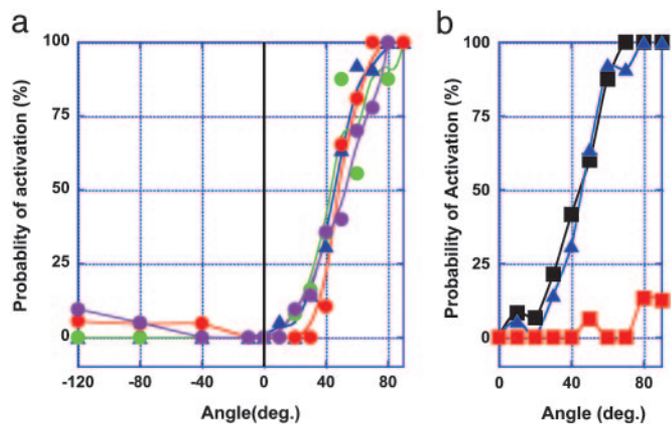


Fig. 2. Angle dependency of the probability of mechanical activation. (a) Probabilities of the mechanical activation were determined in the presence of 0.2 μM ATP (blue); 0.2 μM ATP, 30 mM P_i (green); 0.2 μM ATP, 1 mM ADP (red); and 0.2 μM ATP, 30 mM P_i, 1 mM ADP (purple). Each data point was determined from 17 observations on average. (b) Probabilities were determined in the presence of 0.2 μM ATP (blue), 2 mM ATP (black), and 33 μM ATP, 20 mM ADP (red). Each data point was determined from 15 observations on average.

creased to reach up to nearly 100% at +70°. However, when pulled up to -120°, the F₁ motor mostly returned to 0° (Fig. 1c). Thus, both the rotary direction and angle amplitude were important for the mechanical activation. The activation probability was also determined in the presence of 1 mM ADP and/or 30 mM P_i. No major difference in the presence of ADP was found, except for the lower probability at 30° and 40°. Similar probabilities were also obtained at 2 mM ATP (Fig. 2b). These findings suggest that the mechanical activation is ascribable to the release of the inhibitory ADP, as

opposed to the binding of the substrate/product from the solution. In previous biochemical experiments (24, 25), the ADP-inhibited F₁ was activated by ADP depletion, for example, through gel filtration. To verify this finding, the mechanical activation was examined at 20 mM ADP, where it was completely suppressed, thus confirming the above explanation (Fig. 2c). At 20 mM ADP is much higher than the previously reported K_d values for ADP (26, 27), probably because the affinity of ADP decreases with the rotary angle, as shown below. Therefore, much higher ADP was needed for the suppression of the mechanical activation >40°, whereas the γ-subunit in the biochemical experiments would mostly be 0°, maintaining a higher affinity to ADP. Indeed, 2 mM ADP was enough to inhibit the mechanical activation <30°, indicating that the $K_{d(ADP)}$ for <30° is <2 mM. The $K_{d(ADP)}$ between 40° and 60° was expected to lie between 2 and 20 mM.

Time Course of Mechanical Activation. We examined how the mechanical activation depends on the stalling time. A typical time course of the experiment is shown in Fig. 3a, where the ADP-inhibited F₁ was stalled at 40° for 5, 5, and 15 s. Magnetic tweezers were turned off after each stalling to observe the activation. In this example, the F₁ was activated after 15 s of stalling. From these experiments, we obtained saturation curves for the activation probability, which exponentially increased with the stalling time (Fig. 3b and c). These were well fitted with a single exponential equation that assumes a first-order reaction, ADP-inhibited form → activated form. Each fitting gave a rate constant for the mechanical activation, which exponentially increased with the rotary angle. Interestingly, the activation at 0° was too slow for this fitting, whereas the spontaneous activation occurred at the same rate as the mechanical activation at 20°. These results mean that the affinity of the inhibited F₁ to ADP at 0° is very high, and it decreases with the angle to accelerate ADP release. This finding is consistent with the observation that higher ADP was needed to suppress the

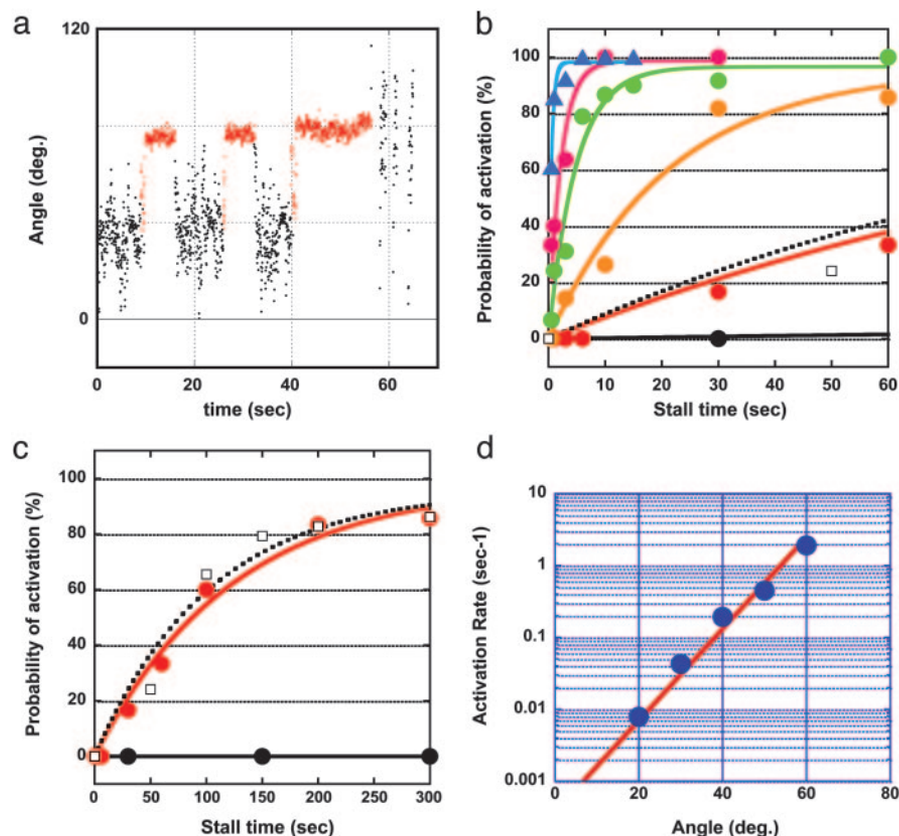


Fig. 3. Rate constant of the mechanical activation. (a) Time course of the mechanical activation with different stalling times. The ADP-inhibited F₁ was stalled and released at +40° after 5 s twice and after 15 s. (b) The probabilities of the mechanical activation by stalling at 0° (black), 20° (red), 30° (orange), 40° (green), 50° (purple), and 60° (blue) were plotted against the stalling time. Each data point was determined from 12 observations on average. (c) Shown is an expanded graph for probabilities of the spontaneous activation (□). (d) Rate constants (blue circles) were determined by fitting the curves in b and c with single exponential equations. The slope of linear fitted line (red) for these rates in semilogarithmic scale gives the angle dependency of the activation energy as $-1.3 k_B T/10^\circ$.

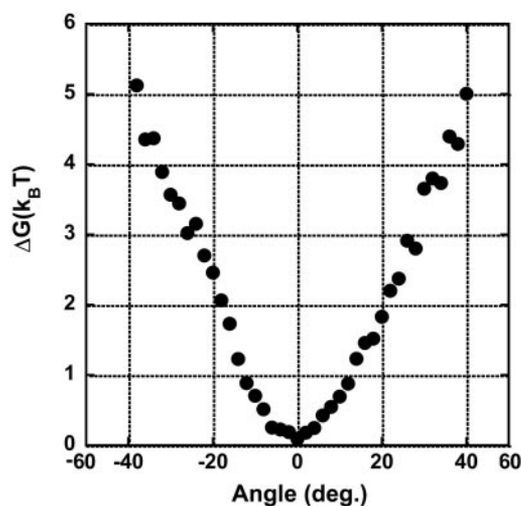


Fig. 4. The rotary potential of the ADP-inhibited form F_1 . The probability density of the angle of the magnetic beads from the six molecules was transformed into potential according to the Boltzmann distribution law.

activation in forward angles. Each fitting gave a saturation value of $\approx 100\%$, suggesting its irreversibility. To verify this idea, the following experiments were performed. The ADP-inhibited F_1 was stalled at $>60^\circ$ for 6 s to activate it, and it was rotated back to 0° to stall for 6 s to inactivate it, should it occur, and then it was released from the magnetic tweezers. In most of the cases (93%), released F_1 showed its active rotation (data not shown). Thus, it was confirmed that F_1 is not reversibly inactivated again in this condition. Once F_1 is activated by stripping inhibitory ADP, the reactivation would need either ADP binding from the solution or production of ADP in catalysis.

Activation Energy of Mechanical Activation. The rate constants of the mechanical activation, k_{act} , determined from Fig. 3 *b* and *c*, are plotted against stall angles (Fig. 3*d*). It was clearly shown that the rate increases exponentially with the angle. This finding means that the activation energy of the mechanical activation, ΔG_a , given in the Arrhenius equation ($\ln k_{act} = \ln A - \Delta G_a/k_B T$), decreases with the angle. The activation energy change was given as $-1.3 k_B T/10^\circ$ from the slope in Fig. 3*d*. This finding strongly suggests that, upon the mechanical energy input through the γ -subunit, the energy state of the inhibitory ADP binding shifts to a higher state, so that the activation energy becomes relatively lower. To determine the amount of energy transferred to the F_1 by the magnetic tweezers, the work for pushing the F_1 in the ADP-inhibited form was estimated. The Brownian motion of the magnetic beads attached to the F_1 in the ADP-inhibited form was observed for >30 s without applying magnetic field, and the probability density of angle was determined to calculate the potential shape according to the Boltzmann distribution law (Fig. 4). The determined potential $>+20^\circ$ had a linear slope, corresponding to the linear decrement of the activation energy of the mechanical activation. The energy increment of $1.5 k_B T$ per 10° coincides well with the activation energy decrement of $1.3 k_B T$ per 10° , suggesting that 85% of the mechanical energy input was transmitted to the catalytic site to weaken the binding energy of ADP. This finding is consistent with the tight coupling model between the catalytic reaction and the γ -rotation in the F_1 motor. The residual energy would remain stored as elastic energy for the other elements in our experimental system (for example, the γ -subunit itself, the $\alpha_3\beta_3$ ring, His tag, or streptavidin).

Angle Deviation Between the γ -Subunit and the Magnetic Beads. When the experimental system has highly flexible components such as the γ -subunit, the $\alpha_3\beta_3$ ring, His tags, or streptavidin, the rotary

angle of the magnetic beads could deviate by a large amount from the γ -angle. Even though the angle deviation was not definitely determined in the present work, some of our experimental data suggested that the deviation is probably not so large. First, the mechanical activation was observed when the magnetic beads were rotated $>20^\circ$, suggesting that the γ -subunit certainly rotates against the $\alpha_3\beta_3$ ring, which implies that the deviation is $<20^\circ$. The rotary potential of the ADP-inhibited F_1 is also suggestive on this point. The determined potential comprises a loose spring-like potential within $\pm 10^\circ$ and linearly increasing potentials for over $\pm 10^\circ$, which have slopes (= torque) of ≈ 35 pNnm for both sides. These values are consistent with the reported torque of ≈ 40 pNnm (16, 24, 28), implying that the potential $>20^\circ$ certainly reflects the interaction between the $\alpha_3\beta_3$ ring and the γ -subunit. Thus, the angle deviation would be at most 20° . One of the most likely candidates for elastic components is the γ -subunit, because its protruding part from the $\alpha_3\beta_3$ ring was twisted by 11° in the crystal structure of the mitochondrial F_1 modified by dicyclohexyl-carbodiimide (8, 9). Further studies on its elasticity remain to be done. It should be noted that the determined torque from the rotary potential is equal to the external torque by the magnetic tweezers because these two torque are equilibrated.

Backward Activation. Although the mechanical activation was not observed when the γ -subunit was pulled backward, up to -120° , preliminary experiments showed that the F_1 can be activated when pulled over by -160° . However, the results have been quite irregular so far. It is only certain that the activation probability was the highest in the presence of both ADP and P_i . A similar finding was reported by Vinogradov and colleagues (29, 30), that is, a submitochondrial particle containing the Fo F_1 -ATPase showed a higher ATP-hydrolysis activity after transient charging of the membrane potential. The most plausible explanation is that when F_1 is rotated in the ATP-synthetic direction in the presence of the ADP and P_i , the F_1 follows the ATP synthesis pathway to be efficiently activated.

Discussion

Mechanical Activation of Enzymes. The catalytically active F_1 spontaneously fails to dissociate ADP with some probability, and its rotation pauses at an angle after an 80° substep. This catalytically inactive state is called the ADP-inhibited form. In this study, we showed the mechanical activation of the F_1 motor in the ADP-inhibited form by pushing it with magnetic tweezers. When the ADP-inhibited F_1 was pushed $>+40^\circ$ with the magnetic tweezers, the F_1 regained its catalytic activity to restart the rotation, whereas it was not readily activated when pulled back. This mechanical activation was completely suppressed by ADP, but not by ATP and/or P_i . These findings showed that the mechanical activation is triggered by the dissociation of the inhibitory ADP. Also, in biochemical experiments, ADP removal is necessary for the activation of the ADP-inhibited F_1 , for example, using gel filtration. The model for the mechanical activation in the catalysis and rotation scheme is shown in Fig. 5*a*. The scheme starts from the ATP-waiting state, A, where the β -subunit with an empty catalytic site waits for the ATP to bind, and the two are bound with ATP and ADP, respectively. This state approximately corresponds to the crystal structure of MF $_1$ solved in 1994, except for P_i binding in the model. Upon ATP binding to the empty β , the F_1 makes an 80° substep, which triggers the following catalytic reactions on the other β s, ATP hydrolysis ($B \rightarrow C$) and ADP and/or P_i release ($C \rightarrow A'$). When ADP releases from the F_1 , the γ -subunit rotates 40° to reset the F_1 in a new ATP-waiting state, A'. With some probability, the F_1 in the state C releases phosphate and isomerizes to the ADP-inhibited form, state D. This scheme accords with a recent biochemical study using a mutant F_1 with an inserted tryptophan residue near the phosphate binding region, which showed that F_1 releases phosphate and then slowly isomerizes to the ADP-inhibited

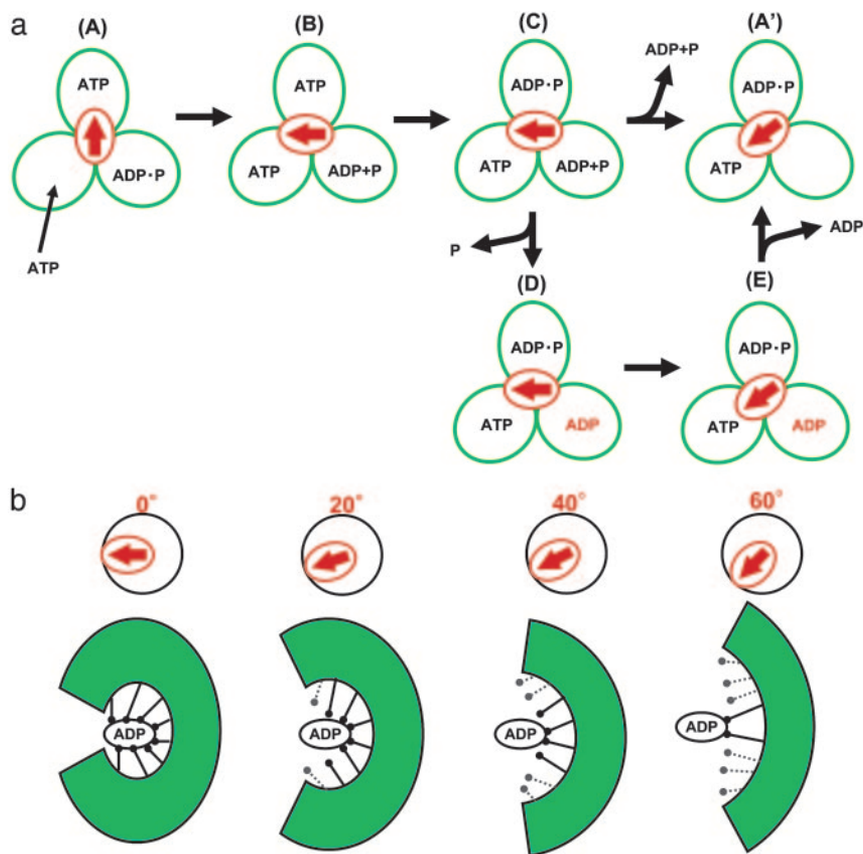


Fig. 5. The proposed model for conformational change around the catalytic site. (a) Reaction scheme in the catalysis and rotation model. Three green circles represent the catalytic state, and the red arrow indicates the angle of the γ -subunit. States D and E represent the ADP-inhibited form and forward-rotated form, respectively. (b) Schematic model of the conformational change around the catalytic site with the inhibitory ADP coupled with mechanical rotation. Upon forced rotation, the catalytic site progressively opens its conformation-breaking interactions with ADP.

form (31). Even though the biochemical experiments were conducted in an unisite condition where the number of catalytic sites occupied with a nucleotide was one at most, it is assumed that a similar isomerization occurs in F_1 that operates in a multisite catalysis mode (32, 33) (here, the trisite model is used). When the γ -subunit of the ADP-inhibited F_1 (D) is rotated $>40^\circ$ with the magnetic tweezers, the mechanical stress on the F_1 changes the conformation of the β -subunit carrying the inhibitory ADP, enforcing the F_1 to strip the inhibitory ADP molecule. After releasing it, the F_1 returns to a new ATP-waiting state, A', on the catalytic active pathway. This scheme explains our experimental data well; however, it should be noted that this scheme is provisional because the angle at which the active F_1 releases ADP and P_i has not been conclusively identified yet. Additionally, the mostly same model can be proposed standing on bisite mode in which site occupancy alternates between one and two.

Angle Dependency of ADP Affinity. The most important result obtained in the present study is that the angle of the γ -subunit determines the rate constant of the ADP dissociation from the β -subunit. The time course of the mechanical activation gave the rate constant of the ADP dissociation that exponentially increases with angles of the γ -subunit in the forward direction. This finding explains well the directed rotation coupled with ADP release. Similar findings on the kinesin motor have been reported by Uemura and Ishiwata (34). They determined the ratio of the ADP-bound kinesin to the nucleotide-free from the measurement of the unbinding force of the kinesin-microtubule complex to reveal that the direction of the external force determines the affinity of kinesin with ADP. Thus, the modulation of the ADP affinity in

response to the external force would be a common feature for molecular motors to give rise to unidirectional motion.

This finding is very suggestive for the ATP synthesis reaction model, in which we assume similar angle dependency of the ADP dissociation rate for active F_1 . For ATP hydrolysis, the F_1 motor has to dissociate ADP in the solution, whereas it also has to efficiently bind ADP from the aqueous solution for ATP synthesis. If the ADP binding/dissociation would occur merely at a specific angle, the F_1 would not manage these two reactions. However, when active F_1 has a similar angle dependency of ADP affinity, F_1 would readily catch ADP from the solution when the F_0 motor pulls F_1 back. Thus, the angle dependency of ADP release accords with a property needed for F_1 to operate its physiological function of ATP synthesis.

Spontaneous Activation. The angle dependency of the activation implies that the ADP remains difficult to strip off until the γ -subunit is moved. It was actually observed that when the F_1 was clamped at 0° , the angle for the ADP-inhibited state, ADP did not dissociate even after 300 s. In contrast, when the F_1 motor was allowed to make the rotary Brownian motion without an external force, the F_1 was spontaneously activated at the same rate as the mechanical activation at 20° . This finding means that when the thermal energy occasionally pushes the γ -subunit $>20^\circ$, the inhibited F_1 dissociates ADP to regain its activity. In this sense, the spontaneous activation is also a mechanical activation caused by the thermal energy. It would be also interesting to examine ADP release in the unisite catalysis from the same view point. In the unisite catalysis, ADP release is known to be very slow, whereas the ATP hydrolysis step and the following phosphate release are fast (31, 35). The F_1 is also in a kind of ADP-inhibited form, although it has a sole bound

nucleotide. Therefore, similar mechanism could occur in the ADP release; F_1 waits for the thermal fluctuation to push forward to release the tightly bound ADP.

Structural Model of Mechanical Activation. As shown in Fig. 3*d*, the rate constant of the mechanical activation increased upon mechanical rotation of the γ -subunit, implying that the interaction between the bound ADP and the catalytic site on the β -subunit was weakened to accelerate ADP release. The analysis of the rate constants according to the Arrhenius equation suggested that the binding energy decreases in proportion to the rotary angle, by $1.3 k_B T$ per 10° . We discuss the structural features of the ADP-inhibited form and models to explain the mechanical activation below.

Structural studies on bovine mitochondrial F_1 (MF_1) by the Walker group (7–9) have revealed that the β -subunit can essentially use three types of conformation. The β -subunit with tightly bound ADP or ATP analog lifts the C-terminal domain up to the nucleotide binding domain enclosing the nucleotide in the crevice between the two domains. In contrast, the β -subunit without a nucleotide pulls down the C-terminal domain opening the catalytic site. Recently, an intermediate conformation was found, which weakly binds ADP in half-opened conformation. Thus, the F_1 closes the catalytic site upon ATP binding and opens its conformation after ADP release via the half-closed conformation. This conformational change is a typical induced-fit. The C-terminal domain has direct contact with the γ -subunit, thereby, this domain movement is thought to mechanically link the catalytic reactions and the γ -rotation. What are the structural features specific for the ADP-inhibited form? Considering that ADP binds tightly to the inhibited F_1 , the ADP-inhibited β would be in the closed conformation (36). Further detailed structural features distinct from a catalytically active F_1 remain to be revealed. However, an interesting point is given by the recent crystal structures of MF_1 with dicyclohexyl carbodiimide (9) and complexes of ADP and beryllium fluoride (37). The catalytic site that corresponds to the ATP-binding site in other crystal structures does not bind ATP but does bind ADP. Because of the absence of the γ -phosphate, the side chain of the $\alpha R373$, a catalytically crucial residue, is in a largely different conformation from other structures and forms a hydrogen bond to the 2'OH of the ribose of the ADP. Considering that isomerization to ADP inhibition occurs after phosphate release, it is possible that this structural rearrangement is a structural base of the ADP-inhibited form.

Kinetic analysis of the mechanical activation suggests that the mechanical forward rotation of the γ -subunit decreases the binding

energy of ADP and the catalytic site. Therefore, we propose a simple structural model in which the mechanical forward rotation enforces the closed- β -subunit with the inhibited ADP to transform into the opened conformation, breaking the interactions between ADP and the catalytic site (Fig. 5*b*). When the interactions are unzipped sequentially with the forced rotation, it explains the linear drop of the ADP binding energy well. As an alternative model, a conformational rearrangement of a few amino acid residues, for example the $\alpha R373$, which is decisive for the ADP inhibition, can be a trigger of mechanical activation. The analysis of the rotary Brownian motion showed that 85% of the energy given for forced rotation was transmitted to the catalytic site, supporting the tight mechanical linkage between the γ and the catalytic site. The structural model shown in Fig. 5*b* coincides well with progressive ATP-binding models proposed by Oster and Wang (38) and Cross (39) independently. These models explain well the efficient energy conversion and the reversibility of F_1 -ATPase. In Oster and Wang's "binding-zipper" model, it is proposed that hydrogen bonds of ATP and the catalytic site are sequentially formed to drive the γ -rotation to explain the F_1 's constant torque against the rotary angle (38, 40). In the near future, the angle dependency of the affinity of F_1 with ATP also will be elucidated by using this method.

Conclusion

The F_1 catalyzes ATP synthesis when the F_0 enforces it to rotate in the reverse direction. It means that catalytic parameters of F_1 can be modulated by rotating the γ -subunit. In this work, we successfully activated the ADP-inhibited F_1 molecules by rotating the γ -subunit with magnetic tweezers. From the analysis of this phenomenon, it was revealed that F_1 decreases its affinity to the inhibitory ADP with the γ -rotation by using external energy with high efficiency. These properties of F_1 well explain its unidirectional nature of rotation and physiological function: synthesis of ATP. The molecular mechanism regarding the manner in which F_1 tunes elementary chemical steps (ATP binding, ATP hydrolysis/synthesis, and P_i release) in response to an external torque would be challenged by this method.

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- Boyer, P. D. (1997) *Annu. Rev. Biochem.* **66**, 717–749.
- Yoshida, M., Muneyuki, E., & Hisabori, T. (2001) *Nat. Rev. Mol. Cell Biol.* **2**, 669–677.
- Kinosita, K., Jr., Yasuda, R., & Noji, H. (2000) *Essays Biochem.* **35**, 3–18.
- Senior, A. E., Nadanaciva, S., & Weber, J. (2002) *Biochim. Biophys. Acta* **1553**, 188–211.
- Itoh, H., Takahashi, A., Adachi, K., Noji, H., Yasuda, R., Yoshida, M., & Kinosita, K. (2004) *Nature* **427**, 465–468.
- Rondelez, Y., Tresselt, G., Nakashima, T., Kato-Yanada, Y., Fujita, H., Takeuchi, S., & Noji, H. (2005) *Nature* **433**, 773–777.
- Abrahams, J. P., Leslie, A. G., Lutter, R., & Walker, J. E. (1994) *Nature* **370**, 621–628.
- Menz, R. L., Walker, J. E., & Leslie, A. G. (2001) *Cell* **106**, 331–341.
- Gibbons, C., Montgomery, M. G., Leslie, A. G., & Walker, J. E. (2000) *Nat. Struct. Biol.* **7**, 1055–1061.
- Ariga, T., Msaiké, T., Noji, H., & Yoshida, M. (2002) *J. Biol. Chem.* **277**, 24870–24874.
- Nishizaka, T., Oiwa, K., Noji, H., Kimura, S., Muneyuki, E., Yoshida, M., & Kinosita, K., Jr. (2004) *Nat. Struct. Mol. Biol.* **11**, 142–148.
- Boyer, P. D. (1993) *Biochim. Biophys. Acta* **1140**, 215–250.
- Duncan, T. M., Bulygin, V. V., Zhou, Y., Hutcheon, M. L., & Cross, R. L. (1995) *Proc. Natl. Acad. Sci. USA* **92**, 10964–10968.
- Sabbert, D., Engelbrecht, S., & Junge, W. (1996) *Nature* **381**, 623–625.
- Noji, H., Yasuda, R., Yoshida, M., & Kinosita, K., Jr. (1997) *Nature* **386**, 299–302.
- Yasuda, R., Noji, H., Kinosita, K., Jr., & Yoshida, M. (1998) *Cell* **93**, 1117–1124.
- Yasuda, R., Noji, H., Yoshida, M., Kinosita, K., Jr., & Itoh, H. (2001) *Nature* **410**, 898–904.
- Shimabukuro, K., Yasuda, R., Muneyuki, E., Hara, K. Y., Kinosita, K., Jr., & Yoshida, M. (2003) *Proc. Natl. Acad. Sci. USA* **100**, 14731–14736.
- Hirono-Hara, Y., Noji, H., Nishiura, M., Muneyuki, E., Hara, K. Y., Yasuda, R., Kinosita, K., Jr., & Yoshida, M. (2001) *Proc. Natl. Acad. Sci. USA* **98**, 13649–13654.
- Jault, J. M., Matsui, T., Jault, F. M., Kaibara, C., Muneyuki, E., Yoshida, M., Kagawa, Y., & Allison, W. S. (1995) *Biochemistry* **34**, 16412–16418.
- Matsui, T., Muneyuki, E., Honda, M., Allison, W. S., Dou, C., & Yoshida, M. (1997) *J. Biol. Chem.* **272**, 8215–8221.
- Lu, H. P., Xun, L., & Xie, X. S. (1998) *Science* **282**, 1877–1882.
- Forde, N. R., Izhaky, D., Woodcock, G. R., Wuite, G. J., & Bustamante, C. (2002) *Proc. Natl. Acad. Sci. USA* **99**, 11682–11687.
- Noji, H., Bald, D., Yasuda, R., Itoh, H., Yoshida, M., & Kinosita, K., Jr. (2001) *J. Biol. Chem.* **276**, 25480–25486.
- Vasilyeva, E. A., Minkov, I. B., Fitin, A. F., & Vinogradov, A. D. (1982) *Biochem. J.* **202**, 9–14.
- Weber, J., Wilke-Mounts, S., Lee, R. S., Grell, E., & Senior, A. E. (1993) *J. Biol. Chem.* **268**, 20126–20133.
- Dou, C., Fortes, P. A., & Allison, W. S. (1998) *Biochemistry* **37**, 16757–16764.
- Noji, H., Hasler, K., Junge, W., Kinosita, K., Jr., Yoshida, M., & Engelbrecht, S. (1999) *Biochem. Biophys. Res. Commun.* **260**, 597–599.
- Galkin, M. A., & Vinogradov, A. D. (1999) *FEBS Lett.* **448**, 123–126.
- Zharova, T. V., & Vinogradov, A. D. (2004) *J. Biol. Chem.* **279**, 12319–12324.
- Msaiké, T., Muneyuki, E., Noji, H., Kinosita, K., Jr., & Yoshida, M. (2002) *J. Biol. Chem.* **277**, 21643–21649.
- Muneyuki, E., Makino, M., Kamata, H., Kagawa, Y., Yoshida, M., & Hirata, H. (1993) *Biochim. Biophys. Acta* **1144**, 62–68.
- Msaiké, T., Mitome, N., Noji, H., Muneyuki, E., Yasuda, R., Kinosita, K., & Yoshida, M. (2000) *J. Exp. Biol.* **203**, 1–8.
- Uemura, S., & Ishiwata, S. (2003) *Nat. Struct. Biol.* **10**, 308–311.
- Grubmeyer, C., Cross, R. L., & Penefsky, H. S. (1982) *J. Biol. Chem.* **257**, 12092–12100.
- Tsunoda, S. P., Muneyuki, E., Amano, T., Yoshida, M., & Noji, H. (1999) *J. Biol. Chem.* **274**, 5701–5706.
- Kagawa, R., Montgomery, M. G., Braig, K., Leslie, A. G., & Walker, J. E. (2004) *EMBO J.* **23**, 2734–2744.
- Oster, G., & Wang, H. (2000) *Biochim. Biophys. Acta* **1458**, 482–510.
- Cross, R. L. (2000) *Biochim. Biophys. Acta* **1458**, 270–275.
- Antes, I., Chandler, D., Wang, H., & Oster, G. (2003) *Biophys. J.* **85**, 695–706.