Robust rotary catalysis of F$_1$-ATPase

Rikiya Watanabe$^{1,2,4}$, Yuki Matsukage$^{2,4}$, Ayako Yukawa$^1$, Kazuhito V Tabata$^{1,2}$ & Hiroyuki Noji$^1$

$^1$Department of Applied Chemistry, The University of Tokyo, Bunkyo-ku, Tokyo 113-8656, JAPAN
$^2$PRESTO, JST, Bunkyo-ku, Tokyo 113-8656, JAPAN
$^3$Institute of Scientific and Industrial Research, Osaka University, Ibaraki, Osaka 567-0047, JAPAN
$^4$These authors contributed equally to this work.

To whom correspondence should be addressed: Hiroyuki Noji, Department of Applied Chemistry, School of Engineering, The University of Tokyo, Tel. +81-3-5841-7252; E-mail: hnoji@appchem.t.u-tokyo.ac.jp

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**Background:** Three catalytically charged-residues of F$_1$-ATPase; p-loop-lysine, general-base, and arginine-finger, are thought to be indispensable for catalysis.

**Results:** Alanine-substituted mutants of the catalytic residues of F$_1$-ATPase drove rotations.

**Conclusion:** The catalytic residues contribute to efficient catalysis but are not indispensable to chemo-mechanical energy coupling of F$_1$-ATPase.

**Significance:** The chemo-mechanical coupling mechanism of F$_1$-ATPase is far more robust than previously thought.

**ABSTRACT**

F$_1$-ATPase (F$_1$) is the rotary motor protein fueled by ATP-hydrolysis. Previous studies have suggested that three charged residues are indispensable for catalysis of F$_1$: the p-loop lysine in the phosphate-binding-loop, GXXXGK(T/S), a glutamic acid that activates water molecules for nucleophilic attack on the $\gamma$ phosphate of ATP (general base), and an arginine directly contacting the $\gamma$ phosphate (arginine-finger). These residues are well conserved among p-loop NTPases. In the present study, we investigated the role of these charged residues in catalysis and torque generation by analyzing alanine-substituted mutants in the single-molecule rotation assay. Surprisingly, all mutants continuously drove rotary motion, even though the rotational velocity was at least 100,000 times slower than...
that of wild-type. Thus, although these charged residues contribute to highly efficient catalysis, they are not indispensable to chemo-mechanical energy coupling, and the rotary catalysis mechanism of $F_1$ is far more robust than previously thought.

Molecular machines, fueled by nucleotide triphosphate (NTP), play pivotal roles in a wide range of cellular activities, such as gene regulation, organelle transport, membrane transport, protein unfolding, signal transduction and energy synthesis. Many of these exert mechanical force on their substrates or associated proteins through conformational changes that occur during NTP hydrolysis (1,2). Although there are some variations in structure, NTP-driven molecular machines share well-conserved structural features, especially around the nucleotide binding domain (1). The most highly conserved primary structure among the NTPases is the phosphate-binding loop (p-loop) comprised of the amino acid sequence GXXXXGKT/S (where X varies). The p-loop sequence motif was first reported by Walker et al. (3), and is therefore referred to as a Walker A motif. Additionally, the lysine is the most crucial residue for NTP hydrolysis among the p-loop residues (2).

Some subfamilies of NTP-driven molecular machines share other catalytically crucial residues that are also electrically charged. Catalytic glutamic acid is found in the nucleotide-binding pocket of many subfamilies of NTPases, such as the AAA$^+$ proteins, ABC transporters, and RecA-type proteins, although the catalytic glutamic acid of RecA-type proteins occupies a different position in the secondary structure compared with other subfamilies (4). The glutamic acid binds to NTP via a coordinated water molecule at the distal end of the $\gamma$ phosphate of bound NTP. Because this residue seems to induce an in-line attack of the water molecule to $\gamma$ phosphate and initiate the hydrolysis reaction by activating the water molecule, the carboxylate residue is termed the ‘general base’. Note that recent theoretical studies have revised the actual working mechanism of the general base in catalysis (5,6).

A catalytic arginine (Arg) residue is also widely distributed among many NTPases (2). The most-studied catalytic arginine is the ‘arginine-finger’ of the G-protein activating protein (GAP), which triggers GTP hydrolysis of the G-protein (7). Several studies have reported that the arginine-finger stabilizes the transition state of hydrolysis to enhance catalysis. Both AAA$^+$ proteins and RecA-type proteins such as RecA and $F_1$-ATPase also carry the corresponding arginine (or lysine in RecA) in the same region of the arginine finger of
GAP (8,9).

Mutation at these charged residues is fatal for NTPase catalysis, and is particularly so when substituted with either a non-charged or an oppositely charged residue, which reduces the catalytic power to undetectable levels in biochemical assays (10-12). Substitution of alanine (Ala) or another non-charged residue for the p-loop lysine (Lys), the most highly conserved of these residues, is the approach used most frequently to knock down the catalytic power of NTPases. However, an explanation as to how the electric charge of these catalytically crucial residues contributes to force generation has remained elusive due to the difficulty of investigating these effects while simultaneously retaining hydrolytic activity. In this study, we evaluated the effects of alanine mutation of the catalytically crucial charged residues on catalytic efficiencies and force generation in a highly sensitive single-molecule assay using the F$_1$-ATPase rotary motor as a model NTP-driven molecular machine.

F$_1$-ATPase (F$_1$), the water-soluble portion of the F$_0$F$_1$-ATP synthase, is a member of the RecA-type protein family and possesses all of the catalytically crucial charged residues: p-loop Lys, the general base, and the Arg-finger. When isolated from the ATP synthase complex, F$_1$ acts as a rotary motor protein, where it rotates the inner rotary subunit against the stator ring, hydrolyzing ATP (13,14). The bacterial type F$_1$ is composed of $\alpha_3\beta_3\gamma\delta\epsilon$, with the $\alpha_3\beta_3\gamma$ sub-complex representing the minimum complex necessary to function as a rotary motor. The $\alpha_3\beta_3$ subunits form the cylindrical stator ring. The $\gamma$ subunit is the rotary shaft that penetrates the center of the cylinder (8,15-17). The catalytic sites for ATP hydrolysis reside on each $\alpha$-$\beta$ interface, mainly on the $\beta$ subunit (8). Therefore, conformational changes of the $\beta$ subunit are responsible for torque generation.

The rotary torque of F$_1$ has repeatedly been reported to be 40 pN·nm for F$_1$ from thermophilic bacteria and the Bacillus PS3 that was investigated in the present study (18,19), while F$_1$ torque from Escherichia coli was reported to be 30–61 pN·nm (14,15,20).

Among motor proteins, F$_1$ is unique for its high reversibility in a chemo-mechanical coupling reaction, i.e., F$_1$ catalyzes ATP synthesis when the rotary shaft is forcibly rotated in the reverse direction (21,22). In cells, F$_1$ binds to F$_0$, the membrane-embedded part of ATP synthase, to form the whole ATP synthase complex. Under physiological conditions, F$_0$, powered by the proton-motive force across the membranes, generates larger torque than F$_1$, thereby reversing F$_1$ to induce the ATP synthesis reaction. This high reversibility means that each reaction step is tightly
coupled with a rotary motion of the γ subunit. The tight coupling feature also allows us to elucidate kinetic analysis of individual catalytic reactions from observations of the rotary motion.

To establish a basis for studying the chemo-mechanical coupling mechanism of F₁, the reaction scheme of F₁ was extensively researched. While most aspects of the scheme have been resolved, some uncertainties remained (23). Rotations reportedly occur in discrete 120° steps, each coupled to a single turnover of ATP hydrolysis (18). The 120° step is further divided into 80° and 40° sub-steps (24,25). The 80° sub-step is triggered by ATP binding and ADP release, each of which occurs on different β subunits (26). The 40° sub-step is triggered by ATP hydrolysis and release of inorganic phosphate (Pᵢ), which also occurs on different β subunits (26,27). The angular positions of the dwell before the 80° and 40° sub-steps are referred to as the ATP-binding and catalytic angles, respectively.

Establishment of the basic reaction scheme has led to the more fundamental question of determining how the locally occurring chemical reaction induces the large conformational change of the motor protein. Previous molecular genetics and biochemical studies have identified the catalytic residues of F₁ based on sequence homology and chemical modifications with ATP-analogues (28). The crystal structures of F₁ elucidated the atomic details of the catalytic reaction centers (8,29-31), providing a foundation for structure-based theoretical studies (32-34). All of these studies have identified the abovementioned charged residues as the catalytically critical residues, although some additional residues were also found to be involved in catalysis. Interestingly, when these charged residues are substituted with non-charged or oppositely charged residues, F₁ catalytic activity is reduced to undetectable levels in biochemical ATPase assays (10,11,35-37), suggesting that these residues are indispensable for catalysis. However, no data have been presented that clearly explain how the electric charges of these residues contribute to force generation of F₁. Recently, a single-molecule assay of F₁ provided quantitative analysis of very low catalytic activity (i.e., less than 0.005 s⁻¹), which was lower than the detection limit of the biochemical ATPase assay (about 0.3 s⁻¹) (35). Therefore, in the present work we re-assessed the catalytic competence of the alanine mutant of the conserved charged residues using this single-molecule rotation assay. All of these mutants exhibited unidirectional rotation, indicating unexpectedly high robustness of catalytic competency. Additionally, we analyzed the impact of the mutation on torque-generation.
EXPERIMENTAL PROCEDURES

Rotation assay - Wild-type $F_1$, and $F_1$ mutants $F_1(\alpha R364A)$, $F_1(\beta K164A)$, $F_1(\beta E190A)$ and $F_1(\beta E190Q)$ were prepared as previously reported (38). In order to visualize the rotation of $F_1$, the stator region ($\alpha_3\beta_3$) was fixed onto a glass surface, and magnetic beads (Seradyn, Indianapolis, USA) were attached to the rotor ($\gamma$) as a rotation probe, as previously reported (27). The rotating beads were observed under a phase-contrast microscope (IX-70 or IX-71, Olympus, Tokyo, Japan). The rotation assay was performed at 25°C. The images of rotary motion were recorded at 30–2,000 frames/s (FASTCAM 1024PCI-SE, Photron, Tokyo, Japan; FC300M, Takex, Kyoto, Japan). Images were stored on the HDD of a computer as AVI files and analyzed using custom-made software.

RESULTS

Rotary motion of alanine-substituted mutants: $\alpha R364A, \beta K164A, \beta E190A$ - The arginine-finger ($\alpha R364$), the p-loop lysine ($\beta K164$), and the general base ($\beta E190$) of $F_1$ from thermophilic Bacillus PS3 (TF$_1$) were substituted with alanine to produce the $F_1$ mutants, $F_1(\alpha R364A)$, $F_1(\beta K164A)$, and $F_1(\beta E190A)$ (Fig. 1a, b). $\beta E190A$ was also substituted with glutamine to produce $F_1(\beta E190Q)$. The ATPase activity of all mutants was undetectable with biochemical assays, which was consistent with the results of previous studies (10,11,35-37). The rotation assay was conducted at a saturating ATP concentration (1 mM) by using magnetic beads as rotation markers. Surprisingly, all mutants exhibited continuous rotary motions in an anticlockwise direction (Fig. 1c), although rotation was extremely slow, in some instances taking as long as 5 min to confirm. Nonetheless, the observed rotation indicated that all of the mutants retained the catalytic power of ATP hydrolysis and their chemomechanical coupling nature. The rotational rate of these mutants ranged from 0.002 to 0.1 rps (Fig. 1c). The slowest rotation was observed for $F_1(\beta E190Q)$. As seen in the time courses in Figs. 1c and 1d, all the mutants exhibited distinct pauses every 120° throughout the rotation, which limited the overall rotation rate. This observation is significant, as it indicates that one or more catalytic reaction steps are distinctively slowed.

The rotational rates of mutants were measured at various ATP concentrations ranging from 1 $\mu$M to 30 mM (Fig. 1c). The mutant $F_1(\alpha R364A)$ rarely exhibited continuous rotations at low ATP concentrations (less than 1 $\mu$M) due to severe inhibition, which did not allow for the assay to detect rotation at an ATP concentration less than
1 μM. The other mutants were observed to follow Michaelis-Menten kinetics, yielding maximum rotation rates ($V_{\text{max}}$) and Michaelis constants ($K_m$). Assuming that the chemomechanical coupling ratio of $F_1$ was 3 ATPs/turn, $3\times V_{\text{max}}$ represented the catalytic turnover rate ($k_{\text{cat}}$), and these values were calculated to be $3.3\times10^1$, $2.2\times10^2$, $1.3\times10^1$, and $6.6\times10^3$ s$^{-1}$ for $F_1(\alpha R364A)$, $F_1(\beta K164A)$, $F_1(\beta E190A)$, and $F_1(\beta E190Q)$, respectively (Table 1). The rate constants of ATP binding ($k_{\text{on}}^{\text{ATP}}$) that were determined as $3\times V_{\text{max}}/K_m$, were $9.0\times10^2$, $3.7\times10^6$, and $4.5\times10^6$ M$^{-1}$s$^{-1}$ for $F_1(\beta K164A)$, $F_1(\beta E190A)$, and $F_1(\beta E190Q)$, respectively. Compared with $k_{\text{cat}}$ and $k_{\text{on}}^{\text{ATP}}$ from the wild-type (387 s$^{-1}$ and $2.6\times10^7$ M$^{-1}$s$^{-1}$, respectively), the impact of the depletion of the conserved charge was significant in the suppression of $k_{\text{cat}}$, while the p-loop lysine mutant also resulted in remarkable suppression of $k_{\text{on}}^{\text{ATP}}$. This indicated that the p-loop lysine is involved in both the binding process and catalysis.

Analysis of stepping rotation - It is known that $F_1$ has two stable states, pausing at either the binding angle or the catalytic angle, with the latter angle calculated at $+80^\circ$ from the binding angle, in the anticlockwise direction. We therefore attempted to determine the angles at which these pauses occurred for the mutants.

For the analysis of $F_1(\beta E190A)$, we conducted a buffer exchange experiment where the ATP concentration was changed from 1 mM to 50 nM, which was comparable with the $K_m$ of this mutant (Fig. 2a). When the buffer was changed to 50 nM, the mutant exhibited sub-steps, characterized by pauses at 6 distinct positions (Fig. 2b). For statistical analysis, we compared the angular position of the pauses appearing at 50 nM ATP from the original pauses at 1mM ATP for individual rotating molecules ($\Delta\theta_2$ in Fig. 2b, c). For comparison, the relative angular positions of the original intervening pauses at two ATP concentrations ($\Delta\theta_1$ in Fig. 2b, c) were also analyzed. The results of this analysis indicated that $\Delta\theta_1$ was $-1.7\pm 6.4^\circ$, which validated this experimental approach, while $\Delta\theta_2$ was $33\pm 9.2^\circ$. These results indicated that the intervening pauses of $F_1(\beta E190A)$ occurred at the same catalytic angle as that previously reported for an aspartic acid mutant at $\beta E190$ (25). When we conducted the same buffer exchange experiment for $F_1(\beta E190Q)$, we observed that the intervening pause of $F_1(\beta E190Q)$ also occurred at the catalytic pauses (Fig. 2d, e, f). Thus, the mutation at the general base caused the distinctly long pause at the catalytic angle. We also analyzed the dwelltime of the intervening pause of $F_1(\beta E190A)$ and $F_1(\beta E190Q)$, as well as the dwelltime of the ATP-
waiting pause (Fig. 3). The histograms of the dwelltimes were fit to single exponential functions 
$y = C \cdot \exp(-k_t)$, allowing us to determine $k_{cat}$ and $k_{on^{\text{ATP}}}$ (Fig. 3a, c). $k_{on^{\text{ATP}}}$ was proportional to [ATP], but $k_{cat}$ was not (Fig. 3b, d). $k_{on^{\text{ATP}}}$ was $4.9 \times 10^6$, $1.7 \times 10^6$ M$^{-1}$s$^{-1}$, and $k_{cat}$ was $0.13$, $9.3 \times 10^{-3}$ s$^{-1}$ for $F_1(\beta E190A)$ or $F_1(\beta E190Q)$ (Fig. 5a, b). These values were consistent with the aforementioned Michaelis-Menten analysis (Fig. 1e).

The mutant $F_1(\alpha R364A)$ had a tendency to lapse into a long inhibitory state (Fig. 4a); i.e., it paused during the rotation, soon after making turns, and did not spontaneously resume rotation during extensive periods of observation. This inhibitory state was attributed to the ADP-inhibited form based on our previous observations of the lysine-substituted mutant of the arginine finger, $F_1(\alpha R364K)$, that also lapsed into a long ADP inhibition state, leading to pauses at the catalytic angle (39). Additionally, as was observed with the ADP-inhibition of the wild-type and other mutant $F_1$’s (40), $F_1(\alpha R364A)$ could also be activated by forcible rotation with magnetic tweezers that allowed us repeated observation of $F_1(\alpha R364K)$. We analyzed the relative angular positions of pauses occurring during rotation from ADP-inhibition ($\Delta \theta$ in Fig. 4b, c). $\Delta \theta$ was $0 \pm 6.0^\circ$, suggesting that $F_1(\alpha R364A)$ also paused at the catalytic angles. To confirm this, we compared the intervening pause with the ATP waiting angle by conducting a rotation assay at low ATP concentration (1 µM). Although the ADP-inhibition of $F_1(\alpha R364A)$ was more pronounced at low ATP concentrations, $F_1(\alpha R364A)$ was observed to rotate only a few steps after forcible activation with magnetic tweezers, allowing the analysis. At low [ATP], $F_1(\alpha R364A)$ exhibited sub-step rotation in each $120^\circ$ rotation. We analyzed the relative ATP waiting angle from the original intervening pause ($\Delta \theta_1$ in Fig. 4d, e). The angle measured for $\Delta \theta_1$ was $41 \pm 12^\circ$, indicating that the original intervening pause occurred at the catalytic angle. We analyzed the dwelltime of pauses at both the ATP-binding and catalytic angles. Dwelltime histograms were fit with single exponential functions, giving the rate constants of ATP binding or catalysis (Fig. 4f). Values calculated for $F_1(\alpha R364A)$ were $k_{on^{\text{ATP}}} = 2.4 \times 10^7$ M$^{-1}$s$^{-1}$, and $k_{cat} = 0.45$ s$^{-1}$ (Fig. 5a, b). It is interesting that $k_{cat}$ for this mutant was ten-fold smaller than that of the lysine-substituted mutant, $F_1(\alpha R364K)$ (39); however, $k_{on}$ was almost the same, which suggests that the positive charge of $\alpha R364$ is necessary for rate-enhancement of hydrolysis, but not for efficient binding of ATP.

We also attempted to identify the pausing position of $F_1(\beta K164A)$. This mutant showed large
rotary fluctuations during the pauses (Fig. 1d), which implied a significantly looser interaction between the stator and the rotor. When observed at low concentrations of ATP, some molecules exhibited rotation with sub-steps at 80° and 40°. Although the reduced probability of observing rotation at low [ATP] did not allow for statistical analysis, it is highly probable that the intervening pause of βK164A also occurred at the catalytic angle.

**Measurement of torque** - To investigate the roles of the electrical charge of the p-loop lysine, general base, and Arg-finger in torque-generation, we measured the torque of the Alanine mutants by using a recently developed method for torque measurement based on the fluctuation theorem (FT) (19). Rotation at 1 mM, where all of the mutants exhibited the 120° stepped rotation, was recorded at 1,000 Hz. For precise torque determination, molecules were selected that displayed symmetric 120° stepping rotation without obvious preferential pausing angles due to surface interactions. From the time trajectory, the portions of rotations were extracted; i.e. the portions of dwells were omitted. It should be noted that F₁ showed the backward movements due to thermal fluctuations during rotations while it showed a net unidirectional movement due to rotary torque. Ratios of the forward and backward movement probabilities [\(P(\Delta\theta)/P(-\Delta\theta)\)] over a set time period were then calculated. The ratio of probabilities was then plotted against \(\Delta\theta\), and the torque was determined from the slope (Fig. 6a, b, c). Results of this analysis indicated that the torque values for the mutants were evidently smaller than those for wild-type F₁; 28, 10, and 24 pN·nm for F₁(αR364A), F₁(βK164A), and F₁(βE190A), respectively (Fig. 6d). In particular, we observed that the torque of F₁(βK164A) was one-fourth that of wild-type F₁.

**Rotary potential** - We also examined the effect of mutations on the rotary potential during catalytic pauses. The probability densities of γ-orientation during catalytic pauses were measured, and the rotary potentials were determined according to Boltzmann’s Law (Fig. 7a). The determined potentials were well fitted with a harmonic function \(\Delta G = \frac{1}{2} \cdot \kappa \cdot \theta^2\), where \(\kappa\) is the torsion stiffness. The determined values of stiffness were 79, 42, 22, and 38 pN·nm for wild-type F₁, F₁(αR364A), F₁(βK164A), and F₁(βE190A), and these values correlated well with those for the rotary torque (Fig. 7b). The lower stiffness values of the mutants suggested that their rotary potentials became more gradual than those of wild-type F₁, which indicated that γ was not tightly held in the cavity of the \(\alpha_3\beta_3\) stator ring of the mutants.
especially for \( F_1(\beta K164A) \). This result concurs with the contention that \( F_1(\beta K164A) \) cannot fully induce the open-closed conformational change of the stator, and, thus, weakens the interaction between the stator and rotor.

**DISCUSSION**

*Robustness of rotary catalysis mechanism of \( F_1 \) -*

All mutants exhibited unidirectional rotation, meaning that all of the charged residues that are highly conserved among p-loop ATPases (p-loop lysine, the general base, and the arginine-finger) are dispensable for catalysis. The unidirectionality of the rotation was also comparable to that of wild-type \( F_1 \), and the frequent backward step was not observed, meaning that the nature of the chemomechanical coupling was fundamentally retained in the absence of the conserved charged residues. This observation suggests that the catalytic power of \( F_1 \) is much more robust than previously assumed.

The impact of the charge depletion was apparent in the kinetics, as the rotational rate was extremely slow (Fig. 1c, Fig. 5). In particular, the \( \beta E190 \) mutant exhibited the slowest rotation (\( V_{\text{max}}=2.2 \times 10^{-3} \)), which was approximately 10⁶-fold slower than that of wild-type \( F_1 \) (\( V_{\text{max}}=1.3 \times 10^{3} \)). Analysis on the stepping rotation of the mutants proved that the mutations were responsible for the lengthened pauses at the catalytic angle, suggesting that all of the mutants decelerated the ATP hydrolysis step. In this sense, these findings support the theoretical prediction that these residues remarkably contribute to the rate-enhancement of the ATP hydrolysis step (19,33).

Considering the conserved structural features observed among p-loop ATPases, it is likely that the conserved, charged residues are also dispensable for other ATPases. To confirm the robustness of other ATPases, a highly sensitive assay is required, such as the single-molecule rotation assay of \( F_1 \). The robustness of catalytic power has not previously been reported for other p-loop motors that have established highly sensitive single-molecule assays, and this is likely due, at least in part, to the relatively low processivity of other molecular motors. Molecular motors translocate their subunit or substrate protein/nucleic acid polymers towards the motor domain with ATPase activity. Because translocation necessarily accompanies the affinity modulation between a motor domain and a substrate, molecular motors dissociate from their substrates at some given probability. If the catalytic turnover rate is largely reduced, the probability of dissociation relatively increases, lowering the processivity of the motors. This could explain why
linear motors such as myosin, kinesin, and dynein do not exhibit clear motion when the ATPase is decelerated by a factor of over 100 (41-43).

We also evaluated a set of double mutants: F₁(αR364A, βK164A), F₁(αR364A, βE190Q), and F₁(βK164A, βE190Q). However, we did not observe any rotating particles in the rotation assay. When the roles of the conserved charged residues in rate-enhancement are additive, the estimated rotational rate approaches the order of 10⁻⁵ revolution/s, and the single-molecule rotation assay might not be sufficiently sensitive to detect such a slow rotation. It should be noted that even if the turnover rate of ATPase is on the order of 10⁻⁵ s⁻¹, it is still faster than the rate of spontaneous ATP hydrolysis in aqueous conditions.

Torque generation - Another impact of the alanine-substitution was observed in torque generation. All mutants decreased rotary torque (Fig. 6d). In particular, the rotary torque of F₁(βK164A) was significantly reduced to 10 pN·nm. This is evidently lower than that of wild-type F₁, and is the lowest among the reported torque values for the F₁ mutants. It should be emphasized that although a wide variety of mutations were introduced into F₁ in an attempt to identify the residues or structural elements that are crucial for torque generation, several mutations have been reported that evidently reduce the torque of F₁. In addition, all of the mutations critical for torque were not found at the catalytic sites, but mainly at the β–γ interfaces, such as the DELSEED loop of the β subunit (16,44). Other nucleotides such as GTP, ITP, and UTP have been previously investigated to determine the effect of base recognition on energy transduction. While GTP and ITP were found to be slow-binding substrates, GTP and ITP apparently supported torque generation at levels comparable with ATP (45). It should be noted that Martin et al proposed in a recent paper that ITP-supported rotation decreases angular velocity. However, because the condition was not viscous-friction limiting, it was not possible to determine if this correlated to a decrease in torque. (46). Thus, the impact of alanine-substitution for the charged residues on torque-generation is remarkable. Considering that all of these residues are involved in γ- or β-phosphate binding, the interaction between the catalytic site of F₁ and the phosphate region of nucleotides is significantly crucial for torque generation.

Interestingly, alanine-substituted mutants did not significantly impact kₐᵣₑₐ₃ values, except for βK164A, although both βE190A and αR364A reduced kₑₐ₃ by a factor of less than 10, while kₑₐ₃ was reduced by a factor between 10⁵ and 10⁶. Thus, it turned out that the apparent binding rate of the substrate was not significantly relevant to torque-
generation. Our recent work on the mechanical modulation of the kinetic power of F₁ indicated that the affinity change process following the first substrate-F₁ docking process (substrate recognition process) is responsible for torque generation. The affinity change process would mainly occur at the phosphate-binding region.

The rotary torque of the mutants positively correlated well with the stiffness of rotary potential, as estimated from the catalytic dwell (Fig. 7b). The stiffness of the rotary potential is a good barometer of how strongly the αβ stator ring holds the γ subunit. This correlation was also observed in a previous study of a mutant with extensive alanine-substitution at the DELSEED loop of the β subunit (44,47,48). Results of the present study also indicated that the p-loop lysine, general base, and arginine-finger, all play a role to tighten the interaction between β and γ, which likely stabilizes the closed conformation of the β subunit.

Role of p-loop lysine - The alanine mutation on p-loop lysine caused drastic suppression of the rate constants for both ATP binding and hydrolysis; 500,000 and 14,000 times smaller than those of wild-type, respectively (Fig. 5, Table 1). Unlike the general base and the arginine-finger, the p-loop lysine is involved in both substrate-recognition and the subsequent affinity change process. Thus, it is highly likely that the p-loop lysine is the most catalytically important residue among the ATP-binding residues in F₁. In the crystal structures of F₁ the p-loop lysine forms hydrogen bonds with β and γ phosphate (Fig. 1b). This residue would therefore be responsible for phosphate binding, and for maintaining the conformation of the phosphate region of ATP to facilitate the binding of other catalytic residues. An NMR study by Yagi et al. also revealed that this residue is crucial for ATP-binding, as well as the conformational transition from the open state to the closed state of the β subunit (10). Taking these findings into account, it is highly likely that hydrogen bond formation between the p-loop lysine and β and γ phosphate of ATP drives the open-to-closed conformational transition that is responsible for torque generation.

Role of general base - The alanine and glutamine mutations on the general base caused drastic suppression of the ATP hydrolysis rate constants to 2,900 and 41,000 times lower than that of wild-type, respectively, while the ATP-binding rate was slightly slower than that of wild-type (Fig. 5, Table 1). Coincidentally, the rotational rate of the glutamine mutant was $2.2 \times 10^{-3}$ rps, equivalent to an ATPase activity of $6.6 \times 10^{-3}$ s⁻¹. To our knowledge, this is the lowest ATPase activity value ever measured for F₁. Thus, the negative charge at
the general base position principally contributes to the acceleration of the ATP hydrolysis step. There are apparent differences in the ATP hydrolysis rates between alanine- and glutamine substituted mutants, as the alanine mutant hydrolyzes ATP faster (Fig. 5, Table 1). In order to retain ATPase activity, it seems likely that similar spatial rearrangements of side chains of other amino acids located at the catalytic sites are required in these mutants, so that the side chain carboxyl groups of other amino acids should occupy spatial positions close to the glutamic acid residues, as indicated in a previous study (35). In view of this, the glutamine, which occupies the aforementioned spatial position with an amide group, prevents spatial rearrangements more readily than the shorter alanine, and hence, reduces the efficiency of ATP hydrolysis.

The general base has been thought to activate the water molecule between the γ phosphate and itself, and thereby induce nucleophilic attack of water on the γ phosphate. However, this prevailing view has been challenged by a recent QM/MM study on the catalysis of F₁ (5), which suggested that γ phosphate dissociation occurs before protonation of the general base, and that the general base-facilitated proton-transfer is the kinetic bottleneck in the hydrolysis step. Our findings that mutants F₁(βE190A or Q) still retain catalytic activity agree with the QM/MM study (5). Therefore, it is likely that proton-relay occurs after γ phosphate dissociation, via the surrounding water molecules, and in the absence of the general base.

Role of arginine finger - The alanine mutation on the arginine-finger reduced the rate constant of the ATP hydrolysis step by a factor of 950, compared with that of the wild-type, while the rate of ATP binding was slightly suppressed (Fig. 5, Table 1). Thus, the positive charge at the arginine-finger position principally contributed to rate-enhancement of the ATP hydrolysis step. This is consistent with results reported from a structural study on F₁ (29). Another prominent impact of the arginine-finger mutation is that it caused severe ADP inhibition (Fig. 4a). The enhancement of ADP inhibition was also reported in the lysine-substituted mutant (39). The inhibition of the alanine-substituted mutant was the most severe inhibition, which occurred after only a few turns of rotations, and could not be activated by thermal agitation within 10 min. We also recently learned that irregular Pᵢ release from the catalytic site prior to ADP leads to an ADP-inhibited state (49). Considering that the positive charge of the arginine-finger electrostatically interacts with the γ phosphate of ATP, it is highly probable that the αR364A mutant reduces the binding affinity to the product Pᵢ, which promotes severe ADP inhibition.
Although arginine finger and p-loop lysine possess a positively charged group that interacts with β- and γ-phosphate of bound ATP, the impacts of the Ala-substitution are different from each other. The p-loop lysine mutant showed drastic suppression of both nucleotide binding and hydrolytic activity while the arginine-finger mutant mainly decreases the hydrolytic activity. This is attributable to the positional difference of the residues in the tertiary structure of F$_1$-ATPase; p-loop lysine is located in the Walker A motif on the β subunit while arginine finger resides on the α subunit. It is evident from the crystal structure of F$_1$-ATPase (29) that F$_1$-ATPase closes the α–β interface upon the rotation to trigger the hydrolytic reaction. The widely accepted scenario is that after ATP binds to the catalytic site that is mainly formed with the residues on the β subunit, F$_1$-ATPase closes the α–β interface. This conformational change accompanies with approximation of arginine finger toward bound ATP. Several lines of theoretical works (5,33) strongly suggested that this positional shift of arginine finger is thought to be crucial for the initiation of the hydrolytic reaction. On the other hand, p-loop lysine forms hydrogen bonds with ATP before the closing of the α–β interface. These structural features would be basis for the distinct difference in the role of these residues.
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FOOTNOTES
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FIGURE LEGENDS
FIGURE 1. Rotary motion of mutant F\textsubscript{1}
\textbf{a}: Side view of the crystal structure of F\textsubscript{1} (PDB code: 2HLD): interface between \(\alpha\)DP and \(\beta\)DP. The \(\alpha\), \(\beta\), and \(\gamma\) subunits are shown in blue, green, and gray, respectively. The arginine-finger (\(\alpha\)R364 in TF\textsubscript{1}; \(\alpha\)R373 in MF\textsubscript{1}), p-loop lysine (\(\beta\)K164 in TF\textsubscript{1}; \(\beta\)K162 in MF\textsubscript{1}), general base (\(\beta\)E190 in TF\textsubscript{1}; \(\beta\)E188 in MF\textsubscript{1}), and ATP are shown by orange, green, blue, and red space-filling models, respectively. Amino acid residues are numbered according to the TF\textsubscript{1} sequence. \textbf{b}: Schematic diagrams of the interaction between the catalytic site (\(\beta\)DP) and ATP. \textbf{c}: Time courses of rotary motion in the presence of 1 mM ATP; red, orange, green, blue, and yellow represent the time courses of wild-type F\textsubscript{1}, F\textsubscript{1}(\(\alpha\)R364A), F\textsubscript{1}(\(\beta\)K164A), F\textsubscript{1}(\(\beta\)E190A), and F\textsubscript{1}(\(\beta\)E190Q), respectively. \textbf{d}: Histogram of the angular position during rotation calculated from Fig. 1c. \textbf{e}: The rotational velocity (\(v\)) at various ATP concentrations. The solid curves represent Michaelis-Menten fits with \(V = V_{\text{max}}[\text{ATP}]/([\text{ATP}]+K_m)\), where \(V_{\text{max}}^{\text{wild-type}} = 5.6 \text{ s}^{-1}\), \(V_{\text{max}}^{\alpha\text{R364A}} = 1.1 \times 10^{-1} \text{ s}^{-1}\), \(V_{\text{max}}^{\beta\text{K164A}} = 7.4 \times 10^{-3} \text{ s}^{-1}\), \(V_{\text{max}}^{\beta\text{E190A}} = 4.4 \times 10^{-2} \text{ s}^{-1}\), \(V_{\text{max}}^{\beta\text{E190Q}} = 2.2 \times 10^{-3} \text{ s}^{-1}\), \(K_m^{\text{wild-type}} = 1.2 \mu\text{M}\), \(K_m^{\beta\text{K164A}} = 2.5 \times 10^2 \mu\text{M}\), \(K_m^{\beta\text{E190A}} = 3.6 \times 10^{-2} \mu\text{M}\), and \(K_m^{\beta\text{E190Q}} = 1.5 \times 10^{-3} \mu\text{M}\). From these, the following rate constants for ATP binding were calculated as \(k_{\text{on}} = 3 \times V_{\text{max}}^{\text{wild-type}}/K_m\), with \(k_{\text{on}}^{\text{wild-type}} = 1.4 \times 10^7 \text{M}^{-1}\text{s}^{-1}\), \(k_{\text{on}}^{\beta\text{K164A}} = 9.0 \times 10^2 \text{M}^{-1}\text{s}^{-1}\), \(k_{\text{on}}^{\beta\text{E190A}} = 3.7 \times 10^6 \text{M}^{-1}\text{s}^{-1}\) and \(k_{\text{on}}^{\beta\text{E190Q}} = 4.5 \times 10^6 \text{M}^{-1}\text{s}^{-1}\). The dashed curve represents the rotational velocity of wild-type F\textsubscript{1} with a gold colloidal bead, measured in the previous study (44).

FIGURE 2. Stepping rotation of \(\beta\)E190 mutants
\textbf{a}: Examples of the time course of rotation by F\textsubscript{1}(\(\beta\)E190A). Buffer was exchanged during observation, and ATP concentration was reduced from 100 \(\mu\text{M}\) (red) to 50 nM (blue). The insets are x-y trajectories during rotation in the presence of 100 \(\mu\text{M}\) (left) or 50 nM ATP (right). \textbf{b}: Histogram of the angular position of F\textsubscript{1}(\(\beta\)E190A) in the presence of 100 \(\mu\text{M}\) (bottom) or 50 nM ATP (top). \(\Delta\theta_1\) and \(\Delta\theta_2\) represent the angle
distances of the pausing angles at 50 nM ATP compared with those at 100 μM ATP. c: Distributions of angle distances of F₁(βE190A) (Δθ₁, Δθ₂). The average values of Δθ₁ and Δθ₂ are -1.8±6.4° and 33±9.2° (mean ± S.D., n = 33). d: Examples of the time course of rotation by F₁(βE190Q). Buffer was exchanged during observation, and ATP concentration was reduced from 10 μM (red) to 10 nM (blue). e: Histogram of the angular position of F₁(βE190Q) in the presence of 1 mM (bottom) or 1 nM ATP (top). Δθ₁ and Δθ₂ represent the angle distances of the pausing angles at 1 nM ATP from that at 1 mM ATP. f: Distributions of angle distances of F₁(βE190Q) (Δθ₁, Δθ₂). The average values of Δθ₁ and Δθ₂ are 9.9±10° and 54±12° (mean ± S.D., n = 21).

FIGURE 3. Kinetic analysis of βE190 mutants

a: Histograms of the dwelltime of F₁(βE190A) at ATP binding angle (top) or catalytic angle (bottom) in the presence of 1 μM (right), 100 nM (middle) and 10nM ATP (left). Curves represent the fittings with a single-order reaction scheme, y=C•exp(-kt). b: Rate constants of ATP binding and catalysis of F₁(βE190A) determined from a. Solid lines represent the linear fittings; k_{on}^{βE190A} = 4.9×10^{6} M^{-1}s^{-1}, k_{cat}^{βE190A} = 0.13 s^{-1}. c: Histograms of the dwelltime of F₁(βE190Q) at ATP binding angle (top) or catalytic angle (bottom) in the presence of 1 μM (right), 100 nM (middle) and 1nM ATP (left). Curves represent the fittings with a single-order reaction scheme, y=C•exp(-kt). d: Rate constants of ATP binding and catalysis of F₁(βE190Q) determined from c. Solid lines represent the linear fittings; k_{on}^{βE190Q} = 1.7×10^{6} M^{-1}s^{-1}, k_{cat}^{βE190Q} = 9.3×10^{-3} s^{-1}.

FIGURE 4. Stepping rotation of αR364A mutant

a: Examples of the time course of rotation by F₁(αR364A). This variant lapsed into severe ADP inhibition (gray), and after forcible rotation with magnetic tweezers (green), exhibited stepwise counterclockwise rotary motion (red). The insets are x-y trajectories during ADP inhibition (left) or stepwise rotation (right). b: Histogram of the angular position during ADP inhibition (top) or stepwise rotation (bottom). Δθ represents the angle distance of the pausing angle at 1 mM ATP from that of ADP inhibition. c: Distributions of angle distances (Δθ). The average value of Δθ was 0±6.0° (mean ± S.D., n = 13). d: Time course of the stepping rotation of F₁(αR364A) in the presence of 1 μM ATP. Δθ₁ represents the relative angular position.
of original pause at ATP saturating condition to ATP binding pause. $\Delta \theta_2$ represents the relative angular position of the original pauses. e: Distributions of angle distances ($\Delta \theta_1$, $\Delta \theta_2$). The average values of $\Delta \theta_1$ and $\Delta \theta_2$ are $42 \pm 12^\circ$ and $109 \pm 28^\circ$ (mean ± S.D., n = 36). f: Histograms of the dwelltime at catalytic angle (top) or ATP binding angle (bottom) in the presence of 1 mM (right) and 1 $\mu$M ATP (left). Curves represent the fittings with a single-order reaction scheme, $y=C\cdot\exp(-kt)$. The rate constants of ATP binding and catalysis were determined as $k_{on}^{\alpha R364A} = 2.4\times10^7$ M$^{-1}$s$^{-1}$, and $k_{cat}^{\alpha R364A} = 0.45$ s$^{-1}$.

**FIGURE 5.** Rate constants of ATP binding and catalysis

a: Rate constants of ATP binding for wild-type and $F_1$($\beta$K164A) were determined from Fig. 1e, where $k_{on}^{ATP} = 3\times V_{max}/K_m$. Those for $F_1$($\beta$E190A), $F_1$($\beta$E190Q), and $F_1$($\alpha$R364A) were determined from the dwelltime analyses of Figs. 3a, 3c, 4f. b: Rate constants of catalysis ($k_{cat}$). The rate constants for wild-type and $F_1$($\beta$K164A) were determined from Fig. 1e, where $k_{cat} = 3\times V_{max}$. Those for $F_1$($\beta$E190A), $F_1$($\beta$E190Q), and $F_1$($\alpha$R364A) were determined from the dwelltime analyses of Figs. 3a, 3c, 4f.

**FIGURE 6.** Rotary torque

a, b, c: The fluctuation theorem was employed for the torque measurement of $F_1$($\beta$E190A), $F_1$($\beta$K164A), and $F_1$($\alpha$R364A). The ln$[P(\Delta \theta)/P(-\Delta \theta)]$ is plotted against $\Delta \theta/k_B T$. The slope of this plot represents the rotary torque generated by $F_1$. The average torque was determined from a linear approximation of all data points (solid line). d: Torque ($N$) amplitudes generated by wild-type $F_1$, $F_1$($\beta$E190A), $F_1$($\beta$K164A), and $F_1$($\alpha$R364A) are represented as red, blue, green and orange bars, respectively. Light gray symbols represent wild-type $F_1$ in the presence of UTP and ribose-triphosphate (RTP) (Arai and Yukawa et al, submitted).

**FIGURE 7.** Rotary potential

a: Rotary potential of the $F_1$ catalytic waiting state. Probability densities of angular positions during the pause from the five molecules were transformed into rotary potentials according to Boltzman’s law: wild-type $F_1$ (red), $F_1$($\beta$E190A) (blue), $F_1$($\beta$K164A) (green), and $F_1$($\alpha$R364A) (orange). Determined potentials were fitted with the harmonic function $\Delta G = 1/2 \cdot \kappa \cdot \theta^2$, where $\kappa$ is the torsion stiffness. Determined stiffness values were 79, 42, 22, and 38 pN·nm for wild-type $F_1$, $F_1$($\alpha$R364A), $F_1$($\beta$K164A), and $F_1$($\beta$E190A),
respectively. b: Rotary torque plotted against rotary potential stiffness.

Table I. Kinetic parameters, Stiffness and Torque

|         | $k_{on}$ (s$^{-1}$ M$^{-1}$) | $k_{cat}$ (s$^{-1}$) | Stiffness (pN·nm) | Torque (pN·nm) |
|---------|-----------------------------|----------------------|-------------------|----------------|
| Wild-type* | 2.6×10$^7$                  | 3.9×10$^2$           | 78                | 43             |
| βE190A  | 4.9×10$^6$                  | 1.3×10$^{-1}$        | 38                | 25             |
| βE190Q  | 1.6×10$^6$                  | 9.3×10$^{-3}$        | N.D.              | N.D.           |
| βK164A  | 5.2×10$^1$                  | 2.8×10$^{-2}$        | 21                | 10             |
| αR364A  | 2.4×10$^4$                  | 4.1×10$^{-1}$        | 42                | 28             |

*Measured in previous studies (Tanigawa et al, 2012; Okuno et al, 2010)
Fig. 1
Fig. 2
Fig. 3
Fig. 4
Fig. 5
Fig. 6
