A Conformational Change of the γ Subunit Indirectly Regulates the Activity of Cyanobacterial F₁-ATPase*†∆‡§

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Background: A conformational change of the γ subunit of ATP synthase may be critical for enzyme regulation.
Results: A conformational change of γ controls both ADP inhibition and ε inhibition.
Conclusion: The γ subunit indirectly regulates the activity by way of ADP inhibition and ε inhibition.
Significance: Regulation system of ATP synthase based on the unique molecular structure of the γ subunit was revealed.

The central shaft of the catalytic core of ATP synthase, the γ subunit consists of a coiled-coil structure of N- and C-terminal α-helices, and a globular domain. The γ subunit of cyanobacterial and chloroplast ATP synthase has a unique 30–40-amino acid insertion within the globular domain. We recently prepared the insertion-removed αβγ complex of cyanobacterial ATP synthase (Sunamura, E., Konno, H., Imashimizu-Kobayashi, M., and Hisabori, T. (2010) Plant Cell Physiol. 51, 855–865). Although the insertion is thought to be located in the periphery of the complex and far from catalytic sites, the mutant complex shows a remarkable increase in ATP hydrolysis activity due to a reduced tendency to lapse into ADP inhibition. We postulated that removal of the insertion affects the activity via a conformational change of two central α-helices in γ. To examine this hypothesis, we prepared a mutant complex that can lock the relative position of two central α-helices to each other by way of a disulfide bond formation. The mutant obtained showed a significant change in ATP hydrolysis activity caused by this restriction. The highly active locked complex was insensitive to N-dimethyldecylamine-N-oxide, suggesting that the complex is resistant to ADP inhibition. In addition, the lock affected ε inhibition. In contrast, the change in activity caused by removal of the γ insertion was independent from the conformational restriction of the central axis component. These results imply that the global conformational change of the γ subunit indirectly regulates complex activity by changing both ADP inhibition and ε inhibition.

The F₀F₁-ATP synthase (F₀F₁) catalyzes synthesis of ATP from ADP and inorganic phosphate using the electrochemical proton gradient formed across chloroplast and cyanobacterial thylakoid membranes, mitochondrial inner membranes, and bacterial plasma membranes by photosynthetic or respiratory electron transfer reaction (1, 2). F₀F₁ consists of a water-soluble F₁ part, which contains catalytic sites for ATP synthesis and hydrolysis, and a membrane-embedded part F₀ which is involved in proton translocation. F₁ consists of five different subunits with a stoichiometry of αβγδε (3) and F₀ consists of three different subunits with stoichiometry of αβγδε (4–6). F₀ solely catalyzes the ATP hydrolysis reaction and the minimum catalytic core of F₁ is αβγ complex (7). In 1994, the first crystal structure of mitochondrial αβγ complex was determined (8) revealing an alternating hexagonal arrangement of three α and three β subunits around an α-helical domain containing the N- and C-terminal regions of the γ subunit. Because three β subunits in the structure showed different conformations due to different nucleotide binding situations, the reported structure pointed to the idea of rotation of the γ subunit against the αβ subunit ring during catalysis, first proposed by Boyer and co-workers (9). In 1997, rotation of the γ subunit coupled with ATP hydrolysis was directly visualized under an optical microscope by attaching a fluorescent-labeled actin filament to the γ subunit of the αβγ complex fixed on a glass surface (10).

When the electrochemical proton gradient across the membrane is insufficient for ATP synthesis, F₀F₁ can potentially hydrolyze ATP. To prevent this wasteful reverse reaction, the enzyme possesses multiple regulatory mechanisms. The most common regulatory mechanism is ADP inhibition irrespective of the origin of F₁-ATPase. Tightly bound MgADP at the catalytic site strongly prevents a ATP hydrolysis reaction but not ATP synthesis (11–15). Recovery from ADP inhibition is accelerated by ATP binding to the noncatalytic sites on the α subunits (16). Single molecule analysis of the catalytic turnover of this enzyme indicates that ADP inhibition can be observed as

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3 The abbreviations used are: F₀F₁, F₀F₁-ATP synthase; CF₁, chloroplast F₁; LDAO, N-dimethyldecylamine-N-oxide.
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long pauses during rotation (17). In addition, the ε subunit works as an intrinsic inhibitor of ATP hydrolysis in bacterial and chloroplast enzymes. This inhibition is referred to as ε inhibition. The C-terminal helix-turn-helix domain of ε is important for the inhibition, which adopts a large conformational change from a “retracted” form to an “extended” form (18–21). In the case of chloroplast-type F₁, the ε subunit seems to inhibit ATP hydrolysis activity more strongly than the bacterial ε (22–24). Single molecule observations of ε inhibition in cyanobacterial and bacterial F₁ showed that the ε subunit of cyanobacterial F₁ completely stops the rotation of the γ subunit (25), whereas that of bacterial F₁ decreases the average rotation speed and increases pause duration (26, 27). Again, the ε inhibition is thought to be the regulatory mechanism that prevents futile ATP hydrolysis, because this subunit may not inhibit the ATP synthesis reaction (20).

Chloroplast FₐF₃ possesses another regulatory mechanism: the redox regulation system mediated by thioredoxin (28). This is thanks to the chloroplast F₁γ containing an additional ~35 amino acid insertion in the middle of the sequence (supplemental Fig. S1) compared with other bacterial and mitochondrial F₁γγ (29). Two critical Cys residues are located in this insertion sequence and regulate ATP hydrolysis activity via disulfide bond formation (30).

Compared with the bacterial and mitochondrial F₁γγ, the γ subunit of cyanobacterial FₐF₃ also bears the inserted sequence like spinach CF₁γγ, although the sequence lacks nine amino acids including two regulatory Cys residues for redox regulation (31). In our previous studies, the mutant α₃β₃γ complex was prepared in which the γ insertion sequence of cyanobacterium Thermosynechococcus elongatus BP-1 F₁ was deleted (25, 32). The mutant complex obtained showed a large increase in ATP hydrolysis activity as a consequence of the low tendency to lapse into ADP inhibition, and lower sensitivity to ε inhibition. Furthermore, the mutant strain of cyanobacterium Synechocystis sp. PCC 6803, whose F₁γγ insertion was deleted, showed lower intracellular ATP levels due to insufficient prevention of the ATP hydrolysis activity in the dark (32, 33). These results strongly suggest that the insertion plays an important role in the regulation of ATP hydrolysis activity of cyanobacterial FₐF₃ in vivo.

However, the location of the insertion seems to be at the bottom of the γ molecule and over 60 Å away from catalytic sites on the α₃β₃ ring (34, 35). How does the conformational change by removal of the inserted sequence affect the catalytic activity despite the catalytic sites being distant from the inserted sequence in the enzyme complex? Since little contact region is reported in the crystal structure of F₁ between the γ subunit and the α₃β₃ ring (Fig. 1A), we postulated that relative slippage of N- and C-terminal α-helices may connect between the conformational change of the bottom part of γ and a conformational change in the upper part of γ, as suggested by our previous study on the redox regulation of this enzyme complex (36).

To examine our hypothesis, we introduced two Cys residues at an appropriate position in each of the N- and C-terminal α-helices of cyanobacterial F₁γγ, allowing us to lock or unlock the relative movement of these two central α-helices. Here we defined the oxidized complex as the locked one and the reduced complex as the unlocked one. The mutant complexes obtained showed significant changes in their ATP hydrolysis activities and sensitivities to ε inhibition. In addition, we prepared several lock mutants lacking a γ insertion, and examined the change in ATP hydrolysis activities by locking or unlocking the two central α-helices. Very interestingly, these mutants showed higher ATP hydrolysis activities irrespective of the lock status of the central α-helices. Based on these results, the relationship between the removal of the insertion and the accompanying conformational change of γ is discussed.

EXPERIMENTAL PROCEDURES

Materials—Aldrich™-2, ATP, phosphoenolpyruvate, and pyruvate kinase/lactate dehydrogenase were purchased from Sigma. Diamide was obtained from MP Biomedicals (Santa Ana, CA). NADH was purchased from Roche Diagnostics. Other chemicals were of the highest grade commercially available.

Strains—Escherichia coli strains used were DH5α for cloning and BL21(DE3) uncΔ702 (Tcr, ATPase mutant, BL21(DE3) uncΔ702, asn::Tn10) (37, 38) for expression of α₃β₃γ complex of T. elongatus BP-1. The latter strain was a kind gift from Dr. C. S. Harwood (University of Iowa).

Construction of Expression Plasmids for Lock Mutants—The expression plasmid for the α₃β₃γ complex of T. elongatus BP-1 and the expression plasmid of Cys-less α₃β₃γ complex (all native Cys residues were mutated to Ser) for the single molecule experiment were constructed in the previous study (25). Using the plasmid for the Cys-less complex as template, the expression plasmids for lock mutants were prepared and two Cys residues were introduced into γ. Site-directed mutagenesis was performed by the overlap extension method (39). The primers used for mutagenesis are shown in supplemental Table S1. To prepare the additional lock mutants lacking the γ (Leu₁⁹₈–Val₂²²) insertion, both expression plasmids for the Cys-less complex and the mutant complex lacking the insertion sequence (25) were digested with SacI and NheI. Ligation then yielded the expression plasmid for the Cys-less mutant complex lacking the insertion (hereafter referred as Cys-less/ΔIns). Using the plasmid as a template, Cys residues that allow locking of the central α-helices were introduced as described, and lock mutant plasmids lacking the insertion were obtained.

Expression and Purification of α₃β₃γ Complex and the ε Subunit—Expression and purification of the α₃β₃γ complex were performed as described (25) with some modifications. After nickel-nitrilotriacetic acid chromatography, proteins were stored at 4°C in solution containing 20 mM potassium phosphate (pH 8.0), 100 mM KCl, 0.1 mM ADP, 0.1 mM MgCl₂, 1 mM DTT, and 55% (w/v) ammonium sulfate before further purification. Then, the ammonium sulfate precipitate was redissolved in “HPLC buffer,” which contains 50 mM HEPES/KOH (pH 8.0), 100 mM KCl, 0.1 mM ADP, and 0.1 mM MgCl₂, and then purified by gel filtration chromatography on a Superdex™ 200 column (GE Healthcare) equilibrated with HPLC buffer in advance. The purified complex was stored at ~80°C in 10% glycerol. The ε subunit was expressed and purified as
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Results

Preparation of Lock Mutant γ to Control the Slippage of α-Helices—In the previous study, we prepared a Cys-less α$_3$β$_3$γ complex of thermophilic cyanobacterial F$_o$F$_1$, for single molecule experiments in which all native Cys at positions 144 and 194 on α, 53 on β, and 90 on γ were substituted with Ser (25). In this study, this complex was referred to as the Cys-less complex and used as a template for preparing lock mutants for two central α-helices in γ. Because no crystal structure information on cyanobacterial or chloroplast F$_1$, α$_3$β$_3$γ complex is available to date, we applied the bovine mitochondrial F$_1$ structure (Protein Data Bank code 1E79) (18) (Fig. 1) instead to design lock mutants for two central α-helices in γ. Residue numbers are those for bovine mitochondrial F$_1$ and the numbers in parentheses correspond to the γ subunit of T. elongatus BP-1. The figures were generated with PyMOL.

Figure 1. Positions of Cys residues introduced for locking of the central α-helices of the γ subunit. A, structures of β$_{up}$ (yellow), and γ (blue) subunits in bovine F$_1$ structure (Protein Data Bank code 1E79) are shown. β$_{up}$ and β$_{up}$ are based on the first crystal structure of F$_1$ (8). N and C termini of α-helices in γ are indicated. Residues of γ, Ala$^{27}$, Tyr$^{31}$, Ala$^{34}$, Leu$^{38}$, Thr$^{222}$, Gln$^{225}$, and Ser$^{229}$ are shown as a stick model and colored in orange. In β$_{up}$ bound MgADP is shown as a sphere model. The apparent position of the inserted sequence of cyanobacterial and chloroplast F$_1$, γ is circled in red. B, magnified structure of the Cys introduced region is marked by a square in A. Residue numbers are those for bovine mitochondrial F$_1$ and the numbers in parentheses correspond to the γ subunit of T. elongatus BP-1.

Preparation of Lock Mutant after Oxidation—The ammonium sulfate precipitant as mentioned was re-dissolved in HPLC buffer with a protein concentration of 0.5 mg/ml. The V32C/A264C complex was oxidized by 100 μM Aldrithiol-2 for 60 min at 25 °C, A28C/L267C, V32C/A264C, and A35C/A264C complexes were oxidized by 500 μM Aldrithiol-2 for 60 min at 25 °C, and the V39C/A264C complex was oxidized by 500 μM diamide for 60 min at 25 °C. Oxidized complexes were further purified by gel filtration and stored as described.

Measurement of ATP Hydrolysis Activity—ATP hydrolysis activity was measured by using an ATP regenerating system. The assay mixture contained 50 mM HEPES/KOH (pH 8.0), 100 mM KCl, 2 mM MgCl$_2$, 2 mM ATP, 6.8 to 11 milliunits/ml of lactate dehydrogenase, 2 mM phospho-enolpyruvate, and 0.2 mM NADH. The assay was carried out at 25 °C. After addition of the ATPase complex, the activity was determined from the steady state slope by monitoring the decrease in NADH absorption at 340 nm with a Jasco spectrophotometer model V650 (Jasco, Tokyo, Japan). To investigate the effect of LDAO on the ATP hydrolysis activity, LDAO (final concentration of 0.1% w/v) was added to the assay mixture before addition of the enzyme complex. In this case, the activity was determined from maximum slope.

Inhibition of ATP Hydrolysis Activity of α$_3$β$_3$γ Complex by the e Subunit—The experimental conditions were the same as described above. The assay was initiated by adding the α$_3$β$_3$γ complex followed by addition of various concentrations of the e subunit. The extent of inhibition of ATP hydrolysis activity was determined by the steady state slope in the presence of absence of the e subunit (25). The titration curve was fitted with the hyperbolic equation, $y = A \times [\epsilon]_{free}/(K_D + [\epsilon]_{free})$, where $y$ represents the percentage of inhibition, $A$ is the maximum inhibition (%), and $K_D$ is an apparent equilibrium dissociation constant for the e subunit. To calculate the concentration of the free e subunit, we assumed the complex that bound the e subunit is completely inhibited, thus $[\epsilon]_{free} = [\epsilon]_{add} - [\text{complex}] \times y/100$.

RESULTS

Preparation of Lock Mutant γ to Control the Slippage of α-Helices—In the previous study, we prepared a Cys-less α$_3$β$_3$γ complex of thermophilic cyanobacterial F$_o$F$_1$ for single molecule experiments in which all native Cys at positions 144 and 194 on α, 53 on β, and 90 on γ were substituted with Ser (25). In this study, this complex was referred to as the Cys-less complex and used as a template for preparing lock mutants for two central α-helices in γ. Because no crystal structure information on cyanobacterial or chloroplast F$_1$, α$_3$β$_3$γ complex is available to date, we applied the bovine mitochondrial F$_1$ structure (Protein Data Bank code 1E79) (18) (Fig. 1) instead to design lock mutants. Finally positions Ala$^{28}$, Val$^{32}$, Ala$^{35}$, and Val$^{39}$ on the N-terminal α-helix, and Ala$^{264}$, Leu$^{267}$, and Ala$^{268}$ on the C-terminal α-helix (residue numbers are those for F$_1$ of T. elongatus BP-1) were selected as candidate residues for Cys substitution. We then prepared nine mutants by a combination of a set of mutations (for example, A28C/L267C, V32C/L267C, and A35C/L267C), which potentially induces N-terminal α-helix slip-up or down by one turn against the C-terminal α-helix after disulfide bond formation. Primers used for the mutation are described (25). Protein concentrations were measured by the Bradford method with bovine serum albumin as a standard.

Determination of the Optimal Oxidation Conditions—Purified complex (50 μl) was oxidized by varying oxidizing reagents (Aldrithiol-2 and diamide), the concentration of the reagent (50, 100, 500, and 1000 μM), reaction time (30 and 60 min), and protein concentration (0.5 and 1.0 mg/ml). The reaction was stopped by adding 5% (w/v, final concentration) trichloroacetic acid. After centrifugation, the supernatant was removed and the remained oxidant was washed away with 500 μl of acetone. Then, the mixture was centrifuged, and air dried after removing the supernatant. The pellet was dissolved in 50 mM Tris/HCl (pH 7.5) and 1% (w/v) SDS and then electrophoresed on 12% polyacrylamide gel. The optimal oxidation condition was determined by comparing the band intensities of the oxidized γ subunit on the gel.

Purification of Lock Mutant after Oxidation—The ammonium sulfate precipitant as mentioned was re-dissolved in HPLC buffer with a protein concentration of 0.5 mg/ml. The V32C/A264C complex was oxidized by 100 μM Aldrithiol-2 for 60 min at 25 °C, A28C/L267C, V32C/A264C, and A35C/A264C complexes were oxidized by 500 μM Aldrithiol-2 for 60 min at 25 °C, and the V39C/A264C complex was oxidized by 500 μM diamide for 60 min at 25 °C. Oxidized complexes were further purified by gel filtration and stored as described.
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listed in supplemental Table S1. Site-directed mutagenesis was performed as described under “Experimental Procedures,” and all mutants were successfully expressed in \( E. \) coli.

Dynamics of the Mutant \( \gamma \) Subunit in the Complex—Mutant complexes were basically prepared as the unlocked form after gel filtration chromatography. We then determined disulfide bond formation conditions to lock two central \( \alpha \)-helices by changing the concentrations of oxidizing reagents, protein concentrations, and reaction periods (For more detail, see “Experimental Procedures”). Aldrithiol-2 and diamide were used as oxidizing reagents. To examine the efficiency of disulfide bond formation, nonreducing SDS-PAGE analysis, in which 2-mercaptoethanol was omitted from the sample preparation buffer, was performed. When the \( \gamma \) subunit was locked by a disulfide bond, a band shift of \( \gamma \) on the gel was clearly observed as a consequence of conformational constraints. After optimizing oxidation conditions, A28C/L267C, A35C/A264C, and V39C/A264C mutants were mostly oxidized (Fig. 2A), whereas A35C/L267C, A28C/A268C, and A35C/A268C were not (Fig. 2B). The other mutants V32C/L267C, V32C/A268C, and V32C/A264C were partially oxidized by 55, 68, and 38%, respectively, as estimated based on their band intensities (Fig. 2, A and B). Incomplete disulfide bond formation between two Cys residues may be attributed to an excessive distance between thiol groups might be slightly different from the reported crystal structure. Consequently, the molecular composition of some of the complexes used in this study was heterogeneous due to their incomplete oxidation, somewhat complicating the resulting interpretation. Nevertheless, our key objective was to obtain a series of Cys mutants that could induce slippage of central \( \alpha \)-helices after oxidation, and we therefore mainly focused on the series of mutants, V32C/A264C, A35C/A264C, and V39C/A264C, and A28C/L267C and V32C/A268C to assess the effect of slippage of the \( \alpha \)-helices.

Change in the ATP Hydrolysis Activity by Unlocking and Locking of Two Central \( \alpha \)-Helices—We then measured the change in ATP hydrolysis activities of unlocked and locked mutants (Fig. 3). Compared with wild type (WT) and Cys-less complexes, unlocked mutants except A28C/L267C showed remarkably higher ATP hydrolysis activities (Fig. 3, open bars), although mutations applied were only Cys substitution of \( \gamma \), which may not directly affect the catalytic sites. Disulfide bond formation in V32C/A268C and V39C/A264C mutants greatly increased their ATP hydrolysis activities, by about 4- and 2.5-fold, respectively. Although the locked A35C/A264C mutant also showed 4-fold higher ATP hydrolysis activity than the unlocked one, the specific activity obtained for this complex was not much higher than the other two mutants. Irrespective of disulfide bond formation, V32C/A264C showed a stable activity (at around 10 units/mg). The activity of A28C/L267C was much lower than that of WT and the Cys-less complex, and slightly decreased by disulfide bond formation.

Stimulation of ATP Hydrolysis Activity by LDAO—To determine the cause of the strong activation by locking the two central \( \alpha \)-helices in the case of V32C/A268C, we examined the effect of LDAO on the activity of the mutant complexes (Table 1). As mentioned, \( F_1 \) has the intrinsic regula-
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FIGURE 3. ATP hydrolysis activities of unlocked and locked mutants. ATP hydrolysis activities of wild type (WT), Cys-less, and lock mutants were determined at 25 °C in the presence of 2 mM MgATP. The hatched and light gray bars represent the activities of WT and Cys-less, respectively. Open and closed bars represent the activities of unlocked and locked mutants, respectively. The results of three experiments were averaged. Error bar represents S.D. The conditions of α-helices in the mutant complexes were shown as U (unlocked) and L (locked), respectively. In the bottom, the change in the activity by lock is indicated based on the activity of each unlocked mutant as 100%.

TABLE 1
Effects of LDAO on ATP hydrolysis activities of lock mutants
The activity in the presence of LDAO was measured as described under “Experimental Procedures.” The activity measured in the absence of LDAO was obtained from Fig. 3. The results of three independent experiments were averaged. The results obtained from LDAO less sensitive complexes are in bold.

| Complex          | ATP hydrolysis activity | Units/mg | Units/mg |
|------------------|-------------------------|----------|----------|
|                  | Unlocked | LDAO       | Locked | LDAO     |
| WT               | 0.27 ± 0.03 | 9.4 ± 0.5 | 0.26 ± 0.03 | 11.0 ± 0.3 |
| Cys-less         | 0.76 ± 0.05 | 21.4 ± 0.9 | 0.75 ± 0.05 | 11.6 ± 0.3 |
| A28C/L267C       | 0.18 ± 0.02 | 15.4 ± 2.1 | 0.18 ± 0.02 | 12.0 ± 0.3 |
| V32C/A268C       | 6.7 ± 1.8  | 30.4 ± 3.7 | 6.7 ± 1.8  | 31.5 ± 2.3 |
| V33C/A268C       | 6.7 ± 1.8  | 30.4 ± 3.7 | 6.7 ± 1.8  | 31.5 ± 2.3 |
| V34C/A268C       | 10.4 ± 0.8 | 31.0 ± 1.2 | 10.4 ± 0.8 | 31.5 ± 2.3 |
| V35C/A264C       | 1.9 ± 0.1  | 26.7 ± 4.7 | 1.9 ± 0.1  | 35.7 ± 3.3 |
| V36C/A264C       | 7.9 ± 1.0  | 35.7 ± 1.5 | 7.9 ± 1.0  | 34.1 ± 5.0 |

The ATP hydrolysis activity of unlocked and locked mutants was determined at 25 °C in the presence of 2 mM MgATP. The hatched and light gray bars represent the activities of WT and Cys-less, respectively. Open and closed bars represent the activities of unlocked and locked mutants, respectively. The results of three experiments were averaged. Error bar represents S.D. The conditions of α-helices in the mutant complexes were shown as U (unlocked) and L (locked), respectively. In the bottom, the change in the activity by lock is indicated based on the activity of each unlocked mutant as 100%.

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FIGURE 4. Time course of ATP hydrolysis by WT, Cys-less, and V32C/A268C lock mutant complexes. The complexes were incubated at 25 °C for 10 min in the presence of 50 μM MgADP in advance. 10 μg of the complexes were added into the assay mixture at 25 s after starting the measurement (down arrow); black line, WT; red line, Cys-less; blue line, unlocked V32C/A268C; purple line, locked V32C/A268C.
Change in the ATP Hydrolysis Activity by Unlocking and Locking of Insertion Removal Mutants—In the previous study, we found that the insertion region of H9253 plays a critical role in conferring the ADP inhibition property to the enzyme complex and this is important in prevention of futile ATP hydrolysis activity in vivo (32). To examine the relevance between the removal of the insertion and the relative slippage of two -helices found in this study, we prepared lock mutants lacking the insertion region in H9253 subunit. For this purpose, we deleted the insertion region from three lock mutants, A28C/L267C, V32C/A268C, and V39C/A264C. These mutants showed a significant change in their activities by lock status as shown in Fig. 3. Hereafter, the mutants were referred as A28C/L267C/Ins, V32C/A268C/Ins, and V39C/A264C/Ins, respectively. All mutants were successfully expressed in E. coli, and purified.

Using the same oxidation conditions applied for the mutants containing the insertion, all the mutants were oxidized to the same degree (Fig. 7A). We then measured the change in ATP hydrolysis activity of the mutants by locking and unlocking them (Fig. 7B). Although the Cys-less complex and A28C/L267C complex showed very low ATPase activities (Fig. 3),

TABLE 2
Apparent dissociation constants for the subunit with lock mutants

| Complex            | Unlocked | Locked |
|--------------------|----------|--------|
| Cys-less           | 0.93 ± 0.29 |        |
| V32C/A268C        | 2.0 ± 0.6  | 1.6 ± 0.1 |
| V32C/A264C        | 1.2 ± 0.2  | 2.2 ± 0.2 |
| A35C/A264C        | 1.2 ± 0.2  | 1.4 ± 0.1 |
| V39C/A264C        | 1.6 ± 0.5  | 2.4 ± 0.2 |

Change in the ATP Hydrolysis Activity by Unlocking and Locking of Insertion Removal Mutants—In the previous study, we found that the insertion region of γ plays a critical role in conferring the ADP inhibition property to the enzyme complex and this is important in prevention of futile ATP hydrolysis activity in vivo (32). To examine the relevance between the removal of the insertion and the relative slippage of two -helices found in this study, we prepared lock mutants lacking the insertion region in the γ subunit. For this purpose, we deleted the insertion region from three lock mutants, A28C/L267C, V32C/A268C, and V39C/A264C. These mutants showed a significant change in their activities by lock status as shown in Fig. 3. Hereafter, the mutants were referred as A28C/L267C/ΔIns, V32C/A268C/ΔIns, and V39C/A264C/ΔIns, respectively. All mutants were successfully expressed in E. coli, and purified. Using the same oxidation conditions applied for the mutants containing the insertion, all the mutants were oxidized to the same degree (Fig. 7A). We then measured the change in ATP hydrolysis activity of the mutants by locking and unlocking them (Fig. 7B). Although the Cys-less complex and A28C/L267C complex showed very low ATPase activities (Fig. 3),
deletion of the insertion significantly accelerated their activities, and Cys-less/ΔIns and unlocked insertion removal mutants showed almost the same activities (15 to 20 units/mg) (Fig. 7B). In contrast, the locked A28C/L267C/ΔIns and V32C/A268C/ΔIns showed a slight increase in their activities (about 1.2-fold) when locked, although Student’s t test indicated that the activity of V32C/A268C/ΔIns mutant was not significantly different (p < 0.05). Activity of the V39C/A264C/ΔIns mutant was not affected irrespective of disulfide bond formation.

**DISCUSSION**

ATP synthase is a very unique enzyme because the reaction catalyzed at the catalytic sites located on the β subunit is coupled to rotation of the central axis γ subunit (10). In addition, this enzyme complex is equipped with multiple systems that work to regulate its activity. Although the major mechanism that acts to regulate its activity is ADP inhibition of the catalytic sites (11, 12, 14, 42), the reason that sites of redox regulation and the e inhibition are structurally distant from the catalytic site is poorly understood. We postulated that the relative slippage of the two central α-helices of the γ subunit, which connect the two active domains for catalysis and regulation in the structure of the complex (8), must be a main cause of the change in the enzyme activity (36). To examine this hypothesis, we prepared several mutant complexes in which the relative slippage of two central α-helices can be locked at a certain position by disulfide bond formation. Accordingly, some of the unlocked and locked mutants prepared in this study showed significant changes in activities caused by disulfide bond formation or reduction. Because the mutated residues for Cys substitution are not supposed to interact with the αβγ ring directly in the referenced crystal structure of bovine F1 (18), the relative slippage of two central α-helices was expected to be a major cause of any observed change in activity. However, introduction of Cys residues for disulfide formation to the central α-helices also changed the ATP hydrolysis activity compared with the WT and Cys-less complex (Fig. 3), irrespective of the redox conditions of these cysteines. This may imply that the position of these Cys residues introduced into the complex is critical in causing relative slippage of α-helices due to substitution of the side residue moiety. Because the pK_	ext{a} value of the thiol moiety of Cys is around 8.0, the thiol group of the introduced Cys residue is thought to be half-deprotonated, which might affect the slippage of α-helices and alter the activity.

To confirm the cause of the variability of the mutant activities, we examined the effects of LDAO, which can release the enzyme from the ADP inhibition state (Table 1). Consequently, the activities obtained in the presence of LDAO were not significantly different; 22 to 36 units/mg in the case of the unlocked form and 25 to 36 in the case of the locked form. Only the mutant complex A28C/L267C showed very poor activity and the activity did not recover completely even in the presence of LDAO (Table 1). Variability of the mutant activities examined might be attributed to different sensitivities of the complex against ADP inhibition, which is affected by the relative α-helices’ position of the γ subunit in the complex.

Where is the significant contact point between γ and the αβγ ring, which is important in transferring the signal of the conformational change via the slippage of α-helices of γ to the catalytic site(s)? In the reported crystal structure of F1, the contacts mainly consist of a “catch” region and a region near the “DELSSEED” sequence (8). Catch is formed by some charged residues, and in the C-terminal α-helix of γ. He et al. (43) reported that substitution of the highly conserved Arg with Leu in the C-terminal α-helix, which forms a catch, stimulated ATP hydrolysis activity and, substitution of Gln close to this Arg with Ala, decreased the sulfite activation in chloroplast ATP synthase. Sulfite is recognized to release the complex from ADP inhibition like LDAO. Thus, these residues seem to be important in transferring the conformational change of the γ subunit to the catalytic site. In addition, the corresponding mutation analysis in E. coli F1 showed a drastic decrease in activity (44). As the catch region is formed by βe and γ and no nucleotide binds in βe, there are some cooperative interactions among
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three $\beta$ subunits as suggested (9). The second contact point between $\beta$ and $\gamma$ is the region near the DELSEED sequence on the $\beta$ subunit. There are a couple of studies that examined the role of this region on catalysis (45–47). Mnatsakanyan et al. (46) and Usukura et al. (47) recently found that the region is important to generate torque and achieve mechanical coupling. Although it is still difficult to assign residues that are responsible to torque generation, this region must also be important for transfer of the signal of the conformational change of the $\gamma$ subunit to the catalytic site(s).

Richter and co-workers (48) recently prepared several cross-linked mutants of $\gamma$ to examine the possibility of helical domain movement during rotational catalysis (49). Although they postulated the twisting movement of the $\alpha$-helices, the position where Cys residues were introduced were very close to those employed in this study. However, all of their mutants showed no significant changes in the activity by disulfide bond formation. In addition, it was difficult to estimate the yield of the oxidized complex by their method, which evaluates accessibility of fluorescent chemicals (see Fig. 3 of Ref. 48). One possible reason for the discrepancy between their study and our study is that the interaction between $\alpha_1$, $\beta_1$, and $\gamma$ is not exactly the same as that of the native enzyme, in their case because their enzyme is a hybrid complex comprised of $\alpha$ and $\beta$ subunits from *Rhodospirillum rubrum* $F_1$ and the $\gamma$ subunit from spinach chloroplast $F_1$. One of the mutants in their study, the $\gamma V31C/A276C$ complex, did not appear any activity by the oxyanion sulfite, implying the enzyme was not recovered from ADP inhibition. However, our complex, V32C/A268C, which corresponds to their $\gamma V31C/A276C$ complex based on the sequence homology, was almost completely released from ADP inhibition and did not show remarkable activation by LDAO when oxidized (Table 1).

When V32C/A268C and A35C/A264C complexes were locked by oxidation, not much constraint was expected to be induced because the expected relative position of $\alpha$-helices seemed to be similar as those of the unlocked form, which is the position of two $\alpha$-helices in the crystal structure as mentioned. Their activities were inhibited almost completely by the $\epsilon$ subunit irrespective of the locked or unlocked forms of the $\gamma$ subunit (Fig. 5, B and D). In contrast, the extent of $\epsilon$ inhibition was lower when V32C/A264C and V39C/A264C complexes were in the locked states. These results indicate that the possible slippage of $\alpha$-helix by disulfide bond formation largely affects the extent of $\epsilon$ inhibition. Because the $\epsilon$ subunit can bind to the locked V39C/A264C complex as well as the unlocked one (Fig. 6), we assumed that the decrease of $\epsilon$ inhibition can be attributed to the constraint of the conformational change of the C-terminal domain in $\epsilon$ from the extended to retracted form. In the latest *E. coli* $F_1$ structure, the $\epsilon$ subunit is in the extended form, and the C-terminal domain of $\epsilon$ can interact with both N- and C-terminal $\alpha$-helices of $\gamma$ (21). The relative slippage of two central $\alpha$-helices may weaken the interaction of $\gamma$ and $\epsilon$, and consequently the $\epsilon$ subunit falls into the retracted form, which cannot inhibit the enzyme (50). Although the inhibitory properties of $\epsilon$ inhibition and ADP inhibition appear to be similar, recent single molecule analyses clearly show that these two inhibitions are mechanically different (41, 51).

We finally sought to investigate whether significant activation of the enzyme by removal of the insertion region of cyanobacterial $F_1$-$\gamma$ could be attributed to the relative slippage of two $\alpha$-helices found in this study. To address this question, we prepared additional lock mutants lacking the $\gamma$ insertion region. By deletion of the insertion, basal activities of Cys-less/$\Delta$Ins and unlocked deletion mutants were drastically increased (Fig. 7) as observed in the previous studies (25, 32). In contrast, their activities remained relatively unchanged by locking. In the case of A28C/L267C, change in the activity by locking was reversed upon removal of the insertion. Two other mutants, V32C/A268C and V39C/A264C, showed full activities by removal of the insertion and the activities did not change after locking. These results clearly indicate that removal of the insertion region of $\gamma$ induces another conformational change in $\gamma$ rather than relative slippage of two $\alpha$-helices. In the crystal structure of bovine $F_1$, the upper part of the globular domain in $\gamma$ is in contact with the DELSEED region in $\beta$ and $\beta_D$. Therefore, removal of the insertion may potentially result in a conformational change that affects the activity by affecting the structure of the globular domain itself, and this conformational change is transferred to the catalytic site(s) via interaction between the $\gamma$ and $\beta$ subunits at the bottom of $\beta$. Finally, ADP inhibition must be controlled at the catalytic sites by the induced conformational change of the $\beta$ subunit.

In the case of the chloroplast ATP synthase, $F_1$ possesses the additional regulation, redox regulation in the $\gamma$ subunit. The two functional Cys residues Cys$^{399}$ and Cys$^{205}$ (spinach $F_1$ numbering) involved in this redox regulation are supposed to be located in the bottom of the globular domain of $\gamma$ close to the c-ring (34). Although we are not currently able to provide a definitive conclusion, redox regulation of chloroplast $F_1$ might be achieved by transferring the conformational change of the disulfide bond formation or reduction to the catalytic site(s) via slippage of the central $\alpha$-helices or change in the interaction between the upper part of the globular domain of $\gamma$ and the DELSEED region in $\beta_D$. On this point, further analysis is required to understand how the conformational change caused by redox regulation is transferred to the catalytic site(s) in the enzyme molecule. Previously, we constructed a chimeric cyanobacterial $F_1$ complex in which the 9-amino acid region of spinach CF$_1$-$\gamma$ including two regulatory Cys was introduced into the cyanobacterial $\gamma$ subunit (36). Based on the single molecular observation of rotation of this chimeric complex, we concluded that redox regulation is achieved by controlling the probability to lapse into ADP inhibition. In addition, we have reported the reverse of the redox regulation of ATPase caused by deletion of the Glu-Asp-Glu sequence from the insertion region of the $\gamma$ subunit of spinach CF$_1$ (52, 53). Hence the regulation must certainly be achieved by a slight conformational change of the $\gamma$ subunit.

In this study, we clearly showed that the conformational change of the $\gamma$ subunit can regulate ATP hydrolysis activity by controlling other regulatory mechanisms, ADP inhibition, and $\epsilon$ inhibition. Further studies, especially analysis of the whole molecular structure of the chloroplast-type $F_1$ including regulatory subunits will lead to a more complete understanding of
the complicated regulatory mechanisms of this enzyme complex.

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