Redox Regulation of Rotation of the Cyanobacterial F₁-ATPase Containing Thiol Regulation Switch*

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 F_1 -ATP synthase (F_1 -ATPase) is equipped with a special mechanism that prevents the wasteful reverse reaction, ATP hydrolysis, when there is insufficient proton motive force to drive ATP synthesis. Chloroplast F₁-ATPase is subject to redox regulation, whereby ATP hydrolysis activity is regulated by formation and reduction of the disulfide bond located on the γ subunit. To understand the molecular mechanism of this redox regulation, we constructed a chimeric F_1 complex ($\alpha_3 \beta_3 \gamma_{redox}$) using cyanobacterial F_1 , which mimics the regulatory properties of the chloroplast F₁-ATPase, allowing the study of its regulation at the single molecule level. The redox state of the γ subunit did not affect the ATP binding rate to the catalytic site(s) and the torque for rotation. However, the long pauses caused by ADP inhibition were frequently observed in the oxidized state. In addition, the duration of continuous rotation was relatively shorter in the oxidized $\alpha_3 \beta_3 \gamma_{\rm redox}$ complex. These findings lead us to conclude that redox regulation of CF1-ATPase is achieved by controlling the probability of ADP inhibition via the γ subunit inserted region, a sequence feature observed in both cyanobacterial and chloroplast ATPase y subunits, which is important for ADP inhibition (Sunamura, E., Konno, H., Imashimizu-Kobayashi, M., Sugano, Y., and Hisabori, T. (2010) Plant Cell Physiol. 51, 855-865).

The ATP synthase complex is ubiquitously found in energytransducing membranes such as bacterial plasma membranes, mitochondrial inner membranes, and chloroplast thylakoid membranes, where its basic architecture is highly conserved. ATP synthase consists of two motor units: the F_1 portion which is powered by ATP, and the F_o portion powered by the proton motive force formed across the energy transducing membranes (1). Fuelling of ATP synthase by ATP and proton motive force results in rotational motion of F_1 and F_o , respectively, though they rotate in opposite directions. On the membranes, these two motors are directly connected by protein-protein interaction, and their functions are coupled to each other: the F_1 portion catalyzes ATP hydrolysis and ATP synthesis, and the F_o portion catalyzes proton translocation. Upon formation of a proton motive force across the membranes as a result of respiratory or photosynthetic electron transport, the F_o portion is forced to rotate by the proton motive force. Rotation of F_o induces rotation of the central axis subunits, γ and ϵ , of F_1 , and finally ATP is formed at the catalytic sites on β subunits, presumably due to forced conformational change at the catalytic sites. *Vice versa*, rotation of F_1 forced by ATP hydrolysis induces rotation of F_o and consequently protons are transferred in the opposite direction, where a proton motive force is generated. This enzyme may therefore hydrolyze ATP and transport protons in the opposite direction when there is insufficient proton motive force to drive ATP synthesis.

However this reverse reaction catalyzed by the enzyme is highly restricted *in vivo* because the reaction results in the wasteful consumption of ATP. To avoid this unfavorable reaction *in vivo*, several regulatory systems are known to maintain the ATP synthesis reaction. The bacterial and chloroplast ATP synthases are equipped with an intrinsic inhibitory subunit ϵ , which strongly suppresses the ATP hydrolysis reaction (2–4). The C-terminal α -helical part of the ϵ subunit has been shown to be critical for this inhibitory function (5). Another regulatory mechanism is known as ADP-inhibition (6). The ATP hydrolysis reaction of F₁-ATPase is strongly inhibited by tight binding of ADP to the catalytic site(s). This inhibition is particularly strong under low concentrations of substrate ATP (7).

In addition to those described above, higher plant chloroplast ATP synthase displays an additional regulatory system, a thiol modulation system (8, 9), whereupon it receives reducing equivalents from the photosynthetic electron transport system via thioredoxin-*f*, a disulfide-dithiol exchange mediator protein (10). The γ subunit of the chloroplast ATP synthase contains two regulatory cysteines (see Fig. 1*A*), which are able to form a disulfide bond (8, 11), and the reduction of this disulfide bond by the reduced form thioredoxin (9) activates the enzyme. Inversely, the formation of the disulfide bond on the γ subunit renders the enzyme inactive (11). This regulation is likely to be critical for photosynthetic organisms, because photosynthesis is subject to highly variable environmental conditions, fuelled by an energy source which is both oscillatory (day and night) and fluctuating (irregular weather changes).

The regulatory cysteines on the γ subunit of chloroplast F_0F_1 , Cys-199 and Cys-205 (see Fig. 1*A*), are conserved in different photosynthetic organisms, from green algae such as *Chlamydomonas reinhardtii* (12) to higher plants (8). Although at present nothing is known about the potential γ subunit conformational changes associated with the redox state, the redox switch region seems to be independent from other parts of the γ subunit. Several studies have shown that the insertion of a chloro-



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plast-like regulatory domain including the two cysteines into the ATP synthase γ subunit of non-regulated organisms enables the enzyme to respond to redox change (13–15); removal of the cysteines from this added domain leads to complete lose of redox regulation (16–17). In a previous study, we reported that the capacity for redox regulation could be conferred to non-redox regulated bacterial F₁ by substitution of the central region (residues 92–239) of the γ subunit with the counterpart of the γ subunit of chloroplast F₁ (14). By using this chimeric protein, we successfully showed redox-regulation of rotation of the γ subunit in the complex, and that an oxidized γ subunit results in an increase in the frequency of long pauses which in turn leads to a decrease in ATPase activity (18).

To gain a better understanding of the redox regulation of γ subunit rotation, we introduced the redox-regulation segment of the higher plant ATP synthase into the γ subunit of cyanobacterial F₁ (19) (see Fig. 1*B*). The enhanced stability demonstrated by this new chimeric complex provides the chance to thoroughly characterize the change in rotational motion conferred by a change in the redox state of the γ subunit.

EXPERIMENTAL PROCEDURES

Materials—Maleimide-PEG₁₁-biotin was purchased from Thermoscientific Co. Ltd (Yokohama, Japan). Carboxylated polystyrene beads were from Polysciences. Other chemicals were of the highest grade commercially available.

Preparation of the Chimeric Complex-The redox sensitive chimera complex (termed as $\alpha_3\beta_3\gamma_{\rm redox}$)² was constructed using an expression plasmid for $\alpha_3\beta_3\gamma$ complex of *Thermosyn*echoccus elongatus BP-1 (19). The redox region of the γ subunit of CF $_1$ from Spinacia oleracea (CF $_1$ $\gamma_{187-210}$, IHTLLPLSPKGE-ICDINGKCVDA) was inserted into the corresponding position of the γ subunit of *T. elongatus* BP-1 (*T. elongatus* BP-1 $\gamma_{188-201}$, VQTLLPLDPQGLET) using the mega-primer method (20) with the following mutation primer: 5'-GGCGG-AAAATTTCATCATCGGCTGCGTCGACACATTTACCA-TTGATGTCGCAAATTTCTCCTTTGGGTGAGAGGGGG-AGCAGGG-3'. The $\alpha_3\beta_3\gamma_{redox}$ was then expressed and purified by Ni-NTA chromatography as described (19). During the purification steps, 2 mM DTT was added to prevent unexpected disulfide bond formation between the four cysteines introduced. Prior to the specific oxidation process as described, DTT was removed by gel filtration using SuperdexTM-200 column (GE Healthcare) equilibrated with 50 mM MOPS-KOH (pH 7.5), and 100 mM KCl. 4 μ M $\alpha_3 \beta_3 \gamma_{redox}$ was incubated with 40 μM thioredoxin-f, 200 μM GSSG, 2 mM GSH, 50 mM MOPS-KOH (pH 7.5), and 100 mM KCl at 25 °C for 10 min to form a disulfide bond between the two cysteines located within the redox region of the chimera complex. This procedure allowed specific biotinylation of Cys-112 and Cys-125 on the γ subunit for observation of $\alpha_3\beta_3\gamma_{\rm redox}$ rotation. Following removal of the oxidizing reagents, 9 μ M $\alpha_3\beta_3\gamma_{redox}$ was biotinylated with 5 mM maleimide-PEG₁₁-Biotin, 50 mM MOPS-KOH, pH 7.5, and 100 mм KCl at 25 °C for 20 min. Unreacted maleimide-PEG₁₁-biotin was removed from the solution by gel-filtration, and the resulting chimera complex was stored at -80 °C in 10% glycerol.

Confirmation of the Specific Biotinylation of Cys-112 and Cys-125 of the γ Subunit by Mass Spectrometry and SDS-PAGE-30 μ l of oxidized and reduced F₁ (0.5 mg/ml) complexes were precipitated with trichloroacetic acid (final 5%), washed with ice-cold acetone, and finally dissolved in 1 mM maleimide-PEG₁₁-biotin, 100 mM Tris-HCl, pH 7.5, and 1% (w/v) SDS. The protein samples were then analyzed by 12% non-reducing SDS-PAGE. Gels from the non-reducing SDS-PAGE analysis were stained with Coomassie Brilliant Blue R-250 and the γ subunit bands manually excised. The proteins in the gel were digested by "in-gel" cleavage (21) at 37 °C with 10 μ g/ml sequencing grade modified trypsin (Promega, Tokyo) in 50 mM NH₄HCO₃, pH 8.0. Gels were reduced with 10 mM DTT at 56 °C for 1 h and alkylated with 1% (w/v) ICH₂CONH₂ in 50 mM NH₄HCO₃, pH 8.0 at room temperature for 45 min. Peptides were extracted from the gel with 5% (w/v) trifluoroacetic acid and 50% (v/v) acetonitrile solution. All digests were analyzed in a MALDI-TOF mass spectrometer (AXIMA-CFR Plus, Shimazu, Kyoto) using α -cyano-hydroxy-*trans*-cinnamic acid as the matrix. The instrument was calibrated with angiotensin II (m/z values, 1046.542) and adrenocorticotropic hormone fragment 18–39 (m/z value, 2465.199). The MS data obtained were analyzed with the data set of expected digested products of the γ subunit with allowance of missed cleavages of 3 and with carbamidomethyl as a variable modification.

ATP Hydrolysis Activity Assay—ATP hydrolysis activity was measured by an ATP regenerating method as described (22). The reaction was initiated by adding 20 nM (final) of enzyme solution into the 1.2 ml of the assay solution. The assay solution contained 50 mM MOPS-KOH (pH 7.5), 100 mM KCl, 2 mM MgCl₂, 200 μ M, or 2 mM MgATP, 0.2 mM NADH, 2 mM phosphoenolpyruvate, 50 μ g/ml pyruvate kinase, and 50 μ g/ml lactate dehydrogenase. The rates of ATP hydrolysis were measured at the steady state phase, 200 s after addition of the enzyme.

Rotation Assay—The rotation assay was performed as described (22) with some modifications. Previously oxidized or reduced $\alpha_3\beta_3\gamma_{redox}$ complex was fixed on the glass surface and then the streptavidin-coated duplex beads ($\phi = 291$ nm) attached to the immobilized $\alpha_3\beta_3\gamma_{redox}$ complex. Rotation of the complex was initiated by addition of 60 μ l of 50 mM MOPS-KOH (pH 7.5), 100 mM KCl, 2 mM MgCl₂, 200 μ M MgATP, 2 mM phosphoenolpyruvate, and 100 μ g/ml pyruvate kinase. To prevent re-oxidation of the complex during the measurement of the reduced complex, 0.5 μ M reduced thioredoxin-*f* and 5 μ M of DTT was added into the assay buffer. Rotation of the duplex beads attached to the immobilized $\alpha_3\beta_3\gamma_{redox}$ complex was observed by dark field microscopy at a frame rate of 200 frames/s, and the recorded data were analyzed by custom-made software (created by Yusung Kim).

Other Proteins—Recombinant thioredoxin-*f* of *S. oleracea* was expressed in *E. coli* and purified as described (23). Chloroplast F_1 -ATPase ($\alpha_3\beta_3\gamma\delta\epsilon$) was purified from fresh market spinach leaves (24) and the δ and ϵ subunits were removed by the following procedures. DEAE-Sephacel (12 ml, Amersham



² The abbreviations used are: $\alpha_3\beta_3\gamma_{redox'}$ redox sensitive chimera complex derived from the $\alpha_3\beta_3\gamma$ complex of *T. elongatus* BP-1; LDAO, lauryl dimethylamine-*N*-oxide.



FIGURE 1. Sequence of the γ subunits and the construction of the redox sensitive γ subunit. *A*, sequences around the redox region of the γ subunit are displayed. The redox region of 9 amino acids unique to the chloroplast γ subunit is shown in green and two cysteines involved in redox reaction in red. So-called insertion region in cyanobacteria (27) is shown in *blue*. The introduced region from *S. oleracea* for construction of $\alpha_3\beta_3\gamma_{redox}$ is *underlined*. *C. reinhardtii*; *Syn. Sp. PCC 6803*, *Synechococcus sp. PCC6803*; *Anaba*. *sp. PCC 7120*, *Anabaena sp. PCC 7120*. *B*, γ subunit (*black*) of *T. elongates* BP-1 ATP synthase was used as the basis for the chimera subunit. The redox region on the γ subunit of CF₁ from *S. oleracea* (*underlined*) was inserted into the corresponding position. To attach streptavidin-coated-beads via biotin-avidin interaction for single molecule experiments, two additional Cys-112 and Cys-125 (*purple*) were introduced in the appropriate position (19).

Biosciences) was previously equilibrated with 25 mM Tris-HCl (pH 8.0), 2 mM ATP, and 5 mM DTT. The crude extract dissolved in the same buffer was loaded on the column and the δ and ϵ subunits were eluted from the column with 25 mM Tris-HCl (pH 8.0), 2 mM ATP, 5 mM DTT, 30% glycerol, 20% ethanol, and 100 mM NaCl. The column was then washed with 25 mM Tris-HCl (pH 8.0) and 2 mM ATP. CF₁ ($\alpha_3\beta_3\gamma$) was eluted with 25 mM Tris-HCl (pH 8.0), 2 mM ATP. CF₁ ($\alpha_3\beta_3\gamma$) was eluted with 25 mM Tris-HCl (pH 8.0), 2 mM ATP. and 400 mM NaCl. The purified CF₁ was stored as ammonium sulfate precipitates at 4 °C. Before use, the preparation was desalted by gel-filtration (NAPTM-5 column, GE Healthcare) using 50 mM Tricine-KOH (pH 8.0) and 100 mM KCl.

RESULTS

*Redox Regulation of Cyanobacterial F*₁-ATPase—To investigate redox regulation of chloroplast-type F₁-ATPase, we first designed a new chimera F_1 complex, $\alpha_3\beta_3\gamma_{redox}$ complex, consisting of cyanobacterial F₁ subunits obtained from *T. elongatus* BP-1, and the redox regulatory region of the spinach $CF_1 \gamma$ subunit as the counterpart (see Fig. 1B). The biochemical properties of cyanobacterial F₁-ATPase are very close to that of higher plant chloroplasts, though the enzyme complex of the former lacks the specific redox regulatory region (see Fig. 1A) (19). By introduction of the 24 amino acid sequence of the spinach CF_1 - γ subunit, including the nine missing amino acid residues EICDINGKC, into the cyanobacterial counterpart, we successfully conferred redox regulatory capacity to the complex. After specific biotinylation of cysteines required for the probe fixation as described below, the activities of the oxidized enzyme and the reduced enzyme were measured. The ratio between the enzyme activity of the reduced form (Act_R) and that of the oxidized one $(Act_O) (Act_R / Act_O = 2.8)$ was similar to that of spinach CF_1 (Act_R/Act_O = 2.7). In addition, our new

chimeric enzyme was rapidly reduced by recombinant thioredoxin-*f* obtained from spinach chloroplasts (Fig. 2*A*), implying that the chimeric enzyme complex successfully mimics the redox regulatory system of CF₁. Compared with the previously synthesized chimeric F₁, $\alpha_3\beta_3\gamma_{\rm TCT}$ (14), the present chimeric complex, $\alpha_3\beta_3\gamma_{\rm redox}$, showed higher stability for purification and assay experiments.

Specific Biotinylation of the $\alpha_{3}\beta_{3}\gamma_{redox}$ Complex for Single Molecule Observations—The $\alpha_3\beta_3\gamma_{redox}$ complex has 4 cysteines on its γ subunit: two are originally located in the redox region and two were introduced to enable optical probe fixation. For single molecule experiments, two cysteines (Cys-112 and Cys-125) located on the γ subunit had to be biotinylated to attach them to the avidin-coated-plastic beads for observation of rotation with high angular resolution; the two cysteines (Cys-200 and Cys-206) located within the γ subunit redox region are expected to have remained unmodified for redox regulation. To carry out this specific modification, we first introduced a disulfide bond between Cys-200 and Cys-206 to protect these residues from biotin modification. This disulfide bond was successfully formed between these residues alone by using the oxidized form of thioredoxin-f and a glutathione pool as an electron acceptor (Fig. 3, *lane d*). The specific labeling of the desired cysteines was confirmed by MALDI TOF-Mass spectrometry following fragmentation of the γ subunit by trypsin (Fig. 4, A-C). Under the conditions in which two cysteines were biotinylated, two isotropic signals containing Cys-112 and Cys-125 were detected as biotinylated fragments (Fig. 4, A and B). The biotinylated fragment containing Cys-206 was only observed when four cysteines were biotinylated without oxidation (Fig. 4C). The γ subunit located within the complex remained redox sensitive after biotinylation of Cys-112 and Cys-125, and full reduction and oxidation of the complex was confirmed by SDS-





FIGURE 2. **Redox sensitivity of biotinylated** $\alpha_3\beta_3\gamma_{redox}$. After biotinylation of the $\alpha_3\beta_3\gamma_{redox}$ complex, the redox sensitivity was confirmed in both regulation of the activity (A) and disulfide bond formation ability by oxidation (B). A, activation by reduction of the γ subunit of $\alpha_3\beta_3\gamma_{redox}$. ATPase activity was measured using an ATP-regenerating system as a decrease in NADH absorption at 340 nm. A final concentration of 200 μ M ATP was used, and the reaction was initiated by adding oxidized $\alpha_3\beta_3\gamma_{redox}$. When the reaction reached the steady state (210 s from the start of reaction), a final concentration of 10 μ M DTT with (*black trace*) or without (*gray trace*) 5 μ M of reduced spinach thioredox. *f* was added. *B*, the disulfide bond formation ability of the biotinylated $\alpha_3\beta_3\gamma_{redox}$ complex was confirmed by further labeling with 4-acetamido-4'-maleimidylstilbene-2,2'-disulfonic acid following oxidation (a) or reduction (b) of the redox sensitive Cys-200 and Cys-206.



FIGURE 3. **Biotinylation of** $\alpha_3\beta_3\gamma_{redox}$. To prevent biotinylation of the cysteines involved in redox regulation (Cys-200, Cys-206), 4 μ M F₁ was oxidized with 40 μ M thioredoxin-*f*, 200 μ M GSSG (oxidized form of glutathione), 2 mM GSH (reduced form of glutathione), 50 mM MOPS-KOH, pH 7.5, and 100 mM KCI at 25 °C for 10 min. Reduced (c) and oxidized (d) F₁ was then biotinylated using maleimide-PEG₁₁-biotin (M.W.: 922.1, net mass addition: 921.5). Reduced (a) and oxidized (b) form F₁s without biotinylation are shown as controls.

PAGE following the thiol modification of Cys-200 and Cys-206 using 4-acetamido-4'-maleimidylstilbene-2,2'-disulfonic acid (Fig. 2*B*).

Effect of LDAO: Relief from ADP Inhibition—By reduction, both the $\alpha_3\beta_3\gamma_{redox}$ complex and the purified CF₁ showed



FIGURE 4. **Peptide analysis of the biotinylated** $\alpha_{3}\beta_{3}\gamma_{redox}$. A–C, mass spectrometry of fragmented γ subunit. The γ subunit separated by SDS-PAGE as shown in Fig. 3, was digested by trypsin in the gel and the resulting peptide fragments were obtained. The molecular weights of the fragments were measured by MALDI-TOF-MS. The fragments containing biotinylated (Bt) cysteines (A; Cys-112, B; Cys-125, and C; Cys-206) of the γ subunit were shown. Molecular weights of each fragment were 2091.7, 1837.0, and 2175.0, respectively. The non-cysteine-containing fragment (M.W. 2176.4) of γ is shown in C.

~2.7-fold activation, as mentioned. We then studied the extent of reduction induced ATPase activation under various assay conditions. As expected, this was affected by addition of lauryl dimethylamine-*N*-oxide (LDAO) (Fig. 5). LDAO is known to be effective in recovering F₁-ATPase from ADP inhibition (25); the $\alpha_3\beta_3\gamma_{redox}$ complex ATPase activity also increased remarkably when LDAO was added to both the reduced and oxidized forms of the complexes. However, the activity ratio between the reduced and oxidized form of the enzyme decreased remarkably depending on the concentration of LDAO, indicating that ADP-inhibition of the oxidized form is more pronounced than the reduced form (Fig. 5).

Redox Regulation of Rotation in the Presence of 200 μ M ATP— To characterize the substep responsible for CF₁ redox regulation during rotation, the rotational behavior of the $\alpha_3\beta_3\gamma_{redox}$



complex was analyzed by single molecule observation. For this purpose, polystyrene duplex beads with a diameter of ~291 nm were attached to the γ subunit of the $\alpha_3\beta_3\gamma_{redox}$ complex, and rotation of the γ subunit in the complex observed for 10–30 min by dark field microscopy. Fig. 6, *A* (oxidized enzyme) and *B* (reduced enzyme) show the typical time course of rotation of the γ subunit in the presence of 200 μ M ATP at room temperature. Torques calculated based on the fluctuation theorem as described (26) were 47.3 ± 6.2 pNnm (n = 3) and 48.6 ± 4.1 pNnm (n = 5), for the oxidized and reduced $\alpha_3\beta_3\gamma_{redox}$ complex, respectively. The maximal rotational speeds, determined from 100 continuous rotations, were 18.5 ± 2.8 rps (n = 19) for



FIGURE 5. Relationship between redox regulation of the complex and LDAO activation. The $\alpha_3\beta_3\gamma_{redox}$ complex was biotinylated in advance and ATP hydrolysis rates were measured using an ATP regenerating system (see "Experimental Procedures"), and the results obtained from three independent experiments were averaged. The ATP hydrolysis activity of the reduced $\alpha_3\beta_3\gamma_{redox}$ complex (open circle) and that of oxidized $\alpha_3\beta_3\gamma_{redox}$ (closed circle) are plotted against various concentrations of LDAO. LDAO was added 200 s after reaction initiation and the effect of LDAO on the activity was assessed for 50–100 s after addition of LDAO. The ratio of activities obtained from the data for the reduced $\alpha_3\beta_3\gamma_{redox}$ complex over those for the oxidized $\alpha_3\beta_3\gamma_{redox}$ complex is shown (triangle).

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oxidized complex and 19.5 \pm 4.2 rps (n = 17) for reduced one, respectively (Fig. 6, A and B), due to large bead friction, though the averaged rotation speeds during 600 s observation showed substantial difference; 2.7 \pm 2.0 rps (n = 16) for the oxidized complex and 7.0 \pm 2.6 rps (n = 10) for the reduced one, respectively. The pause duration was then analyzed using the pause length histogram, which was obtained from pauses longer than 1 s (Fig. 6, C and D). These relatively long pauses are unlikely to originate from the ATP binding state, because ATP binding takes in the order of submilliseconds in the presence of 200 μ M ATP according to the binding rate constant for ATP, k_{on} of $1.8 \times 10^7 \sim 1.9 \times 10^7$ M⁻¹ s⁻¹ (Fig. 7). For curve fitting to the histograms of pause durations, double exponential equations gave adequate results, implying that the observed rotation behavior contained at least two independent pauses. Consequently, the results gave two independent lifetimes (τ), the time constants of the marked events. The short lifetimes (τ_{sp}) for the oxidized and reduced states of the chimeric complex were 6.6 and 8.6 s, respectively. These short pauses were, however, not sufficiently reliable since the bin width (= 10 s) of these histograms was longer than the results obtained. Several studies have been conducted on ADP inhibition including analysis of the relevance between ADP inhibition and rotation of TF1 ATPase (7), and an exploration of the correlation between the inserted region of the γ subunit and ADP inhibition of cyanobacterial F₁ (27). Though the conclusions drawn from these reports were very similar, the origin of the short pause remains to be determined. In contrast, the long pauses with longer lifetimes ($\tau_{\rm lp}$) of oxidized and reduced state of $\alpha_3\beta_3\gamma_{redox}$ complex were significantly different, 124 and 78 s, respectively (Fig. 6, C and D). These long pauses can be interpreted as the ADP-inhibited state, which occurs at the 80° position (7) when the ATP binding position was settled as 0°. The long pause position during



FIGURE 6. **Single molecule analysis of rotation of the** γ **subunit under redox conditions.** *A* and *B*, the typical time courses of the revolutions of the oxidized (*A*) and the reduced (*B*) $\alpha_3\beta_3\gamma_{redox}$ complex. Rotation of the γ subunit was recorded as the movement of the 291-nm diameter beads attached to γ and the data obtained were analyzed using custom-made software. Rotation observation was performed at 25 °C in the presence of 200 μ M ATP. The histogram of pauses (>1 s) of the oxidized $\alpha_3\beta_3\gamma_{redox}$ complex (*C*) and the reduced $\alpha_3\beta_3\gamma_{redox}$ complex (*D*) were constructed from data for 27 and 20 molecules, respectively. In the *inset*, the vertical axis is drawn with a finer scale, and the bars for the data less than 20 s are omitted. The histograms are fitted with a double exponential equation, $N_{sp} \times \exp(-t/\tau_{sp}) + N_{ip} \times \exp(-t/\tau_{ip})$. The histograms of rotation duration between a pause and a next pause of the oxidized (*E*) and the reduced (*F*) $\alpha_3\beta_3\gamma_{redox}$ complex. Complexes were constructed from data for 28 and 20 molecules, respectively. The histograms were fitted with a single exponential equation, $N_{rot} \times \exp(-t/\tau_{rot})$.





FIGURE 7. **Rotation of** $\alpha_3\beta_3\gamma_{redox}$ **at 200 nm ATP.** *A*, typical time courses of rotation of the oxidized (*gray trace*) and reduced (*black trace*) $\alpha_3\beta_3\gamma_{redox}$. Average rotating speeds were 0.87 \pm 0.44 rps (n = 15) for oxidized and 0.81 \pm 0.32 rps (n = 7) for reduced $\alpha_3\beta_3\gamma_{redox}$. *Insets* represent the traces of centroids of the beads. *B*, the dwell time periods during rotation observed in the trace shown in *A* were collected and constructed into histograms. The data for each histogram are from 13 molecules for the oxidized complex (*gray*) and 6 molecules for the reduced one (*black*). Each histogram was fitted with a single exponential equation, Const. $\times \exp(-kt)$. The binding rate constants obtained ATP were $1.9 \times 10^7 \text{ m}^{-1} \text{ s}^{-1}$ for the oxidized complex and $1.8 \times 10^7 \text{ m}^{-1} \text{ s}^{-1}$ for the reduced complex.

rotation was then determined (Fig. 8) by using a medium exchange method from 0.2 μ M to 200 μ M ATP. Consequently, the pause position was determined as 79.3 \pm 12.4° (n = 22) ahead from the ATP binding position, which is close to the position for ADP inhibition. These pause duration analyses suggest that the oxidized $\alpha_3\beta_3\gamma_{redox}$ complex is more likely to be dropped into an ADP inhibited state. Data relating to the duration of rotation which appeared between the two pauses, were also collected and analyzed (Fig. 6, *E* and *F*). The histograms relating to the continuous rotating duration were fitted to a single exponential equation. The lifetime (τ_{rot}) for oxidized and reduced chimera were 29 and 36 s, respectively. Reduction of $\alpha_3\beta_3\gamma_{redox}$ is likely to result in a relatively longer continuous rotation.

Taken together, we conclude that the oxidized $\alpha_3\beta_3\gamma_{redox}$ complex remains in the ADP-inhibited state for longer, and takes longer to recover from the ADP-inhibited state than the reduced $\alpha_3\beta_3\gamma_{redox}$ complex. The difference in the lifetimes



FIGURE 8. **Stop angular position of long pauses.** *A*, after observation of rotation at 0.2 μ m ATP (*blue*), the ATP concentration was changed to 200 μ m ATP (*gray* and *black*). The long pauses (>40 s) are labeled on the traces as p1 to p4. *B* and *C*, the histograms of angular distribution of the beads are shown. *Blue bars* represent the angular distribution of 120° step rotation of the beads at 200 nm ATP. One of the three peak positions at 200 nm ATP is set as 0°. The *black bars* represent the angular distribution of long pauses observed in the data (p1 to p4) after buffer exchange.

between these two states eventually leads to a difference in the total active enzyme population size.

DISCUSSION

In a previous study we prepared a chmieric $\alpha_3\beta_3\gamma_{TCT}$ complex originating from the thermophilic bacteria Bacillus PS3 and spinach chloroplast, and reported that oxidized $\alpha_3\beta_3\gamma_{TCT}$ caused frequent long pauses (> 1 min) (14, 18). However, we could not assign the origin of these long pauses to one of the sub-steps, 80° and further 40° (28), because the significantly modified chimeric mutants had insufficient angular resolution to allow for identification of sub-steps. In this study, a new mutant complex ($\alpha_3\beta_3\gamma_{redox}$) consisting of two distinct benefits was prepared from the thermophilic cyanobacterial F_1 : the capacity to remain stable for extended observation periods, and the ability to achieve high angular resolution by connection of the γ subunit to an optical probe *via* two biotin bonds. Because cyanobacterial and chloroplast F1 share common ancestry, only 24 residues (Fig. 1, insertion of 9 chloroplast specific residues and 15 substitutions near the insertion; within them 8 residues are identical to cyanobacterial γ) were required to confer the redox regulatory capability to the complex. In addition, cysteines in the inserted redox region were reduced by their natural electron donor, thioredoxin-f (Fig. 2). A similar mutant (possessing the same insertion and 4 substitutions of the amino acids near the insertion) has already been prepared by Werner-Grüne et al. using direct transformation of Synechocystis 6803 cells (13). The mutant strain showed redox sensitivity to ATPase activity, as well as physiological differences in the dark and light (low O₂ consumption in the dark and high O₂ production in the light), though this redox switch conferred no growth advantage to the bacteria. Thus the redox regulatory property conferred to the cyanobacterial $\alpha_3 \beta_3 \gamma_{redox}$ must be close to that observed for $\mathsf{CF}_1.$ We therefore sought to determine whether the behavior of $\alpha_3\beta_3\gamma_{redox}$ at the single molecule level reflects the change in activity seen by redox regulation.

By using the newly designed chimeric complex $(\alpha_3\beta_3\gamma_{redox})$ in single molecule studies we successfully confirmed that oxidation of the redox region caused frequent long pauses (Fig. 6, *C* and *D*), which was consistent with a previous study (18). In addition, we observed that duration of rotation under oxidized conditions was relatively shorter than that shown under reduced conditions (Fig. 6, *E* and *F*). The redox-dependent behaviors of the chimeric complex are attributable to insertion of the redox region, since the wild-type complex lacking the redox region did not show any differences in activity and rotation behavior even when the complex had been previously reduced or oxidized (Fig. 9).

As shown in Fig. 8, the stop position of the long pause was determined as 80° ahead of the ATP binding position. This position is known to be where the sub-step of ATP hydrolysis (29) and phosphate release (30) occur; innate ADP inhibition also takes place at this position (7). Based on the results showing that relief from ADP inhibition by LDAO (Fig. 5) reduces redox sensitivity, we concluded that ADP inhibition was involved in the prolonged pauses during rotation, especially those observed under oxidized conditions.

The substrate binding rate at the catalytic sites was calculated by measuring the duration of ATP waiting dwell at 200 nM ATP, where F_1 spends most of its time waiting for ATP (see Fig. 7). When the enzyme activity is regulated by the redox regulatory system, the on-rate of ATP to the $\alpha_3\beta_3\gamma_{redox}$ complex was unchanged, but the ATP binding affinity ($\sim 1/K_m$) was slightly lower in the reduced state (Fig. 10). The difference in ATP affinity between the reduced and oxidized $\alpha_3\beta_3\gamma_{redox}$ complex is unlikely to be a significant element in redox regulation, since the difference in the activity ratio between both states was remarkable and even larger at higher concentrations of ATP (Fig. 10) where the difference of K_m is negligible.

In this study we showed that redox regulation by the redox switch regulates the frequency of ADP inhibition. Considering the position of the redox switch in relation to the inserted region of the γ subunit (Fig. 1*A*), we propose that the redox switch simply takes advantage of the function of the inserted region (27) by changing conformation via a disulfide bond, though as yet there is no crystal structure for the γ subunit of chloroplast F₁.



FIGURE 9. **ATPase activity and rotation of** $\alpha_3 \beta_3 \gamma_{wild}$ **complex under redox conditions.** *A*, ATPase activities of both oxidized (*open square*) and reduced (*closed square*) $\alpha_3 \beta_3 \gamma_{wild}$ complexes were measured (n = 3) in the presence of 200 μ M ATP with or without 0.3% LDAO. *B*, the typical rotational behaviors of oxidized and reduced $\alpha_3 \beta_3 \gamma_{wild}$ complex. Rotation of the complex was observed using 291 nm diameter beads at 200 μ M ATP. Maximum speeds of 10 continuous rotations for the oxidized and reduced complexes were 16.1 \pm 5.7 rps (n = 13) and 15.1 \pm 5.4 rps (n = 11), respectively. Averaged speeds of 10 min rotation for the oxidized and reduced complexes were 3.5 \pm 2.2 rps (n = 10) and 3.6 \pm 3.2 rps (n = 10), respectively.



FIGURE 10. **ATP hydrolysis kinetics of** $\alpha_3\beta_3\gamma_{redox}$. ATPase activities at various MgATP concentrations (*open circle* for reduced, *closed circle* for oxidized $\alpha_3\beta_3\gamma_{redox}$), and the activity ratio of the reduced $\alpha_3\beta_3\gamma_{redox}$ complex over the oxidized one were plotted (*triangle*). Each measurement was conducted under 2 mM of excess Mg²⁺. Activities were fitted with Michaelis-Menten equation. V_{max} values of the reduced and oxidized $\alpha_3\beta_3\gamma_{redox}$ complexes were 60 s⁻¹ and 23 s⁻¹, respectively and K_m s of the reduced and oxidized complexes 35 μ M and 22 μ M, respectively.



Redox Regulation of Rotation of CF₁-ATPase

As shown in this and our previous study (27), the evolutionary tuning of the γ subunit central region is related to ADP inhibition. However the active site on the $\alpha\beta$ surface and the expected position of the regulatory region of the γ subunit are located some distance apart, and no direct contact between these two regions has been reported to date. In our previous study using $\alpha_3 \beta_3 \gamma_{TCT}$, deletion of three negatively charged residues at the regulatory region caused the inverse regulation, higher activity at the oxidized state (31, 32), implying that the activity of the enzyme is very sensitive to small conformational changes in this region. One possible mechanism for the regulatory region is that the redox change of this region slides or results in a change in the angle between the N-terminal α -helix and the C-terminal α -helix of the γ subunit. Since these terminal helices of the y subunit consist of a coiled-coil structure and have direct contact with the inside of the $\alpha_3\beta_3$ ring (33), distortion between two helices could affect its reaction steps, in this case, the tightness of the ADP-inhibited state, or in other words, the probability of dropping in and reverting out of the ADP inhibition state. Moreover, the redox switch located next to the inserted region could regulate the level of distortion via disulfide bond.

In photosynthetic organisms, the ability to reduce the rate of wasteful hydrolysis activity in the dark confers a significant advantage. However, despite multiple dedicated studies, the physiological significance of this regulatory mechanism has to date not been clearly proven. Several experiments in which the redox regulatory property is abolished by removal of the relevant cysteines have been performed in vivo in Arabidopsis thaliana (16) and Chlamydomonas reinhardtii (17). In each of these studies, ATP synthase was converted to a redox insensitive enzyme resulting in increased ATPase activity (16), but growth was not significantly affected. Furthermore, introduction of the redox region into F₁ in Synechocystis had no effect on growth, though the mutant showed reduced oxygen consumption levels in the dark (13). Although to date there has been no strong evidence for viability of these mutants, this regulatory mechanism must play a physiologically important role in evolution, in particular providing a survival advantage under strongly oscillating (day and night) or fluctuating (e.g. irregular weather changes) light conditions. This is likely to be the case because the redox regulation may help minimize energy waste, with redox regulation becoming beneficial to chloroplast type ATP synthesis under insufficient energy conditions such as low light.

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