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REACTION MECHANISM OF F_1 -ATPASE

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GENERAL INTRODUCTION

In oxidative phosphorylation and photophosphorylation, exergonic oxidation-reduction reactions are coupled to the endergonic synthesis of ATP. The chemiosmotic hypothesis of Peter Mitchell($\underline{1}$) best describes the manner in which energy is conserved during ATP synthesis; that is, a transmembrane electrochemical gradient serves as a required intermediate in the following energy conversion processes (Reaction 1 & 2).

$$A_{red} + B_{ox} \xrightarrow{A_{ox}} A_{ox} + B_{red} + \Delta \mu H^{+} \qquad 1$$

$$ADP + P_{i} + \Delta \mu H^{+} \xrightarrow{ATP} + H_{2}O \qquad 2$$

The vectorial transport of protons across the membrane associated with electron transport gives rise to a electrochemical gradient, $\Delta \mu H^+$ (Reaction 1). The energy stored in this gradient is used by membrane-bound coupling factors, $F_1 - F_0$, to drive the net synthesis of ATP(Reaction 2). As this reaction is reversible, the proton gradient is also formed during the net hydrolysis of ATP(<u>1</u>).

The complex of coupling factors that catalyzes Reaction 2 has two empirically defined components. F_0 is an integral membrane complex of three distinct subunits, and has shown to play an essential role in the translocation of protons(2). F_1 can be detached from the membrane as a water soluble complex of five subunits, O_{c} to E(2). The enzyme, F_1 , contains catalytic site for ATP synthesis and hydrolysis. However, the major question still to be answered is, How does the F_1 - F_0 complex catalyze the synthesis of ATP? When physically separated from membrane, soluble F_1 is only capable of catalyzing the net hydrolysis of ATP. However, if we assume the mechanism of ATP synthesis is the

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reversal of that of ATP hydrolysis, we can elucidate the mechanism of ATP synthesis from the detailed analysis on the mechanism of ATP hydrolysis catalyzed by soluble F_1 . Thus, the reaction mechanism of F_1 -ATPase is the subject of this thesis.

I attempted to clarify the reaction mechanism of F_1 -ATPase using a fluorescent analog of ATP. Thus, 2'-(5-dimethylaminonaphthalene-1-sulfonyl) amino-2'-deoxyATP (DNS-ATP) was synthesized by Watanabe and coworkers(<u>3</u>). DNS-ATP has the following advantages for the present purpose. (1) The fluorescence intensity of DNS-ATP incneases markedly on its bunding to F_1 , and it reflects the environment of nucleotide binding site of F_1 . (2) DNS-ATP is hydrolyzed by F_1 very slowly, making kinetic measurements easier than with ATP. (3) DNS-AT³²P can be easily synthesized enzymatically.

In this thesis, following conclusions were drawn from the analysis of the reactions of DNS-ATP with F_1 (isolated from beef heart mitochondria) and EF_1 (isolated from <u>E. coli</u> plasma membrane): (1) F_1 contains two identical DNS-ATP binding sites. The affinity of the sites for DNS-ATP is very high, and the-DNS-ATP bound to this site is hydrolyzed according to the following reaction scheme:

DNS-ATP $E \longleftrightarrow DNS-ATP \Rightarrow E^{DNS-ATP} \Rightarrow E^{DNS-ADP} \Rightarrow E^{DNS-ADP} \Rightarrow E^{DNS-ADP} \Rightarrow E^{DNS-ADP} \Rightarrow E^{DNS-ADP} \Rightarrow E^{DNS-ADP} \Rightarrow E^{DNS-ADP}$ This scheme is quite resembles that of myosin ATPase. (2) High concentrations of nucleotide, such as ATP, accelerate the steps (III), (IV), and (V) in the above scheme. Thus, the low affinity regulatory site(s) is suggested to exist, besides the high affinity catalytic sites. (3) DNS-ATP binds to α subunit of EF_1 with very high affinity and not to β subunit. The chemical modification of

 β subunit of EF₁ with DCCD does not change the binding of DNS-ATP to EF₁ (steps (I) and (II)), and does not inhibit the single turnover of DNS-ATP cleavage (step (III)). However, this modifi= cation leads to the inhibition of accelerating effect of ATP on the DNS-nucleotide release from EF₁ (step (V)), and leads to the inhibition of EF₁-ATPase activity at steady state. Therefore, it is suggested that the high affinity catalytic site and low affinity regulatory site exist in the α subunit and β subunit, respectively.

- 1. Mitchell, P. (1977) Ann. Rev. Biochem. 46, 996-1005
- 2. Kagawa, Y. (1978) Biochim. Biophys. Acta 505, 45-93
- 3. Watanabe, T., Inoue, A., Tonomura, Y., Uesugi, S., & Ikehara, M. (1981) J. Biochem. 90, 957-965

ABBREVIATIONS

ATP	adenosinetriphosphate
ADP	adenosinediphosphate
AT ³² P	[χ ⁻³² P] ATP
DNS-ATP and DNS-ADP	2'-(5-dimethylaminonaphthalene-l-sulfonyl) amino- 2'-deoxyATP and -deoxyADP
DNS - AT ³² P	[γ- ³² P] DNS - ATP
СТР	cytidinetriphosphate
GTP	guanosinetriphosphate
ITP	inosinetriphosphate
NTP and NDP	nucleoside tri- and diphosphate
AMPPNP	adenylyl-5'-imidodiphosphate
F ₁	coupling factor 1 (soluble mitochondrial ATPase)
EF1	F ₁ of <u>E. coli</u>
EDTA	ethylenediaminetetraacetic acid
DCCD	dicyclohexylcarbodiimido
PEP	phosphoenolpyruvate
SDS	sodium dodecyl sulfate
TCA	trichloroacetic acid
Tris	tris(hydroxymethyl)aminomethane
P.i	inorganic phosphate
PPi	inorganic pyrophosphate

PART I.

Reaction Mechanism of the ATPase Activity of Mitochondrial F_1 , Studied by Using a Fluorescent ATP Analog, 2'-(5-Dimethylaminonaphthalene-1-Sulfonyl) Amino-2'-DeoxyATP: Its Striking Resemblance to That of Myosin ATPase.

SUMMARY

The reaction mechanism of beef heart mitochondrial F_1 -ATPase was studied by using a fluorescent ATP analog, 2'-(5-dimethylamino= naphthalene-1-sulfonyl)amino-2'-deoxyATP (DNS-ATP), as a substrate. The following results were obtained.

1. In the presence of Mg^{2+} , 2 mol of DNS-ATP bind to 1 mol of F_1 with an apparent dissociation constant of 0.44 μ M. Upon binding, the fluorescence emission spectrum of DNS-ATP shifted to shorter wavelengths, and the fluorescence intensity at 520 nm increased 4.8 fold. On the other hand, in the absence of Mg^{2+} , the apparent dissociation constant of F_1 -DNS-ATP was 16 μ M, and the extent of fluorescence enhancement was 1.3- to 1.9-fold.

2. The initial rate of fluorescence enhancement, v_f , upon addition of Mg²⁺ to a mixture of F₁ and DNS-ATP was given by the following equation:

$$v_{f} = \frac{V_{f}}{(1 + \frac{K_{f}}{[DNS-ATP]})}$$

where $V_f = 0.34 \text{ s}^{-1}$ and $K_f = 3.3 \,\mu\text{M}$. Following the fluorescence enhancement, TCA-P_i and free P_i were liberated consecutively. The fluorescence intensity maintained its enhanced level even after the free P_i liberation and, on addition of an excess amount of EDTA, decreased slowly to the original level From these findings, the following reaction mechanism for F_1 -DNS-ATPase was proposed: $E+DNS-ATP \rightleftharpoons E^{DNS-ATP} \rightarrow E^{DNS-ATP*} \rightleftharpoons E_{P}^{DNS-ADP*} \xrightarrow{} E^{DNS-ADP*} \xrightarrow{} E_{P_{i}}^{DNS-ADP*} \xrightarrow{} E_{DNS-ADP} \xrightarrow{} E_{D$

where an asterisk indicates the state of bound DNS-nucleotide with enhanced fluorescence, and $E_P^{DNS-ADP*}$ is a TCA-unstable reaction intermediate. This reaction mechanism resembles that of myosin ATPase. In accordance with the reaction mechanism and the stoichiometry of DNS-ATP binding, an initial burst of TCA-P_i liberation of 2 mol/mol F₁ was observed.

The addition of an excess of ATP to F_1 -DNS-nucleotide 3. in the presence of ${\rm Mg}^{2+}$ accelerated markedly the liberation of both TCA-P_i and free P_i. ATP also accelerated the fluorescence decrease of F₁-DNS nucleotide. Direct measurements of released free DNS-nucleotide indicated that the fluorescence decrease was due to the release of DNS-ADP from F₁. ADP, AMPPNP, ITP, and GTP accelerated the F1-DNS-ATPase reaction in a similar manner. On addition of these NTP, the release of DNS-ADP from F₁ proceeded in rapid and slow steps. CTP and PPi largely or completely failed to accelerate the conversion of ${ t E}^{ extsf{DNS-ATP}*}$ into $E_p^{DNS-ADP*}$. Furthermore, CTP and PP_i induced only the slow release of DNS-ADP. Therefore, we concluded that several nucleotides including ATP bind to the regulatory site(s) on F_1 to induce a conformational change at the catalytic sites and accelerate each of the following steps: $E^{DNS-ATP*} \rightleftharpoons$ $E_{P}^{DNS-ADP} \xrightarrow{*} E_{P} \xrightarrow{*} E_{P}$ E. On the other hand, PP_{i} and DNS-ADPCTP mainly accelerate the release of DNS-ADP from $\textsc{b}^{\textsc{DNS-ADP}*}$ by displacement. AMP did not affect the F_1 -DNS-ATPase reaction.

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4. The steady-state rate of F_1 -DNS-ATPase, v_0 , increased linearly with increase in the concentration of DNS-ATP and was 1 s⁻¹ at 200 μ M DNS-ATP exceeding than V_f (0.34 s⁻¹). The time course of ATP hydrolysis after addition of ATP to F_1 -DNS-nucleotide showed no lag phase which corresponds to the release of DNS-ADP. These two findings indicate that F_1 -NTPase can also occur through a reaction path in which $E_p^{\rm NDP}$ is not formed.

5. P_i enhanced the fluorescence of F_1 -DNS-nucleotide and inhibited the displacement of bound DNS-ADP with ATP or PP_i . The extent of fluorescence enhancement by P_i was maximal at 1 mol DNS-ATP/mol F_1 . This finding implies that the two DNS-ATP binding sites on F_1 are heterogenous at least with respect to the interaction with P_i .

INTRODUCTION

It is widely accepted that a mitochondrial F_1 - F_0 complex catalyzes the ATP synthesis in oxidative phosphorylation (for reviews see <u>1,2</u>). When isolated from membrane, F_1 catalyzes only the hydrolysis of ATP, although the reaction center of ATP synthesis is supposed to be remaining in the isolated enzyme.

Recently important information has been accumulated on the structure and function of each subunit in F_1 -ATPase [EC 3.6.1.3] from thermophilic bacteria (2) and <u>E. coli</u> (3) by use of a reconstitution technique. Furthermore, many features of nucleotide binding to F_1 were reported by Slater (4) and others (for review see <u>5</u>). Boyer and coworkers (<u>6</u>) investigated the reaction mechanism of F_1 -ATPase mainly utilizing the oxygen exchange reaction, and Slater and coworkers (<u>7</u>) analyzed kinetic properties of the presteady state of ATP hydrolysis. Several authors (<u>6-10</u>) suggested regulatory effects of ATP and ADP on the reaction rate, nucleotide binding, and P_1 binding to F_1 . Boyer and coworkers (<u>11</u>) have proposed a possible catalytic cooperativity among subunits in the reaction of F_1 -ATPase.

Boyer and coworkers (<u>11</u>) suggested that there was a similarity between F_1 -ATPase and myosin ATPase in their reaction mechanisms. In myosin ATPase, each reaction step is identified and conformational change of the enzyme induced by nucleotide binding is directly measured (<u>12-15</u>). On the other hand, details of the reaction mechanism of F_1 -ATPase are not known despite numerous efforts as mentioned above. Therefore, we attempted to clarify the reaction mechanism of F_1 -ATPase by using the methods developed

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in the research of myosin ATPase (<u>12-15</u>). One of them was to use a fluorescent analog of ATP, 2'-(5-dimethylaminonaphtalenel-sulfonyl) amino-2'-deoxyATP (DNS-ATP) which was synthesized by substituting the 2'-hydroxy group of ATP with an amino group and introducing a dimethyl amino naphtalene sulfonyl group (<u>16</u>). DNS-ATP has the following advantages serving the present purpose: (1) The fluorescence intensity of DNS-ATP increases markedly on its binding to F_1 , and the extent of enhancement depends on experimental conditions. (2) DNS-ATP is hydrolyzed by F_1 very slowly, enabling easier kinetic measurements than using ATP. (3) DNS-AT³²P can be easily synthesized enzymatically. <u>Materials</u> — DNS-ATP and DNS-ADP were synthesized as described previously (16). DNS-AT³²P was synthesized by the same enzymatic method as described by Glynn and Chapell for $AT^{32}P$ synthesis (<u>17</u>). Specific radioactivity of DNS-AT³²P was about 2 µCi/nmol. Adequate purity of DNS-ATP, DNS-ADP, and DNS-AT³²P was confirmed by paper electrophoresis run in 50 mM triethylamine bicarbonate at pH 7.6 (16).

Pyruvate kinase [EC 2.7.1.40] was prepared according to the method of Tietz and Ochoa (<u>18</u>). Luciferase [EC 1.13.12.7] was prepared from firefly lantern as described previously (19). ATP, ADP, and AMP were purchased from Khojin Ltd.(Tokyo). AMPPNP, ITP, GTP, PEP, and luciferin were purchased from Sigma Chemicals Co.(St. Louis). [³H]Glucose was purchased from New England Nuclear Co.(London). All other chemicals were of reagent grade purity.

<u>Purification of F_1 </u> Submitochondrial particles were prepared from beef heart mitochondria as described by Beyer (<u>20</u>). F_1 was extracted from submitochondrial particles with chloroform as described by Beechy <u>et al</u>. (21). After 30-60% ammonium sulfate fractionation, the concentrated extract was subjected to gel filtration on a Sephacryl S-300 Superfine column (3.2x75 cm) equilibrated with 100 mM Tris-HCl, 5mM EDTA, and 50% glycerol at pH 8.0 and 20°C. The eluted enzyme was concentrated by ultrafiltration on an Amicon Diaflo XM-100A membrane, and stored at 0°C. The protein concentration of purified F_1 was determined by the biuret method (<u>22</u>) with bovine serum

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albumin $(A_{279 \text{ nm}} = 0.667 \text{ cm}^2 \cdot \text{mg}^{-1})$ as a standard. The concentration of F₁ was calculated by using a molecular weight of 360,000 (<u>1</u>). The purified enzyme was subjected to SDS-gel electrophoresis on 10% polyacrylamide gel according to the method of Weber and Osborne (23).

The nucleotide content of purified F_1 was measured as follows: To 0.5 ml of F_1 solution at 22 mg/ml, 0.5 ml of 8% HClO₄ was added. After centrifugation at 2000 x g for 20 min, the supernatant was neutralized to pH 7.6 with 4 N KOH. Total nucleotide content was determined spectrophotometrically by using $\xi_{259 \text{ nm}} = 15.4 \text{ mM}^{-1} \text{cm}^{-1}$ for adenine nucleotides, since the UV absorption spectrum of the supernatant was the same as that for adenine nucleotides. The contents of ATP and ADP were determined by the firefly luciferase system, as described by Furukawa <u>et al.</u> (<u>19</u>), after convertion of ADP to ATP by 0.1 mg/ml pyruvate kinase and 4 mM PEP.

Fluorescence Measurement — Usually measurement solutions were prepared by addition of reagents in 5-20 µl portions to 2.0 ml of 50 mM Tris-HC1, pH 8.0 containing 2 mM EDTA. The measurements were performed at 30°C. To initiate a reaction, a reagent was stirred into a cuvette containing the rest of a reaction mixture with the aid of a magnetic stirrer (mixing time ≤ 0.2 s). Fluorescence intensity was measured on a Hitachi MPF-4 spectrofluorometer equipped with a thermostated cuvette hodler. The slit widths on the excitation and emission monochrometers were 6 and 8 nm, respectively. The fluorescence intensity at 520 nm of DNS-ATP and DNS ADP was measured with excitation at 340 nm.

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Partial Reaction of F_1 -DNS-ATPase — All the concentrations were expressed in terms of final concentrations, unless otherwise stated, throughout the text and figure legends. The composition of a reaction mixture was 5 mM MgCl₂, 2 mM EDTA, 50 mM Tris-HCl at pH 8.0 and appropriate concentrations of DNS-ATP and F_1 . After adding 0.1 ml of F_1 solution to 0.85 ml of a solution containing EDTA and DNS-AT³²P in Tris-HCl, it was incubated for 30 s at 30°C. The F_1 -DNS-ATPase reaction was started by addition of 0.05 ml of 100 mM MgCl₂. At appropriate intervals the reaction was stopped by addition of 1 ml of 10% TCA. After centrifugation, ${}^{32}P_1$ in the supernatant (TCA- ${}^{32}P_1$) was measured as described by Nakamura and Tonomura (24).

The concentration of total free ${}^{32}P$ (free ${}^{32}P_i$ + free DNS-ATP 32 P) was measured by a double membrane filtration method developed by Yamaguchi and Tonomura (25). By using a set of two membrane filters (upper Amicon Diaflo XM-100A membrane; lower Millipore filter, pore size 0.8 µm), this technique enabled the separation of free ³²P remaining on the Millipore filter from the F_1 -bound form on the Amicon membrane. The diameter of both membranes was 10 mm. The reaction mixture was the same as used in the TCA- ${}^{32}P_i$ measurement, except that 10 mM [${}^{3}H$]glucose was added to measure the volume of a filtrate. At appropriate times, 50 µl of the reaction mixture was withdrawn and applied on the upper membrane and filtrated by vacuum sucking for 4 s. A small volume (0.5-3 µl) of filtrate remained on the Millipore filter was dried, and the residues were solubilized in 1 ml of dimethyl formamide. The concentration of total free 32 P in the filtrate was determined from the ratio of radioactivities, ${}^{32}P/{}^{3}H$ (25).

The reaction mixture for measurement of the concentration of free DNS-nucleotide was the same as that for the TCA- ${}^{32}P_1$ measurement, except that 10 mM [3 H]glucose was added and DNS-ATP was used instead of DNS-AT ${}^{32}P$. The diameter of the membranes was 25 mm, and the volume of a reaction mixture applied on the membranes was 500 µl, leaving 10-30 µl on the Millipore filter after sucking. The dried Millipore filter was solubilized by 2 ml of dimethyl formamide.

The amount of free DNS-nucleotide in the filtrate was determined by measuring the fluorescence intensity at 530 nm after correcting the fluorescence due to Millipore filter solubilized by dimethyl formamide. The volume of the filtrate was determined from the radioactivity of [³H]glucose.

<u>ATPase Activity</u> — The ATPase activity of F_1 was measured in the presence of a pyruvate kinase system (0.1 mg/ml pyruvate kinase and 4 mM PEP) as a feeder, and the amount of P_i liberated was measured by the method of Fiske and Subbarow (26).

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RESULTS

<u>Properties of Purified F_1 </u> — The specific activity of purified F_1 measured in the presence of 5 mM ATP at pH 8.0 and 30 °C was 120-150 U/mg, which was similar to that already reported (7). When kept in 50% glycerol, F_1 was quite stable. At 0 °C the activity did not change at all for more than three months. As is widely accepted, purified F_1 was composed of 5 subunits (Fig. 1), and the intrinsic inhibitor protein which is usually seen between δ and ξ subunits in SDS-gel electrophoresis (27) was completely removed by gel filtration on Sephacryl S-300. The adenine nucleotide content calculated from ΔA at 259 nm was 2.4 mol/mol F_1 . The ATP content of the sample measured by the luciferin-luciferase method was 2.3 mol/mol F_1 . The ADP content was less than 0.3 mol/mol F_1 .

<u>Fluorescence Change of DNS-ATP on Its Binding to F_1 </u> — Figure 2 shows that the addition of 1.5 µM F_1 to 3.2 µM DNS-ATP in the presence of 2 mM EDTA increased the fluorescence intensity at 520 nm 24% from the level of free DNS-ATP, and that this increase consisted of an initial rapid change and the following slow one. This fluorescence increase was completely reversed by addition of 0.2 mM ATP, 0.2 mM ADP, 1 mM PP₁ or 1 mM P₁ (data not shown). On further addition of 5 mM MgCl₂ to the above mixture of F_1 and DNS-ATP, the fluorescence was markedly enhanced with a halftime of about 10 s, and reached a plateau within about 1 min. The Mg²⁺-dependent fluorescence enhancement was observed only in the presence of F_1 . Subsequent addition of P₁ slowly enhanced the fluorescence intensity as will be

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Fig. 1. SDS-gel electrophoretic pattern of purified F_1 -ATPase. The amount of the applied protein was 5 µg. The length is the migration distance from the origin. The gel was stained by Coomassie Brilliant Blue, and scanned at 550 nm at 0.05 mm slit width.



Fig. 2. Fluorescence intensity change of DNS-ATP on its binding to F_1 . To mixture containing 3.2 μ M DNS-ATP and 2 mM EDTA in 50 mM Tris-HCl, pH 8.0, following additions were made: a, 1.5 μ M F_1 ; b, 5 mM MgCl₂; c, 4 mM K-P_i; d, 0.2 mM ATP.

detailed later. During this series of reaction, fluorescence emission spectra were recorded at various stages as shown in Fig. 3. DNS-ATP in 2 mM EDTA had an emission peak at around 555 nm (trace a) (<u>16</u>), which shifted to 545 nm with a slight increase in fluorescence intensity by addition of F_1 (trace b). Addition of 5 mM MgCl₂ markedly enhanced the fluorescence the maximum shifting to 525 nm (trace c). Finally by addition of 2mM Pi, the fluorescence maximum shifted to 520 nm with an intensity increase (trace d). As reported previously (<u>12</u>), addition of dioxane to an aqueous solution of DNS-ATP enhanced its fluorescence and caused a blue shift reflecting a hydrophobic interaction. Therefore hydrophobicity of the DNS-ATP binding site in F_1 was suggested to be very high (see Table I in DISCUSSION).

When F_1 was added to DNS-ATP in the presence of 2 mM EDTA at pH 6.5 or 7.5 (Fig. 4, arrow a), the fluorescence intensity increased rapidly followed by a slow decrease (Fig. 4). The fluorescence intensities at 1 min after addition of F_1 were 1.8and 2.1-fold greater than those of free DNS-ATP at pH 7.5, and 6.5, respectively. At pH 8.0 this increase was 1.3-fold. The extent and time course of the fluorescence enhancement on addition of 5 mM MgCl₂ (arrow b) were almost the same at these pHs. All the following experiments were carried out at pH 8.0, where the fluorescence of DNS-ATP changed only slightly on addition of F_1 in the absence of Mg²⁺.

The stoichiometry of DNS-ATP binding to F_1 was determined by fluorometric titration of 0.75 or 1.5 uM F_1 with DNS-ATP in the presence of Mg²⁺ (Fig. 5.). The data were analyzed based on the quadratic equation:

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Fig. 3. Fluorescence emission spectra recorded during a course of reaction of DNS-ATP with F_1 . Spectra were recorded in the following order; curve a, 2.1 μ M DNS-ATP alone; curve b, 1 min after addition of 2.3 μ M F_1 ; curve c, 2 min after addition of 5 mM MgCl₂; curve d, 2 min after addition of 2 mM K-P₁. The excitation wavelength was 340 nm.





Fig. 4. pH Dependence of reaction of DNS-ATP with F_1 . The pH of a mixture containing 2.1 μ M DNS-ATP and 2 mM EDTA at 30°C was adjusted to 6.5 with 50 mM Tris-acetate (curve 1), 7.5 with 50 mM Tris-HC1 (curve 2) and 8.0 with 50 mM Tris-HC1 (curve 3). Following additions were made; a, 1.5 μ M F_1 ; b, 5 mM MgCl₂; c, 50 μ M K-P₁.



Fig. 5. Fluorometric titration of F_1 by DNS-ATP in the presence of MgCl₂. The extent of fluorescence enhancement 2 min after addition of 5 mM MgCl₂ was plotted against the amount of DNS-ATP added. The reaction mixture contained 0.75 (•) or 1.5 μ M (O) F_1 in 2 mM EDTA, 50 mM Tris-HCl at pH 8.0. The solid lines are the theoretical curves (see text). The upper is for 1.5 μ M F_1 , and the lower for 0.75 μ M F_1 .

 $\Delta F = \frac{R}{2} \left\{ (nE_0 + S_0 + \phi_{DNS-ATP}) - [(nE_0 + S_0 + \phi_{DNS-ATP})^2 - 4nE_0S_0]^{1/2} \right\} (1)$ where ΔF is the extent of fluorescence enhancement, and is assumed to be proportional to the amount of the F_1 -DNS-nucleotide. R is the fluorescence parameter dependent upon experimental conditions, n is the number of independent and equivalent binding sites of DNS-ATP on F_1 , E_0 and S_0 are the total concentrations of F_1 and DNS-ATP, respectively, and $\phi_{DNS-ATP}$ is the dissociation constant for the complex. It was also postulated that the fluorescence intensity of DNS-ATP during the course of interaction did not change. The curves in Fig. 5 are the best fits of the data to Eqn. 1 with n = 2 and $\phi_{DNS-ATP} = 0.44 \ \mu$ M. With n values other than 2, the fit was poor.

When PP_i had been added to the F_1 -DNS-ATP system in advance, the extent of fluorescence enhancement on addition of Mg²⁺ decreased with increasing concentration of PP_i (Fig. 6). The results were explained by a competitive binding of PP_i to the DNS-ATP binding site on F_1 :

$$\phi'_{\text{DNS-ATP}} = \phi_{\text{DNS-ATP}} \left(1 + \frac{[PP_i]}{\phi_{PP_i}}\right)$$
(2)

where $\phi'_{\text{DNS-ATP}}$ is the apparent dissociation constant for F_1 -DNS-nucleotide in the presence of PP_i and ϕ_{PP_i} is the dissociation constant for F_1 -PP_i. The data in Fig. 6 were analyzed based on Eqn. 1, where $\phi_{\text{DNS-ATP}}$ was replaced by $\phi'_{\text{DNS-ATP}}$ of Eqn. 2. The curve in Fig. 6 is the best fit to the data obtained with $\phi_{\text{PP}_i} = 39 \ \mu\text{M}$.

DNS-ADP was also bound to F_1 and brought about similar fluorescence changes as DNS-ATP. In Fig. 7, the extents of

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Fig. 6. Competitive binding of PP_i to the DNS-ATP binding site on F_1 . The extent of fluorescence enhancement of DNS-ATP after addition of 5 mM MgCl₂ was plotted against the concentration of PP_i . The reaction mixture contained 3.2 μ M DNS-ATP and 0.75 μ M F_1 . The solid line is the theoretical curve (see text).



Fig. 7. Fluorometric titration of F_1 with DNS-ADP in the presence of MgCl₂. The extent of fluorescence enhancement 2 min after addition of 5 mM MgCl₂ was plotted against the amount of DNS-ADP added. The reaction mixture contained 1.5 μ M F_1 . The solid line is the teoretical curve (see text).

fluorescence enhancement on addition of 5 mM MgCl₂ to mixtures containing 1.5 μ M F₁ and various amounts of DNS-ADP were plotted against the amounts of DNS-ADP added. The theoretical curve was obtained by using Eqn. 1, where parameters for DNS-ATP were taken as those for DNS-ADP, with n = 2 and $\phi_{\text{DNS-ADP}} = 0.4 \ \mu$ M. However, the measured values deviated somewhat from the curve.

Reaction Sequence of F_1 -DNS-ATPase — We measured the initial rates of fluorescence enhancement, v_f , on addition of 5 mM MgCl₂ to mixtures containing 1.5 μ M F_1 and various concentrations of DNS-ATP. The double reciprocal plot of v_f against the concentration of DNS-ATP gave a straight line as shown in Fig. 8. This result may be explained by the following reaction scheme, $E + DNS-ATP \xrightarrow[K_f]{K_f} E \cdot DNS-ATP \xrightarrow[V_f]{K_f} E \cdot DNS-ATP^*.$

By assuming that the first step is in rapid equilibrium (see DISCUSSION), we obtain

$$v_{f} = \frac{V_{f}}{1 + \frac{K_{f}}{[DNS-ATP]}}$$
(3)

where K_f is the dissociation constant of E·DNS-ATP, V_f is the rate constant of the fluorescence enhancement, and an asterisk indicates a complex with an enhanced fluorescence intensity. The values of K_f and V_f were estimated as 3.3 μ M and 0.34 s⁻¹, respectively, from the results in Fig. 8.

In Fig. 9, the initial phase of fluorescence enhancement on addition of 5 mM MgCl₂ to a mixture of 4.28 uM DNS-AT³²P and 6.78 uM F₁ is compared with the TCA-³²P₁ liberation

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Fig. 8. Double reciprocal plot of the initial rate of fluorescence enhancement against DNS-ATP concentration. The reciprocal of the initial rate of fluorescence enhancement of DNS-ATP on addition of 5 mM MgCl₂, v_f , was plotted against the reciprocal of the concentration of DNS-ATP added. The concentration of F₁ was 1.5 μ M.



Fig. 9. Time course of fluorescence enhancement, $TCA^{-32}P_i$ liberation and total free ${}^{32}P$ liberation in the initial phase of the reaction of DNS-AT³²P with F_1 . The reaction mixture contained 4.28 µM DNS-AT³²P, 6.78 µM F_1 , and 10 mM [3 H]glucose in 2 mM EDTA and 50 mM Tris-HCl at pH 8.0. The reaction was started by addition of 5 mM MgCl₂, and stopped by addition of 5% TCA (TCA- ${}^{32}P_i$) or filtration through an XM-100A membrane (total free ${}^{32}P$) at the indicated time. The time course of fluorescence enhancement was measured separately under the same conditions.

and release of total 32 P into the filtrate of the reaction mixture (see MATERIALS and METHODS). The percentages of DNS-AT³²P bound to F_1 immediately after addition of Mg²⁺ and at the steady state were calculated by using $K_f = 3.3 \ \mu M$ [Eqn. (3)] and $\phi_{\text{DNS-ATP}}$ = 0.44 μ M [Eqn. (1)] to be 73% and 95%, respectively. Even if the initial concentration of free DNS-AT 32 P is 27%, this is expected to decrease rapidly corresponding to an initial rapid increase in the fluorescence intensity. Therefore, we concluded that the amount of total ³²P in the filtrate was almost equal to or slightly higher than the amount of free ${}^{32}P_{i}$ released from F_1 to the medium. Thus, after the addition of 5 mM MgCl₂ the fluorescence enhancement, liberation of TCA- 32 P; due to hydrolysis of DNS-ATP, and liberation of free $^{32}P_i$ seem to occur sequentially in this order. Since the fluorescence intensity maintained its enhanced level for most of the reaction, it is clear that DNS-ADP formed by hydrolysis of DNS-AT 32 P was still bound to F₁ and that the release of DNS-ADP from F_1 was very slow.

Figure 10A shows the time courses of $TCA-{}^{32}P_i$ liberation after addition of 2.83, 5.65 or 11.3 mol DNS-AT ${}^{32}P/mol F_1$ to 0.75 μ M F₁. In Fig. 10B, the logarithm of the remaining fraction of DNS-AT ${}^{32}P$,([total ${}^{32}P$] - [TCA- ${}^{32}P_i$])/[total ${}^{32}P$], was plotted against time after addition of 5 mM MgCl₂. At all DNS-AT ${}^{32}P$ concentrations examined, the time course of TCA- ${}^{32}P_i$ liberation deviated from a single exponential curve, exhibiting rapid and subsequent slow phases. The amount of TCA- ${}^{\bar{32}}P_i$ liberated during the rapid phase was estimated by extrapolating the second



Fig. 10. Initial phase of TCA- ${}^{32}P_1$ liberation from DNS-AT ${}^{32}P$ by F_1 . A: The reaction mixture contained 2.83 (**a**), 5.65 (O) or 11.3 (X) mol DNS-AT ${}^{32}P$ /mol F_1 and 0.75 μ M F_1 . The reaction was started by addition of 5 mM MgCl₂. B: The data in A were plotted as the logarithm of the fraction of remaining amount of DNS-AT ${}^{32}P$ versus reaction time. The P_i -burst size was obtained by extrapolating the linear portion of a slow change to time 0 (broken line).



REACTION TIME

Fig. 11. Fluorescence decay on addition of EDTA to F_1 -DNS-nucleotide. The reaction mixture contained 2.12 μ M DNS-ATP in 2 mM EDTA, 50 mM Tris-HCl at pH 8.0. Following additions were made; a, 1.36 μ M F_1 ; b, 4 mM MgCl₂; c, 7 mM EDTA.

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Fig. 12. Time course of change in fluorescence on reaction of DNS-ATP with F_1 in the presence of CaCl₂. The reaction mixture contained 1.1 μ M DNS-ATP and 1.1 μ M F_1 . The reaction was started by addition of 4 mM CaCl₂.



Fig. 13. Acceleration by ATP of TCA- ${}^{32}P_1$ liberation from the F_1 -DNS-AT ${}^{32}P$ system. The reaction mixture contained 4.28 μ M DNS-AT ${}^{32}P$ and 6.78 μ M F_1 . The reaction was started by addition of 5 mM MgCl₂. At the arrow (2 min) 0.2 mM ATP was added. The broken line shows the level of 4.28 μ M.



Fig. 14. Acceleration by various phosphate compounds of $TCA^{-32}P_i$ liberation from the F_1 -DNS-AT³²P system. The reaction mixture contained 1.8 µM DNS-AT³²P and 1.22 µM F_1 . The reaction was started by addition of 5 mM MgCl₂. At the arrow a (15 s) 0.2 mM ATP (\blacktriangle) was added. At the arrow b (30 s) 0.2 mM ATP (\bigstar), 0.2 mM AMPPNP (\blacksquare), 0.2 mM ADP (\diamondsuit) or 1 mM PP_i (\asymp) was added. The control (\bigcirc) is also indicated.
calculated by using $\phi_{\text{DNS-ATP}} = 0.44 \ \mu\text{M}$. The rate of TCA- $^{32}P_i$ liberation also increased by addition of 0.2 mM ADP or 0.2 mM AMPPNP but to a lesser extent. Addition of 0.2 mM PP_i showed no effect. The results in Figs. 13-15 clearly demonstrate that DNS-AT³²P bound to F₁ is hydrolyzed very rapidly on addition of 0.2 mM ATP, and that $[\gamma^{-32}P]$ phosphoryl group is rapidly released as free $^{32}P_i$.

Acceleration of DNS-ADP Release from F_1 -DNS-Nucleotide Complex by Addition of High Concentration of ATP — A decrease in fluorescence intensity and release of DNS-nucleotide occurred concomitantly on addition of 0.2 mM ATP to a mixture of 8.6 μ M DNS-ATP, 9.2 μ M F_1 , and 5 mM Mg²⁺, which had been incubated for 2 min (Fig. 16). The fluorescence intensity decreased rapidly to about 50% of the original level within the response time of

apparatus (≤ 0.5 s), followed by a slow decrease approaching the level of free DNS-ATP. Since F_1 -DNS-ATP was readily converted into F_1 -DNS-ADP + P_i by addition of ATP as mentioned above, the observed changes correspond to the release of DNS-ADP from F_1 -DNS-ADP.

When increasing concentrations of ATP were added to F_1 -DNS-nucleotide, the extent of the first rapid decrease of the fluorescence increased with an increase in the rate of slow change (Fig. 17), but did not reach 100% even on addition of 0.2 mM ATP.

Acceleration of DNS-ADP Release from F_1 -DNS-Nucleotide <u>Complex by Addition of Various Phosphate Compounds</u> — ATP, ADP, AMPPNP, ITP, and GTP, 0.2 mM each, added to F_1 -DNS-nucleotide induced a fluorescence decrease consisting of rapid and slow

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Fig. 15. Acceleration by ATP of free 32 P liberation from the F_1 -DNS-AT 32 P system. The reaction was carried out under the identical conditions as described in the legend to Fig. 13 except that the reaction was stopped by filtration through an XM-100A membrane.



Fig. 16. Comparison of the time course of a decrease in fluorescence with that of release of DNS-nucleotide on addition of ATP to F_1 -DNS-ATP. A mixture containing 8.6 µM DNS-ATP, 9.2 µM F_1 , and 10 mM [³H]glucose was incubated with 5 mM MgCl₂ for 90 s, and then 0.2 mM ATP was added. The reaction was stopped by filtration through an XM-100A membrane at the time indicated. The extent of a decrease in fluorescence intensity, $\triangle F1$, is plotted as relative to difference between the intensity before addition of 0.2 mM ATP (1.0) and that of free DNS-ADP (0).



Fig. 17. Fluorescence decay on addition of various concentrations of ATP to F_1 -DNS-nucleotide. The reaction mixture contained 1.1 μ M DNS-ATP and 1.1 μ M F_1 in 2 mM EDTA, 50 mM Tris-HCl at pH 8.0. At the arrow (2 min), following concentrations of ATP were added; 0.5 μ M (a), 1.0 μ M (b), 1.5 μ M (c), 2.0 μ M (d), 5.0 μ M (e), 10 μ M (f), 40 μ M (g), 80 μ M (h), 0.2 mM (i). phases (Fig. 18). CTP at 0.2 mM yielded only a slow phase, and its rate was of the same order of magnitude as the second slow phase observed with 0.2 mM ATP. At the same time, the rate of $TCA-{}^{32}P_{i}$ liberation from F_{1} -DNS-nucleotide (Fig. 19) was accelerated slightly.

On addition of 7 mM EDTA to F_1 -DNS-nucleotide in the presence 4 mM MgCl₂, the fluorescence intensity decreased very slowly as previously shown in Fig. 11 (Fig. 20). Further addition of 0.1 mM ATP or 0.1 mM ADP markedly increased the rate of the fluorescence decrease. The time course of the fluorescence decrease induced by 0.1 mM ADP followed simple kinetics, whereas that induced by 0.1 mM ATP showed both rapid and slow phases as in the presence of Mg²⁺ (Fig. 17).

As previously described, PP_i binds to F_1 competitively with DNS-ATP but with a much lower affinity than that of DNS-ATP. As shown in Fig. 21, the fluorescence intensity of F_1 -DNS-nucleotide decreased on addition of PP_i with a single slow phase, and the extent of the fluorescence change. increased with increasing concentration of PP_i . The rate of the fluorescence decrease in the presence of 10 mM PP_i was almost equal to that induced by 0.2 mM ATP in its slow phase. Furthermore, 1 mM PP_i did not accelerate TCA- $^{32}P_i$ liberation from F_1 -DNS-nucleotide (see Fig. 14). AMP at 0.2 mM had no effect on the fluorescence intensity of F_1 -DNS-nucleotide in the presence of Mg²⁺ (Fig. 22), did not change the rate of TCA- $^{32}P_i$ liberation from F_1 -DNS-nucleotide (Fig. 19).



Fig. 18. Fluorescence decay on addition of various nucleotides to F_1 -DNS-nucleotide. A mixture containing 2.1 µM DNS-ATP and 1.5 µM F_1 was incubated in the presence of 5 mM MgCl₂ for 2 min, and then (at the arrow) following additions were made. A: 0.2 mM ATP (a); 0.2 mM ADP (b); 0.2 mM AMPPNP (c). B: 0.2 mM ATP (a); 0.19 mM GTP (b); 0.22 mM ITP (c); 0.18 mM CTP (d).



Fig. 19. Effect of CTP and AMP on $TCA-{}^{32}P_1$ liberation from the F_1 -DNS-AT ${}^{32}P$ system. The reaction mixture contained 1.8 μ M DNS-AT ${}^{32}P$ and 1.2 μ M F_1 in 2 mM EDTA, 50 mM Tris-HCl at pH 8.0. The reaction was started by addition of 5 mM MgCl₂. At the arrow (30 s), 0.2 mM CTP (**\$\$**) or 0.2 mM AMP (\times) was added. The control (0) is also indicated.



REACTION TIME

Fig. 20. Fluorescence decay of F_1 -DNS-nucleotide on addition of ATP or ADP in the presence of EDTA. To a mixture containing 2 mM EDTA and 50 mM Tris-HCl at pH 8.0, following additions were made; a, 2.1 μ M DNS-ATP; b, 1.4 μ M F_1 ; c, 4 mM MgCl₂; d, 7 mM EDTA (final concentration); e, 0.1 mM ATP; f, 0.1 mM ADP.

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Fig. 21. Fluorescence decay of F_1 -DNS-nucleotide on addition of various concentrations of PP_1 . The reaction mixture contained 4.0 μ M DNS-ATP and 0.75 μ M F_1 . At the arrow (2 min), following concentrations of PP_1 were added; 0 (a), 4 μ M (b); 10 μ M (c); 40 μ M (d); 100 μ M (e); 400 μ M (f); 4 mM (g); 10 mM (h).



Fig. 22. Effect of AMP on fluorescence of F_1 -DNS-nucleotide. The reaction mixture contained 2.1 μ M DNS-ATP and 1.5 μ M F_1 . After addition of 5 mM MgCl₂, following additions were made; a, 0.2 mM AMP; b, 0.2 mM ATP.

Dependence of the Steady-State Rate of F_1 -DNS-ATPase on <u>Substrate Concentration</u>---As shown in Fig. 23, the steady-state rate, v_o , increased in proportion to the DNS-AT³²P concentration up to 200 µM, where the v_o value was 1.0 s⁻¹ accounting for about 0.5% of that of F_1 -ATPase reaction at 200 µM ATP (<u>cf</u>. Fig. 24). If we assume that the F_1 -DNS-ATPase reaction at the steady state follows a simple Michaelis-Menten kinetics, this result indicates that the values of K_m for DNS-ATP and V_{max} are much larger than 200 µM and 1 s⁻¹, respectively.

Comparison of Time Course of ATP Hydrolysis with That of DNS-ADP Release from F₁-DNS-Nucleotide Complex after Addition of ATP — DNS-ATP (6.42 μ M) was allowed to react with F₁ (0.3 μ M) in the presence of ${\rm Mg}^{2+}$, PEP, and pyruvate kinase for 90 s. It was confirmed that ADP but not DNS-ADP can be converted into NTP by pyruvate kinase $(\underline{16})$. Then 0.2 mM ATP was added, and both fluorescence decrease and TCA-P_i liberation were followed in parallel experiments. Under the condition used, almost all TCA-P_i was derived by ATP hydrolysis. At the moment of the ATP addition, more then 95% of F_1 added was estimated to have bound DNS-nucleotide on the basis of $\phi_{DNS-ATP} = 0.44 \ \mu M$. As shown in Fig. 24, the fluorescence intensity decreased in two phases. If we assume that ATP is mainly hydrolyzed at the same site(s) where DNS-nucleotides bind to F₁, the TCA-P_i liberation should have a lag phase corresponding to the release of DNS-ADP (about 3.5 min). However, no such lag phase was observed.

Effect of P_i on Fluorescence of F_1 -DNS-Nucleotide Complex — As already shown in Fig. 2, the fluorescence of F_1 -Mg²⁺-DNSnucleotide was further enhanced by addition of P_i . Figure 25A

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Fig. 23. Dependence on DNS-ATP concentration of the steady-state rate of F_1 -DNS-ATPase reaction. The reaction mixture contained 0.30 µM F_1 , various concentrations of DNS-AT³²P, 2.5 mM MgCl₂, and 50 mM Tris-HCl at pH 8.0 and 30°C. The reaction was started by addition of DNS-AT³²P.



Fig. 24. Comparison of the time course of ATP hydrolysis with that of the DNS-ADP release from F_1 -DNS-nucleotide after addition of ATP. A mixture containing 6.42 μ M DNS-ATP, 0.31 μ M F_1 , 0.48 mg/ml pyruvate kinase, and 4 mM PEP in 2 mM EDTA, 50 mM Tris-HCl at pH 8.0 was incubated with 5 mM MgCl₂ for 90 s, and 0.2 mM ATP was added. The reaction was stopped by addition of 5% TCA at the time indicated. The ti \pm course of the DNS-ADP release was measured separately as a fluorescence change under the same conditions (inset). The arrow in the inset shows the time of addition of 0.2 mM ATP.



Fig. 25. Fluorescence enhancement of F_1 -DNS-nucleotide on addition of P_i . The concentration of DNS-ATP added was 3.2 μ M. A: Following additions were made; a, 1.5 μ M F_1 ; b, 5 mM MgCl₂; c, K-P_i at concentrations of 0 (1), 40 μ M (2), 0.1 mM (3), 0.2 mM (4), 0.4 mM (5), 1 mM (6), 4 mM (7), 10 mM (8). B: The reciprocal of the extent of fluorescence enhancement at 2 min after K-P_i addition (Δ F1) was plotted against the reciprocal of the concentration of K-P_i added.

shows the time courses of the fluorescence enhancement of F_1 -DNS-nucleotide on addition of various concentrations of P_i at pH 8.0. With increasing concentrations of P_i , the extent of the fluorescence enhancement increased, and was saturated at about 4 mM P_i . Figure 25B shows a double reciprocal plot of the extent of the fluorescence enhancement at 2 min after P_i addition and the P_i concentration. A single straight line enabled an estimation of an apparent dissociation constant of P_i from F_1 -DNS-nucleotide (ϕ_{P_i}) as 195 µM. The affinity of F_1 -DNS-nucleotide for P_i increased with decreasing pH (Fig. 4). At pH 6.5 the fluorescence enhancement was saturated at least with 50 µM P_i (data not shown). The rate of DNS-AT³²P hydrolysis in the presence of 4.3 µM DNS-AT³²P, 1.7 µM F_1 , and 5 mM MgCl₂ was unaffected by the addition of 1 mM P_i (data not shown).

We also examined effects of various anions on the fluorescence intensity of F_1 -DNS-nucleotide (Fig. 26). These anions are known to accelerate the F_1 -ATPase activity (28). The fluorescence intensity was almost unaffected by 5 mM HCO₃⁻ or 2 mM maleate, and slightly increased on addition of 0.5 mM HSO₃⁻. The effect of HSO₃⁻ was maximal at 2 mM.

Binding of DNS-Nucleotide to F₁ in the Presence of P_i -

The fluorescence intensity of DNS-ATP after addition of F_1 , MgCl₂, P_i , and ATP, as added in this order, was measured in the presence of different concentrations of DNS-ATP (see Fig. 2), and the fluorescence intensities at each step were plotted against the DNS-ATP concentration (Fig. 27A). The extent of the fluorescence enhancement induced by P_i was replotted in

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Fig. 26. Fluorescence change of F_1 -DNS-nucleotide induced by various anions. The reaction mixture contained 2.1 μ M DNS-ATP and 1.5 μ M F_1 . After addition of 5 mM MgCl₂, following additions were made; a, 5 mM HCO₃⁻; b, 2 mM maleate; c, 0.5 mM HSO₃⁻; d, 2 mM HSO₃⁻.



Fig. 27. Fluorometric titration of F_1 by DNS-ATP in the presence of P_i . A: The fluorescence intensity was measured sequentially as follows; DNS-ATP alone (O); 1 min after addition of 1.5 μ M F_1 (\odot); 2 min after addition of 5 mM MgCl₂ (Δ); 2 min after addition of 8 mM K-P_i (Δ); the level decreased rapidly on addition of 0.2 mM ATP (\times). B: The extent of fluorescence enhancement after addition of 8 mM K-P_i (Δ minus Δ) was plotted against the concentration of DNS-ATP added.

Fig. 27B. With an equimolar concentration of DNS-ATP to F_1 (1.5 μ M), the enhancement was saturated.

Figure 28 shows the effect of 4 mM P_i on the release of DNS-nucleotide from F_1 induced by PP_i (A) or ATP (B). The fluorescence decrease on addition of 1 mM PP_i showed only a slow phase, as already mentioned (<u>cf</u>. Fig. 21), and its rate was remarkably reduced by addition of 4 mM P_i . P_i also inhibited the second slow change of the DNS-ADP release on addition of ATP. After the fluorescence intensity of F_1 -DNS-nucleotide was enhanced by addition of 4 mM P_i , 0.2 mM ATP was added. The fluorescence decreased in two phases as observed in the absence of P_i . However, the extent of the rapid phase was larger and the rate of the second slow phase of P_i .





Fig. 28. Time course of a decrease in fluorescence of F_1 -DNSnucleotide in the presence and absence of P_i on addition of PP_i or ATP. The concentration of DNS-ATP added was 3.2 μ M. A: Following additions were made; a, 1.5 μ M F_1 ; b, 5 mM MgCl₂; c, 1 mM PP_i, d, 4 mM K-P_i; e, 1 mM PP_i. B: Following additions were made; a, 1.5 μ M F_1 ; b, 5 mM MgCl₂; c, 0.2 mM ATP; d, 4 mM K-P_i; e, 0.2 mM ATP.

DISCUSSION

Recently, Slater <u>et al</u>. reported that purified F_1 contains 2 mol of unexchangeable ATP/mol F_1 and proposed that these ATP are indispensable in maintaining the conformation of F_1 (<u>4</u>). The amount of ATP tightly bound to our purified F_1 preparation was 2.3 mol/mol F_1 . In this paper, we assumed that these tightly bound ATP were not directly involved in the F_1 -DNS-ATPase reaction (<u>4</u>). Our purified F_1 preparation contained no intrinsic ATPase inhibitor protein (Fig. 1, <u>27</u>).

So far there are several reports that the nucleotide binding sites on energy-transducing ATPases form hydrophobic clefts (<u>29-31</u>). Upon binding to F₁, DNS-ATP showed a similar change in fluorescence emission spectrum as when it was brought into dioxane (see Fig. 3 and also Table I). This change was reversed by adding several nucleotides including ATP. Therefore, we concluded that the nucleotide binding site of F₁ was in a hydrophobic cleft. The present fluorometric titration studies of F₁ by DNS-ATP in the presence of Mg²⁺ indicated that 2 mol of DNS-ATP were bound to 1 mol of F₁ with an apparent dissociation constant of 0.44 µM (Fig. 5). This finding is consistent with the $\alpha'_2 \beta_2$ stoichiometry of F₁ subunits reported by Vershoor <u>et</u> <u>al.</u> (<u>32</u>). However, we cannot deny the possibility that our purified F₁ preparation was partially inactivated, so that the $\alpha'_3 \beta_3$ stoichiometry (<u>33</u>) cannot be ruled out.

Binding of various phosphate compounds to F_1 are affected markedly by divalent cations and pH (<u>34,35</u>). The properties of fluorescence emission spectra of various complexes of DNS-ATP

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with F₁ are summarized in Table I. The apparent dissociation constant of DNS-ATP bound to F_1 in the absence of Mg^{2+} was about 16 μM (Fig. 15), which was much higher than the ϕ value of 0.44 μ M in the presence of Mg²⁺ In Table I, the dioxane concentrations giving the same fluorescence emission peak and intensity at 520 nm that were obtained with free DNS-ATP and F₁-DNS-ATP in aqueous solution are also included. Thus, the hydrophobicity of environment around the DNS group in E.DNS-ATP and $E \cdot Mg^{2+} \cdot DNS$ -ATP was equivalent to that of 20-30 and 58-76% dioxane, respectively. It is also suggested that ${\rm Mg}^{2+}$ increases both the affinity of F_1 for DNS-nucleotide and the hydrophobicity of the DNS-nucleotide binding site on F_1 . A further increase in the extent of the fluorescence enhancement of DNS-ATP by reaction with F_1 in the absence of Mg²⁺ by lowering the pH (Fig. 4) suggests that the conformation around the nucleotide binding site on F_1 is altered by changing pH.

DNS-ADP was also bound to F_1 with a high affinity in the presence of Mg²⁺, but the titration curve showed some heterogeneity (Fig. 7). One possibility is that two molecules of DNS-ADP bound to F_1 are not equivalent in terms of fluorescence intensity even if there is no interaction between them.

In this study, various elementary steps in the F_1 -DNS-ATPase reaction were identified. The initial rate of fluorescence enhancement on addition of Mg²⁺ to F_1 increased with increasing concentration of DNS-ATP, following typical Michaelis-Menten kinetics (Fig. 8). Thus, we assumed the following reaction scheme to explain the formation of a complex of DNS-ATP and F_1 having enhanced fluorescence intensity (E^{DNS-ATP*}):

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Table I. Properties of fluorescence emission spectra of various complexes of DNS-ATP and F_1 . All the indicated values are for the bound forms of DNS-ATP except for free DNS-ATP.

Complex	Fluor. Max.	Fluor. Intensity at 520 nm (relative)	ø	ø Dioxane (v/v %) ^b		
	(nm)		(µM)	Position of Peak	Intensity at 520 nm	
Free DNS-ATP	555	1.0		0	0	
E.DNS-ATP ^a	545	1.3-1.9	16 ^a	20	30	
E·Mg ²⁺ ·DNS-ATP	525	4.8	0.44	58	76	
P _i ·E·Mg ²⁺ ·DNS-ATP	520	5.8		66	100	

- ^a The ϕ value was calculated from the data of Fig. 15. Fluorescence intensity at 520 nm was calculated from the data of Figs. 2 and 15.
- ^b Concentration of dioxane which induces the same change in fluorescence properties of free DNS-ATP as that induced by formation of complex with F_1 .

$$E + DNS-ATP \xrightarrow{K_f} E^{DNS-ATP} \xrightarrow{V_f} E^{DNS-ATP*}$$

On addition of Mg²⁺ to a mixture containing DNS-ATP and F₁ the enhancement of fluorescence intensity of DNS-ATP, liberation of TCA-³²P_i and free ³²P_i occurred consecutively (Fig. 9). Even after that the fluorescence intensity still maintained its enhanced level. All these findings are easily explained by the proposed reaction scheme shown in Fig. 29. $E_P^{DNS-ADP*}$ is a TCA-unstable intermediate, and its formation is measured as TCA-P_i. The rate of liberation of DNS-ADP from $E^{DNS-ADP*}$ is assumed to be very small, since the enhanced fluorescence level was maintained even after the release of free P_i.

The mechanism is supported by the following findings. (1) TCA-P_i burst of 2 mol/mol F₁ was observed with various concentrations of DNS-ATP (Fig. 10). The burst size agreed with the binding stoichiometry of 2 mol DNS-ATP/mol F₁ obtained from the fluorometric titration of F₁ (Fig. 5). (2) Boyer and coworkers (<u>6</u>) found ¹⁸0/P ratio of more than three for P_i formed during ATP hydrolysis by F₁ at low concentrations of ATP, suggesting a rapid equilibrium of $E^{ATP} \xrightarrow{} E^{ADP}_{P}$. Notably the present reaction scheme resembles strikingly that of myosin ATPase (<u>12-15</u>). This similarity was originally pointed out by Boyer and coworkers (<u>36</u>). When Ca²⁺ was used as a divalent cation instead of Mg²⁺, the release of DNS-ADP was accelerated (Fig. 12). This is also found with myosin ATPase.

There are several pieces of evidence to indicate the regulatory effect of ATP on ATP hydrolysis by F_1 (<u>6-10</u>). Boyer and coworkers (6) found that an increase in the ATP concentration

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ADDITION OF ATP



Fig. 29. The reaction mechanism of F_1 -DNS-ATPase. Although two identical catalytic sites are postulated on F_1 , only one of them is indicated here.

during ATP hydrolysis by F_1 decreased the extent of water oxygen incorporated into each P_i formed, as if ADP and P_i were released at one site when ATP was bound to another. In this study, we found that ATP accelerated markedly the formation of $E_P^{DNS-ADP*}$ from $E^{DNS-ATP*}$ (Fig. 13), release of P_i from $E_P^{DNS-ADP*}$ (Fig. 15), and also release of DNS-ADP from $E^{DNS-ADP*}$ (Fig. 16). This acceleration will be explained by a mechanism that a conformational change is induced by binding of ATP to regulatory site(s) which are different from the catalytic site(s) binding DNS-ATP.

When ATP was added to F_1 -DNS-nucleotide under conditions where most of the DNS-ATP was bound to F_1 , all the DNS-ATP was liberated rapidly as TCA-P_i (Fig. 13). However, when ATP was added at the initial phase of the F_1 -DNS-ATPase reaction, the TCA-P_i liberation was only slightly accelerated, and then stopped (Fig. 14). These findings indicate that the rate of the reverse reaction, $E^{DNS-ATP*} \rightarrow E^{DNS-ATP}$, is very small.

The rapid fluorescence decrease observed on addition of ATP, ADP, AMPPNP, ITP, or GTP to the reaction mixture containing DNS-ATP, F_1 , and Mg²⁺ (Fig. 18) is suggested to be derived by the release of DNS-ADP induced by a conformational change of the catalytic site on F_1 . The second slow phase will be due to the displacement of bound DNS-ADP by other nucleotides. ADP and AMPPNP also accelerated the TCA-P_i liberation, although the extent of the acceleration was smaller than that by ATP. Thus, the effects of nucleotides to accelerate the TCA-P_i liberation and to release DNS-ADP in the first rapid phase can be summarized as follows in the decreasing order of their efficiency:

ATP \geq ADP \approx ITP \geq GTP \approx AMPPNP

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CTP accelerated slightly the step, $E^{DNS-ATP*} \longrightarrow E_P^{DNS-ADP*}$ (Fig. 19), and induced only a slow release of DNS-ADP from F_1 (Fig. 18B). These findings suggest that the affinity of the regulatory site for CTP is very low or a conformational change of the catalytic site is not induced by the CTP binding to the regulatory site. AMP accelerated neither the TCA-P₁ liberation (Fig. 19) nor the DNS-ADP release (Fig. 22).

Recently, interactions of nucleotides with isolated subunits of F₁-ATPase from thermophilic bacteria were investigated by Kagawa and coworkers (37). They found that ATP were bound to of the latter. They also found that CTP was bound to $extsf{C}
extsf{S}$ subunit but not to β subunit, or that the conformational change of β subunit was not induced by CTP. Neither \swarrow nor β subunit bound AMP. If we assume that beef heart F_1 and thermophilic bacterial F_1 have essentially the same structure and function, our results strongly suggest that the catalytic site and the regulatory site exist on \bigwedge and β subunits, respectively. This propose is apparently contrast to a widely accepted view that the catalytic site and the non-catalytic nucleotide binding site of F_1 -ATPase are located on β and α subunits, respectively (38-42). However, it should be noted that this hypothesis was derived mainly by analyzing the effects of chemical modifications of F_1 only on the over-all reaction of F_1 -ATPase. But this discrepancy will be removed by postulating that the hydrolysis of NTP by F_1 occurs at different catalytic sites.

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If we assume that hydrolysis of DNS-ATP occurs only through the pathway <u>via</u> $E_P^{DNS-ADP^*}$ in the steady state, V_{max} has to be smaller than or equal to the maximum rate, V_f , for formation of $E^{DNS-ATP^*}$. However, V_f was 0.34 s⁻¹ (Fig. 8), which is much smaller than V_{max} (\gg 1 s⁻¹) (Fig. 23). Therefore, another pathway is also suggested for hydrolysis of DNS-ATP. Furthermore, when ATP was added under the conditions where most of F_1 was in the form of $E^{DNS-ADP^*}$, ATP hydrolysis occurred without a lag phase (Fig. 24). If ATP is mainly hydrolyzed <u>via</u> E^{ADP} , a lag phase corresponding to the release of DNS-ADP should be observed contrary to the present finding. Thus, it is highly probable that hydrolysis of NTP occurs <u>via</u> different catalytic pathways and at different catalytic sites in F_1 as is observed with myosin ATPase (12,13).

 E_p^{ADP} is widely accepted as a key intermediate in the energy-transducing process of myosin ATPase, and the present investigation clearly demonstrated that this type of intermediate is involved in the F_1 -ATPase reaction. Therefore, it is speculated that the reverse process of the present reaction mechanism functions in the ATP synthesis. Differential effects of inhibitors on ATP synthesis on hydrolysis have been interpreted by several workers (43,44) as showing that the catalytic route for ATP synthesis is separate from that involved in ATP hydrolysis, although this view has been disputed by Boyer (<u>11</u>) from several lines of consideration.

The fluorescence enhancement of F_1 -DNS-nucleotide induced by P_i in the presence of Mg²⁺ suggests that the environment of the nucleotide-binding sites on F_1 becomes more hydrophobic by

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the action of P_i , thus diminishing the rate of the DNS-ADP release (Table I, Fig. 28). An apparent dissociation constant of this complex at pH 8 was 195 μ M (Fig. 25) and this value decreased on lowering the pH. These results are in good agreement with those of Kasahara and Penefsky (<u>45</u>) who obtained the dissociation constant of 285 μ M. Another interesting feature of the P_i effect is the distinction of the two DNS-ATP binding sites on F_1 which otherwise seem to be equivalent; that is, the saturation of the fluorescence enhancement induced by $P_{\underline{i}}$ at 1 mol DNS-ATP/mol F_1 implies the heterogeneity of the two binding sites, at least with respect to the interaction with P_i .

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PART II.

Reactions of a Fluorescent ATP Analog, 2'-(5-Dimethylaminonaphthalene-l-Sulfonyl) Amino-2'-DeoxyATP, with <u>E. coli</u> F_1 -ATPase and Its Subunits: Evidence Suggesting the High Affinity Catalytic Site in (X Subunit and Low Affinity Regulatory Site in β Subunit.

SUMMARY

We performed kinetic studies on the reactions of fluorescent ATP analog, 2'-(5-dimethyl-aminonaphthalene-l-sulfonyl) amino-2'-deoxyATP (DNS-ATP), with <u>E</u>. <u>coli</u> F_1 -ATPase (EF_1) and its subunits, to clarify the role of each subunit in the ATPase reaction. The following results were obtained.

1. In the presence of ${\rm Mg}^{2+}$, 3 mol of DNS-ATP binds to 1 mol of ${\rm EF}_1$ with an apparent dissociation constant of 0.23 μ M. Upon binding, the fluorescence intensity of DNS-ATP at 520 nm increased exponentially with t_{1/2} of 35 s and reached 3.5 times the original fluorescence level. Following the fluorescence increase, DNS-ATP was hydrolyzed, and the fluorescence intensity maintained its enhanced level.

The addition of an excess of ATP over EF1-DNS-nucleotide complex 2. in the presence of ${Mg}^{2+}$ decreased the fluorescence intensity rapidly, indicating the acceleration of DNS-nucleotide release from EF1. ADP and GTP also decreased the fluorescence intensity. DCCD inhibited markedly the accelerating effect of ATP on DNS-3. nucleotide release from EF_1 and the EF_1 -ATPase activity at steady state. On the other hand, DCCD inhibited only slightly the fluorescence increase of DNS-ATP due to its binding to EF_1 and the rate of single turnover of DNS-ATP hydrolysis catalyzed by EF1. 4. In the presence of ${\rm Mg}^{2+}$, 0.65-0.82 mol of DNS-ATP binds to 1 mol of isolated \propto subunit of EF, with an apparent dissociation constant of 0.06-0.07 μ M. Upon binding, the fluorescence intensity of DNS-ATP at 520 nm increased 1.55 fold very rapidly $(t_{1/2} < 1 s)$. The fluorescence intensity of DNS-ATP was unaffected by the addition of the isolated $\boldsymbol{\beta}$ subunit.

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5. The kinetic properties of the fluorescence change of DNS-ATP's reaction with EF_1 -ATPase reconstituted from $\boldsymbol{\alpha}$, $\boldsymbol{\beta}$ and $\boldsymbol{\delta}$ subunits were quite similar to that of native EF_1 .

These findings strongly support the previous proposal that the high affinity catalytic site and low affinity regulatory site exist in the α and β subunit, respectively

INTRODUCTION

 F_1 -ATPase [EC 3.6.1.3] is composed of five subunits, designated \measuredangle through $\boldsymbol{\mathcal{E}}$ in order of decreasing molecular weight $(\underline{1},\underline{2})$. The stoichiometry of either $\alpha_3 \beta_3 \tau$ or $\alpha_2 \beta_2 \tau$ is proposed for the major subunits of $F_1(3-5)$. Recent reconstitution studies on F_1 's from thermophilic bacterium PS3 (TF_1) $(\underline{6}, \underline{7})$ and \underline{E} . <u>coli</u> (EF_1) $(\underline{8}, \underline{9})$ shed. light on the role of each subunit of F $_1$. The complex of \measuredangle with β and $\pmb{\tau}$ subunits possesses the ATPase activity, and the $\pmb{\delta}$ and $\pmb{\epsilon}$ subunits are essential for the binding of the lphaeta γ complex to the membranous component, F_o. The \varkappa and β subunits each have nucleotide-binding sites, which are different in their affinity and specificity for nucleotides (9,10), and the total number of nucleotide-binding site in F_1 is 4-7 (<u>11,12</u>). The different nucleotide-binding sites in F_1 have been revealed by steady-state kinetics of the ATPase reaction (13), binding measurements (11, 14), and the affinity labelling of nucleotide-binding sites (15-17). The F₁-ATPase activity was found to be inhibited by the modification of the $oldsymbol{eta}$ subunit which has a low affinity binding site for nucleotides (15,16,18-23). This finding suggests that the low affinity binding site in $m{eta}$ subunit is the calatytic site. This suggestion is supported by the finding that the steady-state F_1 -ATPase activity has a high K_m value for ATP $(\underline{13}, \underline{24})$.

Recently, we have synthesized a fluorescent ATP analog DNS-ATP, 2'-(5-dimethyl-aminonaphthalene-l-sulfonyl) amino-2'-deoxyATP (25), and studied the mechanism of the DNS-ATPase reaction of beef heart F_1 (26). We found that the reaction mechanism of F_1 -ATPase is very similar to that of myosin ATPase, and that the nucleotides at high

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concentrations accelerate the rates of three elementary steps in F_1 -ATPase (<u>26</u>). These findings were interpreted as showing the existence of low affinity regulatory site(s) as well as high affinity catalytic sites (<u>26</u>). Together with the results on the direct nucleotide binding measurements of the isolated \measuredangle and β subunit (<u>9,10</u>), this view can be extended to the subunit localization of the high affinity catalytic site in the \measuredangle subunit and the low affinity regulatory site in the β subunit. However, this view is quite opposite to the widely accepted hypothesis as mentioned above.

To determine whether the catalytic site exists in the \measuredangle or β subunit, we performed kinetic studies on the reactions of DNS-ATP with EF₁ and isolated active subunits. The results obtained strongly supported our hypothesis that the high affinity catalytic site and low affinity regulatory site exist in the \measuredangle and β subunits, respectively.
<u>Materials</u> — DNS-ATP, DNS-ADP and $[\delta^{-32}P]$ DNS-ATP (DNS-AT³²P) were synthesized as described previously (<u>25</u>). Pyruvate kinase [EC 2.7.1.40] was prepared according to the method of Tietz and Ochoa (<u>27</u>). ATP, ADP and AMP were purchased from Khojin Ltd. (Tokyo). AMPPNP, ITP, GTP, CTP and phosphoenolpyruvate were purchased from Sigma Chemicals Co. (St. Louis, MO). All other chemicals were of reagent grade purity.

Preparation of EF_1 , α and β Subunit, and Reconstituted EF_1 -<u>ATPase</u> — EF₁ was purified as described previously (<u>28</u>) from <u>E</u>. <u>coli</u> KY7485 (λ asn -5) after thermoinduction of the transducing phage carrying a set of structural genes for the proton translocating ATPase $(\underline{29})$. Purified EF₁ was precipitated by 55% ammonium sulfate and dissolved in 50mM Tris-acetate (pH 7.0) and 2mM EDTA, at a concentration of 5-10 mg/ml. The EF_1 solution was dialyzed against the same buffer solution at 20°C for 20 h before use. The \measuredangle , β and \overleftarrow{J} subunits were purified from EF_1 as described previously (9). Reconstituted EF_1 -ATPase was prepared from a mixture of the lpha , eta and $\overline{\sigma}$ subunits as described previously (8), and concentrated to 2 mg/ml by Diaflo XM-100A membrane purchased from Amicon Ltd. (Lexington, MA). The concentrated enzyme was dialyzed against the above buffer solution at 20°C for 20 h before use. The protein concentration was determined by biuret method (30) or by the method of Bradford (31) with bovine serum albumin as a standard. The molecular weights of EF1, lpha , etasubunits, and reconstituted EF₁-ATPase were taken as 376,000, 58,000, 52,000, and 364,000, respectively (9,28).

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<u>Fluorescence Measurements</u> — Measurement solutions were prepared by addition of reagents in 5-20 μ l portions to 2 ml of 2 mM EDTA, 100 mM NaCl and 50 mM Tirs-acetate at pH 7.0 and 30°C. The fluorescence intensity of DNS-ATP at 520 nm with excitation at 340 nm was measured as described previously (26).

<u>DNS-ATPase and ATPase Activity</u> — EF_1 at 0.87-1.6 μ M was allowed to react with 2.8 μ M DNS-AT³²P in 2 mM EDTA, 100 mM NaCl and 50 mM Tris-acetate at pH 7.0 and 30°C for 1 min. Then, the hydrolysis was started by the addition of 5 mM MgCl₂. The amount of ${}^{32}P_i$ liberated when the reaction was terminated by TCA (TCA- ${}^{32}P_i$) was measured as described previously (<u>26</u>). The steady-state ATPase activity of EF_1 was measured in the presence of an ATP-feeder system (0.1 mg/m1 pyruvate kinase and 5 mM phosphoenolpyruvate) at pH 8.0 and 30°C (<u>26</u>). The amount of TCA-P₁ liberated was measured by the method of Fiske and Subbarow (<u>32</u>).

Binding of DNS-AT³²P to \not{a} Subunit — To measure the binding of DNS-AT³²P to \not{a} subunit, the equilibrium dialysis was performed at room temperature using chambers consisting of two wells separated by a layer of dialysis membrane (9). One side of the chamber contained subunit and both wells contained 2 mM EDTA, 100 mM NaCl and 50 mM Tris-acetate at pH 7.0.

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RESULTS

Fluorescence Change of DNS-ATP on Its Binding to EF1 ---- Figure 1(A) shows a typical time course of change in the fluorescence intensity of DNS-ATP at 520 nm during its reaction with EF_1 in the presence of 3 mM free ${\rm Mg}^{2\, +}\,.$ The fluorescence intensity of 1.1 μM DNS-ATP in the presence of 2 mM EDTA increased only slightly on addition of 0.99 μ M EF₁ [arrow <u>a</u> in Fig. 1(A); compare curves <u>a</u> and <u>b</u> in Fig. 2]. On further addition of 5 mM MgCl₂ [arrow <u>b</u> in Fig. 1(A)], the fluorescence increased exponentially with a $t_{1/2}$ of 35 s, and reached a plateu level, which was 3.2 fold the fluorescence intensity of free DNS-ATP. Being observed only in the presence of EF1, the Mg^{2+} -dependent fluorescence increase of DNS-ATP reflects the binding of DNS-ATP to the nucleotide-binding sites in EF_1 (<u>cf</u>. DISCUSSION). On a further addition of 2 mM P_i [arrow <u>c</u> in Fig. 1(A)], the fluo= rescence intensity increased rapidly to the level of 5.1 fold that of free DNS-ATP. When 2 mM ATP was added to the reaction mixture [arrow d in Fig. 1(A)], the fluorescence intensity decreased rapidly to about 50% of the original level, then decreased slowly. When a small amount of DNS-ATP was added to EF_1 (<0.5 mol/mol EF_1), the fluorescence intensity decreased to a level almost identical to that of free DNS-ATP within 5 min after the addition of ATP. However, when a large amount of DNS-ATP was used, the second phase of the decrease occurred very slowly, and even 10 min after the addition of ATP, the fluorescence intensity was not the same as that of free DNS-ATP.

Figure 1(B) shows a time course of change in the fluorescence intensity of DNS-ATP in the presence of 3 mM Ca^{2+} . When 5 mM $CaCl_2$

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Fig. 1. Time course of fluorescence intensity change of DNS-ATP in the presence of EF_1 . To a mixture containing 1.1 µM DNS-ATP, 2 mM EDTA, 0.1 M NaCl and 50 mM Tris-acetate at pH 8.0 and 30°C, the following additions were made at time indicated by arrows (a-d). (A): a, 0.99 µM EF_1 ; b, 5 mM $MgCl_2$; c, 2 mM K-P_i; d, 2 mM ATP. (B): a, 0.94 µM EF_1 ; b, 5 mM $CaCl_2$; c, 2 mM K-P_i; d, 2 mM ATP. The emission and excitation wavelengths were 520 and 340 nm, respectively. was added to a reaction mixture containing 1.1 μ M DNS-ATP, 0.94 μ M EF₁ and 2 mM EDTA (b), the fluorescence intensity of DNS-ATP increased exponentially with almost the same value of t_{1/2} as that in the case of 5 mM MgCl₂ addition. The plateu level of fluorescence in the presence of Ca²⁺ was 2 fold that of free DNS-ATP. Further addition of 1 mM P_i (c) did not cause any fluorescence increase, but caused a slow decrease. On further addition of 2 mM ATP (d), the fluorescence intensity decreased rapidly by about 10%, then approached slowly to the level of free DNS-ATP.

Figure 2 shows the fluorescence emission spectra of free DNS-ATP and various complexes of DNS-ATP with EF_1 . As previously described (25), DNS-ATP had an emission peak at 555 nm (trace a). On addition of EF_1 , the emission peak shifted to around 550 nm with a slight increase in the fluorescence intensity (trace b). Addition of 5 mM MgCl₂ markedly increased the fluorescence intensity, and the maximum shifted to 530 nm (trace c). Further addition of 2 mM P₁ caused a further increase of the fluorescence and the shift of the maximum to around 520 nm (trace d). Addition of 2 mM ATP reversed these fluorescence changes almost backward to the state of free DNS-ATP (trace e). These fluorescence changes of DNS-ATP on its reactions with EF_1 are almost equal to those observed for beef heart F₁.

Stoichiometry of DNS-ATP Binding to EF_1 — The stoichiometry of DNS-ATP binding to EF_1 was determined by fluorometric titration of 1.1 μ M DNS-ATP with EF_1 in the presence of 3 mM free Mg²⁺ (Fig. 3). Several kinds of reaction intermediates are produced by the reaction of EF_1 with DNS-ATP, as will be mentioned in "DISCUSSION." However, we (<u>26</u>) previously found that in the case of beef heart F_1 , the fluorescence intensity of one intermediate is only slightly higher

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Fig. 2. Fluorescence emission spectra recorded during the course of reaction of DNS-ATP with EF_1 . Spectra were recorded in the following order; curve a, 1.1 μ M DNS-ATP alone; curve b, 1 min after addition of 1.6 μ M EF_1 ; curve c, 2 min after addition of 5 mM MgCl₂; curve d, 1 min after addition of 2 mM K-P_i; curve e, 1 min after addition of 2 mM ATP. Other conditions were the same as described in Fig. 1 or in the text.



Fig. 3. Fluorometric titration of DNS-ATP with EF_1 in the presence of MgCl₂. The extent of fluorescence increase 2 min after addition of 5 mM MgCl₂ was plotted against the amount of EF_1 added. The reaction mixture contained 1.1 μ M DNS-ATP in 2 mM EDTA, 0.1 M NaCl and 50 mM Tris-acetate at pH 7.0 and 30°C. The solid line is the theoretical curve (see the text).

than that of free DNS-ATP and the intensities of other intermediates are much higher than that of free DNS-ATP and are equal to each other. Therefore, the data were analyzed based on the quadratic equation:

$$\Delta F = \frac{\Delta F}{2} \left\{ (nE_0 + S_0 + \phi_{DNS-ATP}) - [(nE_0 + S_0 + \phi_{DNS-ATP})^2 - 4nE_0S_0]^{\frac{1}{2}} \right\} (Eq.1)$$

where ΔF is the extent of fluorescence increase, and is assumed to be proportional to the amount of EF_1 -DNS-nucleotide complex. ΔF_{max} is the maximum value of ΔF , and is obtained when all the DNS-ATP binds to EF_1 . Both the ΔF and ΔF_{max} values are expressed as p'ercentage of the fluorescence intensity of free DNS-ATP in the absence of Mg²⁺. E_0 and S_0 are the total concentrations of EF_1 and DNS-ATP, respectively. n is the number of independent and equivalent binding sites of DNS-ATP in EF_1 , and $\phi_{DNS-ATP}$ is their dissociation constant. The solid line in Fig. 3 is the best fit of the data to Eq. 1 with $\Delta F_{max} = 250$ %, n = 3, and $\phi_{DNS-ATP} = 0.23 \ \mu$ M. With n values other than 3, the fit was poor.

<u>Hydrolysis of DNS-ATP by EF_1 </u> — We found that in the absence-of free Mg²⁺, DNS-ATP is not hydrolyzed by EF_1 . Therefore, we measured the time course of TCA-³²P₁ liberation after addition of 5 mM MgCl₂ to a reaction mixture containing 2.8 μ M DNS-AT³²P, 1.6 μ M EF_1 and 2 mM EDTA (Fig. 4). The amount of TCA-³²P₁ liberated at 120 s was 0.47 mol/mol α . The amount of DNS-nucleotide bound to EF_1 under the conditions used was estimated to be 0.53 mol/mol α' by using n = 3 and $\phi_{\text{DNS-ATP}} = 0.23 \ \mu$ M. Under these conditions the fluorescence increase was found to be almost completed after 120 s of reaction. When ATP at 2 mM was added 60 s after starting the reaction (arrow in Fig. 4),

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Fig. 4. Initial phase of TCA-³²P₁ liberation from DNS-AT³²P catalyzed by EF₁ or purified α subunit. The reaction mixture contained 2.8 μ M DNS-AT³²P and 1.6 μ M EF₁ ($\bigcirc, \textcircled{0}$) or 2.3 μ M α subunit ($\triangle, \blacktriangle$) in 2 mM EDTA, 0.1 M NaCl and 50 mM Tris-acetate at pH 7.0 and 30°C. The reaction was started by the addition of 5 mM MgCl₂. At the time indicated by an arrow (60 s after addition of MgCl₂), 2 mM ATP was added ($\textcircled{e}, \bigstar$).

the TCA- ${}^{32}P_1$ liberation ceased immediately, showing no acceleration such as observed on beef heart F₁ (<u>26</u>).

Acceleration of DNS-Nucleotide Release from EF1 - Nucleotide Complex by Addition of Nucleotides and Pyrophosphate — The fluorescence intensity of EF1-DNS-nucleotide complex maintained its enhanced level after liberation of TCA- ${}^{32}P_i$. Because of a very small rate of release of DNS-ADP from the complex, it is clear that DNS-ADP formed by hydrolysis of DNS-AT 32 P was still bound to EF₁. In the presence of 3 mM Mg $^{2+}$, DNS-ATP at 1.1 μM was allowed to react with 0.98 μM EF $_1$ for 2 min. Then, various nucleotides (0.2 mM) or pyrophosphate (2 mM) were added, and the following decrease in the fluorescence intensity was measured (Fig. 5). The addition of ATP caused a fluorescence decrease with first rapid and second slow phases (trace a). In the case of beef heart F_1 , we found that the time course of the fluorescence decrease after addition of ATP was identical to that of the release of DNS-ADP from F_1 -DNS-nucleotide in both rapid and slow phases (26). Therefore, we assumed that the fluorescence decrease after addition of ATP corresponds to the release of DNS-nucleotide from EF_1 -DNS-nucleotide in the case of EF_1 as well. ADP (trace b) and GTP (trace c) also induced a fluorescence decrease, but the rates of the decrease were smaller than that induced by ATP. The rates of the fluorescence decrease on addition of AMPPNP (trace d), PP; (trace e), and CTP (trace f) became smaller in this order. AMP (trace g) had no effect on the fluorescence intensity of EF_1 -DNSnucleotide.

Effects of DCCD on the Reaction of DNS-ATP with EF_1 — Figure 6 shows the time course of the change in the fluorescence intensity of

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Fig. 5. Fluorescence decrease on addition of various phosphate compounds to EF_1 -DNS-nucleotide complex. DNS-ATP at 1.1 μ M was allowed to react with 0.98 μ M EF_1 in the presence of 3 mM Mg²⁺ for 2 min, then at the time indicated by an arrow, the following additions were made; a, 0.2 mM ATP; b, 0.2 mM ADP; c, 0.2 mM GTP; d, 0.2 mM AMPPNP; e, 2 mM PP_i; f, 0.2 mM CTP; g, 0.2 mM AMP. Other conditions were the same as described in Fig. 1.



Fig. 6. Inhibition by DCCD-treatment of decrease in the fluorescence intensity of EF_1 -DNS-nucleotide upon addition of ATP. After incubation of 0.98 μ M EF_1 without (A) or with (B) 0.2 mM DCCD in 2 mM EDTA, 100 mM NaCl and 50 mM Tris-acetate at pH 7.0 and 30°C for 20 min, the following additions were made at the time indicated by arrows (a-c) a, 1.1 μ M DNS-ATP; b, 5 mM MgCl₂; c, 2 mM ATP.

1.1 μ M DNS-ATP during its reaction with 0.98 μ M EF₁, which was preincubated with 0 (A) or 0.2 mM DCCD (B) at pH 7.0 and 30°C for 20 min. The time course of increase in the fluorescence intensity after addition of 5 mM MgCl₂ (b) was almost unaffected by the DCCDtreatment. On the other hand, the decrease of the fluorescence intensity upon subsequent addition of 2 mM ATP (c) was inhibited markedly by the DCCD-treatment (Fig. 6B).

Figure 7 shows the effects of DCCD on the steady-state rate of EF_1 -ATPase and the initial rate of EF_1 -DNS-ATPase. The steady-state rate of the Mg²⁺-ATPase reaction of EF_1 was inhibited almost completely by the DCCD-treatment [Fig. 7(A)], as already reported by Satre <u>et al</u>. (<u>21</u>). The initial rate of TCA-³²P₁ liberation in the presence of a low concentration of DNS-ATP was only slightly inhibited by the DCCD-treatment of EF_1 [Fig. 7(B)]. It must be noted that in both experiments EF_1 of the same concentration range was preincubated with DCCD (legend of Fig. 7).

Fluorescence Change of DNS-ATP on Its Binding to α Subunit of <u>EF</u>₁ — Fig. 8(A) shows a typical time course of change in the fluorescence intensity of DNS-ATP at 520 nm during its reaction with α subunit of EF₁. The fluorescence intensity of 1.1 µM DNS-ATP in the presence of 2 mM EDTA did not change on addition of 0.8 µM α subunit [arrow <u>a</u> in Fig. 8(A), <u>cf</u>. Fig. 9]. On further addition of 5 mM MgCl₂ [arrow <u>b</u> in Fig. 8(A)], the fluorescence intensity increased very rapidly (t_{1/2} < 1 s) by 19%. No fluorescence increase was observed in the absence of this subunit or in the presence of an excess amount of ATP with the subunit. On further addition of 2 mM P_i [arrow <u>c</u> in Fig. 8(A)], the fluorescence intensity changed only slightly and maintained its level. When 2 mM ATP was finally added [arrow d in

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Fig. 7. Effects of DCCD-treatment of EF_1 on its steady-state ATPase and single turnover of DNS-ATP hydrolysis. (A) EF_1 at 0.87 µM was preincubated with (O) or without (\bigcirc) 0.2 mM DCCD in 2 mM EDTA and 50 mM Tris-acetate at pH 7.0 and 30°C for 20 min. Then, EF_1 solution was diluted to 0.087 µM in 5 mM phosphoenolpyruvate, 0.11 mg/m1 pyruvate kinase, 2.5 mM MgCl₂, 0.2 mM EDTA and 50 mM Tris-HCl at pH 8.0 and 30°C. The reaction was started by addition of 5 mM ATP. (B) EF_1 at 1.6 µM was preincubated with (O) or without (\bigcirc) 0.2 mM DCCD under the same conditions as A, then allowed to react with 2.8 µM DNS-AT³²P for 1 min. The reaction was started by addition of 3 mM Mg²⁺.



Fig. 8. Time course of fluorescence intensity change of DNS-ATP in the presence of α subunit. (A) To a reaction mixture containing 0.8 μ M α subunit in a buffer solution (2 mM EDTA, 0.1 M NaCl and 50 mM Tris-acetate at pH 7.0 and 30°C), the following additions were made at the time indicated by arrows (a-d); a, 1.1 μ M DNS-ATP; b, 5 mM MgCl₂; c, 2 mM K-P_i; d, 2 mM ATP. (B) To a reaction mixture containing 4 μ M α subunit in the buffer solution, the following additions were made at the time indicated by arrows (a-d); a, 1.1 μ M DNS-ATP; b, 5 mM MgCl₂; c, 0.2 mM AMP; d, 0.2 mM ATP.



Fig. 9. Fluorescence emission spectra recorded during the course of reaction of DNS-ATP with α' subunit. Spectra were recorded in the following order; curve a, 1 min after addition of 4 μ M α' subunit to 1.1 μ M DNS-ATP; curve b, 1 min after addition of 5 mM MgCl₂; curve c, 1 min after addition of 2 mM ATP. The fluorescence emission spectra of 1.1 μ M DNS-ATP alone in 2 mM EDTA and in 2 mM EDTA + 5 mM MgCl₂ were the same as the curve a and c, respectively. Other conditions are the same as described in Fig. 8.

Fig. 8(A)], the fluorescence intensity decreased exponentially with a $t_{1/2}$ of 8 s to a level which was about 5% lower than that before the addition of 5 mM MgCl₂. The final level was found to be equal to that of free DNS-ATP in the presence of free Mg²⁺.

Essentially the same experiment was carried out with 4.0 μ M α subunit; as shown in Fig. 8(B) the fluorescence intensity increased very rapidly by 50% upon addition of 5 mM MgCl₂ [arrow <u>b</u> in Fig. 8(B)]. Addition of 0.2 mM AMP [arrow <u>c</u> in Fig. 8(B)] had no effect on the fluorescence intensity, whereas subsequent addition of 0.2 mM ATP [arrow <u>d</u> in Fig. 8(B)] decreased the intensity as observed in Fig. 8(A). When 5 mM CaCl₂, instead of MgCl₂, was added to the reaction mixture, no change of the fluorescence intensity was observed (data not shown).

Fluorescence emission spectra were recorded during the course of reaction of DNS-ATP with α' subunit (Fig. 9). Addition of 4.0 μ M α' subunit to 1.1 μ M DNS-ATP in the presence of 2 mM EDTA did not change the fluorescence emission spectrum of the free DNS-ATP (trace a). Further addition of 5 mM MgCl₂ increased the fluorescence intensity, and the emission maximum shifted to around 530 nm (trace b). Addition of 2 mM ATP decreased the fluorescence intensity to the level of free DNS-ATP in the presence of Mg²⁺ (trace c), which was slightly lower than that in the absence of Mg²⁺ (trace a).

Stoichiometry of DNS-ATP Binding to \checkmark Subunit — The stoichiometry of DNS-ATP binding to \checkmark subunit was determined by fluorometric titration of 1.1 µM DNS-ATP with \checkmark subunit in the presence of 3 mM Mg²⁺ at 4° or 30°C (Fig. 10). Data were analyzed based on Eq. 1, where the parameters for _EF₁ were taken as those for \checkmark subunit. The solid lines in Fig. 10 are the best fits of the data of the DNS-ATP titration with \checkmark subunit at 4° ($\textcircled{\bullet}$) and 30°C (O). The parameters

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Fig. 10. Fluorescence titration of DNS-ATP with \propto subunit in the presence of Mg²⁺. The extent of fluorescence increase of 1.1 μ M DNS-ATP after addition of 3 mM Mg²⁺ was plotted against the amount of \propto subunit at 30° (O) or 4°C (e). Other conditions are the same as described in Fig. 8. The extent of fluorescence decrease of free DNS-ATP in the presence of free Mg²⁺ (<u>cf</u>. Fig. 9, curve c) was corrected. The solid lines are the theoretical curves (see the text). When 1.1 μ M DNS-ATP was allowed to react with β subunit in the presence of Mg²⁺ at 30°C, no fluorescence change was observed (Δ).

obtained from the data are n = 0.82, and $\phi_{\text{DNS-ATP}}$ = 0.06 μ M at 4°C, and n = 0.65 and $\phi_{\text{DNS-ATP}}$ = 0.07 μ M at 30°C. The value of ΔF_{max} was 55% at both temperatures. When β subunit up to 4 μ M was added to 1.1 μ M DNS-ATP, no Mg²⁺-dependent fluorescence increase was observed (Δ).

The DNS-ATP binding to α' subunit was also measured at room temperature by the equilibrium dialysis. We found that even in the absence of Mg²⁺, DNS-ATP can bind to α' subunit with an apparent dissociation constant of about 5 times higher than that in the presence of Mg²⁺. However, during the equilibration we noticed some denaturation of α' subunit, and therefore could not analyze the data quantitatively.

Effects of Nucleotides and Pyrophosphate on DNS-ATP Bound to α <u>Subunit</u> — The fluorescence intensity of DNS-ATP increased by addition of 5 mM MgCl₂ to the mixture containing 1.1 µM DNS-ATP, 0.8 µM α subunit and 2 mM EDTA. Then, ADP, AMPPNP, GTP, ITP, CTP at 0.2 mM or PP_i at 2 mM was added. All these compounds decreased the fluorescence intensity exponentially with $t_{1/2} = 7-8$ s to a level almost equal to that of free DNS-ATP (data not shown), as was observed upon the addition of 0.2 mM ATP (Fig. 8).

DCCD inhibited the steady-state rate of EF_1 -ATPase activity at high ATP concentrations, but did not inhibit EF_1 -DNS-ATPase activity at low DNS-ATP concentrations, as mentioned above (Fig. 7). On the contrary we found that DCCD at 2 mM did not affect the Mg²⁺-dependent fluorescence increase and the ATP-dependent fluorescence decrease of DNS-ATP in the presence of $\boldsymbol{\triangleleft}$ subunit (data not shown), suggesting that the results obtained above (Figs. 6 & 7) were not due to the reaction of DCCD to this subunit.

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Inability of $\not{\propto}$ Subunit to Hydrolyze DNS-ATP and ATP — No TCA-³²P_i liberation was observed on either addition of 5 mM MgCl₂ to the reaction mixture containing 2.8 μ M DNS-AT³²P, 2.3 μ M $\not{\propto}$ subunit and 2 mM EDTA (Δ), or subsequent addition of 2 mM ATP (\bigstar) (Fig. 4). Even when 5 mM MgCl₂ was added to the mixture containing 7.6 μ M AT³²P and 1.6 μ M $\not{\propto}$ or 1.8 μ M β subunit in 2 mM EDTA, no TCA-³²P_i liberation was observed (data not shown). These results are in agreement with the previous finding that the isolated $\not{\propto}$, β or \not{o} subunit did not have ATPase activity (7,8).

Fluorescence Change of DNS-ATP on Its Binding to Reconstituted EF₁-ATPase — To determine what kind of subunit interactions are needed for the EF1-ATPase activity, we further investigated the reaction of DNS-ATP with the reconstituted EF, -ATPase. Figure 11 shows the time course of change in the fluorescence intensity of DNS-ATP at 520 nm during its reaction with reconstituted EF₁-ATPase. The fluorescence intensity of 1.1 μ M DNS-ATP in the presence of 2 mM EDTA increased slightly on addition of 0.36 mg/ml reconstituted EF_1 -ATPase [arrow <u>a</u>] in Fig. 11. compare curves a and b in Fig. 12]. On further addition of 5 mM MgCl₂ [arrow <u>b</u> in Fig. 11], the fluorescence intensity increased very rapidly $(t_{1/2} < 1 \text{ s})$, then slowly with $t_{1/2}$ of about 30 s. The plateu level was 1.83 fold the intensity of free DNS-ATP. On further addition of 2 mM P_i [arrow <u>c</u> in Fig. 11], the fluorescence increased rapidly to a level of 2.4 fold that of free DNS-ATP. On addition of 2 mM ATP [arrow d in Fig. 11], the fluorescence intensity decreased to the level of free DNS-ATP.

When 5 mM CaCl₂ was added to the mixture containing 1.1 μ M DNS-ATP and 0.36 mg/ml reconstituted EF₁-ATPase in the presence of 2 mM EDTA, the fluorescence intensity increased exponentially with

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Fig. 11. Time course of fluorescence intensity change of DNS-ATP in the presence of reconstituted EF_1 -ATPase. To a mixture containing 0.36 mg/ml reconstituted EF_1 -ATPase in 2 mM EDTA, 100 mM NaCl and 50 mM Tris-acetate at pH 7.0 and 30°C, the following additions were made at the time indicated by arrows (a-d); a, 1.1 μ M DNS-ATP; b, 5 mM MgCl₂; c, 2 mM K-P_i; d, 2 mM ATP. Specific activity of the reconstituted EF_1 -ATPase was 30 U/mg protein. The concentration of the reconstituted EF_1 -ATPase was shown by the protein concentration, because the reconstitution was not complete [note that the specific activity of purified EF_1 is about 100 U/mg protein (28)]. $t_{1/2}$ of 40 s to reach a plateu level, which was 1.5 fold the intensity of free DNS-ATP. Further addition of 2 mM P_i induced a small increase (6%) in the fluorescence intensity (data not shown).

Figure 12 shows the fluorescence emission spectra of free DNS-ATP (trace a) and complexes of DNS-ATP with reconstituted EF_1 -ATPase. When 0.53 mg/ml reconstituted EF_1 -ATPase was added to 1.1 μ M DNS-ATP in the presence of 2 mM EDTA, the fluorescence intensity increased slightly with a shift of the emission maximum from 555 to 550 nm (trace b). Further addition of 5 mM MgCl₂ markedly increased the fluorescence intensity, and the maximum shifted to 535 nm (trace c). The addition of 2 mM ATP reversed these fluorescence changes backward to the state of free DNS-ATP (trace d).

Figure 13 shows the result of fluorescence titration of DNS-ATP with reconstituted EF_1 -ATPase in the presence of 3 mM Mg²⁺. The extent of the Mg²⁺-dependent fluorescence increase of 1.1 μ M DNS-ATP increased with increase in the amount of reconstituted EF_1 -ATPase added. When the amount of reconstituted EF_1 -ATPase added was 1.5 μ M (0.54 mg/ml), the extent of the Mg²⁺-dependent fluorescence increase was 88% of the intensity of free DNS-ATP. However, we could not obtain a good fit to the data based on Eq. 1, since we could not measure the saturation level of the fluorescence increase.

Acceleration of DNS-Nucleotide Release from Reconstituted EF_{1-} ATPase-DNS-Nucleotide Complex by Nucleotides and Pyrophosphate — DNS-ATP at 1.1 µM was allowed to react with 0.42 mg/ml reconstituted EF_1 -ATPase for 2 min in the presence of 3 mM Mg²⁺. Then, 0.2 mM of nucleotide or 2 mM pyrophosphate was added [arrow <u>c</u> in Fig. 14], and the following decrease in the fluorescence intensity was measured

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Fig. 12. Fluorescence emission spectra recorded during the course of reaction of DNS-ATP with reconstituted EF_1 -ATPase. Spectra were recorded in the following order; curve a, l.l µM DNS-ATP; curve b, l min after addition of 0.53 mg/ml reconstituted EF_1 ; curve c, 2 min after addition of 3 mM Mg²⁺; curve d, l min after addition of 2 mM ATP. Other conditions are the same as described in Fig. ll.



Fig. 13. Fluorometric titration of DNS-ATP with reconstitured EF_1 -ATPase in the presence of Mg^{2+} . The extent of fluorescence enhancement of 1.1 μ M DNS-ATP 2 min after addition of 3 mM Mg^{2+} was plotted against the amount of reconstituted EF_1 -ATPase added. Other conditions are the same as described in Fig. 11.



Fig. 14. Fluorescence decrease on addition of various phosphate compounds to reconstituted EF_1 -ATPase-DNS-nucleotide. DNS-ATP at 1.1 μ M was added to 0.42 mg/ml reconstituted EF_1 -ATPase (indicated by arrow a), then 3 mM Mg²⁺ was added (indicated by arrow b). At the time indicated by arrow c, the following additions were made; 0.2 mM ATP (1), 0.2 mM ADP (2), 2 mM PP₁ (3), 0.2 mM CTP (4) or 0.2 mM AMP (5). Other conditions are the same as described in Fig. 11.

(Fig. 14). The addition of ATP caused a fluorescence decrease, first rapidly ($t_{1/2} < 1$ s) and then slowly, and the final level was almost equal to that of free DNS-ATP (trace 1). When ADP (trace 2), PP_i (trace 3) or CTP (trace 4) was added, the fluorescence decreased almost exponentially. The rate of the fluorescence change decreased in the order of ATP > ADP \approx PP_i > CTP. AMP (trace 5) had no effect on the fluorescence intensity of the reconstituted EF₁-ATPase-DNS-nucleotide.

DISCUSSION

We have previously performed kinetic studies on the single turnover of DNS-ATPase of beef heart F_1 (26), and proposed the following reaction scheme for the F_1 -DNS-ATPase reaction:

 $E + DNS - ATP \rightleftharpoons E^{DNS - ATP} \rightarrow E^{DNS - ATP} \rightleftharpoons E_{P}^{DNS - ADP} \xrightarrow{P_{i}} E^{DNS - ADP} \xrightarrow{P_{i}} E^{DNS - ADP} \xrightarrow{P_{i}} E^{DNS - ADP}$

The asterisk designates the reaction intermediate with enhanced fluorescence intensity of DNS-nucleotide. In the presence of Mg^{2+} , DNS-ATP binds to the high affinity site of F_1 to form a loose complex without enhanced fluorescence ($E^{DNS-ATP}$). The loose complex is converted to a tight complex with enhanced fluorescence ($E^{DNS-ATP*}$), then bound DNS-ATP is cleaved on the enzyme ($E_p^{DNS-ADP*}$). The apparent affinity of DNS-ATP to the enzyme is very high. Nucleotides such as ATP at high concentrations markedly accelerate the following three steps in the reaction sequence:

 $E^{DNS-ATP*} \longrightarrow E^{DNS-ADP*}_{P_i} \xrightarrow{P_i} E^{DNS-ADP*}_{DNS-ADP} \xrightarrow{E}$

Therefore, we suggested that there are low affinity regulatory site(s) as well as the high affinity catalytic sites in F_1 . The purified EF_1 contained 2 mol ATP and 0.5 mol ADP per mol enzyme (33). These bound adenine nucleotides were found to be nonexchangeable with nucleotides in the medium (33). Therefore, in analyzing the results on the reaction of EF_1 with DNS-ATP, we neglected the effect of these tightly bound adenine nucleotides.

The first issue investigated in this study is the similarity of the reaction mechanism of EF_1 -DNS-ATPase to that of F_1 -DNS-ATPase. In the presence of Mg^{2+} , DNS-ATP reacted with EF_1 to induce a

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fluorescence increase similar to that observed on beef heart F_1 (Figs. 1 & 2). This finding suggests that the environment of DNS-ATP binding sites in EF_1 is hydrophobic as is the case for F_1 . The fluorescence intensity of EF1-DNS-nucleotide complex increased on further addition of P_i , again as is the case for F_1 -DNS-nucleotide complex. The fluorescence increase by P_i was observed even in the presence of excess amounts of DNS-ATP to EF1. Furthermore, we $(\underline{26})$ previously found that in the case of beef heart F_1 , the affinity of DNS-ATP was unaffected by the addition of P_i. These findings indicate that the DNS-ATP binding site in EF_1 is made more hydrophobic by the action of Pi. The fluorometric titration of DNS-ATP with EF_1 in the presence of Mg^{2+} indicated that 3 mol of DNS-ATP was bound to 1 mol of EF_1 with an apparent dissociation constant of 0.23 μM (Fig. 3). Using the same method of fluorometric titration, we previously found that 2 mol of DNS-ATP binds to 1 mol of beef heart F $_{1}$ with an apparent dissociation constant of 0.44 μM (26). The stoichiometry of 3 mol DNS-ATP bound/mol EF_1 is consistent with the $d_3 \beta_3 \tau \delta \epsilon$ stoichiometry of EF₁ subunits (<u>34</u>). When Mg²⁺ was added to a reaction mixture containing EF1 and a low concentration of DNS-ATP, the fluorescence intensity was increased $(E^{DNS-ATP*};$ Fig. 1), and the increase was followed by liberation of P_i on termination of the reaction with TCA ($E_P^{DNS-ADP*}$ & $E^{DNS-ADP*}$; Fig. 4). The fluorescence intensity maintained its increased level after liberation of TCA-P_i, indicating that the rate of liberation of DNS-ADP from $E^{DNS-ADP*}$ is very small. These findings are quite similar to those of beef heart F_1 (26).

The mechanism of acceleration of EF1-DNS-ATPase by high

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concentrations of nucleotide was investigated using the same procedures as in the case of beef heart F_1 . A biphasic decrease in the fluorescence intensity of EF1-DNS-nucleotide complex was observed upon addition of high concentrations of ATP (Fig. 5). Therefore, it was suggested that ATP binding to low affinity regulatory site(s) in EF₁ induced a conformational change around the catalytic sites and accelerated the release of DNS-nucleotide, as in the case of beef heart F_1 (26). The rapid phase of the fluorescence decrease observed on addition of ATP, ADP, or GTP to EF₁-DNS-nucleotide complex (Fig. 5) is suggested to be due to the release of DNS-nucleotide induced by a conformational change of catalytic site on EF1. The second slow phase will be due to the displacement of bound DNS-nucleotide by other nucleotide. This slow release of DNS-nucleotide was also observed in the case of addition of PP; or CTP. Thus, the ability of phosphate compounds to release DNS-nucleotide from EF₁ was found to follow in the order of

ATP > ADP \approx GTP > AMPPNP > PP_i \approx CTP \gg AMP This order was similar to that observed on beef heart F₁ (<u>26</u>):

ATP \geq ADP \geq GTP \approx AMPPNP > PP₁ \approx CTP \gg AMP In the case of beef heart F_1 , ATP at high concentrations accelerated markedly the TCA-P₁ liberation from E^{DNS-ATP*} (<u>26</u>), whereas no acceleration of the TCA-P₁ liberation by ATP was observed in the case of EF₁-DNS-ATPase (Fig. 4). This may suggests that the reaction mechanism of EF₁-DNS-ATPase is essentially different from that of F₁-DNS-ATPase. However, if we assume the reaction mechanisms of these ATPases are essentially the same, we can explain the discrepancy between the results of the two enzymes as due to the larger rate constant of the cleavage step (E^{DNS-ATP*} \longrightarrow E^{DNS-ADP*}) than the rates of the following steps in the case of EF₁-DNS-ATPase. Then, the

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acceleration of TCA-P_i liberation due to acceleration of the cleavage step is not observed, since the amount of $E^{\text{DNS-ATP}*}$ at steady state is small.

The second issue is the subunit localization of the high affinity catalytic site and low affinity regulatory site. We previously suggested that the high affinity catalytic site and low affinity regulatory site exist in the α and β subunit, respectively (<u>26</u>). This suggestion is based on the information that the isolated α subunit of thermophilic F_1 or EF_1 and β subunit of thermophilic F_1 have high affinity and low affinity nucleotide-binding site, respectively (<u>9,10</u>). In this study, we found that the fluorescence intensity of DNS-ATP increased markedly on its reaction with α subunit but not with β subunit of EF_1 in the presence of Mg²⁺ (Figs. 8-10). The fluorometric titration of DNS-ATP with α subunit in the presence of Mg²⁺ indicated that 0.65-0.82 mol of DNS-ATP bound to 1 mol of α subunit very tightly (Fig. 10). This finding suggests that properties of DNS-ATP binding to α subunit are very similar to those of DNS-ATP binding to EF_1 .

We investigated the possibility of the existence of a low affinity regulatory site in β subunit by using DCCD, a potent inhibitor of F_1 -ATPase activity specific to β subunit (<u>20-23</u>). DCCD had no effect on the binding of DNS-ATP with α subunit. It inhibited only slightly the fluorescence increase of DNS-ATP by its binding to EF₁ (Fig. 6) and the single turnover of DNS-ATP hydrolysis by EF₁ [Fig. 7(B)]. DCCD inhibited markedly both the accelerating effect of ATP on DNS-nucleotide release from EF₁ (Fig. 6) and the EF₁-ATPase activity in the steady state [Fig. 7(A)]. These finding indicate that the modification of β subunit by DCCD blocks the function of the low affinity regulatory site, without affecting the binding of DNS-ATP to the high affinity catalytic site.

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Thus, our findings strongly suggest that the high affinity catalytic site and the low affinity regulatory site exist in α and β subunits, respectively. This concept is apparently opposite to a widely accepted view that the catalytic site and the non-catalytic nucleotide binding site are located in β and α subunits, respectively (<u>15</u>,<u>16</u>,<u>18</u>-<u>23</u>). It should be mentioned that the latter view has been derived mainly from the inhibition of F₁-ATPase activity in the steady state by chemical modification of β subunit. However, we cannot determine the subunit localization of the catalytic site and regulatory site by measuring the effect of the modification of α or β subunit on the steady-state ATPase. This is because the low affinity nucleotidebinding to the regulatory site accelerates markedly the release of product bound to the catalytic site (<u>26</u>), and the F₁-ATPase activity in the steady state is markedly inhibited by the modification of the regulatory site as well.

In the previous paper, it was suggested that hydrolysis of nucleosidetriphosphate in the steady state occurs <u>via</u> different catalytic pathways and at different catalytic sites in F_1 because of the following two reasons (26). (a) The value of V_f (0.34 s⁻¹) for the formation of $E^{DNS-ATP*}$ was smaller than that of V_{max} ($\gg1$ s) for the steady-state rate of F_1 -DNS-ATPase. (b) When ATP was added under conditions where most of F_1 was in the form of $E^{DNS-ADP*}$, ATP hydrolysis occurred without lag phase. For EF_1 , the modification of β subunit by DCCD blocks both catalytic pathways, because DCCD inhibited the EF_1 -ATPase in the steady state almost completely [Fig. 7(A)].

The third issue investigated in this paper is the difference between the structure and function of the isolated $\not\propto$ subunit and those of integrated $\not\propto$ in EF₁. Recently, Dunn and Futai (<u>7</u>), and Dunn

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(35) reported that the isolated χ subunit of EF₁ binds ATP in the absence of Mg²⁺ with very high affinity (K_d = 0.1 µM at 4°C). In this study, we found that the structure and function of the χ subunit are altered when isolated from EF₁ in the following two respects: (<u>a</u>) In the presence of Ca²⁺, DNS-ATP showed the fluorescence increase on its reaction with EF₁ [Fig. 1(B)], but not with isolated χ subunit. The apparent dissociation constant of isolated χ subunit for DNS-ATP

in the presence of Mg²⁺ was 0.07 μ M, which was about 1/3 of the value found with EF₁ (Figs. 3 § 10). The extent of the fluorescence increase of DNS-ATP bound to the isolated α subunit in the presence of Mg²⁺ was 55% of the fluorescence intensity of free DNS-ATP, which was about 1/5 of the value found with EF₁ (Figs. 3 § 10). These differences suggest that on isolation of α subunit, its affinity for DNS-ATP increased and the environment of the nucleotide-binding site became less hydrophobic. (b) P₁ had no effect on the fluorescence intensity of DNS-ATP bound to the isolated α subunit [Fig. 8(A)], wherease P₁ enhanced markedly the fluorescence intensity in the case of EF₁ [Fig. 1(A)]. This finding is consistent with the report that the P₁ binding site exists in the β subunit (36), and it also suggests an interaction between the α and β subunits in EF₁.

We found that the properties of the reaction of DNS-ATP with the isolated α' subunit was unaffected by the addition of the β subunit, and that the fluorescence changes of DNS-ATP on its reaction with EF₁-ATPase reconstituted from α' , β and γ' subunit were quite similar to those on its reaction with native EF₁. On addition of Mg²⁺ to a reaction mixture containing DNS-ATP and the reconstituted EF₁-ATPase, the fluorescence intensity increased in two phases (Fig. 11).

rapid phase $(t_{1/2} < 1 \text{ s})$ of the increase was suggested to be due to the contamination by free \mathbf{d} subunit, since the rapid phase was not observed on addition of Ca^{2+} instead of Mg^{2+} . It is suggested that the hydrophobicity of the nucleotide binding site is recovered to the level of native EF_1 , since the extent of the Mg^{2+} -dependent fluorescence increase was larger than 88% of the intensity of free DNS-ATP (Fig. 13). Furthermore, the decrease in the fluorescence intensity of reconstituted EF_1 -DNS-nucleotide complex after adding a phosphate compound showed a specificity quite aimilar to that of native EF_1 : which is ATP > ADP $\approx PP_i > CTP \gg AMP$ (Fig. 14). These findings clearly indicate that the subunit interactions in reconstituted EF_1 -ATPase are similar to those in the native EF_1 .

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LIST OF PUBLICATIONS

- Calorimetric studies of heat of respiration of mitochondria.
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- 2. Calorimetric studies of the heat of respiration of mitochondria. Matsuoka, I., Watanabe, T., & Nakamura, T. (1978) <u>Frontiers</u> of <u>Biological Energetics</u> Vol. 1, pp. 689-697, Academic Press, Inc., New York
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- 4. Comparative studies on the effects of linoleate and methyl linoleate and their hydroperoxides on the respiration and reactivities of rat heart mitochondria. Shiotani, A. Watanabe, T., Matsuoka, I., & Nakamura, T. (1980) J. Biochem. <u>88</u>, 677-683
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