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Asymmetric Allosteric Signaling in Aspartate Transcarbamoylase

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nzymes responsible for catalyzing the committed step of critical metabolic pathways, such as aspartate transcarbamoylse (ATCase) in pyrimidine nucleotide biosynthesis, are often regulated by allosteric mechanisms. Allosteric regulation provides a rapid means for turning a pathway on or off dictated by the specific needs of the cell at any point in time. Escherichia coli ATCase has been investigated extensively and has come to serve as a paradigm for allosteric enzymes (1). ATCase is regulated by both homotropic and heterotropic effectors involving the cooperative binding of the second substrate, aspartate, at the active site and nucleotide binding at the allosteric site. The active and allosteric sites are located \sim 60 Å apart on different polypeptide chains of the holoenzyme, a dodecamer composed of six catalytic chains (c) and six regulatory chains (r) (Figure 1, panel a), which can be dissociated into two catalytically active trimers (c_3) and three nucleotide-binding regulatory dimers (r_2) (2).

Kinetic studies revealed activation by ATP, inhibition by CTP, and the synergistic inhibition of UTP in the presence of CTP (3). However, the elucidation of the allosteric mechanism of regulation in ATCase has been exceedingly difficult with many conflicting results. The interactions of the regulatory nucleotides with the enzyme have been investigated using a variety of methods including equilibrium dialysis (4-8), continuousflow dialysis (9), nuclear magnetic resonance (10, 11), site-specific mutagenesis (8, 10–13), and X-ray crystallography (14, 15). Some of these experiments indicate two classes of nucleotide binding sites with differing affinities, suggesting either a nonequivalence of binding sites or negative cooperativity (7). However, these methods lack the sensitivity to accurately measure weak interactions such as the binding of UTP, as well as the binding of nucleotides to the low-affinity sites. Also, the **ABSTRACT** Here we use the fluorescence from a genetically encoded unnatural amino acid, L-(7-hydroxycoumarin-4-yl)ethylglycine (HCE-Gly), replacing an amino acid in the regulatory site of Escherichia coli aspartate transcarbamoylase (ATCase) to decipher the molecular details of regulation of this allosteric enzyme. The fluorescence of HCE-Gly is exquisitely sensitive to the binding of all four nucleotide effectors. Although ATP and CTP are primarily responsible for influencing enzyme activity, the results of our fluorescent binding studies indicate that UTP and GTP bind with similar affinities, suggesting a dissociation between nucleotide binding and control of enzyme activity. Furthermore, while CTP is the strongest regulator of enzyme activity, it binds selectively to only a fraction of regulatory sites, allowing UTP to effectively fill the residual ones. Our results suggest that CTP and UTP are not competing for the same binding sites, but instead reveal an asymmetry between the two allosteric sites on the regulatory subunit of the enzyme. Correlation of binding and activity measurements explain how ATCase uses asymmetric allosteric sites to achieve regulatory sensitivity over a broad range of heterotropic effector concentrations.

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current methods do not allow differentiation between the binding of nucleotides to the allosteric site and nonspecific binding at the active site unless the active site is blocked. However, the ligands used to block the active sites are known to induce conformational changes that may indirectly alter the affinity of the allosteric sites (*16*). In addition, allosteric regulation must respond rapidly to the fluctuating conditions in the cell, but current methods do not allow time-dependent measurements to investigate the rates of nucleotide binding.

The work by Schultz and co-workers (17), to incorporate unnatural amino acids into the structure of proteins, has provided new tools to study complex processes in biological systems. Here we use these tools to introduce a fluorescent amino acid in the regulatory site of ATCase, providing us a means to directly monitor and dissect nucleotide binding to the enzyme without the complexities and limitations of current methods. These experiments validate the usefulness of the methods developed by Schultz and co-workers to incorporate an unnatural amino acid into not only ATCase but other systems as well. In particular, we applied the methodology of Wang *et al.* (*18*) to incorporate the amino acid L-(7hydroxycoumarin-4-yl)ethylglycine (HCE-Gly) sitespecifically into the regulatory chain of ATCase at the allosteric binding pocket to create a fluorescent version of the enzyme that is exquisitely sensitive to nucleotide binding. The 7-hydroxycoumarin side chain is ideal to monitor nucleotide binding because it displays a high fluorescence quantum yield and relatively large Stoke's shift and is sensitive to pH and solvent polarity. This fluorescent ATCase provides not only a tool to monitor nucleotide binding to both the high and low affinity sites but also a means to measure time-dependent binding through the use of stopped-flow spectrofluorimetry.

RESULTS AND DISCUSSION

Selection of HCE-Gly Incorporation Site. The ideal location to place HCE-Gly in the allosteric site of ATCase would be one in which its location does not influence the binding of the nucleotides to the allosteric site, but nucleotide binding should perturb the local environment of the HCE-Gly in such a fashion that its fluorescence would be altered. Candidate positions were selected by visual inspection and tested by model building and energy minimization to determine if the HCE-Gly significantly altered the regulatory binding pocket (Figure 1, panel b). From these studies two sites were selected for incorporation of HCE-Gly: Glu52_R or Leu58_R.

Expression and Purification of HCE-Gly ATCase. The expression and suppression system required the use of two plasmids. The first had the E. coli pyrBI operon, coding for the catalytic and regulatory chains of ATCase, in plasmid pBAD/JYAMB-4TAG-Myo (18), replacing the portion containing the myoglobin gene and the codons at position 52 or 58 in the r chain of ATCase sitespecifically altered to TAG, the amber stop codon. This plasmid also carried the gene encoding the special Methanococcus jannaschii tyrosyl amber suppressor tRNA (18). The second plasmid, pBK-CouRS-D8 (18), contains the gene for the mutant M. jannaschii tRNA synthetase specific for HCE-Gly. The expression was performed by growth of Top10 cells (Invitrogen) containing both of the above plasmids in media supplemented by 1.0 mM HCE-Gly, which was synthesized as described previously (18). Purification was accomplished in two steps. The first employed metal affinity chromatography to purify the enzyme. However, this resulted in a mixture of holoenzyme (c_6r_6) and excess catalytic subunit (c_3) (Figure 2, panel a). The holoenzyme was purified to homogeneity by hydrophobic interaction chromatography (Figure 2, panel b). The levels of suppression of the

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TABLE 1. Kinetic parameters of the wildtype and HCE-Gly52_R ATCase^a

	pН	V _{max} ^b	[Asp] _{0.5}	n _H ^c
Wild-type	8.3	16.6 ± 0.4	12.6 ± 0.1	3.1 ± 0.4
	7.0	10.4 ± 0.1	7.5 ± 0.2	2.1 ± 0.2
HCE-Gly52 _R	8.3	16.8 ± 2.2	12.6 ± 0.7	2.7 ± 0.3
	7.0	8.7 ± 0.3	7.7 ± 0.5	1.7 ± 0.3

^{*a*}All experiments were performed at 25 °C in 50 mM Tris-acetate buffer (pH 8.3) or 20 mM bis-Tris, 20 mM Tris, 20 mM CAPS buffer (pH 7.0) at a saturating CP concentration (4.8 mM). The values reported are the average deviation of four determinations. ^{*b*}The maximal observed velocity (mmol h⁻¹ mg⁻¹) ^cHill coefficient

Figure 2. Purification of HCE-Gly52_R ATCase. SDS-PAGE (a) and nondenaturing PAGE (b) visualized by Coomassie blue stain (panels a and b, left) or fluorescence (panels a and b, right) of the purification of the HCE-Gly52_R ATCase. Lanes 1, wild-type ATCase; Lanes 2, *E. coli* ATCase c₃ containing a C-terminal His tag; lanes 3, total soluble protein from cell lysate; lanes 4, HCE-Gly52_R ATCase after column purification employing Chelating Sepharose Fast Flow (GE Healthcare) charged with Ni²⁺; lanes 5, HCE-Gly52_R ATCase after column purification employing Phenyl Sepharose (GE Healthcare).

TAG codon were significantly different between the two sites, with position 52 proving to be better.

Kinetic Properties of HCE-Gly52_R **ATCase.** The HCE-Gly52_R ATCase was subjected to a battery of kinetic measurements that verified that the mutant enzyme exhibited kinetic parameters similar to those of the wild-type enzyme including maximal velocity, half saturation of aspartate, Hill coefficient, and activation and inhibition by ATP and CTP, respectively (Table 1, Supplementary Figures 1 and 2). Thus, the incorporation of the HCE-Gly at position 52 produced a functional ATCase with kinetic properties very similar to the wild-type enzyme, making the HCE-Gly52_R ATCase a model for the wild-type holoenzyme, albeit with the addition of a fluorescence sensor in the regulatory binding site.

HCE-Gly52_R ATCase Fluorescence Sensitivity to Nucleotides. Upon addition of ATP or CTP to the HCE-Gly52_R ATCase there was a dramatic alteration in fluorescence intensity with little or no shift in the emission maximum (Figure 3, panel a). Crystallography studies



Figure 3. Monitoring nucleotide binding to HCE-Gly52_R ATCase by fluorescence. a) Fluorescence spectrum of HCE-Gly52_R ATCase as a function of the CTP concentrations. HCE-Gly52_R ATCase (50 μ g mL⁻¹) in 50 mM Tris-acetate pH 8.3 at 25 °C was excited at 360 nm, and the emission spectrum was measured from 400-550 nm. The open circle represents the spectrum of enzyme in the absence of CTP. As the CTP concentration was increased to 15 mM (solid circle) in 12 aliquots, the emission intensity at 455 nm decreased significantly. b) Alteration in the fluorescence of the HCE-Gly52_R ATCase upon the binding of ATP (green), CTP (red), UTP (blue), and GTP (black). The fluorescence intensity at 455 nm was normalized to the highest value (0 mM NTP) and plotted versus the nucleotide concentration. Conditions similar to those used in panel a.

have shown that the ribose triphosphate portions of ATP and CTP bind in essentially the same part of the nucleotide-binding site. Therefore one might expect to also observe an interaction of the enzyme with other nucleotides such as UTP and GTP, although it has been reported that these nucleotides do not alter the activity of the enzyme alone or in nucleotide combinations (19). Nevertheless, UTP or GTP alone was able to drastically alter the fluorescence intensity of the HCE-Gly52_R ATCase (Figure 3, panel b; Supplementary Table 1). The biphasic nature of nucleotide binding is particularly apparent in the data for ATP, UTP, and GTP. These data clearly establish the binding of UTP and GTP to the enzyme with affinity comparable to that of ATP and suggest two classes of binding sites with differing affinity. The reduction in fluorescence induced by CTP is less than the other nucleotides, and for CTP these data suggest only one class of binding sites; if there are two classes of sites, the affinity of the weaker sites is too low to be detected.

pH Dependence of HCE-Gly. The alteration in the fluorescence of the HCE-Gly52_R ATCase is dramatically different at pH 7 and pH 8.3. Each nucleotide caused a reduction in the fluorescence at 8.3, while each caused an increase in fluorescence at pH 7 (Figure 4, panel a). The altered response due to nucleotide binding at these two pH values can be partially attributed to the pH dependence of the HCE-Gly, the 7-hydroxyl group of which has a $pK_a = 7.8$ (20). The decrease in fluorescence due to NTP binding at pH 8.3 can be attributed to an increase in the pK_a of the HCE-Gly induced by close proximity of the negative charges of the phosphates of the NTPs to the position of the HCE-Gly52_R (Figure 1, panel b). At pH 7, the situation is somewhat different since the 7-hydroxyl group of HCE-Gly should be almost fully protonated. Under these pH conditions, the binding of the nucleotide acts to increase the local hydrophilicity of the environment of the HCE-Gly52_R, which would cause an increase in fluorescence intensity. Thus, different aspects of nucleotide binding to the enzyme can be probed depending upon the pH at which the experiment is performed.

Binding of ATP and CTP to the Isolated Regulatory Subunit. Previous studies of the binding of ATP and CTP to the isolated regulatory subunit (r₂) indicated that only one molecule of nucleotide bound per homodimeric regulatory subunit (*6*). As seen in Figure 4, panel b, when the binding of ATP or CTP to the HCE-



Figure 4. ATP and CTP binding to HCE-Gly52_R ATCase and the isolated regulatory subunit. a) Alteration in the fluorescence of the HCE-Gly52_R ATCase upon the binding of ATP (green) and CTP (red) at pH 7.0 (squares) and 8.3 (circles). HCE-Gly52_R ATCase (50 μ g mL⁻¹) in 50 mM Tris-acetate buffer at pH 8.3 or 20 mM bis-Tris, 20 mM Tris, 20 mM CAPS buffer at pH 7.0 at 25 °C was excited at 360 or 330 nm, respectively. The fluorescence intensity at 455 nm was normalized to the first data point (0 mM NTP) and plotted *versus* the nucleotide concentration. b) Alteration in the fluorescence of the HCE-Gly52_R r₂ species upon the binding of ATP (green) and CTP (red) at pH 8.3. Conditions similar to those in panel a.

Gly52_R r₂ was monitored, the fluorescence response of each nucleotide exhibited a biphasic binding isotherm, consistent with high and low affinity sites. Furthermore, both nucleotides exhibit similar affinity to both classes of binding sites; however, the difference in affinity between the high and low affinity sites is even greater than that observed for the binding of these nucleotides to the holoenzyme (compare Figure 4, panels a and b). Although these experiments do not distinguish between a preexisting or induced asymmetry, the observed asymmetry does exist upon the binding of nucleotides and allows the two binding sites on one regulatory subunit to be distinguished.

Competitive or Synergic Binding of CTP and UTP. The functional significance of the enzyme having two classes of regulatory sites, which are distinguishable by their different affinity, may explain why UTP can only inhibit the enzyme in the presence of CTP (*3*). To test this, the fluorescence response to the binding of CTP and

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Figure 5. Interaction of ATCase with CTP and UTP. a) Alteration in the fluorescence of the HCE-GLy52_R ATCase at pH 8.3 upon the binding of UTP to the enzyme-CTP complex. The red circles represent saturation with CTP only, and the blue circles represent saturation with UTP in the presence of 2 mM CTP. b) Alteration in the fluorescence of the HCE-Gly52_R ATCase upon the binding of CTP to the enzyme-UTP complex. The blue circles represent saturation with UTP only, and the red circles represent the saturation with CTP in the presence of 2 mM UTP. HCE-Gly52_R AT-Case (50 μ g mL⁻¹) in 50 mM Tris-acetate buffer at pH 8.3 at 25 °C was excited at 360 nm. The fluorescence intensity at 455 nm was normalized to the first data point (0 mM NTP) and plotted *versus* the nucleotide concentration.

UTP alone and in combination was measured (Figure 5). The binding of CTP (Figure 5, panel a) or UTP (Figure 5, panel b) alone to the HCE-Gly52_R holoenzyme results in a biphasic reduction in relative fluorescence intensity, with maximal reductions of 49% and 51% for CTP and UTP, respectively, at the highest nucleotide concentration tested, 15 mM. The binding affinity of CTP was higher than that of UTP for the high affinity sites. In separate experiments, the binding UTP in the presence of 2 mM CTP (Figure 5, panel a) and the binding of CTP in the presence of 2 mM UTP (Figure 5, panel b) were determined. The second nucleotide dramatically reduced the fluorescence intensity, with UTP reducing the intensity of the enzyme.CTP complex an additional 22%, and CTP reduced the intensity of the enzyme-UTP complex an additional 14%. The binding of CTP to the enzyme-UTP complex resulted in a less substantial reduction in fluorescence that may also be obtained with much higher saturation of UTP alone. However, the total inhibition with the combination of both nucleotides

was significantly higher than that observed for any one nucleotide binding in isolation at the concentrations tested. Thus, at the concentration that CTP exhibits maximal inhibition (1-2 mM), UTP can still bind and cause a further diminution in fluorescence.

Notably, CTP binds to the enzyme with similar affinity regardless of the presence of UTP and *vice versa*. If CTP and UTP were competing for the same binding site, the affinity would be drastically reduced in the presence of the competing nucleotide. Instead, our results suggest the two allosteric sites on a regulatory dimer are asymmetric. Whether the asymmetry is induced or preexisting, our data indicate that irrespective of which nucleotide binds first, the binding of the second occurs independently. It is tempting to speculate that the two essentially identical allosteric sites are functionally asymmetric.

Relationship between Allosteric Site Saturation and **Enzyme Activity.** Functional asymmetry is supported by our experiments investigating the relationship between nucleotide binding and the corresponding extent of inhibition or activation of the enzyme. By comparing our fluorescent binding and our kinetic data for HCE-Gly52_R ATCase, determined under identical conditions, the percent saturation of regulatory sites compared to the percent of inhibition by CTP or activation by ATP was determined (Figure 6). These data indicate a cooperative relationship between binding and activity. Notably, for CTP (Figure 6, panel a), significantly more inhibition occurs at low saturation of the allosteric sites, whereas for ATP (Figure 6, panel b), significantly more activation occurs at high saturation of the allosteric sites. It is tempting to speculate that the sites causing CTP inhibition are not the same sites that are causing the ATP activation.

The typical cellular concentrations of CTP (0.5 mM) and ATP (6.0 mM) would be expected to be saturating in isolation; however, there is significant sensitivity of ATCase to these nucleotides in their physiological range due to competition with other nucleotides present in the cell (*19*). Our data can be used to explain how ATCase displays sensitivity over a broad range of nucleotide concentrations, since significant activation requires nearly full saturation of the allosteric sites by ATP, while significant inhibition by CTP requires only about half saturation of the allosteric sites. Thus it is possible that the response of the nucleotides may be related to their specific binding to one or the other of the



Figure 6. Relationship between the saturation of allosteric sites by CTP or ATP and inhibition or activation of enzyme activity. a) Percent saturation of regulatory sites *versus* the percent of maximum inhibition by CTP (red circles). The dashed lines delineate the approximate width of a 45° line due to experimental uncertainties. b) Percent saturation of regulatory sites *versus* the percent of maximum activation by ATP (green circles). Kinetic and fluorescence measurements performed at pH 8.3 as described in Methods.

asymmetric allosteric sites within each regulatory dimer of the enzyme.

These experiments present a practical application for unnatural amino acid mutagenesis and help to decipher the complex nature of allosteric regulation. The incorporation of HCE-Gly into the allosteric site of ATCase provides an extremely sensitive method to measure nucleotide binding at both the high and low affinity sites, even for UTP, which binds with relatively weak affinity. Binding measurements revealed similar affinities for all four nucleotides, indicating that strength of binding does not necessarily dictate the extent of allosteric activation or inhibition of enzyme activity. This observation was confirmed by relating binding measurements with kinetic measurements that showed CTP being able to effectively inhibit enzyme activity at low percent binding, whereas ATP effectively enhances enzyme activity only at high percent binding. Because of the multimeric nature of ATCase, allosteric inhibition can be accomplished by CTP binding to only a few of the six regulatory sites, but ATP has to bind to many to enhance it. In addition, CTP binds selectively to only a fraction of regulatory sites with UTP able to fill the residual ones with similar affinity regardless of the presence of CTP. These findings suggest that the two essentially identical allosteric binding sites of each regulatory dimer appear to be functionally asymmetric.

METHODS

Selection of a Site for Incorporation of the HCE-Gly To Monitor the Binding of the Regulatory Nucleotides. Initial evaluation of possible positions for placement of the HCE-Gly was performed visually using the structure of the ATCase CTP complex determined to 2.2 Å resolution (PDB entry 1ZA1) (21). Candidate positions required that the side chain of the HCE-Gly would not interfere with the binding of CTP or cause a shift in the backbone of the structure. Positions 3, 48, 52, and 58 were selected as possible locations for incorporation of the HCE-Gly. For each candidate position the HCE-Gly was built into the structure followed by conjugate-gradient energy minimization using CNS (22). During the minimization the position of the CTP and the backbone were fixed. For comparison, a base structure was determined by minimizing the wild-type structure. Next, each of the minimized HCE-Glv structures was compared to the minimized base structure. An acceptable site for incorporation of the HCE-Gly required that the side chain of the HCE-Gly fit into the structure with no bad contacts, no significant alterations in the positions of nearby side chains, and no direct interactions with CTP

Based upon the examination of the coordinates from these minimized structures, position 52 was selected for incorpora-

tion of the fluorescent amino acid because it is in close proximity to the allosteric site (approximately 5 Å from the δ phosphate of CTP), it is not highly conserved among species, and the space occupied by the large Glu52 side chain could accommodate the side chain of HCE-Gly.

Mutagenesis and Construction of Expression Plasmids. A TAG stop codon was introduced at the position corresponding to Glu52 of the regulatory chain of ATCase on plasmid pEK351 (23) by site-specific mutagenesis employing a Quick-Change Site-Directed Mutagenesis Kit (Qiagen) to produce plasmid pEK664. This plasmid contains the E. coli pyrBI operon with a His-tail added to the C-terminus of the catalytic chain. Site-directed mutagenesis was performed on pEK664 to introduce a Nco I site overlapping the ATG initiation codon of the pyrB gene to create pEK679. Finally, pEK679 and pBAD/JYAMB-4TAG-Myo (24) were digested with Nco I and KpnI. The 1550 base pair Nco I to KpnI fragment from pEK679 carrying the pyrBI gene and the backbone of pBAD/JYAMB-4TAG-Myo (24) were isolated after agarose gel electrophoresis using a QIAquick Gel Extraction Kit from Qiagen. These fragments were mixed and treated with T4 DNA Ligase in 1X T4 DNA ligase buffer supplied by New England Biolabs at 16 °C for 16 h to create plasmid pEK684. To generate a plasmid to express only the fluorescent regulatory chain, the 1000

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base pair *pyrB* region was removed from pEK684 by digesting with Ndel, isolating the backbone after agarose gel electrophoresis using the QlAquick Gel Extraction Kit from Qiagen, and treating with T4 DNA Ligase in 1X T4 DNA ligase buffer supplied by New England Biolabs at 16 °C for 16 h to create plasmid pEK692. The plasmids pEK684 and pEK692 have the *pyrBl* or *pyrl* operon, respectively, under the control of the arabinose promotor, the gene for resistance to tetracycline, and the gene for the *Methanococcus janaschii* amber suppressor tRNA that uniquely specifies HCE-Gly in response to the TAG codon.

Synthesis of L-(7-Hydroxycoumarin-4-yl)ethylglycine. The synthesis of the fluorescent amino acid HCE-Gly was completed as described previously (24). The product was verified using ¹H NMR and LC-MS (ESI).

Expression and Purification of HCE-Gly52_R ATCase and HCE-Gly52_R Regulatory Dimers. Plasmid pEK684 (to express the holoenzyme) or plasmid pEK692 (to express the regulatory chain) was cotransformed with pBK-CouRS-D8 into competent Top10 cells (Invitrogen) and plated on 2 YT plates (25) with 50 μ g mL⁻¹ kanamycin and 12.5 μ g mL⁻¹ tetracycline. A starter culture with the same antibiotics was used to inoculate 500 mL of liquid M9 glycerol minimal media supplemented with 0.5% casamino acids (Bacto), appropriate antibiotics, and 1.0 mM HCE-Gly. Cells were then grown at 37 °C to an OD₆₀₀ of 0.5, and protein expression was induced by the addition of 0.2% arabinose. After further incubation at 37 °C for 12 h, cells were harvested by centrifugation and lysed by sonication. This system has been reported to have a minimum of 95% incorporation purity for HCE-Gly into myoglobin determined by ESI protein mass spectrometric analysis (24).

Because the gene for the regulatory chain (pyrl) follows the gene for the catalytic chain (pyrB) on the pyrBI operon, excess catalytic chain was produced as a result of the inefficiency of suppression of the amber codon in the *pyrl* gene. The large excess of catalytic chain, in conjunction with the location of the His-tag on the catalytic chain, complicated the purification of the HCE-Glv52_P ATCase holoenzyme, requiring a three-step purification to be developed. The HCE-Gly52 $_{\mbox{\scriptsize R}}$ ATCase was first purified by column chromatography employing Chelating Sepharose Fast Flow (GE Healthcare) that was charged with Ni²⁺. The HCE-Gly52_R ATCase holoenzyme and contaminating ATCase catalytic subunit were eluted with 400 mM imidazole. The catalytic subunits were removed by column chromatography employing Phenyl Sepharose (GE Healthcare). The fractions containing the fluorescent HCE-Gly52_R ATCase were pooled and extensively dialyzed against 50 mM Tris-acetate buffer at pH 8.3. Purity was verified by both SDS (26) and nondenaturing (27, 28) PAGE.

The HCE-Gly52_R regulatory dimers were first purified by precipitation with 65% saturated ammonium sulfate. The precipitated material was suspended in buffer containing 50 mM Tris pH 8.3, 2 mM 2-mercaptoethanol, and 50 μ M zinc chloride. The solution was further purified using ion-exchange chromatography employing Quaternary Fast Flow (GE Healthcare) by elution with 0.5 M NaCl. A second step of column chromatography employing Phenyl Sepharose purified the regulatory dimers to homogeneity. Purity was verified by SDS-PAGE (*26*) (Supplementary Figure 1).

Aspartate Transcarbamoylase Activity Assay. The activity of ATCase was measured at 25 °C by the colorimetric method (29). Aspartate saturation curves (Supplementary Figure 2), ATP activation curves, and CTP inhibition curves (Supplementary Figure 3) were performed in duplicate. Assays were performed in 50 mM Tris acetate buffer, pH 8.3, or in a tripart buffer at pH 7.0 containing 20 mM bis-Tris, 20 mM Tris, and 20 mM CAPS buffer, in the presence of a saturating concentration of carbamoyl phosphate (4.8 mM). Data analysis of the steady-state kinetics was carried out as described previously (30). Fitting of

the experimental data to theoretical equations was accomplished by nonlinear regression. The data was analyzed using an extension of the Hill equation that included a term for substrate inhibition (*31*).

Steady-State Fluorescence Measurements for Nucleotide Binding to HCE-Gly52_R ATCase and HCE-Gly52_R Regulatory Dimers. All steady-state measurements were performed using a IASCO spectrofluorometer FP-6300 in a quartz SUPRASIL fluorescence cell (Hellma) with a slit width of 5 nm at 25 °C in 50 mM Trisacetate buffer pH 8.3, or the tripart buffer at pH 7.0, and HCE-Gly52_R ATCase or r_2 at a concentration of 50 μ g mL⁻¹. An excitation spectrum from 300-500 nm was acquired to determine the maximum excitation wavelength. Using an excitation wavelength of 360 nm at pH 8.3 or 330 nm at pH 7.0, a fluorescence spectrum from 400-600 nm was acquired in the absence of nucleotides with maximum emission at 455 nm. Microliter amounts of the nucleotides ATP, CTP, UTP, or GTP were added to the fluorescence cell up to a concentration of 15 mM. At each concentration of nucleotide, a fluorescence spectrum was measured. These tests were repeated using the same volume of buffer in place of the nucleotides to adjust for dilution. The tests were also repeated using free amino acid HCE-Gly and nucleotides in the absence of the HCE-Gly52_R ATCase. The maximum fluorescence intensity at each concentration was determined, and the data fit by nonlinear least-squares to an equation for binding at two classes of sites (Supplementary Tables 1 and 2).

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Supporting Information Available: This material is available free of charge *via* the Internet at http://pubs.acs.org.

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