Residues in the Polar Loop of Subunit c in *Escherichia coli* ATP Synthase Function in Gating Proton Transport to the Cytoplasm*

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Background: H\(^+\) transport through membrane-traversing ATP synthase mechanically drives bond formation.

**Results:** Metal chelation with cysteine residues in cytoplasmic domains near the surface of the membrane blocks proton transport.

**Conclusion:** Gating from the H\(^+\) channel traversing the lipid bilayer occurs in cytoplasmic loops of transmembrane proteins.

**Significance:** The proton transport pathway traversing ATP synthase extends beyond the lipid bilayer into the cytoplasm.

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Rotary catalysis in F\(_{1}\)F\(_{0}\) ATP synthase is powered by proton translocation through the membrane-embedded F\(_{0}\) sector. Proton binding and release occur in the middle of the membrane at Asp-61 on the second transmembrane helix (TMH) of subunit c, which folds in a hairpin-like structure with two TMHs. Previously, the aqueous accessibility of Cys substitutions in the transmembrane regions of subunit c was probed by testing the inhibitory effects of Ag\(^+\) or Cd\(^{2+}\) on function, which revealed extensive aqueous access in the region around Asp-61 and on the half of TMH2 extending to the cytoplasm. In the current study, we surveyed the Ag\(^+\) and Cd\(^{2+}\) sensitivity of Cys substitutions in the loop of the helical hairpin and used a variety of assays to categorize the mechanisms by which Ag\(^+\) or Cd\(^{2+}\) chelation with the Cys thiolates caused inhibition. We identified two distinct metal-sensitive regions in the cytoplasmic loop where function was inhibited by different mechanisms. Metal binding to Cys substitutions in the N-terminal half of the loop resulted in an uncoupling of F\(_{1}\) from F\(_{0}\) with release of F\(_{1}\) from the membrane. In contrast, substitutions in the C-terminal half of the loop retained membrane-bound F\(_{1}\) after metal treatment. In several of these cases, inhibition was shown to be due to blockage of passive H\(^+\) translocation through F\(_{0}\) as assayed with F\(_{0}\) reconstituted into liposomes. The results suggest that the C-terminal domain of the cytoplasmic loop may function in gating H\(^+\) translocation to the cytoplasm.

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F\(_{1}\)F\(_{0}\) ATP synthase utilizes the energy stored in an H\(^+\) or Na\(^+\) electrochemical gradient to synthesize ATP in bacteria, mitochondria, and chloroplasts (1–3). The ATP synthase complex is composed of two sectors, i.e. a water-soluble F\(_{1}\) sector that is bound to a membrane-embedded F\(_{0}\) sector. In eubacteria, F\(_{1}\) is composed of five subunits in an \(\alpha_{3}\beta_{3}\gamma\delta\varepsilon\) ratio and contains three catalytic sites for ATP synthesis and/or hydrolysis centered at the \(\alpha-\beta\) subunit interfaces. F\(_{0}\) is composed of three subunits in an \(a_{1}b_{2}c_{10–15}\) ratio in eubacteria and functions as the ion-conducting pathway (4–8). The c-ring composition in mitochondria varies between species, being eight for bovine and 10 in yeast (9, 10). Ion translocation through F\(_{0}\) drives rotation of a cylindrical ring of c subunits that is coupled to rotation of the \(\gamma\) subunit within the \(\alpha\beta\gamma\) hexamer of F\(_{1}\) to force conformational changes in the three active sites and in turn drive synthesis of ATP by the binding change mechanism (1–3, 11–14).

Subunit c of F\(_{0}\) folds in the membrane as a hairpin of two extended \(\alpha\)-helices connected by a short cytoplasmic loop. In *Escherichia coli*, 10 copies of subunit c pack together to form a decameric cylindrical ring with TMH1\(^2\) on the inside and TMH2 on the periphery (5, 15, 16). In the atomic resolution structures of H\(^+\)-translocating \(c_{14}\)-ring from spinach chloroplast (17, 18) and the \(c_{15}\)-ring from *Spiroplasma platensis* (19), the H\(^+\)-binding Glu, which corresponds to Asp-61 in *E. coli*, is located in TMH2 at the middle of the lipid bilayer. Several other neighboring residues form a hydrogen bonding network between TMHs 1 and 2 of one subunit and TMH2 from the adjacent subunit. A more recent structure of the \(c_{15}\)-ring from *Bacillus pseudofirmus* OF4 (20) revealed a more hydrophobic H\(^+\) binding site in which a water molecule is coordinated by the H\(^+\)-binding Glu and the backbone of the adjacent TMH2, an architecture likely shared by the *E. coli* c-ring. The loop of subunit c, particularly a conserved R/Q/N/P motif, is involved in binding the \(\gamma\) and \(\epsilon\) subunits of F\(_{1}\) and coupling H\(^+\) translocation to ATP synthesis/hydrolysis (21–30).

Subunit a consists of five transmembrane helices, four of which likely interact as a four-helix bundle (31–34). Subunit a lies on the periphery of the c-ring with TMHs 4 and 5 from subunit a and TMH2 from subunit c forming the a-c interface (35, 36). During ion translocation through F\(_{0}\), the essential Arg-210 on TMH4 of subunit a is postulated to facilitate the proto-
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nation/deprotonation cycle at Asp-61 of subunit c and cause the rotation of the c-ring past the stationary subunit a (2, 3, 37).

Chemical modification of cysteine-substituted transmembrane proteins has been widely used as a means of probing the aqueous accessible regions (38–40). The reactivity of a substituted cysteine to thiolate-directed probes provides an indication of aqueous accessibility because the reactive thiolate species is preferentially formed in an aqueous environment. The aqueous accessibility of subunits a and c in E. coli F<sub>0</sub> has been probed using Ag<sup>+</sup>, Cd<sup>2+</sup>, and other thiolate-reactive probes (37, 41–47). The results suggested the presence of an aqueous accessible channel in subunit a in the center of TMHs 2–5 extending from the periplasm to the center of the membrane (34, 36, 37, 44). Protons entering through this periplasmic access channel are postulated to bind to the essential Asp-61 residues of the c-ring with the protonation and deprotonation of Asp-61 driving c-ring rotation. A second putative channel was suggested at the interface of TMH4 of subunit a and TMH2 of subunit c, providing an exit pathway at the a-c interface leading from Asp-61 to the cytoplasm (47).

In previous studies (46, 47), we probed the thiolate reactivity of Cys substitutions in the transmembrane regions of subunit c based upon inhibition of function in response to Ag<sup>+</sup> or Cd<sup>2+</sup> treatment. Here we report on the reactivity of Cys substitutions in the cytoplasmic loop and at the N- and C-terminal ends of subunit c, completing the screening of nearly all residues in the subunit. The sensitivity of function to inhibition by Ag<sup>+</sup> and Cd<sup>2+</sup> proved to be widespread in the loop. Inhibition in this region was surprising because metal chelation to the transmembrane Cys thiolates was thought to inhibit function by blocking proton translocation through F<sub>0</sub>, a function in which the loop was not thought to directly participate. We sought to distinguish between several possible mechanisms of inhibition by Ag<sup>+</sup> and Cd<sup>2+</sup>. Our experiments support the well-characterized role of the loop in F<sub>1</sub> binding, which proved to be disrupted by Ag<sup>+</sup> and Cd<sup>2+</sup> in some substitutions, but also suggest involvement of a distinct section of the loop in gating H<sup>+</sup> transport to the cytoplasm. In light of this and other data, we hypothesize that the aqueous half-channel from the middle of the membrane to the cytoplasm continues through a cytoplasmic domain at the subunit a-c interface formed by cytoplasmic loops from both subunits.

EXPERIMENTAL PROCEDURES

Strain Construction—The mutagenesis procedure of Barik (48) described previously (46, 47) was used to generate the Cys substitutions in subunit c, and all substitutions were expressed in plasmid pCMA113 (41). Fused subunit c dimers containing a single Cys substitution in the C-terminal copy were generated from plasmid pPJ2CR, which encodes two c subunits joined by a short linker (49). For purification of wild type and mutant F<sub>1</sub>F<sub>0</sub> complexes, mutant uncE was excised from the pCMA113 derivative plasmid and transferred into pFV2 between the PflMI and BssHII restriction sites (50). Like pCMA113, pFV2 codes a Cys-less F<sub>1</sub>F<sub>0</sub> complex; however, a His tag is present at the N terminus of subunit β rather than subunit a (50). The presence of modifications in either plasmid was confirmed by DNA sequencing through the ligation sites. Mutant plasmids were transferred into the chromosomal unc (atp) operon deletion strain JWP292 (5) for growth and biochemical characterization or DK8 (51) for purification. Growth yields of the mutant strains on glucose minimal medium and succinate minimal medium agar plates were assayed as described previously (41).

ATP-driven ACMA Fluorescence Quenching—Inside-out membrane vesicles were prepared from JWP292 transformant strains, and protein concentrations were determined as described previously (46). Aliquots of membranes (1.6 mg at 10 mg/ml) were suspended in 3.2 ml of HMK-NO<sub>3</sub> buffer (10 mM HEPES-KOH, 1 mM Mg(NO<sub>3</sub>)<sub>2</sub>, 10 mM KNO<sub>3</sub>, pH 7.5) or HMK-Cl buffer (10 mM HEPES-KOH, 5 mM MgCl<sub>2</sub>, 300 mM KCl, pH 7.5) and incubated at room temperature for 10 min. ACMA was added to 0.3 μg/ml, and fluorescence quenching was initiated by the addition of 30 μl of 2.5 mM ATP, pH 7. Each experiment was terminated by adding nigericin to 0.5 μg/ml. The fluorescence measured after the addition of nigericin was used as the base line from which the relative quenching of the mutant membranes was calculated. Treatments with Ag<sup>+</sup> or Cd<sup>2+</sup> were carried out in HMK-NO<sub>3</sub> buffer or HMK-Cl buffer, respectively. Inhibition caused by metal treatment was reversed by adding dithiothreitol (DTT) to 2 mM or β-mercaptoethanol (β-MSH) to 4 mM after initiation of quenching with ATP.

Measurement of ATPase Activity—Typically, 10 μg of membrane protein were diluted into 0.9 ml of TM buffer (55.5 mM Tris-SO<sub>4</sub>, 0.2 mM MgSO<sub>4</sub>, pH 7.8). The assay was initiated by the addition of 100 μl of 4 mM ATP supplemented with 0.8 μCi/ml [γ<sup>32</sup>P]ATP. After incubation for 6 min at 30 °C, the reaction was stopped by the addition of 800 μl of 1.125% (w/v) trichloroacetic acid and immediately placed on ice for 10 min. Liberated phosphate was extracted from aqueous solution by adding 0.3 ml of 5 M H<sub>2</sub>SO<sub>4</sub>, 0.6 ml of 6% (w/v) ammonium molybdate, and 3 ml of benzene:isobutanol (1:1) followed by Vortex mixing. After phase separation, 0.5 ml of the organic phase was removed for scintillation counting. For treatments with DCCD, HMK-NO<sub>3</sub> buffer containing membrane protein at 0.5 mg/ml was supplemented with 50 μM DCCD from a 0.1 M ethanolic stock solution and incubated for 20 min at room temperature prior to additional treatment. For silver and cadmium treatments, membrane protein in HMK-NO<sub>3</sub> buffer was incubated with 40 μM AgNO<sub>3</sub> or 300 μM CdCl<sub>2</sub> for 10 min at room temperature. To measure F<sub>1</sub> dissociation, membranes at 0.5 mg/ml in HMK-NO<sub>3</sub> buffer were incubated at room temperature for 5 min with or without the inclusion of 40 μM AgNO<sub>3</sub> or 300 μM CdCl<sub>2</sub>. Half of this suspension was centrifuged at 541,000 × g for 10 min at room temperature to pellet the membrane vesicles. Membranes were exposed to metal a total of 15 min prior to the ATPase assay. A 10-μg aliquot from the centrifuged suspension and an equivalent volume of supernatant from the centrifuged suspension were transferred to TM buffer and assayed as above.

Purification of F<sub>1</sub>,F<sub>0</sub>—The His-tagged F<sub>1</sub>,F<sub>0</sub> complex was purified as described previously (50, 52). Membrane vesicles from 12 g of plasmid pFV2/strain DK8-transformed cells were suspended at ∼20–25 mg/ml in 10 ml of extraction buffer (50 mM Tris-Cl, 100 mM KCl, 250 mM sucrose, 30 mM imidazole, 5 mM MgCl<sub>2</sub>, 0.1 mM K<sub>2</sub>EDTA, 40 mM 6-aminocaproic acid, 15 mM p-aminobenzamidine, 0.2 mM DTT, 0.8% soybean phos-
Phosphatidylcholine (Sigma, Type IV-S), 1% octyl glucoside, 0.5% cholate, 0.5% deoxycholate, 2.5% glycrol, pH 7.5) and incubated at 4 °C with rocking for 30 min. Insoluble material was removed by centrifugation at 541,000 × g for 30 min. A 1.25-ml nickel-nitritriacetic acid column (His Select, Sigma) was pre-equilibrated with 5 volumes of extraction buffer, and the supernatant fraction of the extract was loaded at a flow rate of 0.18 ml/min. The column was washed with 10 volumes of extraction buffer. Bound protein was eluted with 5 ml of elution buffer (extraction buffer containing 400 mM imidazole) at a flow rate of 1 ml/min. The column was washed with 10 volumes of elution buffer.

Concentrations were determined using a modified Lowry method. Reconstituted ATP synthase was loaded on a chromatography column, which was washed with extraction buffer, and the fractions were analyzed for ATP synthase activity. Protein concentrations were determined using a modified Lowry method.

RESULTS

Phenotype and Metal Sensitivity of Cys Substitutions—We report the phenotype and metal sensitivity of function for Cys substitutions in the cytoplasmic loop and the N terminus and C terminus of subunit c. This study nearly completes the screening of subunit c with 71 of 79 positions tested for function by in vivo oxidative phosphorylation, ATP-driven H⁺ pumping activity, and its sensitivity to treatment with Ag⁺ and Cd²⁺, and the entirety of this data set is tabulated in Table 1. Most of the new Cys substitution mutants generated in this study grew nearly as well as wild type on both glucose and succinate minimal media (Table 1), indicating a functional oxidative phosphorylation system. Only the R41C substitution abolished growth on succinate, and this substitution was not further characterized. Previous studies determined that Arg-41 is essential for F₁ binding and intolerant of even conservative mutations (24, 25). Inverted membrane vesicles from most of the remaining Cys substitution strains were functional in ATP-driven proton pumping as assayed by ACMA fluorescence quenching (Table 1). For mutant F₁F₀ complexes that were functional in this assay, we tested the effects of 40 µM Ag⁺ and 300 µM Cd²⁺ on H⁺ pumping as described previously (46, 47). Sensitivity values are tabulated in Table 1. Notably, substitutions at most positions in the loop (residues 33–49) were strongly inhibited by Ag⁺ (Table 1). Only the G33C, K34C, and E37C substitutions at the C-terminal end of TMH1 showed minimal inhibition. Substitutions at positions 35 and 42–49 were also sensitive to inhibition by Cd²⁺ (Table 1).

Distinguishing the Mechanisms of Inhibition for the Cytoplasmic Loop Cys Substitutions—Ag⁺ and Cd²⁺ react with the thiolate form of Cys, which should predominate in an aqueous versus a lipid-exposed environment. In previous studies of Cys substitutions in transmembrane regions, sensitivity to a metal was thought to be indicative of aqueous exposure, and inhibition was thought to reflect direct obstruction of an aqueous proton translocation pathway (41–44, 46, 47). It is possible that substitution of a hydrophobic residue with Cys could introduce novel aqueous accessibility into subunit c. However, such a Cys residue would not be expected to be sensitive to metal inhibition of ATP-driven H⁺ pumping unless the Cys side chain packed in a position that was proximal to the native H⁺ translocation pathway. Cys substitutions outside the membrane should also have access to the bulk aqueous solvent except for those where the side chain is buried in the protein. Ag⁺ sensitivity is therefore less likely to reflect blockage of an aqueous proton channel unless the channel continues through a protein domain after exiting the lipid bilayer of the membrane. Alternatively, reaction of Cys in the loop region with a metal may disrupt the binding of F₁ or the coupling of F₁ to F₀ and thus prevent ATP-driven quenching while making membranes proton-permeable. Substitutions in this region have been shown previously to reduce the affinity of F₁ binding to F₀ (22–26), and modification of an eH38C mutation at the interface of subunits c and c with N-ethylmaleimide disrupts F₁ binding (56). To determine whether sensitive residues in the cytoplasmic loop of subunit c are directly involved in H⁺ transport and/or gating,
TABLE 1—continued

| Substitution | Colony size on succinate<sup>a</sup> | Growth yield on glucose<sup>b</sup> | ATP-driven quenching<sup>b</sup> | Inhibition of H<sup>+</sup> pumping<sup>c</sup> | Ref. |
|--------------|-----------------------------|-----------------------------|-----------------------------|-------------------------------|-----|
|              | mm                         | %                          | %                          | %                             |     |
| TMH2         |                             |                            |                            |                               |     |
| Q52C         | 2.1                        | 87                         | 61                         | 97 76                         | 46  |
| F53C         | 2.1                        | 91                         | 64                         | 99 21                         | 46  |
| F54C         | 2.1                        | 100                        | 6                          |                               | 46  |
| I55C         | 2.1                        | 95                         | 3                          |                               | 46  |
| V56C         | 2.1                        | 108                        | 68                         | 97 24                         | 46  |
| M57C         | 2.1                        | 80                         | 71                         | 97 78                         | 46  |
| G58C         | 2.1                        | 87                         | 57                         | 99 95                         | 46  |
| L59C         | 2.1                        | 90                         | 48                         | 84 90                         | 46  |
| V60C         | 2.1                        | 103                        | 65                         | 99 8                          | 46  |
| D61C         | 0                          | 65                         |                            | 62                            | 46  |
| A61C         | 1.1                        | 98                         | 78                         | 99 35                         | 46  |
| I63C         | 2.1                        | 100                        | 74                         | 98 93                         | 46  |
| P64C         | 2.1                        | 95                         | 69                         | 95 44                         | 46  |
| M65C         | 2.1                        | 94                         | 66                         | 97 28                         | 46  |
| I66C         | 2.1                        | 105                        | 73                         | 91 7                          | 46  |
| A67C         | 2.4                        | 100                        | 67                         | 91 9                          | 46  |
| M65C         | 2.1                        | 94                         | 66                         | 97 28                         | 46  |
| L68C         | 2.1                        | 100                        | 61                         | 96 87                         | 46  |
| R50C         | 2.0                        | 75                         |                            | 63                            | 46  |
| T51C         | 2.0                        | 100                        |                            | 63                            | 46  |

Irreversibility of Metal Inhibition Suggests Disruption of F<sub>i</sub> Binding to F<sub>0</sub>—If metal binding to a substituted Cys disrupts F<sub>i</sub> binding to F<sub>0</sub>, then removal of the bound metal may not restore function. Conversely, if metal binding simply blocks the H<sup>+</sup> translocation pathway, then function would more likely be restored upon removal of the blocking metal. Binding of Ag<sup>+</sup> or Cd<sup>2+</sup> to a substituted Cys should be readily reversed by the addition of a metal-chelating agent such as DTT or β-MSH. The thiolates of the reduced forms of these reagents compete with protein Cys residues for the Ag<sup>+</sup> or Cd<sup>2+</sup> ions causing the inhibition. Extended incubation with DTT reversed Ag<sup>+</sup> inhibition in previous studies of subunit a (41), and both DTT and β-MSH have been used in electrophysiological experiments of the cystic fibrosis transmembrane regulator to rapidly reverse silver modifications of Cys substitutions (57). To determine whether treatment with metals disrupts the integrity of the F<sub>i</sub>F<sub>0</sub> complex, we attempted to rapidly reverse metal inhibition by treatment with DTT or β-MSH during ATP-driven H<sup>+</sup> pumping.

Inverted membrane vesicles were treated with 40 μM Ag<sup>+</sup> or 300 μM Cd<sup>2+</sup> and assayed for ATP-driven quenching of ACMA fluorescence. During the quenching assay, DTT or β-MSH was added shortly after the addition of ATP. For most transmembrane Cys substitutions in subunit c, addition of DTT or β-MSH immediately reversed inhibition as represented by quenching of ACMA fluorescence for the M57C substitution (Fig. 1A). Addition of DTT also reversed inhibition at several Cys substitutions in the loop as represented by the L48C substitution (Fig. 1B). The reversal effect is consistent with the removal of a blockage of the H<sup>+</sup> translocation pathway and was seen with most transmembrane substitutions as summarized in Table 2. On the other hand, ≤20% reversal of inhibition was observed for substitutions at positions 40, 42, and 49 in the loop region of subunit c (Fig. 1C and Table 2), and only partial reversal was observed for substitutions at positions 25, 35, 36, 38, 39, 43, and 52 (Fig. 1D and Table 2). Persistence of Ag<sup>+</sup> inhibition after the addition of DTT suggested that metal treatment caused an irreversible alteration of the
Cys-substituted F1F0 complex or a slowly reversible alteration where reversal is beyond the time course of the experiment. The lack of reversal is consistent with disruption of the interactions between F1 to F0 and possible loss of F1 from the membrane.

Effect of Ag+/H11001 on Dissociation of F1.—To investigate the possibility that the irreversible inhibition of proton pumping by metal binding at some positions in the cytoplasmic loop was caused by disruption of F1 binding to F0, we measured the effect of Ag+/H11001 treatment on the ATPase activity of the six substitutions in the c loop showing the least reversibility in Ag+/H11001-inhibited ATP-driven H+ pumping. Inverted membrane vesicles from each substitution were incubated in HMK-NO3 buffer with 40 μM Ag+/H11001, and then half of this suspension was centrifuged to remove the membranes and any bound ATPase. Equivalent volumes of the suspension before centrifugation and the supernatant fraction after centrifugation were assayed for ATPase activity. No significant release of ATPase activity from wild type membranes was observed even after metal treatment (Fig. 2).

For the three substitutions where the Ag+/H11001 inhibition of proton pumping was essentially irreversible (i.e. A40C, Q42C, and L49C), only a small fraction of ATPase activity was released from membranes without metal treatment. However, Ag+/H11001 treatment markedly increased the amount of the ATPase activity in the supernatant for each of these substitutions (Fig. 2A).

Similar results were seen for the G38C substitution. For the two substitutions showing more moderate reversibility, F35C and L36C, approximately a third of the ATPase activity measured in the uncentrifuged membrane suspension was present in the supernatant after centrifugation without metal treatment, sug-

| Substitution | Inhibition by Ag+/H11001 | Restoration by DTT |
|--------------|--------------------------|--------------------|
| TM1          |                          |                    |
| A20C         | 97                       | 88                 |
| A24C         | 97                       | 78                 |
| A25C         | 90                       | 47                 |
| L28C         | 97                       | 95                 |
| Loop         |                          |                    |
| F35C         | 96                       | 29                 |
| L36C         | 94                       | 34                 |
| G38C         | 96                       | 24                 |
| A39C         | 91                       | 65                 |
| A40C         | 96                       | 16                 |
| Q42C         | 96                       | 12                 |
| P43C         | 96                       | 70                 |
| D44C         | 97                       | 97                 |
| L45C         | 96                       | 94                 |
| I46C         | 97                       | 100                |
| L48C         | 94                       | 96                 |
| L49C         | 96                       | 20                 |
| TM2          |                          |                    |
| Q52C         | 97                       | 69                 |
| F53C         | 99                       | 90                 |
| V56C         | 97                       | 95                 |
| M57C         | 97                       | 96                 |
| G58C         | 99                       | 90                 |
| L59C         | 84                       | 99                 |
| V60C         | 99                       | 83                 |
| A62C         | 99                       | 88                 |
| I63C         | 98                       | 95                 |
| P64C         | 95                       | 75                 |
| M65C         | 97                       | 97                 |
| I66C         | 91                       | 85                 |
| V74C         | 87                       | 94                 |
| M75C         | 91                       | 79                 |

a Inhibition of quenching of ACMA fluorescence after the addition of ATP but before the addition of DTT.

b Percentage of quenching response restored by DTT relative to the maximum quenching in the absence of Ag+.

c Constructed with an A20P suppressor mutation.

d 4 mM β-mercaptoethanol used in place of 2 mM DTT.

Effect of Ag+ on Dissociation of F1.—To investigate the possibility that the irreversible inhibition of proton pumping by metal binding at some positions in the cytoplasmic loop was caused by disruption of F1 binding to F0, we measured the effect of Ag+ treatment on the ATPase activity of the six substitutions in the c loop showing the least reversibility in Ag+-inhibited ATP-driven H+ pumping. Inverted membrane vesicles from each substitution were incubated in HMK-NO3 buffer with 40 μM Ag+, and then half of this suspension was centrifuged to remove the membranes and any bound ATPase. Equivalent volumes of the suspension before centrifugation and the supernatant fraction after centrifugation were assayed for ATPase activity. No significant release of ATPase activity from wild type membranes was observed even after metal treatment (Fig. 2). For the three substitutions where the Ag+ inhibition of proton pumping was essentially irreversible (i.e. A40C, Q42C, and L49C), only a small fraction of ATPase activity was released from membranes without metal treatment. However, Ag+ treatment markedly increased the amount of the ATPase activity in the supernatant for each of these substitutions (Fig. 2A). Similar results were seen for the G38C substitution. For the two substitutions showing more moderate reversibility, F35C and L36C, approximately a third of the ATPase activity measured in the uncentrifuged membrane suspension was present in the supernatant after centrifugation without metal treatment, sug-
sensitivity of ATPase activity to DCCD, an inhibitor that specifically modifies Asp-61 of subunit c and blocks H⁺ translocation with a consequential inhibition of F₁-ATPase activity. To test the effects of metals (both Ag⁺ and Cd²⁺) on DCCD sensitivity, which is a measure of the coupling of F₁ to F₀, inverted membrane vesicles were incubated with or without 50 μM DCCD for 30 min at room temperature, and for the final 10 min of incubation, samples were treated with or without 40 μM Ag⁺ (or 300 μM Cd²⁺). The ATPase activity of these membrane suspensions was then assayed. For wild type membranes, Ag⁺ slightly stimulated activity (Fig. 3). DCCD inhibited wild type ATPase activity by 70 ± 5%, and inhibition was not significantly affected by metal treatment. The ATPase activities of L36C, G38C, A40C, Q42C, and L49C membrane suspensions were greatly stimulated by treatment with Ag⁺, and metal treatment strikingly reduced the DCCD sensitivity of the reaction, most strikingly for the A40C substitution (Fig. 3A). We observed similar effects after Cd²⁺ treatment of Q42C and L49C (data not shown). The enhancement of ATPase activity can be attributed to the release of F₁-ATPase from the membrane (as shown in Fig. 2) with removal of the drag resistance by the F₀ rotary motor on the F₁ rotary motor. The loss of DCCD sensitivity on metal treatment of the L36C, G38C, A40C, Q42C, and L49C substitutions supports the conclusion that F₁ is uncoupled from F₀. In contrast, the ATPase activity of Cys substitutions thought to be involved in H⁺ translocation (i.e. mutants where metal inhibition was reversed by DTT) was inhibited by Ag⁺ in a manner similar to the inhibition observed after treatment with DCCD (Fig. 3B). In the case of transmembrane residues I28C and G58C and loop residue L48C, DCCD inhibition of ATPase activity was not reversed by Ag⁺ (Fig. 3B).

FIGURE 2. Effect of Ag⁺ on the release of F₁ from the membrane. Inverted membrane vesicles were assayed for ATPase activity. White bars indicate the ATPase activity of untreated, uncentrifuged membranes. The substitutions tested showed inhibition of ACMA quenching by Ag⁺ that was irreversible (A) or reversible (B) by DTT treatment. Prior to the assay, membranes in HMK-NO₃ buffer were incubated at room temperature without metal (light gray bars) or with 40 μM AgNO₃ (dark gray bars). After 1 min, half of the membrane suspension was centrifuged to collect the membrane fraction. The light and dark gray bars indicate the ATPase activity measured in the supernatant fraction expressed as a ratio relative to the total ATPase activity of the untreated membrane suspension. Error bars represent the S.D. for each measurement.

FIGURE 3. Effects of Ag⁺ on total ATPase activity and its sensitivity to DCCD. Inverted membrane vesicles were assayed for ATPase activity. The substitutions tested showed inhibition of ACMA quenching by Ag⁺ that was irreversible (A) or reversible (B) following treatment with DTT. Prior to the assay, membranes were incubated for 30 min at room temperature in HMK-NO₃ buffer with no additions (solid bars) or 50 μM DCCD (hatched bars). These incubations were followed by treatment with or without 40 μM AgNO₃ (gray bars). Activity is expressed as a ratio relative to the activity of untreated membranes. Error bars represent the S.D. for each measurement.

suggesting that the Cys substitutions themselves cause a degree of disruption of F₁ binding to F₀ (Fig. 2A). Treatment with Ag⁺ did not significantly further increase the ATPase activity released from the membranes of these mutants. For substitutions classified as fully reversible (i.e. I28C, D44C, I46C, L48C, and G58C), only a small amount of ATPase activity was released without metal treatment, and treatment with Ag⁺ did not significantly increase the activity released to the supernatant (Fig. 2B).

Effects of Ag⁺ and Cd²⁺ on Total ATPase Activity and Its Sensitivity to DCCD—Disruption of the functional coupling of F₁ and F₀ during Ag⁺ or Cd²⁺ treatment with or without release of F₁ from the membrane surface should reduce the sensitivity of ATPase activity to DCCD, an inhibitor that specifically modifies Asp-61 of subunit c and blocks H⁺ translocation with a consequential inhibition of F₁-ATPase activity. To test the effects of metals (both Ag⁺ and Cd²⁺) on DCCD sensitivity, which is a measure of the coupling of F₁ to F₀, inverted membrane vesicles were incubated with or without 50 μM DCCD for 30 min at room temperature, and for the final 10 min of incubation, samples were treated with or without 40 μM Ag⁺ (or 300 μM Cd²⁺). The ATPase activity of these membrane suspensions was then assayed. For wild type membranes, Ag⁺ slightly stimulated activity (Fig. 3). DCCD inhibited wild type ATPase activity by 70 ± 5%, and inhibition was not significantly affected by metal treatment. The ATPase activities of L36C, G38C, A40C, Q42C, and L49C membrane suspensions were greatly stimulated by treatment with Ag⁺, and metal treatment strikingly reduced the DCCD sensitivity of the reaction, most strikingly for the A40C substitution (Fig. 3A). We observed similar effects after Cd²⁺ treatment of Q42C and L49C (data not shown). The enhancement of ATPase activity can be attributed to the release of F₁-ATPase from the membrane (as shown in Fig. 2) with removal of the drag resistance by the F₀ rotary motor on the F₁ rotary motor. The loss of DCCD sensitivity on metal treatment of the L36C, G38C, A40C, Q42C, and L49C substitutions supports the conclusion that F₁ is uncoupled from F₀. In contrast, the ATPase activity of Cys substitutions thought to be involved in H⁺ translocation (i.e. mutants where metal inhibition was reversed by DTT) was inhibited by Ag⁺ in a manner similar to the inhibition observed after treatment with DCCD (Fig. 3B). In the case of transmembrane residues I28C and G58C and loop residue L48C, DCCD inhibition of ATPase activity was not reversed by Ag⁺ (Fig. 3B).
Effect of Cd\(^{2+}\) on H\(^+\) Transport through F\(_{0}\)—To support the hypothesis that Ag\(^+\) and Cd\(^{2+}\) block H\(^+\) translocation in the Cys substitutions where inhibition was reversed by DTT, we directly measured the effects of metal treatment on H\(^+\) transport by Cys-substituted F\(_{0}\). We reconstituted wild type and mutant F\(_{0}\) into liposomes and measured the quenching of ACMA fluorescence driven by a ΔΨ. As described under “Experimental Procedures,” His-tagged F\(_{1}F_{0}\) was purified by detergent extraction and affinity chromatography and then reconstituted into unilamellar phosphatidylcholine liposomes. F\(_{1}\) was removed under conditions of high pH and low ionic strength, and the resultant F\(_{0}\) liposomes were loaded with K\(^+\). To measure H\(^+\) influx through F\(_{0}\), we diluted K\(^+\)-loaded F\(_{0}\) liposomes into a K\(^+\)-free buffer containing ACMA and added valinomycin to generate a ΔΨ in response to the K\(^+\) diffusion potential, which then drove H\(^+\) import into the liposomes through F\(_{0}\). Ag\(^+\) interfered with the valinomycin-driven quenching assay likely because of its affinity for valinomycin (58), so we only examined Cd\(^{2+}\)-sensitive mutants here. Wild type F\(_{0}\) and several mutant F\(_{0}\) complexes containing Cys substitutions in subunit c were purified and functionally reconstituted; however, complexes containing mutations F35C, D44C, L49C, M57C, G58C, and I63C were not successfully reconstituted. To determine whether Cd\(^{2+}\) when bound to a substituted Cys blocks H\(^+\) permeability, we supplemented the ΔΨ-driven quenching assay with various concentrations of Cd\(_{2}\)SO\(_{4}\). Wild type F\(_{0}\) was minimally sensitive with <15% inhibition being seen at 300 μM Cd\(^{2+}\) (Fig. 4). Cd\(^{2+}\) markedly reduced the quenching response of F\(_{0}\) liposomes with I28C or L59C transmembrane substitutions and markedly inhibited the quenching response of the I46C and L48C loop substitutions (Fig. 4). The Cd\(^{2+}\) inhibition of this quenching response, which directly measures passive H\(^+\) translocation through F\(_{0}\), strongly suggests that Cd\(^{2+}\) also inhibits ATP-driven H\(^+\) pumping in these four mutants by directly blocking H\(^+\) transport through F\(_{0}\). Relative to that seen in wild type, Cd\(^{2+}\) treatment resulted in somewhat greater inhibition of the quenching response with Q42C, L45C, or Q52C F\(_{0}\) liposomes (Fig. 4), which suggested that the structural changes causing uncoupling upon metal treatment may also result in minor effects on H\(^+\) movement through the loop region at the surface of F\(_{0}\). The relatively minor inhibition observed for these substitutions is consistent with the conclusion that uncoupling is the primary mechanism of metal inhibition at these positions.

Flexibility of the Cytoplasmic Loop—The length of the cytoplasmic loop of subunit c (i.e. the number of residues between the conserved GxGxGxGxG motif in TMH1 and the Asp-61 or equivalent residue in TMH2 is invariant among the c subunits of eubacteria and euarkytes (Fig. 5A) and an NMR study of the loop from E. coli subunit c suggested that it forms a rigid structure (59). We tested the structural flexibility of the region of the cytoplasmic loop implicated in H\(^+\) translocation by inserting or deleting 1–3 residues near Ile-46 (Fig. 5B). This region of the loop was chosen for alteration due to its low sequence conservation. Residues were genetically inserted or deleted by oligonucleotide-directed mutagenesis as described under “Experimental Procedures.” We then assayed the insertion or deletion mutants for ATP-driven H\(^+\) pumping to determine whether the alterations disrupted activity. To our surprise, inverted membrane vesicles from mutants in which Ile-46 was deleted or Ala was inserted after Leu-45 were functional in ATP-driven proton pumping as assayed by ACMA fluorescence quenching (Fig. 5C). Mutants in which Ile-46 and Leu-48 were both deleted or Ala-Gly-Ala was inserted after Leu-45 were not functional in this assay (Fig. 5C).

Metal-mediated Cross-linking at Some Cys Positions—A third effect of Ag\(^+\) may contribute to inhibition at several positions. Ag\(^+\) is multicoordinate and has been shown to simultaneously bind multiple thiolates (60). A Cys substituted into subunit c would be present in every subunit making up the decameric ring so that any substituted Cys would be in close proximity to the Cys in subunits on either side. Single Cys substitutions at the C-terminal end of cTMH2 have been shown to form dimers in the presence of Cu\(^{2+}\)-phenanthroline (15), and other cross-linking experiments indicate that cTMH2 is capable of significant mobility (61). To test the possibility that inhibition of H\(^+\) pumping could be caused by metal-mediated cross-linking, we have created genetically fused subunit c dimers that contain a single substituted Cys in only one copy of the two fused subunits. A c-ring assembled from these dimers would not position Cys in adjacent subunits, so formation of a metal-mediated cross-link would be unlikely. Inverted membrane vesicles isolated from these fusion mutants are competent in ATP-driven proton pumping as assayed by quenching of ACMA fluorescence (Table 3). When assayed for Ag\(^+\) sensitivity, most dimer substitutions retained the sensitivity observed in the Cys-substituted c monomer, suggesting that cross-linking is not the source of inhibition for these sensitive substitutions. Marked reductions in Ag\(^+\) sensitivity occurred for several substitutions on the periplasmic half of subunit c, including c’A20C, c’V60C, c’M75C, and c’V78C where the c’ substitution is placed in only the C-terminal subunit of the c-c’ dimer. A reduction in sensitivity to Ag\(^+\) resulting from increased spacing between Cys suggests that metal-mediated cross-linking may contribute significantly to inhibition at these positions. This conclusion for c’M75C and c’V78C is supported by Cu\(^{2+}\)-mediated cross-linking of these residues, suggesting increased flexibility in the C terminus (15). On the other hand, a more likely explanation for the c(wild type)-c’A20C and c(wild type)-c’V60C dimers may be a reduced disruption of structure in rigidly packed regions of the c-ring, especially for the V60C substitution, which lies on the buried side of TMH2 proximal to TMH1.

DISCUSSION

Previous studies demonstrated that Ag\(^+\) and Cd\(^{2+}\) ions and presumably H\(_2\)O can access Cys substitutions in the transmembrane regions of subunit c and defined an aqueous pathway by which protons can travel from the proton-binding Asp-61 in the middle of the membrane to the cytoplasmic surface (46, 47). In the current study, we surveyed the Ag\(^+\) and Cd\(^{2+}\) sensitivity of Cys substitutions in the polar loop and at the N- and C-terminal ends of subunit c and identified several Ag\(^+\)-sensitive regions lying outside of the membrane bilayer with distinctly different properties. Two distinct regions were identified in the cytoplasmic loop. In the first region including residues between...
positions 35 and 43 on the N-terminal side and 49 and 52 on the C-terminal side of the loop (Fig. 6), substitutions became uncoupled upon treatment with Ag\(^{+}\) or Cd\(^{2+}\), suggesting that metal treatment of Cys substitutions in this region disrupts F\(_1\) binding to the c-ring. In a second region of the loop including residues 44–48 (Fig. 6), metals inhibited ATP-coupled H\(^{+}\) transport without uncoupling F\(_1\) from F\(_0\), and Cd\(^{2+}\) was shown to block passive F\(_0\)-mediated H\(^{+}\) transport, suggesting that this was the mechanism of inhibition. Finally, in the C-terminal region at positions 74–78, inhibition appeared to be the result of Ag\(^{+}\)-mediated cross-linking between Cys in adjacent c subunits in the ring. This hypothesis is plausible given that Ag\(^{+}\) is bicordinate (60) and that Cys substitutions at positions 74, 75, and 78 form high yield dimers in the presence of Cu\(^{2+}\)-phenanthroline, suggesting structural flexibility in this region (15).

Most of the metal-inhibited Cys substitutions in the loop, including positions 35, 36, 38–43, 49, and 52, show defects in the binding or coupling of F\(_1\) to F\(_0\) as initially evidenced by the irreversibility of inhibitory effects of Ag\(^{+}\) by subsequent addition of DTT. At the positions studied in greater detail, metal treatment resulted in increased F\(_1\) release from the membrane with a resultant increase in ATPase activity, reduced DCBD sensitivity, and Ag\(^{+}\) inhibition of H\(^{+}\) pumping that was unaffected or only partially reversed by treatment with DTT (Figs. 2 and 3). These effects are consistent with previous mutagenesis results supporting the role of this region in F\(_1\) binding. Arg-41,
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**A)**

\[
\text{ATP-driven quenching by } e_3 \
\text{Inhibition by } Ag^\ast \\
\begin{array}{l|l|l}
\text{Substitution} & \text{ATP-driven quenching by } e_3 & \text{Inhibition by } Ag^\ast \\
A12C & 69 & 32 \\
A20C & 70 & 97^a \\
A24C & 43 & 97^a \\
A28C & 78 & 94 \\
A40C & 47 & 96 \\
A41C & 47 & 94 \\
A49C & 70 & 96 \\
Q52C & 18 & 95^a \\
F53C & 71 & 95^a \\
V56C & 55 & 96^a \\
G58C & 96^a \\
V60C & 97^a \\
I63C & 51 & 96^a \\
M75C & 74 & 91 \\
V78C & 65 & 75 \\
\end{array}
\]

*Reported in Ref. 47.*

*Reported in Ref. 46.*

**B)**

Alterations

\[
\text{del1} & \text{PDL_P_LLR} \\
\text{del2} & \text{PDL_PLLR} \\
\text{ins1} & \text{PDLAPPLLR} \\
\text{ins3} & \text{PDLAPGLLR} \\
\]

**C)**

Relative ACMA Fluorescence

\[
\text{ATP} \\
\text{del2, ins3} & \text{N} \\
\text{del1, ins1} & \text{WT} \\
\]

**TABLE 3**

Sensitivity of fused c dimers to Ag^+**

Gln-42, and Pro-43 form a conserved motif, and in general, mutations in this region are characterized by reduced binding affinity of the c-ring with F₁, increased dissociation of F₁, reduced DCCD sensitivity, and a defect in ATP-driven H⁺ pumping, all of which are indicators of defective coupling of F₁ and F₀ (21–27). The deleterious effect of a Q42E mutation in this motif can be suppressed by a second, compensating mutation at Glu-31 in subunit ε (26). Furthermore, disulfide cross-linking experiments indicate that Cys at positions 40, 42, 43, and 44 can be cross-linked to subunits γ and ε in F₁ (28, 29, 56).

A recent structure of the F₁, c complex from yeast mitochondria revealed the c-γε interface at high resolution and supports the electrostatic interaction of this region of the loop with the F₁ stalk (62). Ag⁺ or Cd²⁺ modification of a Cys at this position might easily alter the chemical environment required for F₁ binding. Despite its role as a binding site for the γε subunits of F₁, the loop is remarkably tolerant of substitution, even at the latter two positions in the highly conserved RQP motif. This tolerance is consistent with the results of a random mutagenesis study where mutants with substitutions at every position in the loop except for Arg-41 still supported growth on succinate (63). In a striking example of loop plasticity, we observed that ATP-driven H⁺ pumping activity is not significantly affected by the insertion of an Ala after Leu-45 or the deletion of Ile-46, even though the length of the loop is invariant across bacterial and eukaryotic c subunits (Fig. 5A).

We have also identified a region of the loop in which binding of metals to Cys block ATP-coupled H⁺ transport in a manner that was totally reversed by DTT as was the case for most transmembrane substitutions. This region includes positions 44–48.
and is continuous with the aqueous accessible cytoplasmic end of TMH2. Two of the loop substitutions also show passive F₀-mediated proton transport activity that was inhibited by Cd²⁺ in a manner akin to the inhibition seen with representative transmembrane substitutions where metals are thought to inhibit function by directly blocking proton transport through F₀. These two Cys substitutions, i.e. L46C and L48C, show no indication of dissociation of F₁ from F₀ on treatment with Cd²⁺ or Cd⁴⁺. Two other Cys substitutions in this region, D44C and L45C, also show complete reversal of inhibition by DTT and no evidence of uncoupling F₁ from F₀. In addition, metal treatment was shown to inhibit ATPase activity for several representative substitutions both in this region of the loop and in TMH substitutions in contrast to the metal-dependent stimulation of ATPase activity in mutants that became uncoupled (Figs. 2 and 3). Although the effects of Ag⁺ on the H⁺ permeability of F₀ could not be tested, the common chemistry shared by Ag⁺ and Cd²⁺ in sulfur chelation as well as the identical effects caused by both metals for many substitutions in the other assays reported here (i.e. inhibition of ATP-driven H⁺ pumping, reversibility of inhibition by DTT, and inhibition of ATPase activity) suggests that Ag⁺ also blocks H⁺ translocation in these regions (64).

These results support our previous interpretations (41–44, 46, 47) that the sensitivity of substitutions in transmembrane regions to Ag⁺ and Cd²⁺ is due to blockage of the H⁺ translocation pathway and that accessibility to the metals is provided by an aqueous channel that functions in F₀-mediated H⁺ transport.

The occurrence of metal sensitivity in the residue 44–48 region of the loop that is likely to be caused by steric or electrostatic blockage of H⁺ transport suggests that the H⁺ translocation pathway does not end at the surface of the lipid bilayer but rather continues through a protein domain into the cytoplasm. Although the structure of the E. coli c-ring has not been determined, several crystal structures of H⁺-translocating c-rings indicate that residues 46 and 48 lie on the end of TMH2 and project toward each other at the interface of two c subunits (17, 19, 20, 62, 65). Furthermore, residue 44 projects into the cytoplasm from the loop. The arrangement of the side chains of residues 46 and 48 suggests that the H⁺ translocation pathway may run at the interface of two c subunits packing against subunit a in this region. Recently, Moore et al. (45) reported on the Ag⁺ sensitivity of two Cys substitutions in the 1-2 loop and substitutions in the 194–199 region of the 3-4 loop of subunit a. Cross-linking experiments showed that the residues in the 1-2 loop, including M93C, lie in close proximity to the highly Ag⁺-sensitive L195C substitution. The authors concluded that these Ag⁺-sensitive regions pack into a single cytoplasmic domain that is somehow involved in the H⁺ translocation pathway, perhaps as part of a gating mechanism. Residues 194–199 in the 3-4 loop lie at the cytoplasmic end of TMH4 just outside of the membrane, and earlier cross-linking studies (35) indicate that an extensive face of the cytoplasmic side of αTMH4 packs against cTMH2 in the membrane. These observations suggest that the Ag⁺-sensitive residues in the 3-4 loop of subunit a lie in close proximity to the portion of the cytoplasmic loop of subunit c proposed in this study to be involved in H⁺ translocation. Given the expected proximity of aM93C to the region of the subunit c loop implicated here in H⁺ translocation, interaction between each of these Ag⁺-sensitive domains is a reasonable expectation. The exit channel at the aTMH4 and cTMH2 interface from the middle of the membrane to the cytoplasm may then be gated by interacting cytoplasmic domains of the polar loop of subunit c and the cytoplasmic loops of subunit a.

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