Visualization of a radical B\textsuperscript{12} enzyme with its G-protein chaperone

The MIT Faculty has made this article openly available. Please share how this access benefits you. Your story matters.

<table>
<thead>
<tr>
<th>Citation</th>
<th>Jost, Marco, Valentin Cracan, Paul A. Hubbard, Ruma Banerjee, and Catherine L. Drennan. “Visualization of a Radical B\textsuperscript{12} Enzyme with Its G-Protein Chaperone.” Proc Natl Acad Sci USA 112, no. 8 (February 9, 2015): 2419–2424.</th>
</tr>
</thead>
<tbody>
<tr>
<td>As Published</td>
<td><a href="http://dx.doi.org/10.1073/pnas.1419582112">http://dx.doi.org/10.1073/pnas.1419582112</a></td>
</tr>
<tr>
<td>Publisher</td>
<td>National Academy of Sciences (U.S.)</td>
</tr>
<tr>
<td>Version</td>
<td>Final published version</td>
</tr>
<tr>
<td>Accessed</td>
<td>Mon Mar 11 14:21:46 EDT 2019</td>
</tr>
<tr>
<td>Citable Link</td>
<td><a href="http://hdl.handle.net/1721.1/98024">http://hdl.handle.net/1721.1/98024</a></td>
</tr>
<tr>
<td>Terms of Use</td>
<td>Article is made available in accordance with the publisher's policy and may be subject to US copyright law. Please refer to the publisher's site for terms of use.</td>
</tr>
<tr>
<td>Detailed Terms</td>
<td></td>
</tr>
</tbody>
</table>
Visualization of a radical B$_{12}$ enzyme with its G-protein chaperone

Marco Jost$^{a,6}$, Valentin Cracan$^{b,1}$, Paul A. Hubbard$^{b,2}$, Ruma Banerjee$^{b}$, and Catherine L. Drennan$^{a,c,3}$

$^{a}$Department of Chemistry and $^{b}$Department of Biology, $^{c}$Howard Hughes Medical Institute, Massachusetts Institute of Technology, Cambridge, MA 02139; and $^{d}$Department of Biological Chemistry, University of Michigan, Ann Arbor, MI 48109-0600

Edited by Markus W. Ribbe, University of California, Irvine, CA, and accepted by the Editorial Board January 21, 2015 (received for review October 14, 2014)

G-protein metallochaperones ensure fidelity during cofactor assembly for a variety of metalloproteins, including adenosylcobalamin (AdoCbl)-dependent methylmalonyl-CoA mutase and hydrogenase, and thus have both medical and biofuel development applications. Here, we present crystal structures of IcmF, a natural fusion protein of AdoCbl-dependent isobutryl-CoA mutase and its corresponding G-protein chaperone, which reveal the molecular architecture of a G-protein metallochaperone in complex with its target protein. These structures show that conserved G-protein elements become ordered upon target protein association, creating the molecular pathways that both sense and report on the cofactor loading stage. Structures determined of both apo- and holo-forms of IcmF depict both open and closed enzyme states, in which the cofactor-binding domain is alternatively positioned for cofactor loading and for catalysis. Notably, the G protein moves as a unit with the cofactor-binding domain, providing a visualization of how a chaperone assists in the sequestering of a precious cofactor inside an enzyme active site.

metallocofactor delivery | vitamin B$_{12}$ | metallochaperone | G protein | crystallography

Metallocofactors are ubiquitous in biology and essential for many cellular processes. Their use comes at a price, however, because metallocofactors are expensive to biosynthesize and transport, can have associated toxicity, and must be correctly assembled into their target enzyme for activity. To address these challenges, nature often employs protein metallochaperones. Chaperone-assisted cofactor delivery limits cellular dilution, minimizes inadvertent reactivity associated with the free cofactor, and ensures proper enzyme assembly (1–3). The importance of metallocofactor trafficking is underscored by the manifestation of diseases when mutations affect trafficking proteins. One important group of metallochaperones comprises the small guanine nucleotide-binding proteins (G proteins) belonging to the SIMB1 class (after signal recognition particle, MinD, and BioD) of P-loop NTPases (4). The human G-protein chaperone, MMAA (methylmalonic aciduria type A protein, mitochondrial; gene product of cblA), is involved in the assembly of adenosylcobalamin (AdoCbl, coenzyme B$_{12}$)-dependent methylmalonyl-CoA mutase (MCM) (5, 6). Mutations in MMAA and MCM result in methylmalonic aciduria, a genetically inherited metabolic disease (7). Bacterial homologs of MCM are involved in assembly of other metalloproteins and include HypB for hydrogenase (8) and UreG for urease (9). These G-protein chaperones bind to their target proteins and then use their GTP hydrolase (GTPase) activity to assist maturation of the target with high specificity (10–12). The mechanisms by which these essential chaperones perform their functions, however, are not well established, in large part because of the absence of structural information on their complexes with target proteins. An extensively studied member of the G-protein chaperones is MeaB$^{12–14}$, a bacterial ortholog of MMAA. Its target MCM requires AdoCbl to catalyze the radical-mediated 1,2-rearrangement of methylmalonyl-CoA (CoA) to succinyl-CoA (Fig. SLA) (13), an essential step in the degradation of odd-chain fatty acids, cholesterol, and branched amino acids. In this reaction and those of related isomerases, AdoCbl serves as a radical reservoir, reversibly generating the working 5'-deoxyadenosyl radical via homolytic cleavage of its cobalt–carbon bond (13–15). It is critically important for these isomerases to be loaded with AdoCbl and not any other cofactor derivative, which would yield inactive enzyme. MeaB performs this gating function for MCM and uses its GTPase activity to enable cofactor delivery (12) from the adenosyltransferase that synthesizes AdoCbl (16, 17). MeaB additionally remains associated with MCM during turnover and reduces the rate of oxidative inactivation (18). Notably, MeaB is structurally and mechanistically distinct from the ATP-dependent chaperones for AdoCbl-dependent eliminating enzymes, such as diol dehydratase: these chaperones, also termed reactivases, bind to inactivated enzymes and use their ATPase activity to eject damaged cofactor but do not affect AdoCbl delivery (19). An ortholog of MeaB, MeaL, is found fused to isobutryl-CoA mutase. Termed IcmF (20), the fusion protein exhibits both GTPase activity and AdoCbl-dependent carbon skeleton isomerase activity, interconverting isobutryl-CoA and n-butyryl-CoA, as well as pivalyl-CoA and isovaleryl-CoA (Fig. S1 B and C) (20, 21). The latter activity is newly discovered and may have relevance to the biodegradation of branched compounds (22).

Author contributions: M.J., V.C., R.B., and C.L.D. designed research; M.J. and V.C. performed research; V.C., P.A.H., and R.B. contributed new reagents/analytic tools; M.J. and C.L.D. analyzed data; and M.J. and C.L.D. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission. M.W.R. is a Guest Editor invited by the Editorial Board.

Data deposition: The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org [PDB ID codes 4K6H (holo-IcmF GDP), 4K7C (apo-IcmF) and 4KCB (apo-IcmF GDP)].

$^{1}$Present address: Department of Molecular Biology, Howard Hughes Medical Institute, Massachusetts General Hospital, Harvard Medical School, Boston, MA 02114.

$^{2}$Present address: Department of Biomedical Sciences, Cedars-Sinai Medical Center, Los Angeles, CA 90048.

$^{3}$To whom correspondence should be addressed. Email: cdrennan@mit.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1419582112/-/DCSupplemental.
Herein, we present the crystal structure of IcmF from *Cupriavidus metallidurans* containing bound AdoCbl and GDP•Mg²⁺, providing what is, to our knowledge, the first visualization of the juxtaposition of a G-protein chaperone and its target AdoCbl-dependent mutase. In addition, the crystal structure of IcmF in the absence of cofactors reveals the conformational changes required for cofactor loading.

**Results**

**Conserved GTPase Elements Are Involved in G-Protein:Mutase Complex Formation.** To visualize the molecular architecture of a G-protein:mutase complex, we determined the crystal structure of *C. metallidurans* IcmF with bound AdoCbl and GDP•Mg²⁺ (holo-IcmF•GDP) to 3.35 Å resolution (Fig. 1, Fig. S2, and Table S1). Each protomer of the IcmF dimer consists of four functional domains: an N-terminal cobalamin (Cbl)-binding Rossmann-fold domain (residues 21–157), a G-domain (residues 169–417), a structured linker region involved in dimer formation (residues 418–579), and a C-terminal acyl-CoA substrate-binding domain with an (α/β)₉-triose phosphate isomerase (TIM) barrel fold (residues 580–1093) (Fig. 1A–C).

Both the Cbl- and substrate-binding domains of IcmF share high similarity to other known AdoCbl-dependent mutases, such as MCM and glutamate mutase (Fig. 1B and D, Fig. S3A–C, and Table S2) (23–26). As with these other enzymes, AdoCbl is bound to a Rossmann-fold domain in a “base-off/His-on” conformation with a His (His39) serving as the lower axial ligand to the cobalt (Fig. S3D and E) and the 5’-deoxyadenosyl group (5’-dAdo) positioned in the active site cavity on the C-terminal face of the substrate-binding TIM barrel. Acyl-CoA substrates are known to access the AdoCbl cofactor by threading through the TIM barrel (Fig. S3F), which can open and close (23, 27). Finally, the dimer architecture of IcmF is similar to other AdoCbl-dependent mutases, with substantial parts of the dimer interface formed by the substrate-binding domains (Fig. S3G–I). The linker region of IcmF, which is not conserved, additionally stabilizes the dimeric arrangement of the subunits.

Remarkably, the two G-domains of an IcmF dimer are located on opposite ends and do not share an interface (Fig. 1A), in contrast to all other G-protein metallochaperones characterized to date, which are dimers (28–31). Apart from this difference, the G-domain exhibits high structural similarity to other G-protein chaperones with a central seven-stranded β-sheet, which is flanked by α-helices on both sides (Fig. 1C and E and Table S2) (29, 30), housing the signature G-protein motifs (Fig. 2A). The catalytically essential Mg²⁺ is coordinated by the side-chains of Ser223, Asp262, and Glu310 and by a β-phosphate oxygen of GDP (Fig. 2A). Arg249 is in proximity to the Mg²⁺-binding site and seems ideally positioned to activate a water molecule for GTP hydrolysis, as predicted previously based on homology to HypB (32). Arg265, which was disordered in structures of MeaB alone, interacts with the phosphate groups of the bound GDP and could help stabilize the negative charge that builds up during GTP hydrolysis, thus playing the role of a cis-acting Arg finger.

This conserved nucleotide-binding site is located directly at the interface of the G-domain and the mutase substrate-binding domain (Fig. 1F), burying 667 Å² and 626 Å² on the G-domain and the substrate-binding domain, respectively. The substrate-binding domain contributes two hydrogen bonds to the GDP ribose, from the side-chain of Asn1092 and the backbone carbonyl of Glu973 (Fig. 2B). Glu973 is also involved in salt bridges with base specificity loop residues Arg361 and Lys358. Interestingly, Glu973 immediately precedes a long α-helix that spans the width of the TIM barrel and interacts with the Cbl-binding domain, thus connecting the nucleotide binding site with both mutase domains (Fig. 2B). Two other residues of the substrate-binding domain, Arg1091 and Asn1092, form hydrogen bonds across the interface via their backbone atoms to the side-chain of the substrate-binding domain, Arg1091 and Asn1092, form hydrogen bonds across the interface via their backbone atoms to the side-chain of the substrate-binding domain.

![Structure of IcmF](image_url)
of Arg265, which is likely involved in GTP hydrolysis as described above (Fig. 2B). It is notable how much of the G-protein:substrate-binding domain interface is comprised of residues from each domain that directly or indirectly contact GDP.

The interface between the G-domain and the Cbl-binding domain buries 969 Å² and 943 Å², respectively. Residues 258–261 of the G-domain, which were disordered in the structures of MeaB (29) and MMAA (33), form a short β-strand (Fig. 2B, dark blue) that hydrogen bonds to the outermost β-strand of the Cbl-binding domain (Fig. 2B, orange), creating an extended β-sheet. Interestingly, this domain-linking β-strand contains part of the GTPase switch I motif (Leu259) (Fig. 2A and B) known to undergo conformational changes during the GTPase cycle. There are a number of interactions made between residues in or near switch I that connect the GDP•Mg⁺⁻-binding site in the G-domain to the Cbl-domain. In particular, Asp262 of switch I and downstream Arg265 resemble a C-clamp, pointing toward and interacting with Mg⁺⁻ and GDP, respectively (Fig. 2B), whereas Arg265 points in the opposite direction and directly contacts the Cbl-domain via a bifurcated interaction with Gln75 and Glu76. Gln75, Glu76, Leu259, Asp262, Arg263, and Arg265 are all fully conserved in IcmF sequences (20).

Insights into MCM:G-Protein Complexes and Methylnalonic Aciduria from the IcmF Structure. Given the high structural similarity between the IcmF domains and the homologous domains of MCM and MMAA/MeaB, the IcmF crystal structure allows us to predict the architectures of MCM:G-protein chaperone complexes by 3D structural alignments. These structure-based docking models place MCM and the G proteins in the same relative positions as in IcmF, with complementary surfaces and no major clashes (Fig. 1G, and Figs. S4 and S5). In the modeled MCM:G-protein complexes (with MMAA or MeaB), GDP resides at the interface of the MCM substrate-binding domains and the G protein, where it can make similar interactions (Fig. S4A and B). At least 10 residues that contribute to the interface between the IcmF Cbl-binding and G-domains are conserved in MCMs and their G proteins (Fig. S6), including the switch I motif and the interface-forming β-strand in the G protein.

Interestingly, a number of residues in MCM and MMAA that are associated with methylnalonic aciduria, map to this predicted interface. These residues include the MCM mutations R616C (R28 in IcmF) and G642R (G54 in IcmF), and G188R in MMAA (G257 in IcmF) (Fig. 2B–D and Table S3). We expect all three of these mutations to substantially alter the MCM/MMAA interface: the MCM R616C mutation would disrupt several hydrogen bonds, whereas the MCM G642R or MMAA G188R mutations would introduce severe clashes. Similarly, two other disease-causing mutations, F573S and G648D, map to IcmF residues G1086 and L60, respectively. Both these residues are located at the mutase:G-protein interface in IcmF (Fig. S4D and E).

Asymmetry Within the Holo-IcmF+GDP Structure Highlights Subunit Flexibility. Comparison of the two protomers of the holo-IcmF+GDP dimer reveals a displacement of the Cbl- and G-domains in one chain (B) relative to the other chain (A), leading to an ∼85 Å² difference in buried surface area and an 8 Å shift of the Cbl cofactor (Fig. 3A and Fig. S7A). Whereas the interactions between the Cbl and its binding domain are unchanged, almost all interactions between the Cbl and the substrate-binding domain are lost in chain B, with reduction of the buried surface area from 839 Å² to 159 Å² and concomitant loss of 5’-dAdo. Although the Cbl-binding domain is farther from the substrate-binding domain in chain B, parts of the G-domain are closer to the substrate-binding domain, increasing the interface between the latter two domains from 646 Å² to 768 Å². A difference distance matrix plot (34) demonstrates that the Cbl- and the G-domains move as a single rigid body that is best described as a swinging motion around a hinge centered at the nucleotide-binding site of the G-domain (Fig. 3A and Figs. S7B and S8). Given its location at the hinge, the guanosine nucleotide moves by only ∼3 Å, whereas the Cbl, which is far from the hinge, is displaced by a larger distance of ∼8 Å.

Inspection of crystal packing suggests that the chain B Cbl-binding domain is held in this extended position by lattice
Comparison of Apo- and Holo-IcmF Structures Support Relevance of Domain Motions to G-Protein-Assisted Cofactor Delivery. To investigate the significance of the motion of the Cbl- and G-domains observed in the holo-IcmF•GDP structure, we determined the structure of cofactor-free IcmF (apo-IcmF) to 3.45 Å resolution (Table S1). This structure lacks both AdoCbl and GDP•Mg²⁺ but has the substrate N-butyryl-CoA bound to the Cbl. Comparison of the holo and apo structures reveals that the Cbl-binding domain of chain A is no longer tightly associated with its substrate-binding domain. The surface area between these domains has decreased from 1,502 Å² to 799 Å², and the Cbl-domain is moved by ~6 Å from its position in holo-IcmF•GDP (Fig. S9A). At the same time, the interface between the Cbl-domain and the G-domain remains intact with an unchanged average buried surface area and no loss of interactions, demonstrating again that the Cbl-domain and the G-domain are moving as one rigid body. This displacement is very similar to that in the holo-IcmF•GDP structure, and as before, it is best described as a swinging motion hinged near the nucleotide-binding site of the G-domain.

Notably, this “open” conformation in apo-IcmF exposes the AdoCbl binding site and the His39-containing loop known to be important for cofactor docking (16) and could therefore depict a structural intermediate in cofactor delivery. To test if this conformation of IcmF is capable of binding AdoCbl, we incubated a preformed apo-IcmF crystal with fresh AdoCbl and determined the resulting structure. Strikingly, the structure reverts back to the active conformation, with AdoCbl bound in the catalytically competent position (Fig. S9B). Thus, the open conformation is competent for cofactor binding, and cofactor binding triggers a conformational change into the active conformation.

The G-Protein Active Site Undergoes Nucleotide-Sensitive Conformational Changes. The structures of holo-IcmF•GDP and apo-IcmF also allow us to observe changes associated with GDP binding. In the absence of GDP in the apo-IcmF structure, three loops near the G-protein active site are disordered or exhibit conformational changes compared with the holo-IcmF•GDP structure (Fig. S9B). These loops include: (i) residues 252–256, which immediately precede switch II; (ii) residues 312–318, which include the last two residues of switch II and are located next to the residues corresponding to the recently identified switch III region in MeaB (36) (residues 333–344); and (iii) residues 281–291, which are packed between loops (i) and (ii). To examine whether these loop differences are a result of the presence of GDP and not other crystal-to-crystal variations, we incubated a preformed apo-IcmF crystal with GDP•Mg²⁺ and determined the resulting apo-IcmF•GDP structure to 3.25 Å resolution (Table S1). Notably, the loops become ordered (Fig. S9B) and adopt the same conformations as observed in the holo-IcmF•GDP structure. Thus, these crystallographic data support the responsiveness to GDP of these three loops in the G protein.

Discussion

G-protein metallochaperones constitute an important enclave of metal homeostasis regulators whose activity in cofactor delivery has been difficult to understand without structural data depicting a metallochaperone:target enzyme complex. The fortuitous fusion of two “stand-alone” proteins into one in IcmF has allowed us to obtain the requisite structural data to interrogate both the signaling pathways and the conformational dynamics related to cofactor delivery. With IcmF serving as a model system, we predict that the human and bacterial MCM:G-protein complexes will form such that their G proteins interact similarly with their mutase domains despite their differences in oligomerization state [bacterial MCM is a heterodimer of an active and an inactive chain (22, 23, 37) and human MCM is a homodimer (33, 38)]. Notably, our structure-based docking model places five pathogenic human mutations—four in human MCM and one in

Fig. 3. Conformational movements in IcmF. (A) Differential positioning of the Cbl-domain and G-domain of holo-IcmF•GDP in chain A (cyan/orange) compared with chain B (light blue/yellow) when superimposed by their substrate-binding domains (gray surfaces). Cbl is shown in pink (chain A) and black (chain B), and GDP is shown in yellow (chain A) and green (chain B). Mg²⁺ is shown as an orange sphere. S'-dado not shown for clarity. (B) Superposition of the TIM barrel β-strands of holo-IcmF•GDP chains A (dark green) and B (gray). Cbl colored as in A and in distances between corresponding Cα-atoms in Å. (C) Superposition of the TIM barrel β-strands of substrate-bound (dark blue) and substrate-free (gray) MCM [PDB ID codes 4R0Q (35) and 2R0Q (27), respectively]. Cbl is shown in pink (substrate-bound) and black (substrate-free).

contacts, whereas chain A makes fewer lattice contacts and is thus free to pack tightly against its substrate-binding domain (Fig. S7 C and D). Chain A represents an active state of the enzyme, with AdoCbl positioned in the barrel ready for catalysis, whereas chain B represents an inactive state with a cleaved Cbl cofactor positioned out of the active site.

The substrate-binding domains of the holo-IcmF•GDP dimer also exhibit structural differences (Fig. 3 and Fig. S8). Whereas the TIM barrel in chain A is intact, with all interstrand hydrogen bonds in place, the TIM barrel in chain B is split into two halves of four strands each (Fig. 3B). Comparing the two conformations of the substrate-binding domain to structures of MCM reveals that the closed barrel of chain A resembles the structure of substrate-bound MCM (27, 35), even though no substrate is bound in our holo-IcmF•GDP structure. The open barrel of chain B closely matches the structure of substrate-free MCM (Fig. 3C) (23, 27). Thus, chain B, which has the cleaved Cbl cofactor swung out of the active position, also exhibits a disrupted substrate-binding barrel. Although opening and closing of the substrate-binding barrel was previously associated exclusively with substrate binding, here we observe both states in the absence of substrate. The opening is instead accompanied by a displacement of the Cbl out of the active position.

The G-Protein Active Site Undergoes Nucleotide-Sensitive Conformational Changes. The structures of holo-IcmF•GDP and apo-IcmF also allow us to observe changes associated with GDP binding. In the absence of GDP in the apo-IcmF structure, three loops near the G-protein active site are disordered or exhibit conformational changes compared with the holo-IcmF•GDP structure (Fig. S9B). These loops include: (i) residues 252–256, which immediately precede switch II; (ii) residues 312–318, which include the last two residues of switch II and are located next to the residues corresponding to the recently identified switch III region in MeaB (36) (residues 333–344); and (iii) residues 281–291, which are packed between loops (i) and (ii). To examine whether these loop differences are a result of the presence of GDP and not other crystal-to-crystal variations, we incubated a preformed apo-IcmF crystal with GDP•Mg²⁺ and determined the resulting apo-IcmF•GDP structure to 3.25 Å resolution (Table S1). Notably, the loops become ordered (Fig. S9B) and adopt the same conformations as observed in the holo-IcmF•GDP structure. Thus, these crystallographic data support the responsiveness to GDP of these three loops in the G protein.

Discussion

G-protein metallochaperones constitute an important enclave of metal homeostasis regulators whose activity in cofactor delivery has been difficult to understand without structural data depicting a metallochaperone:target enzyme complex. The fortuitous fusion of two “stand-alone” proteins into one in IcmF has allowed us to obtain the requisite structural data to interrogate both the signaling pathways and the conformational dynamics related to cofactor delivery. With IcmF serving as a model system, we predict that the human and bacterial MCM:G-protein complexes will form such that their G proteins interact similarly with their mutase domains despite their differences in oligomerization state [bacterial MCM is a heterodimer of an active and an inactive chain (22, 23, 37) and human MCM is a homodimer (33, 38)]. Notably, our structure-based docking model places five pathogenic human mutations—four in human MCM and one in
MMAA—at the MCM:MMAA interface. We now posit that MCM F573S, R616C, G642R, and G648D, as well as MMAA G188R (Table S3) (7), lead to disease because they disrupt the MCM:MMAA interaction, thereby impairing AdoCbl delivery to MCM and causing methylmalonic aciduria. Biochemical data are available to support two of these predictions: the pathogenic G188R mutation in MMAA was shown to inhibit complex formation with human MCM (33) and the pathogenic R616C mutation in MCM was mimicked in a bacterial ortholog and exhibited reduced affinity for MeaB (12).

Our atomic level view of this G-protein:enzyme complex reveals that the switch I region lies at the interface between the G-domain and the Cbl-domain, where it can communicate a signal to regulate Cbl delivery. We have captured multiple structures of both open and closed forms of IcmF and every structure depicts the movement of the Cbl-domain as a concerted displacement of both the Cbl- and the G-domain, with the nucleotide-binding site in the G-domain serving as a hinge. With an interface between the Cbl- and G-domains that appears fairly rigid, the G protein is not just sending a remote signal for the Cbl-domain to move, it is accompanying the Cbl-domain in its movement. Additionally, our structures with and without GDP identify three conformationally flexible loops, which are either near or include residues of the common G-protein elements (switch I and II) and the MeaB/MMAA-specific element known as switch III. Notably, mutations of switch III residues in MeaB affect AdoCbl delivery (36).

Using these structural snapshots and previously published biochemical data (12), we can propose a molecular model for the involvement of G-protein metallochaperones in Cbl delivery (Fig. 4). Enabled by association with the G protein, the Cbl-domain can undergo substantial motions, equilibrating between open and closed conformations. In the open state, the Cbl-domain is partially dissociated from the substrate-binding domain, exposing the His-loop known to be important for cofactor docking (16) and allowing access to the binding pocket for the Cbl di-methylbenzimidazole tail. Indeed, we observe that the open conformation is capable of binding AdoCbl, returning to the catalytically competent closed conformation. AdoCbl delivery (16) should be promoted by GTP binding. Based on the position of GDP, the γ-phosphate of GTP would be positioned between switch I and switch II in a solvent-exposed region, where it could make direct contact with the adenosyltransferase. Alternatively, GTP binding could stabilize the open conformation of the Cbl-domain via the switch I region, but elucidation of the precise role of GTP awaits a GTP-bound structure. In our model, AdoCbl binding and GTP hydrolysis would favor release of the adenosyltransferase and collapse of the open state, sequestering the high value cofactor in its binding pocket. In this catalytically active state, the cofactor is secured by multiple interactions from both the substrate- and Cbl-binding domains, which should inhibit active site reopening and thus prevent inactivation because of loss of the Cbl upper ligand.

In addition to the molecular interactions observed between the G-domain and the Cbl-binding domain that are undoubtedly involved in cofactor delivery, we find that the GTPase active site is located directly at the interface with the mutase substrate-binding domain. This location provides a molecular explanation for the observation that chaperone GTPase activity is affected by binding to a target protein, as observed with MCM: MMAA/MeaB (33, 39). Sizable regions of the GTPase active site are disordered in structures of MeaB and MMAA alone, and likely become ordered upon complex formation to assume the positions observed in IcmF. In particular, complex formation between MCM and MMAA/MeaB is expected to position the conserved Arg (265 in IcmF and 108 in MeaB) at the G-protein:mutase interface such that it can facilitate GTP hydrolysis (Fig. 2B). Thus, by the repositioning of a few key side-chains through a protein:protein binding event, communication is established between the substrate-binding domain of the target protein and the GTPase active site of the metallochaperone.

This molecular communication could play a role in mediating conformational changes of the substrate-binding domain that have been observed crystallographically for MCM, with the TIM barrel split open in the absence of substrate and closed in its presence (23, 27). Here, we observe the same open and closed conformations of the TIM barrel, but the correlation is with the absence/presence of Cbl in the barrel as neither structure has substrate. Although the key role of the G-protein metallochaperone must be in AdoCbl loading, MeaB and MMAA also reduce the rate of inactivation of their target enzyme MCM at no cost to catalytic efficiency (6, 18). It is tempting to consider that this protection arises from the G protein assisting conformational changes of the TIM barrel to prevent formation of inactive enzyme states.

In summary, we have provided atomic resolution structures that depict the interaction between an AdoCbl-dependent mutase and its corresponding G-protein chaperone. These structures suggest a molecular pathway to explain the observed bidirectional signaling (12, 36) and indicate how large conformational changes between cofactor- and substrate-binding domains can be achieved to load one of nature’s most precious metallocofactors.

Materials and Methods

C. metallidurans icmF was amplified from genomic DNA and ligated into a pET28a vector (Novagen) at the NdeI and BamHI sites. C. metallidurans IcmF was purified as described previously (20, 21). IcmF-SeMet was expressed using the methionine biosynthesis inhibition method.

All forms of IcmF were crystallized at 25 °C by hanging-drop vapor diffusion. Purified IcmF at a final concentration of 12 mg/mL was supplemented with 300 μM AdoCbl, 1 mM GDP, and 3 mM MgCl2 to generate holo-IcmF•GDP. For IcmF without AdoCbl and GDP (apo-IcmF), the substrate n-butyl-CoA

---

**Fig. 4.** Schematic of proposed molecular model of Cbl delivery. Only one monomer of the dimer is depicted. See main text for details.
(BcoA, 10 mM) was included to stabilize the protein. To generate crystals of apo-IcmF-GDP, preformed apo-IcmF crystals were soaked with 1 mM GDP and 1.5 mM MgCl₂ for 6 h. To reconstitute crystals of apo-IcmF with AdoCbl (apo-IcmF-reconst), preformed apo-IcmF crystals were soaked with 1 mM AdoCbl for 6 h.

All IcmF crystals belong to space group H2221. A Se anomalous peak dataset for holo-IcmF–SeMet–GDP was collected at the Advanced Light Source (Berkeley, CA) at a wavelength of 0.97914 Å (2.663 eV). All other data were collected at the Advanced Photon Source (Argonne, IL). All data were collected at a temperature of 100 K. Data collection statistics are summarized in Table S1. The holo-IcmF–GDP structure was solved using SSe-single isomorphous replacement with anomalous scattering, whereas the structures of apo-IcmF, apo-IcmF–GDP, and apo-IcmF–reconst were determined by molecular replacement. The models were refined by iterative cycles of manual adjustment in COOT (40) and refinement in phenix (41). Crystallographic refinement of the structures yielded models that possess low free R-factors, excellent stereochemistry, and small rmsd from ideal values for bond lengths and angles. Refinement statistics are summarized in Table S1. Full experimental details are provided in SI Materials and Methods.

ACKNOWLEDGMENTS. We thank Yan Kung and Jeremy W. Setser for help with data collection, and Daniel P. Dowling for help with data processing. This work was supported in part by National Institutes of Health Grants GM069857 (to C.L.D.) and DK45776 (to R.B.), and by a Massachusetts Institute of Technology Poitrax pre-doctoral Fellowship (to M.J.). C.L.D. is a Howard Hughes Medical Institute Investigator. This work is based in part on research conducted at the Advanced Photon Source on the Northeastern Collaborative Access Team beamlines, which are supported by National Institute of General Medical Sciences Grant P41 GM103403 and the National Institutes of Health. This research used resources of the Advanced Photon Source, a US Department of Energy (DOE) Office of Science User Facility operated for the DOE Office of Science by Argonne National Laboratory under Contract DE-AC02-06CH11357. This work was also based in part on research conducted at the Advanced Light Source, which is supported by the Director, Office of Science, Office of Basic Energy Sciences, of the US Department of Energy under Contract DE-AC02-05CH11231.