Structure of Human B<sub>12</sub> Trafficking Protein CblD Reveals Molecular Mimicry and Identifies a New Subfamily of Nitro-FMN Reductases*  

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Background: Mutations in CblD, involved in B<sub>12</sub> (or cobalamin) trafficking, lead to disease.
Results: The first crystal structure of human CblD is reported.
Conclusion: CblD most closely resembles CblC, another cobalamin trafficking protein, and they belong in a new subclass with the nitro-FMN reductase superfamily.
Significance: Disease-causing mutations that impair cobalamin oxidation kinetics can now be localized on the CblD structure.

In mammals, B<sub>12</sub> (or cobalamin) is an essential cofactor required by methionine synthase and methylmalonyl-CoA mutase. A complex intracellular pathway supports the assimilation of cobalamin into its active cofactor forms and delivery to its target enzymes. MMADHC (the methylmalonic aciduria and homocystinuria type D protein), commonly referred to as CblD, is a key chaperone involved in intracellular cobalamin trafficking, and mutations in CblD cause methylmalonic aciduria and/or homocystinuria. Herein, we report the first crystal structure of the globular C-terminal domain of human CblD, which is sufficient for its interaction with MMADHC (the methylmalonic aciduria and homocystinuria type D protein), or CblC, and for supporting the cytoplasmic cobalamin trafficking pathway. CblD contains an α+β fold that is structurally reminiscent of the nitro-FMN reductase superfamily. Two of the closest structural relatives of CblD are CblC, a multifunctional enzyme important for cobalamin trafficking, and the activation domain of methionine synthase. CblD, CblC, and the activation domain of methionine synthase share several distinguishing features and, together with two recently described corrinoid-dependent reductive dehalogenases, constitute a new subclass within the nitro-FMN reductase superfamily. We demonstrate that CblD enhances oxidation of cob(II)alamin bound to CblC and that disease-causing mutations in CblD impair the kinetics of this reaction. The striking structural similarity of CblD to CblC, believed to be contiguous in the cobalamin trafficking pathway, suggests the co-option of molecular mimicry as a strategy for achieving its function.

Cobalamins are B<sub>12</sub> derivatives that serve as essential cofactors for two housekeeping enzymes in mammals: the mitochondrial methylmalonyl-CoA mutase and the cytosolic methionine synthase (1, 2). Following the entry of cobalamin into cells, the concerted action of several enzymes and chaperones result in the interorganellar transport, maturation, and loading of the cofactor into the active sites of its target enzymes (see Fig. 1) (3–5). Dysfunction of proteins involved in the cobalamin trafficking pathway results in either isolated or combined homocystinuria and methylmalonic aciduria, with attendant complications (6). Early clinical genetic studies on patients with inherited disorders of cobalamin metabolism led to the identification of distinct complementation groups; cblA-G, cblI, mut, and cblX (7–15). The genes at the cblI and cblD loci encode proteins that have been named MMACHC (for methylmalonic aciduria cblI type, with homocystinuria) (9) and MMADHC (methylmalonic aciduria cblD type, with homocystinuria) (11). For simplicity, we refer to these proteins as CblC and CblD, respectively.

Both cobalamin-dependent enzymes are impaired in patients with mutations in CblC, indicating that this protein functions early in the trafficking pathway (16). In contrast, mutations in CblD can interrupt the function of either methylmalonyl-CoA mutase or methionine synthase or both enzymes (17). Consequently, patients with CblD dysfunction are classified into three distinct groups with isolated methylmalonic aciduria, isolated homocystinuria, and combined methylmalonic aciduria and homocystinuria. These clinical observations suggest that CblD functions downstream of CblC in the cobalamin trafficking pathway (see Fig. 1A) (17). Subcellular localization studies on CblD revealed its presence in both the cytoplasmic and mitochondrial compartments consistent with its suggested function at a branch point in B<sub>12</sub> processing (18). Four missense mutations have been described in CblD that are associated with isolated homocystinuria; i.e. they affect only the cytoplasmic branch of the trafficking pathway. These mutations are located in the C-terminal half of CblD (see Fig. 1B).

CblC is predicted to be the first protein that binds cobalamin in the cytoplasm as the cofactor exits the lysosome (see Fig. 1A). It is a multifunctional enzyme that catalyzes a range of activities including reductive decyanation of cyanocobalamin (19) and
dealkylation of alkylcobalamins (20, 21). The activity of CblC converts cobalamins to a common cob(II)alamin intermediate, which is subsequently partitioned into the biosynthetic active cofactor forms, methylcobalamin and 5′-deoxyadenosylcobalamin, required in the cytoplasm and mitochondrion, respectively. In contrast to CblC, the biological role of CblD has remained elusive. When first identified, it was speculated, based on weak primary sequence homology, to be an ATPase and/or to bind cobalamin (11). However, subsequent biochemical studies revealed that human CblD alone is incapable of binding cobalamin or hydrolyzing ATP (22, 23). So far, no ligand binding or enzymatic activity has been demonstrated for CblD. The only biochemical clue into CblD function derives from the observation that it binds to CblC, to which cob(II)alamin is bound (22–24). Based on these observations, it has been proposed that CblD serves as an adaptor protein, exerting its function in a protein-protein complex with CblC (4, 23, 25).

Progress in elucidating the function of CblD has been hampered by its lack of any obvious sequence similarity to any protein with known function and by the absence of structural information. To bridge this gap, we have solved the first crystal structure of the human protein truncated at the N terminus (CblDΔN108) to remove a region that is predicted to be highly disordered. The C-terminal domain is sufficient for interaction with CblC and for supporting the cytoplasmic cobalamin-processing pathway (11, 23). We demonstrate that CblD accelerates oxidation of CblC-bound cob(II)alamin to aquocobalamin (OH₂Cbl)⁴ and that missense pathogenic mutations in CblD differentially impact the kinetics of oxidation of CblC-bound cob(II)alamin. The crystal structure provides a framework for locating pathogenic mutations in CblD. Unexpectedly, the structure reveals that CblD exhibits a nitro-FMN reductase (NFR)-like fold despite the lack of sequence similarity to other family members. Within this family, CblD shares several distinguishing structural features with two other proteins involved in mammalian cobalamin metabolism, CblC, and the activation domain of methionine synthase (MSact) and with two recently identified protein family members. Within this family, CblD shares several distin-

**Structure of Human CblD**

The abbreviations used are: OH₂Cbl, aquocobalamin; NFR, nitro-FMN reductase; MSact, activation domain of methionine synthase; CblDΔN108, human CblD variant in which 108 N-terminal residues are deleted; SAD, single-wavelength anomalous dispersion; AdoMet, S-adenosyl methionine.

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**Experimental Procedures**

Chemicals—Reagents used for cloning were purchased from Agilent Technologies (Santa Clara, CA) and New England BioLabs (Ipswich, MA). Tris(2-carboxyethyl) phosphine and iso- propyl β-D-1-thiogalactopyranoside were from GoldBio Technical. All other chemicals were purchased from Fisher Scientific.

Cloning of CblD—CblDΔN108 was cloned into a pET-28b(+) vector (Novagen) using the NdeI and XhoI restriction enzyme sites to generate the N-terminal His₆-tagged construct containing a thrombin cleavage site. Oligonucleotides containing the bases corresponding to Ser¹⁰⁹–Met¹¹⁶ (5′-primer) and the last 24 bases (3′-primer) of the CblD cDNA (with restriction sites underlined) were used as primers: forward, 5′-CATATGAGTGAAAGAC ATGAGTTTGTAG-3′, and 3′-primer: 5′-CTCGAGTAATTTCGCACTTATCTCCATATGAC-3′. Construction of T182N, D246G, Y249C, and L259P pathogenic mutants of human CblD was performed using the QuickChange site-directed mutagenesis kit according to manufacturer’s instructions (Agilent Technologies Inc.). The pET-28b(+) vector encoding the human CblD residues 116–296 (CblDΔN108 background (23)) was used as a template with the following forward primers: 5′-ACTGTAACACAAAAAT- AAGATGATGACT-3′, 5′-ACTCTTTTGGAACTGGTT- GAACGCTACCGACAT-3′, 5′-GAAACTGATAAGCCTGC- CGCATAATTAGGATTGTC-3′, and 5′-HTTCCTGTGATGAC- CCGGGATCTGTAAGTG-3′ to generate T182N, D246G, Y249C, and L259P mutations. The mutagenic codons introducing asparagine, glycine, cysteine, and proline are underlined. The mutations were confirmed by DNA sequencing at the DNA Sequencing Core (University of Michigan).

Expression and Purification of CblD—Wild-type and mutant CblDΔN108 and CblDΔN108 proteins were expressed and purified as previously described (23). Purified protein was subjected to thrombin (GenTrac, Inc., Middleton, WI) treatment (2 units/mg protein) at 4 °C overnight, and the released His₆ tag was removed by purification through a second nickel-nitrilotriacetic acid chromatographic step. The flow-through containing the tagless CblDΔN108 protein was collected and concentrated using a YM-10 (Millipore, Billerica, MA) filter and loaded onto an $200 size exclusion column (120 ml, HiLoad 16/600 Superdex 200 PG; GE Healthcare, Pittsburgh, PA) to exchange the buffer to 50 mM Tris, pH 8, containing 0.5 mM Tris(2-carboxyethyl) phosphine. The selenomethionine derivative of CblDΔN108 (SeMet-CblDΔN108) was expressed like the unlabeled protein with the exception that the cells were grown in minimal medium supplemented with 50 mg/liter SeMet as described previously for CblC (28).

Oxidation of CblC-bound Cob(II)alamin—Cob(II)alamin was generated by photolysis of 5′-deoxyadenosylcobalamin in a sealed anaerobic vial, and full conversion to cob(II)alamin was monitored by UV-visible spectroscopy (conversion of the 525 nm absorbance to 475 nm). An anaerobic CblC stock solution was mixed with stoichiometric cob(II)alamin (300 μM each), and excess cob(II)alamin was removed using a Nanosep 10 K centrifugal device (Pall Life Sciences, Port Washington, NY). CblC-bound cob(II)alamin oxidation was then monitored aerobically at 20 °C in 0.1 M Hepes, pH 7.4, containing 150 mM KCl and 10% glycerol. For this, the CblC-bound cob(II)alamin stock solution was added to a 150-μl cuvette containing the same but aerobic buffer to give a final solution of CblC: cob(II)alamin (30 μM bound cobalamin). For experiments containing CblD, stoichiometric amount of wild-type or T182N, D246G, Y249C, and L259P (30 μM each) CblD was added to the aerobic buffer prior to addition of CblC-bound cob(II)alamin. Oxidation of cob(II)alamin to OH₂Cbl was monitored at 525 nm and was accompanied by the decrease in the cob(II)alamin absorbance at 475 nm. Control experiments were performed by adding bovine serum albumin at 1 mg/ml to the CblC-bound cob(II)alamin solution.
Crystallization Conditions—Stock solutions of CblD<sup>ΔN108</sup> (10 mg/ml) and of SeMet-CblD<sup>ΔN108</sup> in 50 mM Tris, pH 8, containing 0.5 mM Tris-(2-carboxyethyl) phosphate were used for crystallization trials. Crystals of both CblD<sup>ΔN108</sup> and SeMet-CblD<sup>ΔN108</sup> were obtained at 20 °C by the vapor diffusion method from 1:1 mixtures of protein and reservoir solution in sitting drop plates. Typical volumes of well and protein solution used for crystallization varied between 0.5 to 1.5 μl. The reservoir solutions for CblD<sup>ΔN108</sup> and for SeMet-CblD<sup>ΔN108</sup> contained 20% PEG3350, 0.1 M Tris-HCl, pH 7.5, 0.2 M MgCl<sub>2</sub>, 0.1 M NaF (for CblD<sup>ΔN108</sup>), and 20% PEG3350, 0.1 M Tris-HCl, pH 7.5, 0.185 M MgCl<sub>2</sub>, 0.12 M NaF (for SeMet-CblD<sup>ΔN108</sup>), respectively. Harvested crystals were cryo-protected by soaking in 16% PEG 3,350, 0.08 M Tris-HCl, pH 7.5, 0.16 M MgCl<sub>2</sub>, 0.08 M NaF, and 20% (v/v) glycerol for a few minutes prior to flash cooling in liquid nitrogen. Crystals of CblD<sup>ΔN108</sup> were of space group I<sub>2</sub>2<sub>2</sub>2<sub>1</sub> (a = 47.6 Å, b = 67.1 Å, c = 66.2 Å, and β = 110.5°) with two molecules in the asymmetric unit (Matthews’ coefficient VM = 2.3 Å<sup>3</sup>/Da for two molecules per asymmetric unit, and 46.8% solvent content). Crystals of SeMet-CblD<sup>ΔN108</sup> were of space group I222 (a = 65.3 Å, b = 66.2 Å, c = 71.8 Å) with one molecule in the asymmetric unit (Matthews’ coefficient VM = 1.8 Å<sup>3</sup>/Da for one molecule per asymmetric unit, and 32.2% solvent content). Detailed crystallographic information regarding data processing and refinement statistics is provided in Table 1.

X-ray Diffraction and Data Analysis—Diffraction data were collected at 100 K on Beamline GM/CA-CAT 23-ID-B at the Advanced Photon Source, Argonne National Laboratory (Argonne, IL). The data were recorded on a Mar300 detector and processed with HKL2000 (29). Phenix AutoSol (30) was used to identify the selenium sites and calculate density-modified 2.31 Å experimental maps based on a single-wavelength anomalous dispersion (SAD) data set from one SeMet-CblD<sup>ΔN108</sup>. Specifically, four of four selenium sites were located and used for SAD phasing, using phenix.hys. Subsequently, Phaser (31) was used to calculate the experimental phases, followed by density modification with RESOLVE (32) (figure of merit 0.34 before and 0.65 after density modification). The experimental density map showed clear features of the protein backbone and well defined side chains. RESOLVE traced and automatically built 120 residues and their side chains in the experimental electron density. COOT (33) was used to manually correct the incorrectly modeled residues, and through successive iterative rounds of refinement and manual model building, the remaining residues were traced in the electron density to afford the final model. Restained individual atomic refinement and restrained isotropic individual B-factor refinement with maximum likelihood targets using the Babinet model for bulk solvent scaling was performed using REFMAC5 (34) of the CCP4 suite (35). The final SeMet-CblD<sup>ΔN108</sup> model, after the removal of residues with high B-factors and water molecules was used as a starting model for the refinement of CblD<sup>ΔN108</sup> to 1.9 Å resolution. Initial simulated annealing refinement (torsional and Cartesian) was performed with phenix.refine followed by iterative rounds of refinement and model building/correcting with Refmac5 and Coot as described above. In the final CblD<sup>ΔN108</sup> model, residues 109–117, and 123–131 are missing in one monomer and residues 108–132, 169, and 170 in the second monomer in the asymmetric unit. The missing residues were not modeled because no visible electron density associated with these residues was found. The geometric quality of the models was assessed with MolProbity (36). PyMOL (37) was used to create molecular images.

Results

Construct Design—Attempts to crystallize full-length CblD failed despite considerable effort. We therefore analyzed the primary sequence by DisEMBL (38), PrDOS (39), and PSIPRED (40) to detect possible disordered regions at the protein termini. All three servers predicted that CblD is highly disordered and lacks any secondary structure at the N-terminal region spanning approximately the first 100 amino acids. Therefore, we designed a variant in which the first 108 residues of human CblD were deleted, which we refer to as CblD<sup>ΔN108</sup> (Fig. 1B).

Structure Determination—SAD phasing using the SeMet derivative was used to determine the crystal structure of CblD<sup>ΔN108</sup> (Table 1). Phenix Autosol was used to calculate the initial phases, and the initial experimental maps were obtained at 2.31 Å resolution. A complete model was built and refined for the SeMet derivative that was subsequently used for obtaining a solution for the wild-type CblD<sup>ΔN108</sup> structure to 1.9 Å resolution (Table 1). Because the structures of the wild-type and the SeMet derivative of CblD<sup>ΔN108</sup> were identical, the higher resolution wild-type structure is discussed in the remainder of the text. In the wild-type structure, we find two molecules of CblD<sup>ΔN108</sup> in the asymmetric unit, which is involved in dimer formation as discussed below. In the final refined model, we see clear and unambiguous electron density in both monomers for residues corresponding to the C-terminal region that spans from residues 132 to 296. The N-terminal region (residues 108–131), with the exception of a very short polypeptide (118–122) in one of the molecules in the asymmetric unit, was not modeled because no visible electron density corresponding to this region could be discerned. Therefore, the N-terminal region even in the truncated CblD<sup>ΔN108</sup> variant is unstructured and disordered.

Overall Fold and Structure Description—The CblD<sup>ΔN108</sup> structure exhibits a single domain of 152 residues (145–296) with an overall topologic arrangement characteristic of an α + β fold (Fig. 2). Specifically, the core of CblD<sup>ΔN108</sup> is comprised of a central four-stranded anti-parallel β-sheet sandwiched between four α helices and a β hairpin. The anti-parallel β-sheet is enclosed on one side by a very long (αC) and a short (αE) helix (Fig. 2A, left panel) and on the other side by two short helices (αB and αD) and a β hairpin (β1′-β2′) that are arranged perpendicular to the central β-sheet (Fig. 2A, right panel). The loops that connect the various secondary structure elements comprising the core of CblD<sup>ΔN108</sup> are numbered sequentially (Fig. 2B). This structural arrangement is comparable to that found in members of the NFR superfamily, which is discussed in detail later.

Dimer/Subunit Interaction—Although full-length CblD is reported to be a monomer in solution (22, 23), CblD<sup>ΔN108</sup> is found as a dimer in the asymmetric unit (Fig. 3, A and B). Despite its extended interdomain interface (1435 Å<sup>2</sup> buried surface) containing multiple ionic and hydrophobic interac-
tions, the dimer is most likely a crystallization artifact. The N-terminal region (referred to as the "dimerization arm"), comprising a small helix and two loops spanning residues 132–145, promotes dimerization and contributes mostly to the dimer interface. The dimer interface is assembled mostly via two types of contributions, an "arm and arm" and an "arm and body." In the "arm and arm" type of interaction, one dimerization arm and specifically the L2 region, interacts with the corresponding region (L2') in the adjacent monomer (Fig. 3B). In the "arm and body" interface, one dimerization arm and specifically the L1 region and helix and strand β1 in the adjacent monomer. In the crevice formed between the subunits at the "arm and body" interface, a combination of mainly hydrophobic and a few electrostatic interactions are involved (Fig. 3D).

Because both the full-length and the variously truncated forms of CblD reportedly behave as monomers in solution (22, 23), we used ultracentrifugation analysis to investigate the oligo-

FIGURE 1. The cobalamin trafficking pathway and boundaries of the CblDAN1108 construct used in this study. A, cobalamin first enters the lysosome from circulation, and CblC binds alkylcobalamin (R-Cbl) as it enters the cytoplasm via the action of two lysosomal membrane proteins. Following processing of cobalamin to a common cob(II)alamin (Cbl2++) intermediate, it is partitioned into the cytoplasmic and mitochondrial pathways for cofactor assimilation. Although the precise function of CblD is unknown, it functions downstream of CblC. CblE (methionine synthase reductase) provides reducing equivalents needed for the conversion of cob(II)alamin to methylcobalamin (MeCbl), the active cofactor form for CblG (methionine synthase). MS++ binds AdoMet and provides the methyl group needed for MeCbl synthesis. For clarity, details of the mitochondrial pathway, where cob(II)alamin is converted to adenosylcobalam- min (AdoCbl) and loaded into methylmalonyl-CoA mutase (mut) are not shown. The mechanism by which cobalamin enters the mitochondrion and the role of CblD in this process (denoted by a question mark) are unknown. B, boundaries of full-length CblD showing the N-terminal initiation site (Met1) as well as two internal initiation sites (Met62 and Met116). The locations of the missense patient mutations in CblD that result in isolated homocystinuria are indicated. The boundaries of the CblDAN1108 construct used for crystallography are shown below.

TABLE 1
X-ray crystallography data collection and refinement statistics

| Parameter                        | CblDAN1108 | SeMet CblDAN1108 |
|----------------------------------|------------|------------------|
| **Data collection**              |            |                  |
| Beamline                         | APS, GMCA 23-IDB | APS, GMCA 23-IDB |
| Wavelength (Å)                   | 0.979      |                  |
| Resolution (Å)                   | 50.0–1.90 (1.97–1.90) | 50.0–2.31 (2.39–2.31) |
| Space group                      | P2₁       |                  |
| Cell dimensions (Å)              | a = 47.6, b = 67.1, c = 66.2 | a = 65.3, b = 66.2, c = 71.8 |
| Cell dimensions (%)              | α = 90, β = 110.5, γ = 90 | α = β = γ = 90 |
| Unique reflections               | 30,281 (2986) | 6667 (4888) |
| Multiplicity                     | 3.2 (2.8)  | 7.0 (3.5)       |
| Completeness (%)                 | 98.1 (97.0) | 94.1 (70.4)     |
|<I/α>                             | 13.1 (2.5)  | 14.5 (2.5)      |
| Rmerge (%)                       | 7.9 (41.7)  | 9.8 (35.9)      |
| **Refinement**                   |            |                  |
| Resolution range                 | 62.01–1.90 | 48.71–2.31       |
| Number of reflections (work/test set) | 28,735/1523 | 633/320          |
| Number of atoms (protein/water)  | 2666/303   | 1266/21          |
| Mean B-factors Å²                | 26.2/33.6  | 45.2/36.7        |
| Rwork/Rfree (%)                  | 18.8/22.7  | 20.8/26.3        |
| Root mean square deviation from ideal values (bonds, Å) | 0.12 | 0.11 |
| Root mean square deviation from ideal values (angles, °) | 1.39 | 1.37 |
| Ramachandran plot (favored/outliers, %) | 99.4/0.0 | 96.1/0.0 |
| MolProbity Score                 | 1.01 (100th percentile) | 1.75 (97th percentile) |
| Protein Data Bank code           | 5CV0       | 5CUZ             |
meric state of CblD\textsuperscript{N108}. Based on this analysis, CblD\textsuperscript{N108} also behaves as a monomer in solution even at concentrations (10 mg/ml) that were used for crystallization (data not shown). Thus, the unique dimerization mode of CblD\textsuperscript{N108} as observed \textit{in crystallo} is clearly distinct from the dimerization mode of the CblD structural homologs that belong to the NFR family members and will be detailed later.

**Identification of CblD Structural Homologs—** Prior to this study, the cbl\textit{D} gene product was listed as an “uncharacterized conserved protein (DUF2246)” in the NCBI conserved domain database with no available structural homolog. Our initial DALI search (41) revealed that the closest structural homologs of CblD\textsuperscript{N108} are CblC, some members of the NFR superfamily (Conserved Domain Database accession number cl00514), \textit{e.g.} BluB from \textit{Sinorhizobium melloti} and flavin reductase P from \textit{Vibrio harveyi}, and MS\textsuperscript{act}, the activation domain of cobalamin-dependent MS. These proteins exhibit high structural similarity (z score >4) despite their very low sequence similarity (3–18% identity). In addition to these proteins, two corrinoid-dependent dehalogenases (26, 27) that are topologically similar to CblC (28) have been described recently.

CblD, CblC, NpRhdA (a corrinoid-dependent reductive dehalogenase from \textit{Nitratireductor pacificus} pht-3B), and MS\textsuperscript{act} share common structural features (within the blue rectangle), including a characteristic \(\beta\)-hairpin, which distinguishes them from other NFR family members (Fig. 4). In Fig. 4, the red dotted boxes represent the most conserved feature in all proteins described here including those belonging to a typical NFR family member, whereas the blue boxes include features exclusively shared by CblD, CblC, NpRhdA, and MS\textsuperscript{act}, but not...
Structure of Human CblD

FIGURE 3. Structure of the CblD^[4108] homodimer. A and B, two views of the CblD^[4108] dimer shown in ribbon representation with the subunits colored in blue and yellow, respectively. The N-terminal region including the extended α helix (αA) and the L2 loop (up to residue Arg^[145]) is the primary contributor of the dimer interface. C, close-up of the arm to arm subunit interface (pink dotted box in B). Amino acid residues involved in the interface analyzed by PISA are shown in stick representation. Hydrogen bonds and salt bridges are found in the loop region (Glu^[42]–Arg^[145], between αA and β1). Specifically salt bridges between Glu^[42] and Arg^[145], hydrogen bonds between the Glu^[42] side chain carboxylate and the Arg^[145] and Val^[146] backbone amido groups, and the Arg^[145] amine side chain with the Ser^[143] Glu^[142] and Arg^[145] hydrogen bonds and salt bridges are shown in the loop region (Glu^[142]–Arg^[145], between αA and β1). Specifically, the hydrophobic interface is lined up by residues Ile^[134], Ser^[136], Ala^[137], Tyr^[140], and Phe^[141] in the arm of one monomer and the residues Val^[404], Cys^[404], Val^[404], Phe^[404], Gly^[397], Ala^[408], Ile^[397], Ala^[397], Ala^[397], Ile^[397], Ile^[397], and Leu^[397] all found in a crevice formed from helices αC and αE and strand β1 of the adjacent monomer. The electrostatic interactions in the arm and body part are formed through hydrogen bonds between the Ala^[137] backbone amide and Leu^[397] carbonyl, the Glu^[142] side chain carboxylate and the Cys^[404] backbone amide, and a salt bridge between Glu^[143] and Lys^[292]. In C and D, hydrogen bonds and salt bridges are shown by dotted green lines.

by other proteins in the NFR superfamily. These conserved structural features are relevant to the cobalamin-related biological functions of these proteins as discussed later.

CblD Is Similar but Different from Other NFR Family Members—A detailed comparison of CblD was made with the NFR family member, BluB (Fig. 5A). Superposition of CblD^[4108] and BluB monomers (Fig. 5B) with Swiss-Pdb Viewer gives a root mean square deviation of 1.33 Å based on 53 of 152 structurally equivalent residues, 22% identity). The central anti-parallel β-sheet bundle is found in nearly identical positions in both proteins, whereas the arrangement of the surrounding α-helices is very similar. Of particular note is the near ideal superposition of a long α-helix (αC in CblD^[4108] (Fig. 2A) and αE in BluB (Fig. 5A), which is a structural hallmark of the NFR family. The majority of NFR family members exist as homodimers, and the long α-helix is important for dimerization. The flavin- and substrate-binding sites are located at the dimer interface in these proteins. In contrast, CblD and CblC do not form dimers in solution, and the dimerization interface seen in their crystal structures is distinct from that of other NFR family members in that it does not involve the long helix (28, 42).

Substantially more contact area (≈4800 Å²) is buried at the interface of the BluB dimer (Fig. 5C) than the CblD^[4108] dimer (≈1435 Å²). By superimposing two molecules of CblD^[4108] on a BluB dimer, a theoretical homodimerization model for CblD^[4108] was generated (Fig. 5D). The αC helices from both CblD^[4108] monomers in dark blue and yellow can theoretically align and assemble an interface similar to that in BluB. However, other interactions at the interface that are present in BluB and other NFR family members are missing in CblD^[4108]. These include the long N-terminal region including helix αA and the β-strand (β5) in BluB that are important in the dimer interface (Fig. 5C). Based on this analysis, we conclude that neither CblD nor CblC has the capacity to dimerize in the same mode as other members of the NFR superfamily, despite sharing a similar core subunit topology (Fig. 4).

CblC Is the Closest Structural Homolog of CblD—Although CblC and CblD share little sequence similarity, their overall topology is remarkably similar with a root mean square deviation value of 1.5 Å based on 70 Ca atoms (Fig. 6, A and B). Solution of the crystal structure of CblC led to the recognition that it resembles some members in the NFR superfamily (28, 42). CblC and CblD share an identical core of antiparallel β-sheets with similar structural elements in comparable arrangements framing the core. Both proteins exhibit superimposable long helices (αC in CblD and αE in CblC), and the arrangements of the structural components between strands β3 and β4 are very similar. Specifically, following strand β3, a long loop (L7 in CblD spanning residues 226–245 and L9 in CblC spanning residues 103–116) is present, followed by a short helix (αD in CblD and αF in CblC). Furthermore, a β-hairpin (β1’ and β2’) and a loop connecting to strand β4 (L10 in CblD (residues: 267–274), and L13 in CblC (residues: 151–158)) are found in perfectly superimposable positions in both proteins. Although the residues spanning the β1 and β2 strands adopt different conformations in the two proteins, they are nonetheless found in approximately similar locations.

Despite the similarities between the two proteins, there are also a number of differences. CblC has a four-helical cap comprising α1-αL (Fig. 6A) that is situated on top of the core domain and is absent in CblD. The long αE helix that facilitates
dimerization in other members of the NFR family is not continuous in CblC and therefore shorter than the corresponding helix in CblD (Fig. 6B). We had previously suggested that the presence of the four-helical cap in CblC forces what could be a continuous helix, such as /H9251C in CblD, to bend and break into two helices /H9251E and /H9251F (28). Another difference is that the loop following strand /H92523( L7 and L9 in CblD and CblC, respectively), is six amino acids longer in CblD. In CblC, this loop adopts two distinct conformations depending on the presence or absence of cobalamin. The binding of cobalamin induces a conformational change that allows loop L9 to move closer to the four-helical cap and to contribute to the cobalamin-binding pocket. Loop L7 in CblD adopts a conformation similar to that of loop L9 in CblC in the absence of cobalamin. Furthermore, the loop between the two β-strands in the β-hairpin, is approximately three times shorter in CblD. Finally, a functionally significant difference between the two proteins is the presence of a large cavity in CblC that accommodates cobalamin but is largely occluded in CblD (Fig. 6C). Indeed, attempts to dock cobalamin in the CblD cavity were largely unsuccessful. In addition, the four-helical cap forms part of the cobalamin-binding pocket in CblC and, as noted, is missing in CblD. Instead, the long /H9251C helix and the loop connecting to /H92523 occupy a region in CblD corresponding to the cobalamin-binding pocket in CblC. These large structural protrusions along with smaller ones would lead to steric clashes with cobalamin, explaining the observed lack of binding of this cofactor to CblD (22, 23).

**Topological Similarities between CblD, MS<sup> act</sup>, and Corrin-dependent Reductive Dehalogenases**—Cobalamin-dependent methionine synthase is a multidomain protein in which the terminal module constitutes the activation domain (Fig. 7A). MS<sup> act</sup> binds S-adenosylmethionine (AdoMet) and is important
for repairing inactive cobalamin bound in the preceding domain (43, 44). Human MS\textsuperscript{act} is approximately twice the size of CblD\textsubscript{AN108} and therefore exhibits additional features. Nevertheless, topological similarities are evident between them in their core regions (Fig. 7B).

The crystal structures of two corrinoid-dependent reductive dehalogenases, NpRdhA (26) and PceA (27), were recently reported. Both enzymes harbor two iron/sulfur clusters and bind one corrinoid cofactor per monomer. NpRhdA shown in Fig. 7C is a monomer and binds the corrin cofactor at the C terminus, whereas PceA is a homodimer. The corrinoid-binding domains in NpRhdA and PceA are similar to that of CblC and distinct from the classic Rossmann fold-containing motif found in other cobalamin proteins (45, 46). The CblD structure superimposes very well on the corrin-binding domains of NpRhdA (Fig. 7D) and PceA (data not shown). It is interesting that although CblD does not bind a corrin cofactor, its topology is more similar to that of CblC and the reductive dehalogenases than to other member of the NFR superfamily. We propose that these proteins (CblC, CblD, PceA, and NpRdhA) constitute a new subfamily within the NFR superfamily and are distantly related to the other NFR family members.

*Putative Ligand Binding Sites in CblD*—Because CblD does not bind cobalamin, we looked for similarities between CblD and the MS\textsuperscript{act} domain. Like CblD, MS\textsuperscript{act} does not bind cobalamin but binds AdoMet in a pocket (indicated by an arrow in Fig. 7A). The equivalent location in CblD is partially occupied by loop L7 and is thus unlikely to serve as a pocket for binding AdoMet. Interestingly, this equivalent location in CblD corresponds to a putative flavin-binding site in CblC that forms only when cobalamin is not bound (28). When cobalamin binds, structural changes to L9 (equivalent to L7 in CblD) open up the putative flavin-binding pocket. It is thus conceivable that conformational changes occur when CblD interacts with a physiological partner leading to reorganization in the region of the L7 loop and creating a ligand-binding cavity. Nonetheless, CblD alone is unable to bind ATP, GTP, AdoMet, NADP\textsuperscript{+}, or NADPH (data not shown).

*Location of Pathogenic Missense Mutations and Their Effects on CblD Function*—The majority of the disease-causing mutations in CblD result in either frameshifts or premature termination (11), leading to an expected loss of function. The four known missense disease-causing mutations can now be mapped on the CblD structure (Fig. 8A). Three of the four mutations cluster at one surface. D246G resides in loop 7, Y259C is in the aD helix, and L259P is in the \(\beta\)-hairpin. Interestingly, the position comparable to the aD helix and the \(\beta\)-hairpin in MS\textsuperscript{act} is proximal to the AdoMet-binding site or to
the site that interacts with the cobalamin-binding domain in methionine synthase (47). The fourth mutation, T182N, is located at the end of the β2 strand (Fig. 2B).

We have previously reported that CblD interacts with CblC albeit only when the product of the CblC reaction, cob(II)alamin, is bound to CblC (23). When reconstituted with cob(II)alamin, CblC protects it from further oxidation to OH₂Cbl (Fig. 8C, \( v = 0.067 \pm 0.009 \) nmol h⁻¹). Surprisingly, in the presence of stoichiometric CblD, the kinetics of cob(II)alamin oxidation is significantly enhanced (\( v = 1.81 \pm 0.09 \) nmol h⁻¹)).

All four pathogenic mutations diminish the rate of CblC-bound cob(II)alamin oxidation (T182N CblD, \( v = 1.33 \pm 0.19 \) nmol h⁻¹, D246G CblD: \( v = 1.24 \pm 0.10 \) nmol h⁻¹, Y249C CblD: \( v = 0.57 \pm 0.09 \) nmol h⁻¹, and L259P CblD: \( v = 0.47 \pm 0.05 \) nmol h⁻¹).

A phage display study was previously employed to identify putative peptides on CblD that might be important for its interactions with CblC. As shown in Fig. 8D, these peptides are diffusely scattered across the surface of CblD and therefore are of limited utility in delineating the CblD surface that interacts with CblC.

**Discussion**

Herein, we report the first structure of human CblD, a protein involved in intracellular cobalamin trafficking. Based on the clinical phenotypes of patients with the cblD disorder, the protein is predicted to play an important role in intercellular cobalamin delivery to both the cytosolic and mitochondrial targets, methionine synthase and methylmalonyl-CoA mutase (17). However, despite identification of the gene associated with the cblD locus in 2008 (11), insights into its function have been limited primarily because of its lack of sequence similarity to other proteins and its inability to bind cobalamin or other ligands that have been tested.

The cblD gene encodes a protein (from the Met¹ initiation codon; Fig. 1B) with a weak mitochondrial leader sequence at the N terminus, consistent with its proposed role in the mitochondrial branch of cobalamin metabolism (11). Replacement of the CblD leader sequence with a more efficient one from aldehyde dehydrogenase 2 improved adenosylcobalamin synthesis and cobalamin delivery to methylmalonyl-CoA mutase.
Patients carrying nonsense mutations at the 5′-end of the gene express N-terminally truncated CblD variants that are presumably translated by use of internal initiation sites, presumably Met62 or Met116 (Fig. 1B). These truncated variants can support cobalamin-dependent methionine synthase function (25). Based on these observations, it has been proposed that CblD functions in directing cobalamin delivery, with the N- and C-terminal portions of the protein being important for supporting methylmalonyl-CoA mutase and methionine synthase functions, respectively (11, 25). The CblD^{ΔN108} variant used in this study includes the internal Met116 initiation site and the residues where mutations have been reported in cblD patients exhibiting isolated methylmalonic aciduria or combined homocystinuria and methylmalonic aciduria. Hence, whereas the truncated protein used in our study lacks a large N-terminal segment that is presumed to be disordered, it is nevertheless relevant for understanding how CblD functions in the cytoplasmic branch of the cobalamin trafficking pathway.

The most striking feature of the ordered C-terminal half of CblD (residues 145–296) is that it exhibits a structural fold very similar to that found in CblC, which was not expected. We speculate that this architectural similarity despite the lack of sequence similarity could be functionally important for CblD, which is predicted to operate immediately downstream of CblC in the trafficking pathway (Fig. 1). The molecular mimicry might be functionally important if CblC and CblD compete for binding to the same surface on a target protein in the trafficking pathway. It is possible that CblC and CblD arose via a distant gene duplication event followed by loss of the cofactor-binding site, which led to the functional speciation of CblD.

We know that CblD forms a stable complex with CblC albeit only under conditions where the product of the CblC processing reaction, cob(II)alamin, is bound (23). In this study, we demonstrate that the CblC-bound cob(II)alamin becomes more prone to oxidation in the presence of CblD forming OH2Cbl, presumably by increasing access of oxygen to the cobalt ion in CblC (Fig. 8, B and C). The significance of this observation is presently not known and is being investigated. However, this initial observation is noteworthy on two accounts. First, because CblC displays a lower affinity for OH2Cbl than for cob(II)alamin, enhanced oxidation in the CblC-CblD complex might be important for its subsequent transfer to a donor protein. Second, all four pathogenic mutations reduce the rate of CblC-bound cob(II)alamin oxidation. These observations are consistent with the initially proposed adaptor function of CblD (4). The folded and globular C-terminal region is sufficient for complex formation with CblC (23) and for supporting the cytoplasmic branch of the trafficking pathway. In contrast, an additional and seemingly unstructured N-terminal region is needed for the mitochondrial branch of the trafficking pathway and presumably becomes ordered in complex with a partner protein.
Two of the members of the new NFR subfamily that we have described here (CblD and M5act) do not bind cobalamin directly but interact with proteins (CblC) or domains (B12 domain of methionine synthase) that do. The structure of the C-terminal half of methionine synthase comprising the cobalamin- and M5act domains (47) shows that the M5act module interacts via a unique M15-hairpin (B12-B2) with a propionamide side chain of cobalamin in the adjacent B12-binding module. The corresponding M15-hairpin element is found in identical positions in CblD, CblC and in the reductive dehalogenases. Interestingly, in the B12-binding proteins, i.e. CblC and the reductive dehalogenases, the M15-hairpin structure is directly involved in cofactor binding. In CblC, the corrin ring stacks against the hairpin, which also contributes to the wall of the groove into which the dimethylbenzimidazole moiety of cobalamin binds (28). We posit that the M15-hairpin structure in CblD competes with the equivalent structure in CblC in a protein-protein complex and interacts with the cobalamin. Interestingly, three of the disease-causing missense mutations are located in the vicinity of the M15-hairpin (Fig. 8A).

In summary, although the precise role of CblD remains elusive, the structure of its globular C-terminal domain reveals a surprising and unexpected structural similarity to CblC, with which this domain interacts. Together with the newly described structures of other corrinoid proteins that do not use the canonical Rossmann fold for binding B12, they have allowed delineation of a new subfamily within the NFR superfamily that share unique structural features. Finally, the CblD structure allows mapping of disease-causing mutations identified in patients with the cblD disorder.

Author Contributions—K. Y., C. G., R. B., and M. K. wrote the manuscript. C. G. expressed and purified the protein and performed biochemical analysis. K. Y. crystallized the protein, and K. Y. and M. K. collected crystallographic data, solved the structure, and performed structural analysis. All authors edited and approved the final version of the manuscript.

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