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Structural insight into metalloc cofactor maturation in carbon monoxide dehydrogenase

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Running title: Crystallographic studies of C-cluster assembly

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Abstract

The nickel-dependent carbon monoxide dehydrogenase (CODH) employs a unique heterometallic Ni-Fe-S cluster, termed the C-cluster, to catalyze the interconversion of CO and CO₂. Like other complex metalloenzymes, CODH requires dedicated assembly machinery to form the fully intact and functional C-cluster. In particular, nickel incorporation into the C-cluster depends on the maturation factor CooC; however, the mechanism of nickel insertion remains poorly understood. Here, we compare X-ray structures (1.50–2.48 Å resolution) of CODH from Desulfovibrio vulgaris (DvCODH) heterologously expressed in either the absence (DvCODH−CooC) or presence (DvCODH+CooC) of co-expressed CooC. We find that the C-cluster of DvCODH+CooC is fully loaded with iron but does not contain any nickel. Interestingly, the so-called unique iron ion (Feₙ) occupies both its canonical site (80% occupancy) and the nickel site (20% occupancy), with addition of reductant causing further mismetallation of the nickel site (60% iron occupancy). We also demonstrate that a DvCODH variant that lacks a surface-accessible Fe-S cluster (the D-cluster) has a C-cluster that is also replete in iron but lacks nickel, despite co-expression with CooC. In this variant, all Feₙ is in its canonical location and the nickel site is empty. This D-cluster-deficient CODH is inactive despite attempts to reconstitute it with nickel. Taken together, these results suggest that an empty nickel site is not sufficient for nickel incorporation. Based on our findings, we propose a model for C-cluster assembly that requires both CooC and a functioning D-cluster, involves precise redox-state control, and includes a two-step nickel-binding process.
Introduction

Anaerobic carbon monoxide dehydrogenases (CODHs) catalyze the reversible oxidation of CO to CO$_2$, enabling certain microbes, such as *Rhodospirillum rubrum* and *Carboxydothermus hydrogenoformans*, to live on CO as a sole source of carbon and energy (1,2). This microbial activity accounts for the removal of an estimated 10$^9$ tons of CO from the lower atmosphere each year, making CODHs an important part of the global carbon cycle (3). Given that CO is a toxic pollutant and a component of fossil fuel emissions, CODH has attracted attention as a possible bioremediation catalyst. Similarly, CODHs also have potential applications in the capture and removal of CO$_2$ via the Wood-Ljungdahl pathway of carbon fixation, in which the CO that is generated is incorporated into the acetyl group of acetyl-CoA (4). Use of CODH in such capacities would benefit from an ability to produce large amounts of active enzyme. In particular, CODH activity requires a complex heterometallic Ni-Fe-S cofactor (termed the C-cluster), the biogenesis of which is poorly understood.

The CODH C-cluster is housed within a homodimeric protein scaffold that contains two additional Fe-S clusters, termed the B- and D-clusters, that are used for electron transfer during catalysis (Figure 1A). The D-cluster, depending on the bacterial species, is either a [4Fe-4S] or [2Fe-2S] cluster that resides at the CODH dimer interface and serves as an electron conduit to external redox partners, such as ferredoxins; whereas the B-cluster is a [4Fe-4S] cluster that mediates electron transfer between the C- and D-clusters (5-9). The C-cluster is a structurally unique metallocluster composed of a distorted [Ni-3Fe-4S] cubane linked through a sulfide ion (S$_i$) to a mononuclear iron site (Fe$_n$) (Figure 1B) (5,6). This canonical C-cluster architecture is essential for catalysis, as it allows for binding of CO at the nickel ion of the cubane, activating it for nucleophilic attack by a water molecule ligated in immediate proximity at Fe$_n$ (10-13). To access this chemistry, organisms require dedicated cellular machinery for C-cluster assembly, similar to the requirements of other complex metalloclusters (14). Our understanding of the C-cluster assembly process, however, remains incomplete. It is still unknown what the biosynthetic origin of the Fe-S scaffold of the cluster is, how the Ni-Fe-S cluster is assembled, and what roles individual accessory proteins play in this assembly process.

Limited insight into the process of C-cluster assembly has been gleaned from the co-operonic expression of accessory proteins that appear to play roles in cluster maturation, in particular incorporation of nickel. Previous studies have shown that integration of nickel into the C-cluster is dependent on the accessory protein CooC (15-17). Certain organisms express additional proteins, CooJ and CooT, that have been implicated in C-cluster maturation; however, CooC appears to be the only dedicated and essential maturation factor expressed by all CODH-containing organisms (15,18-22). CooC is a P-loop ATPase with sequence similarity to UreG and HypB, maturation factors involved in nickel transfer to the active sites of urease and Ni-Fe hydrogenase, respectively (15,23). In analogy to UreG and HypB, CooC has been proposed to use ATP hydrolysis to facilitate nickel insertion into CODH (15,16,23,24). Alternatively, CooC has been proposed to fold or otherwise mediate formation of the proper nickel binding site in CODH (16,17,25).

To gain a further understanding of C-cluster maturation and the role of CooC, we have recently developed a means to heterologously express *Desulfovibrio vulgaris* CODH (DvCODH) in either the presence (DvCODH$^{CooC}$) or absence (DvCODH$^{−CooC}$) of the *D. vulgaris* CooC maturase (DvCooC) using *D. fructosovorans* as an expression host (17). This differential expression results in substantially different enzymatic phenotypes (Table 1). As-isolated DvCODH$^{CooC}$ binds about half of the expected nickel content and exhibits a lag phase in activity followed by a relatively low rate of CO oxidation (160 µmol·min$^{-1}$·mg$^{-1}$) as compared to the previously published activities of monofunctional CODHs from other species, which range from ~4400–16,000 µmol·min$^{-1}$·mg$^{-1}$ (17,26,27). Incubation of the as-isolated DvCODH$^{−CooC}$ with both NiCl$_2$ and the reductant sodium dithionite results in elimination of the lag phase and a 10-fold increase in CO oxidation activity; however, activation does not occur in the presence of NiCl$_2$ or sodium dithionite alone (17). In contrast, as-isolated DvCODH$^{−CooC}$ contains low amounts of nickel (0–0.2 Ni/monomer), has nearly no activity (4% of as-isolated DvCODH$^{CooC}$), and undergoes limited
activation with NiCl₂ and sodium dithionite (17), suggesting that DvCooC is involved in constructing the appropriate nickel binding site in DvCODH (25).

Interestingly, our previously published crystal structures of DvCODH-CooC revealed that the C-cluster adopts an alternative conformation upon exposure to oxygen in which the Ni, Feₚ, and Sₚ ions shift by as much as 3 Å and the Ni and Feₚ ions adopt new coordination environments (Figure 1C) (9). Notably, this oxidized conformation of the C-cluster can be converted back to the canonical, reduced conformation by incubation with reducing agent (9). The oxidized conformation involves ligation by a cysteine residue that is strictly conserved in CODHs but that does not serve as a ligand to the active, reduced conformation of the cluster (9). Mutation of this cysteine residue (Cys 301) in DvCODH-CooC to serine (C301S) in DvCODH⁺CooC to serine (C301S) was shown to result in inactive enzyme that does not bind nickel (Table 1) (9), similar to previous results on the CODH from Moorella thermoacetica (28). The crystal structure of DvCODH(C301S)⁺CooC revealed a partially assembled C-cluster in which Feₚ adopted a split conformation: at 70% occupancy Feₚ was in its canonical binding site; and at 30% occupancy Feₚ was incorporated into the cubane portion of the cluster, taking up the canonical Ni binding site (Figure 1D) (9). This split C-cluster conformation combined with the inability of the DvCODH(C301S)⁺CooC variant to incorporate nickel led us to propose that the oxidized conformation of the cluster could be an intermediate in C-cluster maturation, although how this conformation may participate in the assembly process remained unclear (9).

To further interrogate the process of C-cluster assembly, we have now determined crystal structures of DvCODH produced in the absence of CooC (DvCODH-CooC−) and of a DvCODH variant produced in the presence of CooC and engineered to not contain the surface-accessible D-cluster (DvCODH(ΔD)-CooC). Comparison of the DvCODH⁻CooC structure to that of DvCODH(C301S)⁺CooC (9) suggests a possible link between CooC-dependent cluster assembly and the ability to adopt the alternative, oxidized cluster arrangement. Furthermore, removal of the D-cluster leads to formation of an incomplete C-cluster, highlighting the importance of this redox active iron-sulfur cluster for C-cluster maturation. Combined, these results expand our understanding of C-cluster biogenesis, with an emphasis on the importance of accessing different cluster conformations and redox states.

Results

The C-cluster expressed in the absence of CooC is a [3Fe-4S] cluster with a mobile fourth iron. The CODH from D. vulgaris was expressed heterologously in D. fructosovorans in the absence of the C-cluster maturation factor CooC (DvCODH⁺CooC), as described previously (17). The preparation of protein that was used for crystallization displayed no detectible CO oxidation activity and contained 0 Ni/monomer and 10 Fe/monomer (Table 1). The crystal structure of DvCODH⁺CooC was determined to 1.50 Å resolution (Table 2). The structure aligns well (Cα r.m.s.d. of 0.18 Å for 1250 Cα atoms within the CODH dimer) with our previously determined structure of DvCODH⁻CooC (9) and the B- and D-clusters of the enzyme are both present and fully intact. Thus, the overall structure of DvCODH is retained when expressed in the absence of CooC.

At the C-cluster of DvCODH⁻CooC, the [3Fe-4S] partial cubane portion of the canonical C-cluster is intact and present at full occupancy (Figure 2), indicating that CooC is not necessary for formation of this part of the C-cluster. Modeling of the Feₚ ion, however, was more complicated. When the C-cluster was modeled as a [3Fe-4S]-Feₚ cluster at full occupancy, residual positive difference electron density was observed in the open cubane position, indicating the presence of an additional atom (Figure 2A). Further, iron anomalous difference maps (Table 2) reveal a shoulder extending from the canonical Feₚ binding site into the cubane position, suggesting the presence of iron at partial occupancy (Figure 2B). Given the lack of nickel in the sample that was crystallized, the positive difference density in the electron density maps, and the shoulder in the iron anomalous maps, we rationalized that Feₚ could be present in a split conformation. Refinement of Feₚ with a split conformation revealed that at 80% occupancy Feₚ is in its canonical binding site, ligated by His 266, Cys 302, and a water molecule; whereas at 20% occupancy, Feₚ is incorporated into the cubane...
portian of the cluster and ligated by Cys 519, forming a distorted [4Fe-4S] cluster (Figure 2C). Similar split Fe₄ conformations were observed across multiple crystal structures of DvCODH CooC samples that lacked nickel. Interestingly, this split Fe₄ conformation is similar to what was observed previously in our structure of DvCODH(C301S) CooC (9) (see Figure 1D), a CODH variant that is unable to adopt the alternative, oxidized C-cluster conformation due to the absence of the Cys 301 thiol for coordination to Fe₄. Together, the structural similarity between the C-clusters in DvCODH CooC and DvCODH(C301S) CooC (9) suggest a link between Cys 301 and the role of CooC in cluster assembly, perhaps due to a CooC-induced conformational change in which Fe₄ becomes ligated by Cys 301 (see Discussion). Additionally, these data suggest that it is not the lack of an open coordination site for nickel that prevents nickel incorporation into the C-cluster. Although there is some Fe₄ in the Ni binding site, there is not enough to explain the inability to reconstitute the C-cluster with nickel.

Reduction of DvCODH CooC induces movement of Fe₄ into the cubane position. The presence of Fe₄ at partial occupancy in the Ni binding site of the C-cluster in both our new structure, DvCODH CooC, and previous structure, DvCODH(C301S) CooC (9), is intriguing. Notably, the [3Fe-4S] clusters of aconitase and ferredoxins, as well as synthetic model compounds, are well known to incorporate exogenous metal into their open cubane site upon reduction due to the increased nucleophilicity of the open sulfide ions in the reduced state (29-33). Therefore, we hypothesized that reduction of the immature pre-C-cluster, which in part resembles a [3Fe-4S] cluster, could have led to movement of Fe₄ from its canonical binding site into the cubane position in some CODH molecules. To test this hypothesis, crystals of as-isolated DvCODH CooC were soaked in the reductant sodium dithionite prior to cryo-cooling and X-ray data collection. The structure of reduced DvCODH CooC was determined to 1.72-Å resolution (Table 2) and reveals greater incorporation of Fe₄ into the cubane position relative to the structure of the as-isolated enzyme (Figure 3A,B). Here, the Fe₄ ion resides in its canonical position at 40% occupancy and in the cubane portion at 60% occupancy (Figure 3C). Together, these data suggest that reduction of the pre-C-cluster before Ni is inserted can lead to mismetalation of the Ni site, and therefore careful control of cluster redox state is likely essential during the C-cluster maturation process in vivo.

The D-cluster is necessary for proper C-cluster assembly in the presence of CooC. To test the hypothesis that control of redox state is essential to C-cluster maturation, we sought to disrupt electron transfer between the C-cluster and external redox partners by removal of the solvent-exposed D-cluster, which serves as an electron conduit during CO/CO₂ interconversion. Towards this goal, we designed a DvCODH double-mutant variant in which the D-cluster-ligating cysteine residues (Cys 42 and Cys 45) were replaced with alanine residues to abolish binding of the D-cluster (DvCODH(DΔ)). This variant was expressed in the presence of CooC (DvCODH(DΔ) CooC) and purified to homogeneity. Similar to DvCODH CooC and DvCODH(C301S) CooC, DvCODH(DΔ) CooC is inactive as-isolated and does not contain appreciable amounts of Ni (Table 1). No increase in activity is observed upon incubation with nickel (Table 1). These observations indicate that the D-cluster is essential for C-cluster maturation.

To characterize the impact of a D-cluster deletion on C-cluster architecture, the crystal structure of DvCODH(DΔ) CooC was determined to 2.48-Å resolution (Table 2). The overall structure aligns well (Ca r.m.s.d of 0.29 Å for 1242 Ca atoms within the CODH dimer) with that of DvCODH CooC. The structure contains both the B- and C-clusters and confirms that the D-cluster is not present in this protein variant (Figure 4A). The absence of the D-cluster leads to local disorder, and residues 41–44 could not be modeled (Figure 4A inset). At the C-cluster of DvCODH(DΔ) CooC, we observe an intact [3Fe-4S]-Fe₄ scaffold with Fe₄ present at 100% occupancy in its canonical binding site (Figure 4B,C). This result is consistent with the above-mentioned idea that movement of Fe₄ into the cubane is induced by reduction and that the D-cluster mediates that reduction. Additionally, it is notable that the C-cluster of DvCODH(DΔ) CooC, in which 100% of Fe₄ is in the canonical location, cannot be activated by incubation with nickel. Further, the fact that the structure of DvCODH(DΔ) CooC is largely unchanged by D-cluster loss suggests that
it is the D-cluster’s redox role, rather than a structural role, that is required for nickel insertion.

**Discussion**

Here we present a series of crystal structures of *DvCODH* to provide insight into the process of C-cluster assembly and maturation, the mechanisms of which remained largely elusive. Our structures suggest that the C-cluster maturase CooC is primarily involved in nickel insertion rather than in formation of the [3Fe-4S]-Fe₄ scaffold and reveal that nickel insertion is additionally dependent on the D-cluster, likely due to its role in mediating electron transfer. Together, these findings allow us to propose a model for C-cluster assembly and maturation involving multiple cluster conformations and redox states.

In our structure of as-isolated *DvCODH*-CooC, we observe a largely (80%) intact Fe-S scaffold that contains 4 Fe ions and 4 S ions arranged as a [3Fe-4S]-Fe₄ “pre-cluster” that lacks nickel. The presence of this prearranged Fe-S scaffold in the absence of dedicated C-cluster assembly machinery suggests that the [3Fe-4S]-Fe₄ cluster arrangement can be formed using general Fe-S cluster biogenesis pathways, such as the SUF or NIF systems, both of which are present in the *D. fructosovorans* expression host as well as *D. vulgaris* itself. Two possibilities for the formation of the Fe-S scaffold can be envisioned (Figure 5A). First, the pre-C-cluster could be inserted in two pieces: a single iron ion inserted into the unique His 266/Cys 302 site and a [3Fe-4S] cluster inserted into the cubane site. Linkage of Fe₄ and the [3Fe-4S] cluster via the cubane sulfide (S₈) could occur subsequently (Figure 5A, upper pathway). Alternatively, the C-cluster binding site of CODH could become loaded with a [4Fe-4S] cluster that is distorted by CODH concomitant with the insertion step or is acted upon by an unknown maturation factor to remove an iron ion from the cubane, forming Fe₄ (Figure 5A, lower pathway). Regardless of the exact assembly mechanism, our data indicate that CooC is not necessary for formation of a 4-Fe containing Fe-S scaffold and that its primary role is likely in facilitating nickel insertion.

Once the Fe-S framework of the C-cluster has been assembled in CODH, nickel insertion can occur to form the fully mature and active cluster. Here we consider two possibilities for nickel insertion. In the first, C-cluster maturation in vivo involves the CooC-dependent insertion of nickel into a preformed [3Fe-4S]-Fe₄ scaffold that resembles our *DvCODH*-CooC structures with Fe₄ in its canonical site coordinated by His 266 and Cys 302 (Figure 5B). In the second, nickel is inserted into a [3Fe-4S]-Fe₄ scaffold in which Cys 301 coordinates Fe₄ (Figure 5C), a state that is reminiscent of the metal positions observed in our previous structure of fully oxidized *DvCODH*-CooC (Figure 1C) (9).

For scenario I (Figure 5B), the key role of CooC, in addition to nickel insertion, may be to control the redox state of the pre-C-cluster, allowing for nickel insertion without mismetallation of the nickel site. In analogy to metal capture by [3Fe-4S] clusters in other systems, nickel insertion into the scaffold as shown in Figure 5B would likely require that the [3Fe-4S] framework be in a reduced state to increase the nucleophilicity of the open cubane site, allowing for binding of exogenous metal (29-33). In the case of the C-cluster, however, addition of exogenous nickel is likely complicated by the presence of Fe₄, which we have shown can migrate into the open coordination site of the reduced cubane (Figures 2, 3). In the context of C-cluster assembly, this observation indicates that the redox state of the pre-C-cluster must be tightly regulated to avoid mismetallation. One strategy for ensuring Ni incorporation in vivo could be to couple binding of CooC with cluster reduction, such that cluster reduction occurs just prior to nickel insertion. For example, binding of CooC could in some way facilitate interaction of CODH with a low-potential electron transfer protein, such as a reduced ferredoxin.

Although control of cluster reduction provides one route to prevent mismetallation, the previously reported structure of a fully oxidized C-cluster (Figure 1C) (9) suggests another possible strategy for avoiding incorporation of Fe₄ into the cubane in vivo (Figure 5C). In particular, the position of Fe₄ in the oxidized cluster, ligated by Cys 301, could represent an alternative binding site in which Fe₄ is positioned prior to nickel insertion, such that Fe₄ is not ligated in immediate proximity to the remainder of the [3Fe-4S] scaffold (Figure 5C, state II). In this model, CooC could be involved in inducing a conformational change in the C-cluster prior to nickel insertion such that Fe₄...
becomes ligated by Cys 301 (Figure 5C, state I to II). Given the inability of the C-cluster to incorporate nickel in the absence of the D-cluster, this conformational change could additionally be redox-dependent. In any case, nickel could then be inserted into the His 266/Cys 302 binding site that is normally occupied by Fe₆, resulting in formation of the oxidized C-cluster conformation (Figure 5C, state IIIa). Subsequent reduction, possibly facilitated by a change in reduction potential as a result of nickel binding, would then trigger formation of the active C-cluster via the three-atom migration of Fe₆, S₁, and Ni that we have described previously and that occurs upon reduction of the oxidized cluster conformation (Figure 5C, state IIIa to IV) (9).

With these two proposals in mind (Figure 5B and 5C), we revisited the CODH literature. In addition to our previous characterization of DvCODH(C301S)₄CooC (9), several additional mutagenesis studies on the CODHs from Moorella thermaacetica (MtCODH) (28), Rhodospirillum rubrum (RrCODH) (34-36), and Carboxydothermus hydrogenofermum (ChCODH-II) (37) are better explained by the mechanism shown in Figure 5C than Figure 5B. First, the proposal in Figure 5B does not explain why substitution of the non-canonical C-cluster ligand Cys 301 in DvCODH and MtCODH results in inactive CODH variants that lack nickel (9,28), whereas the mechanism in Figure 5C provides a role for Cys 301 in nickel insertion. Second, substitution of the C-cluster-ligating histidine residue with valine (in RrCODH) or alanine (in ChCODH-II) resulted in CODHs with iron contents that were indistinguishable from wild-type, but that were impaired in their ability to incorporate nickel in vivo (34,37). Additional mutagenesis experiments in which each of the canonical C-cluster-ligating cysteine residues were mutated to alanine or serine, revealed that His 266 and Cys 302 (D. vulgaris numbering) are in fact the only protein-based ligands to the canonical C-cluster that are necessary for nickel incorporation (34-37). Together, these data support the hypothesis that the His 266/Cys 302 site serves as the binding site for nickel during nickel incorporation (Figure 5C). Second, the kinetics of nickel activation in nickel-deficient RrCODH (produced in the presence of RrCooC) suggest a two-step mechanism in which nickel first binds to the enzyme reversibly and then is seated into its active and stable site (26,36).

Collectively, these findings support a model in which Cys 301 binds Fe₆ while nickel is first inserted into the His 266/Cys 302 site, followed by rearrangement to form the active C-cluster (Figure 5C, upper pathway), consistent with structures of the C-cluster that we have observed experimentally (9). We note that an alternative assembly pathway could also involve coordination of Fe₆ by Cys 301 while nickel is inserted into its canonical site in the Fe-S cubane, although such a state has not been observed crystallographically (Figure 5C, state IIIb) and does not explain the RrCODH mutational data mentioned above. One caveat of this model is that we have only observed Fe₆ coordinated to Cys 301 in the oxidized state of the C-cluster (9), whereas the presence of reducing agent is known to be essential for nickel-dependent activation in vitro (17,26). That being said, it has not yet been possible to visualize a nickel-deficient form of DvCODH⁴CooC in either an oxidized or reduced state to know whether Fe₆ movement occurs and/or is redox-dependent in the absence of nickel.

Overall, our data begin to reveal the requirements for assembly of a fully intact and activatable C-cluster: 1) the C-cluster maturase CooC (this work)(15-17), 2) the D-cluster (this work), and 3) the non-canonical Fe₆ ligand Cys 301 (9,28). Together, these observations begin to expand our understanding of the complex and tightly-regulated process of C-cluster biogenesis. In particular, given the varied metal binding sites that we have observed within the C-cluster scaffold, the insertion of nickel is not a straightforward process and appears to be more complicated than originally thought.

**Experimental Procedures**

**Cloning and purification of DvCODH⁴CooC and DvCODH(ΔD)⁴CooC**. Protein was expressed and purified as described previously (17). Briefly, the D. vulgaris gene encoding CODH (cooS) was cloned into a modified pBGF4 shuttle vector under the control of the promoter of the D. fructosovorans Ni-Fe hydrogenase operon and included an N-terminal strep-tag. For DvCODH(ΔD)⁴CooC, the expression vector also contained the gene for the CooC maturase (cooC), and mutations encoding C42A and C45A were
introduced into the cooS gene by site-directed mutagenesis. To perform mutagenesis by PCR, the HindIII-SacI fragment of the modified pBGF4 plasmid, containing the 5’ end of cooS, was subcloned into pUC19 to serve as a DNA template. The primers GACACGGCGGCGCGGCGCAA ATTCGGCAATTGGGACACCC (forward; mutations underlined) and GGTGTTGGCCTA ATTCGGCAATTGGCCGGCGGCCTGCAC C (reverse; mutations underlined) were used to generate the C42A/C45A variant. The mutated HindIII-SacI fragment was then reintroduced into the HindIII-SacI digested expression vector. The final mutated plasmid was verified by DNA sequencing. Protein was expressed in D. fructosovorans and purified under strictly anaerobic conditions in a Jacomex anaerobic chamber (100% N₂ atmosphere) by affinity chromatography on Strep-Tactin Superflow resin. Protein concentrations were determined by amino acid analysis at the Centre for Integrated Structural Biology (Grenoble, France). Metal content was analyzed by inductively coupled plasma optical emission spectroscopy (ICP-OES). The as-isolated samples contained Ni and Fe as follows: \(DvCODH^{+CooC}\): 0 Ni/monomer, 10 Fe/monomer; \(DvCODH(\Delta D)^{+CooC}\): 0.02 Ni/monomer, 8.5 Fe/monomer. CO oxidation activity was assayed at 37 °C by monitoring the reduction of methyl viologen at 604 nm (\(ε = 13.6 \text{ mM}^{-1} \cdot \text{cm}^{-1}\)), as described previously (17). Neither \(DvCODH^{-CooC}\) nor \(DvCODH(\Delta D)^{-CooC}\) exhibited detectable CO oxidation activity. Reconstitution of either sample with NiCl₂ under reducing conditions did not lead to an increase in activity.

**Crystallization of \(DvCODH\) variants.** \(DvCODH^{+CooC}\) and \(DvCODH(\Delta D)^{+CooC}\) were crystallized in an N₂ atmosphere at 21 °C by hanging drop vapor diffusion in an MBraun anaerobic chamber. A 1-µL aliquot of protein (10 mg/mL in 100 mM Tris-HCl pH 8) was combined with 1 µL of a precipitant solution (200–275 mM MgCl₂, 14–20% PEG 3350) on a glass cover slip and sealed over a reservoir containing 500–700 µL of precipitant solution. Diffraction quality crystals grew in 4–7 days. Crystals were soaked in a cryo-protectant solution containing 200 mM MgCl₂, 20–30% PEG 3350, and 10–16% glycerol and cryo-cooled in liquid nitrogen. For structures of reduced \(DvCODH^{-CooC}\), crystals were soaked in 250 mM MgCl₂, 18% (w/v) PEG 3350, 5 mM sodium dithionite for 30 min prior to cryo-protecting and cryo-cooling in liquid nitrogen.

**Data collection, model building, and refinement.** Data were collected at the Advanced Photon Source (Argonne, IL) on beamline 24-ID-C using a Pilatus 6M pixel detector and at a temperature of 100 K. Native and Fe peak data were collected on the same crystal for each sample, where applicable. The \(DvCODH(\Delta D)^{+CooC}\) structure was determined and refined using data collected at the Fe peak wavelength. Data for \(DvCODH^{-CooC}\) (as-isolated and reduced) were integrated in XDS and scaled in XSACE (38). Data for \(DvCODH(\Delta D)^{-CooC}\) were integrated and scaled in HKL2000 (39). All data collection statistics are summarized in Table 2.

Structures were determined by molecular replacement (MR) in the program Phaser (40) using our previously published structure of \(DvCODH\) (PDB ID: 6B6V) as a search model. Following MR, 10 cycles of simulated annealing refinement were performed in Phenix (41) to eliminate existing model bias. Refinement of atomic coordinates and atomic displacement parameters (B-factors) was performed in Phenix and models were completed by iterative rounds of model building in Coot (42) and refinement in Phenix. Metal cluster geometries were restrained during refinement using custom parameter files. In advanced stages of refinement, water molecules were added automatically in Phenix (41) and modified in Coot (42) with placement of additional water molecules until their number was stable. For the \(DvCODH^{-CooC}\) structures, final stages of refinement included translation, libration, screw (TLS) parameterization with one TLS group per monomer (43). Side chains without visible electron density were truncated to the last atom with electron density and amino acids without visible electron density were not included in the model. Final models contain the following residues (of 629 total): as-isolated \(DvCODH^{+CooC}\): 4–628 (chain A), 4–629 (chain B), 4–629 (chain C), 3–628 (chain D); reduced \(DvCODH^{-CooC}\): 4–629 (chain A), 4–629 (chain B), 4–629 (chain C), 3–629 (chain D); \(DvCODH(\Delta D)^{-CooC}\): 4–40, 45–628 (chain A).
Final refinement yielded models with low free $R$-factors, excellent stereochemistry, and small root mean square deviations from ideal values for bond lengths and angles. Models were validated using simulated annealing composite omit maps calculated in Phenix (41). Model geometry was analyzed using MolProbity (44). Analysis of Ramachandran statistics indicated that each structure contained the following percentages of residues in the favored, allowed, and disallowed regions, respectively: as-isolated $DvCODH^{\text{COoC}}$: 96.7%, 3.0%, 0.3%; reduced $DvCODH^{\text{COoC}}$: 96.9%, 2.8%, 0.3%; $DvCODH(\Delta D)^{\text{COoC}}$: 95.8%, 4.0%, 0.2%. Refinement and geometry statistics are summarized in Table 2. Figures were generated in PyMOL (45). Crystallography packages were compiled by SBGrid (46).
Acknowledgements
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§The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Conflict of Interest
The authors declare that they have no conflicts of interest with the contents of this article.

Author Contributions
ECW and SEC performed the crystallographic experiments and analyzed the crystallographic data with CLD. MM and SD purified protein and performed activity assays. ECW and CLD wrote the manuscript with critical contributions from CL, VF, and SD.
Crystallographic studies of C-cluster assembly

References


Abbreviations
CODH: carbon monoxide dehydrogenase
S_L: linking sulfide
Fe_u: unique iron
Table 1. Metal content and activity of DvCODH variants

<table>
<thead>
<tr>
<th>DvCODH sample</th>
<th>as-isolated Ni/monomer</th>
<th>as-isolated Fe/monomer</th>
<th>as-isolated CO oxidation activity (µmol·min⁻¹·mg⁻¹)</th>
<th>Ni-reconstituted CO oxidation activity (µmol·min⁻¹·mg⁻¹)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT⁺CooC</td>
<td>0.4–0.9</td>
<td>8–10.5</td>
<td>160</td>
<td>1660</td>
<td>(17)</td>
</tr>
<tr>
<td>WT⁻CooC</td>
<td>0–0.2</td>
<td>7.5–8.5</td>
<td>&lt;5</td>
<td>4–60</td>
<td>(17)</td>
</tr>
<tr>
<td>C301S⁺CooC</td>
<td>0</td>
<td>13</td>
<td>N/D</td>
<td>N/D</td>
<td>this work</td>
</tr>
<tr>
<td>ΔD⁺CooC</td>
<td>0.02</td>
<td>8 ± 1</td>
<td>N/D</td>
<td>&lt;5</td>
<td>this work</td>
</tr>
</tbody>
</table>

Crystallographic studies of C-cluster assembly
Table 2. Crystallographic data collection and refinement statistics

<table>
<thead>
<tr>
<th>Data collection</th>
<th>WT−CooC as-isolated</th>
<th>WT−CooC reduced</th>
<th>WT−CooC Fe peak†</th>
<th>(AD)−CooC Fe peak†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wavelength (Å)</td>
<td>0.9792</td>
<td>0.9792</td>
<td>1.7398</td>
<td>1.7379</td>
</tr>
<tr>
<td>Space group</td>
<td>P2₁</td>
<td>P2₁</td>
<td>P2₁</td>
<td>C2</td>
</tr>
<tr>
<td>Cell dimensions</td>
<td>a, b, c (Å)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>64.8, 144.1, 123.4</td>
<td>64.7, 143.8, 123.1</td>
<td>64.7, 143.7, 123.2</td>
<td>110.5, 124.7, 124.7</td>
</tr>
<tr>
<td>β (°)</td>
<td>98.5</td>
<td>98.6</td>
<td>98.6</td>
<td>124.7</td>
</tr>
<tr>
<td>Resolution (Å)</td>
<td>100–1.50</td>
<td>100–1.72</td>
<td>100–2.32</td>
<td>50.0–2.48</td>
</tr>
<tr>
<td></td>
<td>(1.53–1.50)</td>
<td>(1.75–1.72)</td>
<td>(2.37–2.32)</td>
<td>(2.57–2.48)</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>95.0 (93.1)</td>
<td>95.9 (95.6)</td>
<td>94.2 (93.0)</td>
<td>93.4 (88.6)</td>
</tr>
<tr>
<td>Redundancy</td>
<td>6.5 (6.0)</td>
<td>4.3 (4.9)</td>
<td>4.3 (4.2)</td>
<td>3.0 (2.0)</td>
</tr>
<tr>
<td>Unique reflections*</td>
<td>338890</td>
<td>225437</td>
<td>178944</td>
<td>38000</td>
</tr>
<tr>
<td></td>
<td>(24577)</td>
<td>(16599)</td>
<td>(13009)</td>
<td>(3637)</td>
</tr>
<tr>
<td>R_{sym} (%)</td>
<td>9.2 (90.4)</td>
<td>9.3 (91.3)</td>
<td>11.6 (87.5)</td>
<td>22.0 (78.3)</td>
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<tr>
<td>CC{12}</td>
<td>99.9 (71.2)</td>
<td>99.8 (68.5)</td>
<td>99.6 (65.0)</td>
<td>95.8 (47.2)</td>
</tr>
<tr>
<td>&lt;I/σI&gt;</td>
<td>11.5 (1.9)</td>
<td>7.9 (2.0)</td>
<td>11.3 (2.0)</td>
<td>6.1 (2.1)</td>
</tr>
</tbody>
</table>

Refinement

| Resolution (Å) | 93.3–1.50          | 93.3–1.72       | 45.4–2.48        |
| No. reflections| 338812             | 225376          | 37976            |
| R_{work}/R_{free} | 0.154/0.178      | 0.149/0.176     | 0.207/0.248      |
| Monomer/asu    | 4                   | 4                | 1                 |
| No. atoms      | protein 18977       | 18682           | 4481             |
|                | B-cluster 32        | 32              | 8                |
|                | C-cluster 36        | 36              | 8                |
|                | D-cluster 8         | 8               | 8                |
|                | water 2673          | 1899            | 25               |
| B-factors      | protein 19.6        | 22.6            | 41.5             |
|                | B-cluster 15.8      | 16.7            | 40.3             |
|                | C-cluster 21.0       | 24.4            | 45.6             |
|                | D-cluster 17.9      | 18.9            | 40.7             |
|                | water 33.9          | 34.0            |                  |
| R.m.s. bond deviations | Lengths (Å) 0.007 | 0.007           | 0.003            |
|                | Angles (°) 0.932    | 0.919           | 0.737            |
| Rotamer outliers (%) | 0.15              | 0.52            | 0.67             |

†Bijvoet pairs were not merged during data processing.
*Values in parentheses are for the highest-resolution shell.
Figure 1. The metalloclusters of CODH. A) The overall homodimeric structure of \( Dv\text{CODH} \) (PDB ID 6B6V). Metalloclusters are shown as spheres and labeled. Note that the B-cluster of one monomer completes the electron transfer pathway of the opposing monomer. B) The C-cluster in its canonical, reduced state (PDB ID 6B6V). C) The oxidized C-cluster (PDB ID 6B6W). A lysine residue that completes a distorted tetrahedral coordination geometry around the Ni ion has been omitted for simplicity. D) The C-cluster of \( Dv\text{CODH}(C301S)^{\text{CooC}} \) (PDB ID 6DC2). Residue numbers correspond to the sequence of \( Dv\text{CODH} \). Protein is shown in ribbon representation in pink with metalloclusters shown as spheres and sticks with Ni in green, Fe in orange, S in yellow; in panels B–D, ligating amino acid residue side chains are shown as sticks with S in yellow, N in blue, and O in red. Structures shown in this figure are described in Reference (9).
Figure 2. The DvCODH C-cluster is a [3Fe-4S] cluster with a mobile Feₚ. A) Refinement of a [3Fe-4S]-Feₚ C-cluster results in positive Fₒ−Fₑ electron density (green mesh, contoured to +3σ) at the Ni binding site. 2Fₒ−Fₑ electron density (blue mesh) contoured to 1σ. A water molecule (red sphere) is bound to Feₚ. Cys 519 adopts alternative conformations. B) Fe anomalous difference map (orange mesh, contoured to 6σ) indicates the presence of Fe at partial occupancy in the canonical Ni binding site. The Feₚ-ligating water molecule has been omitted for simplicity. C) The C-cluster refined with an alternative conformation of Feₚ. At 80% occupancy, Feₚ is ligated by His 266 and Cys 302 in its canonical binding site. At 20% occupancy, Feₚ is incorporated into the cubane portion of the cluster and ligated by Cys 519. 2Fₒ−Fₑ electron density (blue mesh) contoured to 1σ. Protein is shown in ribbon representation in teal with ligating amino acid residue side chains in sticks; cluster ions shown as spheres and sticks; Fe in orange, S in yellow, N in blue, O in red.

Figure 3. Reduction of DvCODH C-cluster induces movement of Feₚ into the Fe-S cubane portion of the C-cluster. A) Isomorphous difference map (Fₑ(reduced)−Fₑ(as-isolated)) reveals increased electron density at the canonical Ni binding site of the cubane (green mesh, contoured to +5σ) and decreased electron density at the canonical Feₚ binding site (red mesh, contoured to −5σ) in the structure of reduced DvCODH C-cluster relative to as-isolated DvCODH C-cluster. B) Fe anomalous difference map (orange mesh, contoured to 6σ) reveals a strong peak of Fe anomalous signal in the canonical Ni binding site of the cubane (compare to Figure 2B). C) The C-cluster of DvCODH C-refined with an alternative conformation of Feₚ. At 60% occupancy, Feₚ is incorporated into the cubane portion of the cluster and ligated by Cys 519. At 40% occupancy, Feₚ is ligated by His 266 and Cys 302 in its canonical binding site. Cys 302 adopts alternative conformations. 2Fₒ−Fₑ electron density (blue mesh) contoured to 1σ. Protein is shown in ribbon representation in teal with ligating amino acid residue side chains in sticks; cluster ions shown as spheres and sticks; Fe in orange, S in yellow, N in blue.
Figure 4. Removal of the D-cluster does not alter the overall structure but leads to incomplete C-cluster assembly. A) Structural alignment of \( \text{DvCODH(\Delta D)}^{\text{+CooC}} \) (maroon) with \( \text{DvCODH}^{\text{+CooC}} \) (grey; PDB ID 6B6V, Ref. (9)). Inset shows disorder in the vicinity of the D-cluster in \( \text{DvCODH(\Delta D)}^{\text{+CooC}} \). Proteins are shown as the C\( \alpha \) trace of each structure. B- and C-clusters of \( \text{DvCODH(\Delta D)}^{\text{+CooC}} \) are shown as spheres. B) Fe anomalous difference map (orange mesh, contoured to 5\( \sigma \)) suggests the presence of Fe\( \alpha \) at full occupancy in its canonical binding site. C) Refinement of \( \text{DvCODH(\Delta D)}^{\text{+CooC}} \) confirms the location and occupancy of Fe\( \alpha \). \( 2F_o - F_c \) electron density (blue mesh) contoured to 1\( \sigma \). In panels B and C, protein is shown in ribbon representation in maroon with ligating amino acid residue side chains in sticks; cluster ions shown as spheres and sticks; Fe in orange, S in yellow, N in blue, O in red.
Figure 5. Models of C-cluster assembly. A) Formation of the C-cluster Fe-S scaffold. The Fe-S scaffold could be assembled through two different pathways. First, the components of the C-cluster could be inserted as a [3Fe-4S] cluster that combines with a mononuclear Fe ion (upper pathway). Alternatively, the C-cluster site could become loaded with a [4Fe-4S] cluster followed by removal of an Fe ion from the cubane to form Fe_u (lower pathway). In either case, an off-pathway reduction event could (re)convert the [3Fe-4S]-Fe_u scaffold into a [4Fe-4S] cluster. B/C) Two independent models for nickel insertion into the C-cluster. B) Nickel could be inserted directly into a reduced [3Fe-4S]-Fe_u pre-C-cluster. C) Alternative model for nickel insertion involving multiple C-cluster conformations. Starting from the [3Fe-4S]-Fe_u pre-C-cluster (state I), CooC may be involved in inducing a conformational change in the C-cluster in which Fe_u becomes ligated by Cys 301 (state II). Nickel could then bind in either the canonical Fe_u binding site (as observed in structures of the oxidized C-cluster (9); state IIIa) or in the cubane position (state IIIlb). Cluster reduction could then result in formation of the fully mature C-cluster (state IV). Electrons (e^-) indicate reduction events. In panel C, conformations of the C-cluster that have not been characterized crystallographically are shown in faded colors.