AcsF Catalyzes the ATP-dependent Insertion of Nickel into the Ni,Ni-[4Fe4S] Cluster of Acetyl-CoA Synthase*

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Acetyl-CoA synthase (ACS) catalyzes the reversible condensation of CO, CoA, and a methyl-cation to form acetyl-CoA at a unique Ni,Ni-[4Fe4S] cluster (the A-cluster). However, it was unknown which proteins support the assembly of the A-cluster. We analyzed the product of a gene from the cluster containing the ACS gene, cooC2 from Carboxydoterhmus hydrogenofomans, named AcsFCh and showed that it acts as a maturation factor of ACS. AcsFCh and inactive ACS form a stable 2:1 complex that binds two nickel ions with higher affinity than the individual components. The nickel-bound ACS-AcsFCh complex remains inactive until MgATP is added, thereby converting inactive to active ACS. AcsFCh is a MinD-type ATPase and belongs to the CooC protein family, which can be divided into homologous subgroups. We propose that proteins of one subgroup are responsible for assembling the Ni,Ni-[4Fe4S] cluster of ACS, whereas proteins of a second subgroup mature the [Ni4Fe4S] cluster of carbon monoxide dehydrogenases.

The cellular maturation of metalloenzymes, previously considered a spontaneous process in vivo, typically depends on a machinery of uptake, storage, processing, and delivery factors. How metalloenzymes mature has been investigated for some systems, revealing surprisingly complex maturation pathways. Enzymes containing nickel, although still relatively small in number, play critical roles in archaea, bacteria, and eukarya, through which they impact the global hydrogen metabolism. The nickel-enzymes acetyl-CoA synthase (ACS) and carbon monoxide dehydrogenase (CODH) catalyze the reversible reduction of CO2 to CO at the interface (31, 32). Additional proteins support the maturation of these enzymes, now proposed to have a good understanding of how nickel is incorporated into the active site (8, 10).

The nickel-enzymes acetyl-CoA synthase (ACS) and carbon monoxide dehydrogenase (CODH) are found in a variety of anaerobic microbes, including bacterial sulfate reducers, acetogens, and hydrogenogens, as well as archaeal methanogens and sulfate reducers, where they act as the prime CO2 and CO converter (11–14). ACS and CODH can be found as independent monofunctional enzymes in Carboxydoterhmus hydrogenofomans (15) but are typically found in other microorganisms as protein complexes: in acetogens ACS and CODH form a bifunctional (αβ)2 complex, whereas in methanogens they are part of a large (αβγδε)9 multienzyme complex (11, 13, 15, 16). CODHs catalyze the reversible reduction of CO2 to CO at the C-cluster, in which a single nickel ion, embedded within a 3Fe-4S scaffold with an additional iron in exo, binds nickel ions are typically found in other microorganisms as protein complexes: in acetogens ACS and CODH form a bifunctional (αβ)2 complex, whereas in methanogens they are part of a large (αβγδε)9 multienzyme complex (11, 13, 15, 16). CODHs catalyze the reversible condensation of CO, CoA, and a methyl-cation to form acetyl-CoA (13). ACS depends on a Ni,Ni-[4Fe4S] cluster (also called A-cluster) for activity, in which the two nickel ions have distinct coordination: the nickel ion distal to the [4Fe4S] cluster (NiD) is coordinated by two amide nitrogen atoms and two cysteine thiolates within a Cys-X-Cys motif, holding nickel in a stable square-planar coordination environment (15, 20, 21). NiDp is weakly bound by three cysteine thiolates and is removable by 1,10-phenanthroline (22), and may be replaced by zinc and copper, thereby inactivating ACS (15, 20, 21, 23, 24). NiDp likely adopts different oxidation states during turnover and is the presumed place where substrates are activated and converted (13).

Despite their ubiquity and importance in different metabolisms, little is known about the maturation of CODH and ACS, which both limit our understanding of metalloenzyme maturation and hampers their adaptation for biotechnological processes. CODH activity in vivo depends on an ATPase termed CooC (25–28). CooC belongs to the MinD-type ATPases, an enzyme family with diverse functions and a conserved GG signature in their Walker A motif (GGGhGK(T/S)) containing a characteristic lysine residue (signature lysine) in addition to the P-loop lysine (29). Common to the MinD-type ATPases is an ATP-dependent dimerization, which is central to their function and a prerequisite to hydrolyze ATP (30). CooC binds Ni(II) through the Cys-X-Cys motifs of two monomers that combine to form a metal binding site within the dimer interface (31, 32). Additional proteins support the maturation of the CODH in Rhodospirillium rubrum (25); however, these additional proteins are not conserved in bacteria outside the Rhodobacteraceae and may be functionally replaced by non-homologous isofunctional enzymes in other organisms. Although CooC was shown to mature monofunctional CODHs (25–28), it was proposed that the homologous ATPase AcsF supports the assembly of the C-cluster in the bifunctional ACS-CODH complex in Moorella thermoacetica (33).
Maturation of Acetyl-CoA Synthase

In contrast, how the Ni,Ni-[4Fe4S] cluster of ACS is formed in vivo is not known. The purpose of this paper is to provide insights into the role of an ATPase closely related to CooC in this process, which we named AcsF<sub>Ch</sub>. We provide evidence that AcsF<sub>Ch</sub> catalyzes the nickel and MgATP-dependent activation of ACS by forming a complex with apoACS that serves as a platform for nickel binding from which active ACS is generated at the expense of ATP.

Results

CooC Proteins Can Be Divided into Three Subgroups—The genome of Carboxythermus hydrogenoformans contains three genes, annotated as cooC<sub>1</sub>, cooC<sub>2</sub>, and cooC<sub>3</sub>, which encode for proteins of the CooC family (34). The cooC<sub>1</sub> and cooC<sub>2</sub> genes are located in the same gene cluster as the genes encoding CODH-III and ACS (Fig. 1A). The cooC<sub>3</sub> gene is located in proximity to the gene encoding CODH-I.

The InterPro database lists 881 protein sequences as members of the CooC family (35, 36). A sequence similarity network analysis of the non-redundant sequences of the CooC proteins indicates that the sequences may be grouped into three different clusters (Fig. 1B). The two largest clusters, hereafter called “CooC-type” cluster and “AcsF-type” cluster, contain proteins typically associated with gene clusters of CODHases or ACS/CODH. The third cluster contains 116 proteins, mostly found in methanogenic archaea (Methanomicrobia, Methanococci). The CooC-type cluster harbors 298 sequences and includes all CooC proteins shown to be involved in CODH maturation, such as CooC from R. rubrum (CooC<sub>Rr</sub>), 26), CooC<sub>3</sub> from Desulfovibrio vulgaris (CooC<sub>Dv</sub>), 28). The CooC-type cluster also encompasses CooC1 from C. hydrogenoformans (CooC<sub>Ch</sub>), which is a nickel-binding ATPase whose crystal structure has been determined (27, 31).

The AcsF-type cluster encompasses 309 protein sequences, among them AcsF from M. thermoacetica (AcsF<sub>Ma</sub>), AcsF<sub>Ch</sub> has been described as an ATPase of unknown function whose involvement in CODH maturation of bifunctional ACS/CODH was postulated but not shown (27, 33). The AcsF-type cluster also encompasses the gene product of cooC2 from C. hydrogenoformans, which we therefore named AcsF<sub>Ch</sub>. All clusters include proteins from both bacteria and archaea. Most organisms (63%) containing a gene in the CooC-type cluster also harbor a paralogue gene of the AcsF-type cluster in their genome and only 13% of the organisms with an AcsF-type sequence have no CooC-type protein in the Interpro database.

AcsF<sub>Ch</sub> Dimerizes in the Presence of ATP, Does Not Bind Nickel and Has Low ATPase Activity—Typical size exclusion chromatograms of AcsF<sub>Ch</sub> showed peaks at 83.2 and 90.5 ml. The corresponding molecular masses, 57.7 and 28.8 kDa, were determined by comparison to a standard curve and agree with the predicted molecular masses of dimeric (55.4 kDa) and monomeric AcsF<sub>Ch</sub> (27.7 kDa). The monomer:dimer ratio depended on the cultivation batch rather than on individual purifications. Additionally, two variants of AcsF<sub>Ch</sub> were expressed and purified. The K10A-AcsF<sub>Ch</sub> variant, lacking the signature lysine Lys-10, showed similar monomer:dimer ratios for as isolated protein as wild type AcsF<sub>Ch</sub>, whereas the C108A-AcsF<sub>Ch</sub> variant was only present as monomer.

We analyzed by size exclusion chromatography (SEC) whether monomeric AcsF<sub>Ch</sub> dimerizes in the presence of MgADP and MgATP (Fig. 2A). Although the presence of MgADP had no influence on the chromatogram, we found AcsF<sub>Ch</sub> in the presence of MgATP to be predominantly dimeric, as expected for a MinD-type ATPase. The monomeric fraction of the K10A-AcsF<sub>Ch</sub> variant remained in the monomeric state independent of the presence of MgADP or MgATP (Fig. 2B).

![Figure 1](image1.png)

**FIGURE 1.** A, gene clusters containing the gene for bifunctional ACS/CODH in C. hydrogenoformans and M. thermoacetica. B, sequence similarity network of CooC proteins. The network was generated from an all-by-all distance matrix storing Kimura distances calculated from a multiple amino acid sequence alignment. Gray colored lines connect nodes (amino acid sequences) with a Kimura distance below 1.3. Nodes are arranged using the yFiles organic layout of Cytoscape 3.3.0 (59). Larger nodes represent sequences (amino acid sequences) with a Kimura distance above 1.3. Nodes are arranged using the yFiles organic layout of Cytoscape 3.3.0 (59). Larger nodes represent sequences (amino acid sequences) with a Kimura distance above 1.3.

![Figure 2](image2.png)

**FIGURE 2.** Nucleotide dependent dimerization of AcsF<sub>Ch</sub>. A, the elution profile of monomeric AcsF<sub>Ch</sub>, without nucleotide is shown as solid line. Monomeric AcsF<sub>Ch</sub> was incubated with either MgADP or MgATP and the oligomeric state was analyzed by SEC. The elution profiles of AcsF<sub>Ch</sub>, in the presence of MgADP or MgATP are shown as dashed and dotted lines, respectively. B, influence of nucleotide on the oligomeric state of K10A-AcsF<sub>Ch</sub>. The elution profile of monomeric K10A-AcsF<sub>Ch</sub> is shown as solid line and the elution profile of K10A-AcsF<sub>Ch</sub>, in the presence of MgADP or MgATP are shown as dashed and dotted lines, respectively. All dimerization experiments were at least performed with two independent samples.
AcsF<sub>Ch</sub> shares with the other CooC proteins the CXC sequence motif. In CooC<sub>1</sub>Ch, the two Cys residues of this motif are able to bind nickel, which can be followed by an increased absorption in the region 310–350 nm originating from a ligand to metal charge transfer between Cys-S<sup>-</sup> and Ni(II) (31). The UV/visible absorption spectrum of AcsF<sub>Ch</sub> had a maximum at 274 nm. When we added increasing amounts of NiCl<sub>2</sub> to AcsF<sub>Ch</sub>, the UV/visible spectral features remained unchanged. Furthermore, when we titrated NiCl<sub>2</sub> to AcsF<sub>Ch</sub> in an anoxic isothermal titration calorimetry (ITC) experiment, we only observed heats of dilution. Thus, in contrast to CooC<sub>1</sub>Ch, we could not detect specific nickel binding by AcsF<sub>Ch</sub>.

The amino acid sequence of AcsF<sub>Ch</sub> contains the characteristic Walker A signature of the MinD-type ATPase. We determined the specific ATPase activity of AcsF<sub>Ch</sub> with a coupled assay and a colorimetric assay. Determined specific activities varied with each purification batch and averaged at 7.4 nmol min<sup>-1</sup> mg<sup>-1</sup> at 25 °C and 25.5 nmol min<sup>-1</sup> mg<sup>-1</sup> at 45 °C in the coupled assay. In the colorimetric assay the average activities measured were 3.9 nmol min<sup>-1</sup> mg<sup>-1</sup> at 25 °C, 20.4 nmol min<sup>-1</sup> mg<sup>-1</sup> at 45 °C, and 56.5 nmol min<sup>-1</sup> mg<sup>-1</sup> at 60 °C. In contrast, we could not detect ATPase activity above background for the K10A-AcsF<sub>Ch</sub> variant. There is an intrinsic risk that low ATPase activities, such as those determined for AcsF<sub>Ch</sub>, could arise from minor protein impurities of high specific activities and not the protein investigated. However, all AcsF<sub>Ch</sub> preparations tested had substantially higher activities than the preparation of the K10A-AcsF<sub>Ch</sub> variant, which was purified using the same protocol. Furthermore, the ATPase activity increased until 60 °C, which is expected for an ATPase complexed with AcsF<sub>Ch</sub>.

ApoACS Can Be Activated by AcsF<sub>Ch</sub>, NiCl<sub>2</sub>, and MgATP—The UV/visible spectrum of ACS had the characteristic absorption shoulder of a [4Fe4S]<sup>2+</sup> cluster at 420 nm. Metal analysis by inductively coupled plasma optical emission spectrometry (ICP-OES) confirmed the presence of iron in as isolated ACS, whereas nickel could not be detected (Table 1). In agreement with the lack of nickel, the enzyme is inactive. Therefore, the as isolated ACS is designated as apoACS.

The lack of nickel and catalytic activity in apoACS served as a starting point to investigate the maturation of ACS. When we added stoichiometric amounts of nickel, MgATP, and AcsF<sub>Ch</sub> to ACS, it reached within 2 h a specific activity of 70 nmol min<sup>-1</sup> mg<sup>-1</sup> (Fig. 4A). In contrast, when we omitted AcsF<sub>Ch</sub> from the mixture, the activity of ACS increased only marginally. Activation of ACS by AcsF<sub>Ch</sub> strictly depended on MgATP addition. Thus, AcsF<sub>Ch</sub> catalyzes the Ni<sup>2+</sup> and MgATP-dependent maturation of ACS. The process is catalytic as also substoichiometric amounts of AcsF<sub>Ch</sub> (1:20 AcsF<sub>Ch</sub>:apoACS) sufficed to activate apoACS.

In a next step, we investigated how the amino acid exchanges K10A and C108A influence the capability of AcsF<sub>Ch</sub> to activate apoACS. Neither the ATPase-deficient variant K10A-AcsF<sub>Ch</sub>, nor the C108A-AcsF<sub>Ch</sub> variant was able to activate apoACS (Fig. 4B). This indicates that the ATPase activity of AcsF<sub>Ch</sub> as well as the presence of the CXC motif, are essential for the activation process.

AcsF<sub>Ch</sub> is a specific maturation factor of ACS, as neither CooC<sub>1</sub>Ch nor CooC<sub>3</sub>Ch can replace AcsF<sub>Ch</sub> in the activation assay (Fig. 4B). Even though the data may indicate a slight increase of ACS activity over time (Fig. 4B, inset), the determined activities of ACS are within the error of the activity assay. They remained within the range detected in the absence of AcsF<sub>Ch</sub> and are more than 37 times lower than when incubated with AcsF<sub>Ch</sub>. Thus only AcsF<sub>Ch</sub>, not CooC<sub>1</sub>Ch or CooC<sub>3</sub>Ch, acts as a specific maturation factor of ACS.

The ACS-AcsF<sub>Ch</sub> Complex Binds Nickel Stoichiometrically and Can Be Subsequently Activated by MgATP—We determined whether there is a difference between the nickel binding properties of apoACS and the apoACS-AcsF<sub>Ch</sub> complex. Therefore, we titrated the proteins with nickel using ITC and analyzed the metal contents of these samples by ICP-OES after removal of excess nickel. When we titrated apoACS with NiCl<sub>2</sub>, we observed substoichiometric nickel binding: the ICP-OES metal analysis detected only 0.43 nickel atoms per 4 Fe atoms in nickel-titrated apoACS (Table 1). The apoACS-AcsF<sub>Ch</sub> complex, by contrast, binds nickel stoichiometrically; the ICP-OES analysis detected 1.85 nickel atoms. Thus, whereas apoACS alone cannot bind Ni<sup>2+</sup> stoichiometrically, the apoACS-AcsF<sub>Ch</sub> complex is able to bind approximately two Ni<sup>2+</sup> ions (Table 1).

As the apoACS-AcsF<sub>Ch</sub> complex has the full complement of metal ions after titration with nickel, we determined its specific activity. Despite the presence of two nickel ions, the nickel-
bound apoACS-AcsF<sub>Ch</sub> complex lacked ACS activity. But when we removed excess Ni<sup>2+</sup>/H<sub>2</sub>O and incubated the nickel-bound apoACS-AcsF<sub>Ch</sub> complex with 10 mM MgATP (1 h, 60 °C), it gained an activity of 67 nmol min<sup>−1</sup> mg<sup>−1</sup>, the same specific ACS activity as observed when all components were present in our activation assay. Thus, MgATP is needed to convert the active site in the ACS-AcsF<sub>Ch</sub> complex from an inactive nickel-bound to an active nickel-bound state.

**Discussion**

Role of AcsF<sub>Ch</sub> in Ni,Ni-[4Fe4S] Cluster Formation—Our analysis allows first mechanistic conclusions about how AcsF<sub>Ch</sub> supports the assembly of the Ni,Ni-[4Fe4S] cluster of ACS (Fig. 5). In the first step, two AcsF<sub>Ch</sub> monomers or one AcsF<sub>Ch</sub> dimer form a complex with apoACS. Complex formation creates at least one additional high-affinity nickel binding site in the complex, as only the apoACS-AcsF<sub>Ch</sub> complex binds the full complement of nickel when presented with low physiological concentrations: we detected no affinity for AcsF<sub>Ch</sub> to bind nickel; apoACS binds less than one nickel ion when titrated with nickel; however, the apoACS-AcsF<sub>Ch</sub> complex binds two nickel ions under the same conditions. Thus, complex formation likely
induces conformational changes, creating new binding sites for nickel, in apoACS, AcsF<sub>Ch</sub>, or at the interface between both.

When the Cys-X-Cys motif in AcsF<sub>Ch</sub> is altered, the complex does not form, suggesting that it is part of the ACS-AcsF<sub>Ch</sub> interface. It is tempting to assume that the presence of strictly conserved Cys-X-Cys motifs in both AcsF<sub>Ch</sub> and ACS is not purely coincidental (37), but serves to have binding sites with similar affinity for Ni<sup>2+</sup> in both proteins.

Once nickel is bound to the apoACS-AcsF<sub>Ch</sub> complex, MgATP is required to activate ACS. Although MgATP binding and its hydrolysis are key to the activation, their impact on the bound nickel ions remains to be defined. Furthermore, it should be noted that the observed rate constant of apoACS activation of 1.47 h<sup>−1</sup> is substantially lower than expected from the determined rate of ATP hydrolysis by AcsF<sub>Ch</sub> alone.

Two Classes of CooC Proteins Are Responsible for the Maturation of ACS and CODH—Despite the sequence similarity between AcsF and the paralogous CooC1 and CooC3 proteins of <i>C. hydrogenoformans</i>, it is obvious that the CooC proteins, as defined by InterPro, can be grouped in at least three classes (Fig. 1). We suggest that enzymes clustered in the CooC-type group may act in the maturation of the [Ni4Fe4S] cluster of CODHs, whereas those that belong to the AcsF-type group, such as AcsF<sub>Ch</sub> and AcsF<sub>Mt</sub>, catalyze the MgATP and nickel-dependent maturation of the Ni,Ni-[4Fe4S] cluster of ACS. A possible division of selected CooC proteins in two classes has also been proposed based on a phylogenetic tree (27).

CODH and ACS differ not only in their function and overall fold, but their active sites differ in (i) composition: CODH has 1 nickel ion per 4 iron ions, whereas ACS has 2 nickel ions; (ii) architecture: in CODH nickel is integrated into an FeS scaffold, whereas in ACS the 2 nickel ions are bound next to a [4Fe4S] cluster; and (iii) accessibility: in CODH the Ni,Fe-cluster is connected to the surface via small channels, whereas the Ni,Fe-cluster of ACS is surface exposed in at least one conformation of the protein (21). It is surprising that despite all these differences the maturation involved in nickel- and MgATP-dependent activation of the enzymes belong to the same ATPase family, have remarkably similar amino acid sequences, and likely evolved from a common ancestor.

Comparison with Homologous NTPases—AcsF<sub>Ch</sub> is homologous and functionally related to other NTP-hydrolyzing maturation factors, which act, for example, in the assembly of ureases (UreG), Ni,Fe-hydrogenases (HypB), nitrogenases (NifH), and cytosolic Fe/S proteins (Cfd1/Nbp35). UreG and HypB are homologous GTPases that are essential for the maturation of ureases and hydrogenases, respectively. Both are like AcsF<sub>Ch</sub>

![Figure 4. Activation of apoACS. A, time dependent activation of apoACS in the presence of AcsF<sub>Ch</sub>, MgATP, and nickel. 20 μM apoACS were incubated with 40 μM AcsF<sub>Ch</sub>, 10 mM MgATP, and 40 μM NiCl<sub>2</sub> at 60 °C. Aliquots were taken from the activation assay at the indicated time points and the specific activity was measured at 25 °C by following the conversion of Mecob(III)imad to cob(I)imad by UV/visible spectroscopy. Assay conditions included a sample of the activation assay, Ti(III)citrate, Mecob(III)inamid, CO, and CoA in 0.1 M HEPES, pH 7.2. Each data point was measured in triplicate and error bars represent the standard deviation. Four control experiments were performed, in which AcsF<sub>Ch</sub>, MgATP, NiCl<sub>2</sub>, or apoACS was omitted from the activation assay. Data points for the control experiments were measured once. A monoeXponential function was fitted to the data points giving an observed rate constant of 1.47 h<sup>−1</sup> for the activation. B, activation of apoACS after 60 (dark) and 180 min (light) in the presence of AcsF<sub>Ch</sub>, K10A-AcsF<sub>Ch</sub>, C108A-AcsF<sub>Ch</sub>, and CooC3<sub>Ch</sub>, which was measured once. The inset shows a magnification of the data points for K10A-AcsF<sub>Ch</sub>, C108A-AcsF<sub>Ch</sub>, CooC1<sub>Ch</sub>, and CooC3<sub>Ch</sub>.](image1)

![Figure 5. Preliminary model of ACS activation by AcsF<sub>Ch</sub>. ApoACS (1), which contains the [4Fe4S] cluster, but no nickel, can form a complex with either one dimer or two monomers of AcsF<sub>Ch</sub>. The apoACS-AcsF<sub>Ch</sub> complex (2) is then able to bind two nickel ions. The nickel ions may bind either sequentially (3) or simultaneously. However, neither the details of the nickel binding mechanism nor the location of the nickel binding site are known. ATP hydrolysis by AcsF<sub>Ch</sub> converts inactive (4) to active ACS (5).](image2)
SIMIBI NTPases, but belong to the G3E family and carry the signature motif (ESGG) and the guanine specificity loop (NKTD) characteristic for this family (29, 38). In contrast to AcsF(CooC), they act together with other maturation factors to transfer a metal to a target protein: UreG forms a complex with the large subunit of Ni,Fe-hydrogenase (7, 10, 41, 42). In contrast to other metalloenzymes, DNA polymerases, and T4 DNA ligase were purchased from New England Biolabs, Fermentas (Thermo Scientific), or Agilent Genomics. N2, N2/H2 (95/5%), and CO were purchased from Air Liquide. Chromatography columns were purchased from GE Healthcare or IBA. Bottles for metal-free buffers were rinsed with distil HCl and then washed with MilliQ/H2O. Chelex 100 resin (Bio-Rad) was added to all metal-free buffers (5 to 10 g liter⁻¹). All anaerobic solutions were prepared at a Schlenk line in a bottle equipped with a butyl rubber septum by at least four cycles of evacuating and purging with N2. CO-saturated buffer was prepared by bubbling CO through the buffer for at least 10 min. Ti(III) citrate was pre-prepared as previously described (52). Methylcobinamide was synthesized from methylcobalamin using the method from Zou et al. (53) except that it was purified with a SepPak C18 reversed phase column (Waters).

**Chromatography**—All chemicals were at least of analytical grade and purchased from Sigma, AppliChem, or Roth. Restriction enzymes, DNA polymerases, and T4 DNA ligase were purchased from New England Biolabs, Fermentas (Thermo Scientific), or Agilent Genomics. N2, N2/H2 (95/5%), and CO were purchased from Air Liquide. Chromatography columns were purchased from GE Healthcare or IBA. Bottles for metal-free buffers were rinsed with distilled HCl and then washed with MilliQ/H2O. Chemaux 100 resin (Bio-Rad) was added to all metal-free buffers (5 to 10 g liter⁻¹). All anaerobic solutions were prepared at a Schlenk line in a bottle equipped with a butyl rubber septum by at least four cycles of evacuating and purging with N2. CO-saturated buffer was prepared by bubbling CO through the buffer for at least 10 min. Ti(III) citrate was pre-prepared as previously described (52). Methylcobinamide was synthesized from methylcobalamin using the method from Zou et al. (53) except that it was purified with a SepPak C18 reversed phase column (Waters).

**Cloning, Mutagenesis, and Expression**—The cooC2 gene was amplified by PCR from genomic DNA of *C. hydrogenoformans*, Z-2901 using *Pfu* DNA polymerase with primer C2_fw with an Ndel restriction site and C2_rv with an Eco31I restriction site (primer sequences are listed in Table 2), which produces a BamHI compatible ligation site. The PCR product was digested with Eco31I and Ndel and then ligated into an Ndel/BamHI-digested pET11a vector (Novagen). The ligation product was named pET11a_AcsF. From this plasmid, the cooC2 gene was cloned into a modified pET28a vector with an N-terminal streptag and a tobacco etch virus (TEV) protease cleavage site, named pET28aTEVstrep, via polymerase incomplete primer extension (PIPE) cloning (54, 55). For PIPE cloning the vector PCR contained 0.1 ng of template, 1 μM of each primer (PIPE_vec_fw and PIPE_vec_rv; Table 2), and 25 μl of Phusion High Fidelity PCR master mix in a 50-μl reaction volume. The DNA was amplified using the Mastercycler pro PCR machine (Eppendorf) with the following protocol: 2 min initial denaturation at 98 °C, 35 cycles of 98 °C for 10 s, 65 °C for 10 s, and 72 °C for 2 min, followed by immediate cooling to 4 °C. The PCR product was purified and aliquots were stored at −20 °C. Using appropriate primers and templates, the set up for the insert

| Name     | Sequence                                                                 |
|----------|--------------------------------------------------------------------------|
| C2_fw    | 5′-GGA ATT CCA TAT GGC CTT TAA AAT TGC GGT TG-3′                         |
| C2_rv    | 5′-CAT GGC CTC GCG GAT TCA GAT CCA TAT GGC CTT TAA AAT TGC GGT TG-3′      |
| C3_fw    | 5′-GGA ATT CCA TAT GCA GGT GTC TG-3′                                    |
| C3_rv    | 5′-CCC AAG CT TCA ACA ATT CCT CC-3′                                     |
| PIPE_vec-fw  | 5′-GGA TCC GAA TTC TCG CTC GTG CG-3′                                    |
| PIPE_vec-rv | 5′-CAT ATG GCC CTG GAA ATA CAA GTT TTC GTC CG-3′                        |
| PIPE_C2_fw  | 5′-CTT GTA TTA CCA GGG CCA TAT GCC CTT TAA AAT TGC GGT TG-3′             |
| PIPE_C2_rv  | 5′-CGA GGG AGC TCG AAT TCG GAT CCC TAG ATA ATCC GCA TTT TGT AAA ATT TTC-3’ |
| PIPE_ACS_fw  | 5′-CTT GTA TTA CCA GGG CCA TAT GAG CGA AGT TAA TTT TGA TCA AAT TTT TG-3′  |
| PIPE_ACS_rv  | 5′-CGA CCG AGC TCG AAT TCG GAT CCT TAG AGT AGT GGG GCC CGC ATT TG-3′      |

**Experimental Procedures**

**Bioinformatic Methods**—881 sequences were extracted from InterPro entry IPR014433 (35, 36). The sequence of CooC from *D. vulgaris* (CooC2) was added, because it was missing in the InterPro entry. After removal of redundant (identical) sequences 723 sequences remained. These sequences were aligned using MAFFT (49) and a distance matrix was calculated from the multiple sequence alignment using protdist/PHYLIP (50) using the Kimura distance as a criterion for similarity. The distance cutoff used to define an interaction in the sequence similarity plot was 1.3. Cytoscape 3.3.0 was used for clustering and displaying the sequence similarity network using the yFiles organic layout provided in Cytoscape (51).

**Maturation of Acetyl-CoA Synthase**

**TABLE 2**

| Name Sequence                                                                 |
|--------------------------------------------------------------------------|
PCR was the same as for the vector PCR. The reaction mixture was treated as follows: 2 min initial denaturation at 98 °C, 35 cycles of 10 s at 98 °C, 10 s at 62 °C, and 12 s at 72 °C, followed by immediate cooling to 4 °C. After purification of the insert, 1 μl of vector was annealed with 1 μl of insert for 1 min on ice and then directly transformed into *Escherichia coli* DH5α. The plasmid was named pET28a_AcsFstrep. K10A-AcsF<sub>Ch</sub> and C108A-AcsF<sub>Ch</sub> variants were prepared using the QuickChange method following standard procedures. The cooC<sub>3</sub> gene was amplified by PCR from genomic DNA of *C. hydrogenoformans* Z-2901 using Herculase II DNA fusion polymerase with primer C3<sub>fw</sub> with an NdeI restriction site and C3<sub>rv</sub> with a HindIII restriction site (Table 2). The PCR product was digested with the respective restriction enzymes and ligated into the original pET28a vector and the modified pET28aTEVstrep vector. The latter plasmid was named pET28a_C3strep. The *acsB* gene was cloned into the pET28aTEVstrep vector from genomic DNA of *C. hydrogenoformans* Z-2901 (10 ng) using PIPE cloning as described above with primers PIPE_ACS<sub>fw</sub> and PIPE_ACS<sub>rv</sub> (Table 2). The plasmid was named pET28a_ACSstrep. The pET28a_ACSstrep plasmid was also digested with NdeI and BamHI and the digestion product was ligated into a pET28a_twinstrep vector, which has a twin-streptag and a TEV protease cleavage site. The plasmid was named pET28a_ACStwinstrep. All plasmids were transformed into *E. coli* DH5α and verified by nucleotide sequencing (Eurofins Genomics). For expression, all plasmids were transformed into *E. coli* BL21(DE3). Cells containing the pET28a_AcsFstrep or pET28a_C3strep plasmid were grown aerobically in LB medium containing 50 μg ml<sup>-1</sup> of kanamycin at 37 °C. After the optical density at 600 nm (A<sub>600</sub>) reached 0.7 ± 0.1, the culture was induced with 0.2 mM isopropyl β-D-thiogalactopyranoside and the temperature was decreased to 28 °C. Cells were harvested 16 to 24 h after induction and washed with 50 mM Tris-HCl, pH 8.0, 100 mM NaCl, 2 mM tris(2-carboxyethyl)phosphine, and 2 mM EDTA. Cell pellets were frozen in liquid N<sub>2</sub> and stored at −80 °C. Cells with the pET28a_AcsFstrep or pET28a_ACSstrep plasmid were grown anaerobically in LB medium containing 50 μg ml<sup>-1</sup> of kanamycin in 5-liter bottles in a water bath at 37 °C. The culture was stirred aerobically and after 1 h the medium was supplemented with 0.1 mM FeSO<sub>4</sub> and 0.5 mM cysteine. When the A<sub>600</sub> reached 0.6 ± 0.1 cultures were induced with 0.5 mM isopropyl β-D-thiogalactopyranoside and transitioned to anaerobic growth by closing the bottles with a butyl rubber septum. After 24 h cells were quickly harvested aerobically, frozen in liquid N<sub>2</sub>, and stored at −80 °C.

**Purification**—All purifications were carried out at room temperature in an anaerobic glove box (model B; COY Laboratory Products) using metal-free buffers. CooC<sub>1</sub> was prepared as previously described (31). For AcsF<sub>Ch</sub>, 5 to 10 g of frozen cells were resuspended in buffer A (50 mM Tris-HCl, pH 8.0, 100 mM NaCl, 2 mM tris(2-carboxyethyl)phosphine) with a small amount of avidin and lysozyme. The cell suspension was sonicated (Bandelin Sonoplus 2200) three times for 5 min each (5 × cycle, 50% amplitude) in a rosette cooling cell on ice. It was then centrifuged in a polycarbonate bottle with cap assembly at 142,400 × g for 1 h at 12 °C. The supernatant was loaded on a StrepTactin-Sepharose High Performance column (10–30 ml) equilibrated in buffer A. After washing the column with 3 column volumes of buffer A, AcsF<sub>Ch</sub> was eluted with a linear gradient of 0–2.5 mM desthiobiotin in the same buffer. The streptag of AcsF<sub>Ch</sub> was cleaved by incubation with a streptagged TEV protease for ~16 h. The protein solution was applied to a PD-10 desalting column and was then reloaded on the StrepTactin column equilibrated with buffer A. The flow-through was collected, concentrated, and loaded on a Superdex 200 HiLoad 16/60 column equilibrated in buffer B (50 mM Tris-HCl, pH 8.0, 100 mM NaCl). Fractions corresponding to monomeric and dimeric AcsF<sub>Ch</sub> were pooled, concentrated, frozen in liquid N<sub>2</sub> (in glass vials equipped with a PTFE/silicone septum), and stored at −80 °C. For K10A-AcsF<sub>Ch</sub>, C108A-AcsF<sub>Ch</sub>, and CooC<sub>3</sub> variants, the purification procedure was continued as described above. AcsF<sub>Ch</sub> was prepared following the same basic strategy except that buffers did not contain tris(2-carboxyethyl)phosphine and the streptag or the twin-streptag was not cleaved. Cells containing the pET28a_ACStwinstrep or pET28a_ACStwinstrep plasmid were lysed, centrifuged, and captured on the streptactin column as described above. The protein was then eluted with desthiobiotin, concentrated, and directly loaded on the Superdex 200 size exclusion column. The fractions corresponding to monomeric AcsF were collected, concentrated, frozen in liquid N<sub>2</sub> (in glass vials equipped with a PTFE/silicone septum), and stored at −80 °C.

**Analysis of Dimerization of AcsF<sub>Ch</sub> and K10A-AcsF<sub>Ch</sub> in the Presence of Nucleotide**—The dimerization behavior of AcsF<sub>Ch</sub> and K10A-AcsF<sub>Ch</sub> in the presence of nucleotide was analyzed by size exclusion chromatography. Monomeric AcsF<sub>Ch</sub> was incubated in 50 mM Tris-HCl, pH 8.0, 100 mM NaCl, 2 mM tris(2-carboxyethyl)phosphine, 0.25 mM MgCl<sub>2</sub>, and 0.25 mM ADP or ATP for 10 min. The incubation mixture was then loaded on a Superdex 200 HiLoad 16/60 column equilibrated with the incubation buffer containing the corresponding nucleotide. The flow rate was 1 ml min<sup>−1</sup>.

**UV/Visible Spectroscopy**—UV/visible spectra were obtained inside the glove box, using an Agilent 8453 photodiode array spectrophotometer with a Peltier temperature controller at 25 °C. Black walled quartz cuvettes with 1-cm path length were used to record spectra.

**Nickel Binding Assay**—20 μM AcsF<sub>Ch</sub> in 50 mM Tris–HCl, pH 8.0, 100 mM NaCl were titrated with NiCl<sub>2</sub> to yield concentrations of 1, 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, and 40 μM Ni<sup>2+</sup>. After each addition one spectrum was recorded. Titrations and UV/visible spectroscopy were performed inside the glove box.

**Measurements of ATPase Activity**—To measure the ATPase activity of AcsF<sub>Ch</sub> a coupled assay using pyruvate kinase and lactate dehydrogenase was performed (56). In this assay the conversion of one molecule of NADH to NAD<sup>+</sup> by lactate dehydrogenase equals the consumption of one molecule of ATP by AcsF<sub>Ch</sub>. The rate of ATP hydrolysis was determined by measuring the decrease of absorption at 340 nm for 10 min and calculating the reaction rates using the extinction coefficient of NADH ε<sub>340</sub> = 6220 M<sup>−1</sup> cm<sup>−1</sup>. The assay solution was prepared in 50 mM HEPES, pH 7.2, containing 150 μM KCl, 10 mM MgCl<sub>2</sub>, 2.3 mM phosphoenolpyruvate, 140 μM NADH, 9 units of lactate dehydrogenase, 6 units of pyruvate kinase, and 1 mM MgATP. The assay was started by addition of 5 μM AcsF<sub>Ch</sub>. This assay was performed at two different temperatures, at 25 or
Maturation of Acetyl-CoA Synthase

45 °C. The velocities obtained by the coupled assay were confirmed with a modified malachite green ammonium molybdate assay (57). This assay was performed at 25, 45, and 60 °C. In this assay the amount of phosphate released by the hydrolysis of ATP is detected by following the increase of absorption at 630 nm, which is due to the formation of a malachite green phosphomolybdate complex. The rate calculation was based on a standard curve, which was prepared with KH₂PO₄ in the range from 1 to 9 μM. The assay solution contained 5 μM AcsF₄₁₉, 10 mM MgCl₂, and 1 mM MgATP in 0.1 mM HEPES, pH 7.2. Aliquots were taken at different time points, incubated with the malachite green ammonium molybdate mixture and 34% (w/v) sodium citrate for 15 min, and then the absorption was measured at 630 nm. Control experiments in the absence of AcsF₄₁₉ were performed to assess the rate of autophosphorylation of ATP at different temperatures.

Analysis of Complex Formation Behavior between ACS and AcsF and CooC Proteins—ApoACS and AcsF₄₁₉ were incubated for 10 min inside the glove box and loaded on a Superdex 200 HiLoad 16/60 column equilibrated in 50 mM Tris-HCl, pH 8.0, 100 mM NaCl. The flow rate was 1 ml min⁻¹. Elution profiles were recorded by following the absorption at 280 nm. As a control apoACS and AcsF₄₁₉ were also loaded individually on the column. From these three chromatography runs, the fractions were analyzed by SDS-polyacrylamide gel electrophoresis. The stoichiometry of the complex was analyzed with the software GelAnalyzer 2010a. Complex formation of apoACS with K10A-AcsF₄₁₉, C108A-AcsF₄₁₉, CooC1₄₁₉, and CooC3₄₁₉ was also analyzed by size exclusion chromatography as described for wild type AcsF₄₁₉. A calibration of the Superdex HiLoad 16/60 column was performed with the following molecular mass standards: aprotinin (6.5 kDa), RNase A (13.7 kDa), DNase I (31 kDa), ovalbumin (44 kDa), conalbumin (75 kDa), lactate dehydrogenase (140 kDa), aldolase (161 kDa), and catalase (250 kDa). The decadic logarithm of the molecular weight was plotted against the elution volume and the calibration curve was determined by linear regression.

Reconstitution of ApoACS—All incubation and reaction mixtures were set up inside an anaerobic glove box (model B; COY Laboratory Products). ApoACS was reconstituted by incubating 20 μM apoACS with 40 μM NiCl₂, 40 μM AcsF₄₁₉, 10 mM MgCl₂ and 10 mM MgATP in 0.1 mM HEPES, pH 7.2, at 60 °C. Four control experiments were set up; in each case, one component of the incubation mixture was missing. Reconstitution experiments were also performed with K10A-AcsF₄₁₉, C108A-AcsF₄₁₉, CooC1₄₁₉, and CooC3₄₁₉ instead of AcsF₄₁₉. Aliquots were taken at different time points and the rate of acetyl-CoA formation was analyzed by UV/visible spectroscopy.

Measurements of Acetyl-CoA Formation Activity—Acetyl-CoA synthesis activity of ACS from CO, methylcobinamide, and CoA was determined by following the conversion of methylcob(III)inamide (MeCbi) to cob(II)inamide (Cbi). The change of absorption was measured at 387 and 462 nm. The reaction rates were calculated using the differences of the extinction coefficients, Δε₃₈₇ nm(MeCbi-Cbi) = −17,410 M⁻¹ cm⁻¹ and Δε₄₆₅ nm(MeCbi-Cbi) = 7,840 M⁻¹ cm⁻¹, which were calculated from spectra of Robertson et al. (58). The reaction was set up inside the glove box as follows: 0.1 mM HEPES, pH 7.2, 50 μM MeCbi, and 300 μM Ti(III)citrate were placed in a screw cap cuvette; the ACS incubation mixture (see ACS reconstitution) was then added to yield a final concentration of 2 μM ACS. After closing the cap, CO-saturated buffer (10% of the final volume) was injected with a syringe and the reaction was started by adding 200 μM CoA. All measurements were performed in darkness.

Isothermal Titration Calorimetry—ITC experiments were performed inside a glove box (LABstar, MBRAUN) with a MicroCal VP-ITC system (GE Healthcare) at 25 °C in 50 mM Tris, 100 mM NaCl. Three different experiments were performed: (a) 15 μM AcsF₄₁₉ were titrated with 300 μM NiCl₂, (b) 24.7 μM apoACS were titrated with 750 μM NiCl₂, and (c) 24.4 μM ACS-AcsF₄₁₉ complex were titrated with 750 μM NiCl₂. The concentrations of apoACS and ACS-AcsF₄₁₉ complex were normalized to the amount of iron, which was determined after ITC measurements by ICP-OES. The enzyme was present in the reaction cell (1.4 ml) and NiCl₂ was in the syringe. The initial injection was 3 μl (lasting 6 s) and the 47 subsequent injections were 6 μl (lasting 12 s). There was an interval of 300 s between injections. The solution in the reaction cell was stirred at 307 rpm.

Removal of Excess Metal from ITC Samples—When the ITC experiment was finished, samples b and c were used for further experiments, i.e. activity measurements and metal analysis. The protein samples were concentrated to ~300 μl and the excess metal was removed using a PD MiniTrap-G25 column. Only colored fractions were collected. To ensure that all excess metal was removed a buffer sample was prepared for metal analysis: 130 μM NiCl₂ were added to the buffer and the sample was treated exactly the same as the protein samples.

Metal Analysis—Metal contents were analyzed using a PerkinElmer Optima 2100 DV ICP-OES spectrometer. Samples included two different purifications of apoACS and ITC, samples b and c, which were treated as described above. 500-μl protein samples were wet-washed overnight in a 1:1 mixture with 65% nitric acid (Suprapur, Merck, Darmstadt, Germany) at 100 °C. The samples were diluted with 4 ml of H₂O prior to their injection into the ICP-OES. The multielement standard solution XVI (Merck) was used as a reference.

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