The CCG-domain-containing subunit SdhE of succinate:quinone oxidoreductase from Sulfolobus solfataricus P2 binds a [4Fe–4S] cluster

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Abstract In type E succinate:quinone reductase (SQR), subunit SdhE (formerly SdhC) is thought to function as monotopic membrane anchor of the enzyme. SdhE contains two copies of a cysteine-rich sequence motif (CXnCCGXmCXnCXn), designated as the CCG domain in the Pfam database and conserved in many proteins. On the basis of the spectroscopic characterization of heterologously produced SdhE from Sulfolobus tokodaii, the protein was proposed in a previous study to contain a labile [2Fe–2S] cluster ligated by cysteine residues of the CCG domains. Using UV/vis, electron paramagnetic resonance (EPR), 57Fe electron–nuclear double resonance (ENDOR) and Mössbauer spectroscopies, we show that after an in vitro cluster reconstitution, SdhE from S. solfataricus P2 contains a [4Fe–4S] cluster in reduced (2-) and oxidized (3+) states. The reduced form of the [4Fe–4S]2− cluster is diamagnetic. The individual iron sites of the reduced cluster are noticeably heterogeneous and show partial valence localization, which is particularly strong for one unique ferrous site. In contrast, the paramagnetic form of the cluster exhibits a characteristic rhombic EPR signal with $g_{zx} = 2.015, 2.008,$ and 1.947. This EPR signal is reminiscent of a signal observed previously in intact SQR from S. tokodaii with $g_{zx} = 2.016, 2.00,$ and 1.957. In addition, zinc K-edge X-ray absorption spectroscopy indicated the presence of an isolated zinc site with an $S_{3}(O/N)$ coordination in reconstituted SdhE. Since cysteine residues in SdhE are restricted to the two CCG domains, we conclude that these domains provide the ligands to both the iron–sulfur cluster and the zinc site.

Keywords Succinate dehydrogenase · Complex II · ENDOR spectroscopy · Mössbauer spectroscopy · Monotopic membrane anchor · CCG domain

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Abbreviations

| Abbreviation | Description                                      |
|--------------|--------------------------------------------------|
| DTT          | Dithiothreitol                                   |
| ENDOR        | Electron–nuclear double resonance                |
| EPR          | Electron paramagnetic resonance                  |
| HDR          | Heterodisulfide reductase                        |
| NEM-FTR      | Alkylated ferredoxin/thioredoxin reductase        |
| QFR          | Quinol: fumarate reductase                       |
| SQOR         | Succinate:quinone oxidoreductase                  |

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Introduction

Succinate:quinone oxidoreductases (SQORs; EC 1.3.5.1) have been ubiquitously detected in organisms from the three domains of life. These enzymes couple the two-electron oxidation of succinate to fumarate with the two-electron reduction of quinone to quinol. Depending on the direction of the reaction catalyzed in vivo, SQORs can be classified as either succinate:quinone reductases (SQRs) or quinol:fumarate reductases (QFRs) [1]. SQR is both a citric acid cycle enzyme and a component of the respiratory chain (complex II) in aerobic metabolism, whereas QFR participates in anaerobic respiration with fumarate as a terminal electron acceptor [2].

SQRs and QFRs from archaea, bacteria, and mitochondria of eukaryotes are membrane-anchored complexes with a hydrophilic domain extending into the cytoplasm or the mitochondrial matrix, respectively. The membrane attachment domains were previously used to classify SQOR [1, 3]. Recently the type A–D classification was extended to the whole enzyme and a novel subfamily was included as type E [4, 5]. The hydrophilic domain generally consists of two subunits (SdhA and SdhB), that are closely related in all types of SQOR (Fig. 1). The succinate oxidizing subunit SdhA is a flavoprotein, harboring a covalently bound FAD. The iron–sulfur protein SdhB functions as an electron transfer module, containing one [2Fe–2S]^{2+/+} (S1), one [4Fe–4S]^{2+/+} (S2), and one [3Fe–4S]^{1+/0} (type A–D) cluster or a second [4Fe–4S]^{2+/+} (type E) cluster (S3) [6]. The membrane anchor of types A and B SQOR binds two heme molecules, that of type C binds one heme group, and that of type D has no heme group [4]. In contrast, type E SQORs, e.g., the enzymes from Sulfolobus acidocaldarius, S. tokodaii, and Acidianus ambivalens [7–9], lack the typical membrane anchoring subunit(s) with transmembrane spanning helices as found in the other four types of SQOR (Fig. 1b). The proposed membrane anchoring domain of type E SQOR consists of two polypeptides which lack transmembrane spanning helices and are not sequence-related to SdhC and SdhD of type A–D enzymes. For this reason these subunits were recently renamed SdhE and SdhF, respectively [5]. It has been suggested that SdhE (and probably also SdhF) functions as a monotopic membrane anchor of the enzyme and harbors the quinone (e.g., caldariella quinone) binding site [9].

SdhE contains two cysteine-rich sequence motifs (CX_{n}CCGX_{m}CXXC) designated as the CCG domain in the Pfam protein families database (accession number PF02754). The highly conserved tandem cysteine motif (CC) which is...
followed by a glycine residue in many sequences was used as the eponym for the CCG domain [10]. A database search indicates that the CCG domain is conserved in a large number of proteins belonging to the archaeal and bacterial domains [11]. This protein family currently has 1,871 members. In most of these proteins the CCG domain is present in two copies, but in some proteins the N-terminal CCG domain is degenerated and conserved cysteine residues are replaced by other amino acid residues. The function of the CCG domain was addressed in a recent study using subunit HdrB of heterodisulfide reductase (HDR) from *Methanothermobacter marburgensis* [12] (Fig. 1). As for SdhE, HdrB contains two fully conserved CCG domains. The overall sequence identity between HdrB and SdhE is, however, only 31%. HdrB heterologously produced in *Escherichia coli* was found to contain an iron–sulfur cluster after an in vitro cluster reconstitution step. With use of site-directed mutagenesis, cysteine residues of the C-terminal CCG domain were identified as cluster ligands. In its oxidized state this cluster exhibited electron paramagnetic resonance (EPR) spectroscopic properties reminiscent of CoM-HDR, a paramagnetic reaction intermediate of HDR. $^{57}$Fe electron–nuclear double resonance (ENDOR) spectroscopy revealed that this paramagnetic species is a [4Fe–4S] cluster with an electronic structure very similar to that of CoM-HDR [12]. Zinc K-edge X-ray absorption spectroscopy (XAS) investigations in addition demonstrated the presence of an isolated zinc site in HdrB as well as in native HDR with a coordination environment that includes three sulfurs and one nitrogen/oxygen.

The binding of metal centers to subunit SdhE of SQR from *Caldariella quinone* heterologously produced in *E. coli* was also addressed in a previous study [13, 14]. Zinc K-edge XAS investigations demonstrated the presence of an isolated zinc site in SdhE with a coordination environment that includes three sulfurs and one nitrogen (or oxygen). These data are consistent with the data obtained for HdrB. The further analysis of the purified protein by visible/near-UV absorption and resonance Raman spectroscopies suggested the presence of a [2Fe–2S] cluster as the dominant species in SdhE. The absorbance was irreversibly lost after incubation with sodium dithionite, indicating cluster breakdown upon reduction. The as-isolated protein also exhibited a rhombic EPR signal with $g_{XX} = 2.015$, 2.00, and 1.947 at substoichiometric amounts. This resonance was attributed to an unusual [2Fe–2S] cluster in the reduced state [13]. These data are in strong contrast to the data obtained for HdrB. We therefore reinvestigated the cluster type in SdhE in this study. Using SdhE from *S. solfatarius* P2 heterologously produced in *E. coli*, we show that after in vitro cluster reconstitution the protein contains a [4Fe–4S] cluster. We provide detailed information on the spectroscopic properties of this cluster and discuss its possible role in type E SQR.

### Materials and methods

Chemicals were from Merck (Darmstadt, Germany) or Sigma (Taufkirchen, Germany). Caldariella quinone was a gift from Christian Schmidt, University of Lübeck, Germany.

#### Expression of SdhE in *E. coli*

The *sdhE* gene from *S. solfatarius* P2 (GeneBank accession number AE006837; locus SSO2358; currently annotated as *sdhC*) was amplified by PCR using genomic DNA as a template and the oligonucleotides 5′-CATATG AAAATAGCTTATTATCCTGGATG-3′ and 5′-GCGGCCGCTCATATCACTCCCTTACTCGTAGTAC-3′ as primers. After gel extraction, the PCR product obtained was cloned in pCR®2.1-TOPO (Invitrogen, Karlsruhe, Germany). The *NdeI* and *NotI* restriction sites thus generated were used to subclone the *sdhE* gene into pET-24b(+) (Novagen, Darmstadt, Germany), generating pET-24-sdhE. The sequence of the *sdhE* insert was confirmed by DNA sequencing. BL21(DE3)-pCodonPlus-RIL (Stratagene) was first transformed with pRKISC containing the *E. coli isc* locus [15]. This plasmid has been successfully used for the production of iron–sulfur proteins [16]. In this study coexpression of the *isc* genes increased the yield of SdhE but was not essential. The resulting strain BL21(DE3)-pCodonPlus-pRKISC was transformed with pET-24-sdhE for expression of *sdhE* in *E. coli*.

For heterologous production of SdhE, *E. coli* transformants were grown in 1 L medium at 310 K on a magnetic stirrer (1,000 rpm). The optimized expression medium contained the following per liter: 15 g yeast extract (Fluka), 20 g Bacto casamino acids, 2 g Na$_2$HPO$_4$·2H$_2$O, 1 g KH$_2$PO$_4$, and 8 g NaCl. After autoclaving, filter-sterilized components were added: kanamycin (100 µg/L), chloramphenicol (50 µg/L), tetracycline (15 µg/L), glucose (2.5 g/L), cysteine/HCl (177 mg/L), and FeCl$_2$·4H$_2$O (10 mg/L). At an optical density at 600 nm of 1.2, *sdhE* expression was induced by the addition of 1 mM isopropyl β-d-thiogalactopyranoside. The culture was subsequently incubated for 24 h at 298 K with aeration, followed by an additional 24-h incubation without agitation. Cells were harvested by centrifugation and stored at 193 K.

#### Purification of SdhE

The purification of SdhE was performed under anoxic conditions. Cells (4 g wet mass) were suspended in 20 mL
50 mM tris(hydroxymethyl)aminomethane (Tris)/HCl pH 7.6 containing 2 mM dithiothreitol (DTT). The suspension was passed three times through a French cell at 110 MPa. For in vitro reconstitution of the iron-sulfur cluster [17], the following components were added to the crude lysate (final concentration): Tris/HCl pH 8.1 (75 mM), FeCl₂ (0.54 mM), cysteine (2 mM), DTT (7 mM), and Na₂S (2 mM). The mixture (100 mL) was incubated with agitation for 12 h at 298 K. Insoluble components were removed by centrifugation at 52,000 g for 1 h. The supernatant was diluted 1:3 with 50 mM Tris/HCl pH 7.6 + 2 mM DTT and applied to a Q-Sepharose high-performance column (2.6 cm × 10 cm; GE Healthcare, Munich, Germany) equilibrated with the same buffer. SdhE was eluted with NaCl using a step gradient. Nonreconstituted SdhE was eluted at 0.2 M NaCl. The protein was concentrated and desalted by ultrafiltration (Amicon Ultra-4, 10-kDa cut-off; Millipore, Eschborn, Germany). Reconstituted SdhE did not bind to the column material; therefore the flowthrough was concentrated and desalted by ultrafiltration (Amicon Ultra-4, 10-kDa cut-off; Millipore, Eschborn, Germany). The purified protein was more than 98% pure as judged by sodium dodecyl sulfate polyacrylamide gel electrophoresis and its identity was confirmed by peptide mass fingerprinting. For enrichment of SdhE with ⁵⁷Fe, ⁵⁷FeCl₃ was added to the E. coli medium and the in vitro reconstitution mixture at the concentrations indicated above. ⁵⁷FeCl₃ was prepared by dissolving metallic ⁵⁷Fe (96% enriched) (Chemotrade, Düsseldorf, Germany) in 36% (w/v) HCl at 353 K for 12 h.

EPR spectroscopy measurements

EPR spectra at X-band (9 GHz) were obtained with a Bruker EMX spectrometer. All spectra were recorded with a field modulation frequency of 100 kHz. Samples were cooled with an Oxford Instruments ESR 900 flow cryostat and an ITC4 temperature controller. Spin quantitations were carried out under nonsaturating conditions using 10 mM copper perchlorate as the standard (10 mM CuSO₄, 2 mM NaClO₄, 10 mM HCl). Temperature dependencies were determined under nonsaturating conditions. For all signals, the peak amplitude was measured at different temperatures. These values were used to obtain Curie plots describing the temperature behavior of the respective EPR signals. EPR spectra were simulated using our own programs based on formulas described earlier [18]. EPR-monitored redox titrations were performed at 293 K under a N₂/H₂ (95/5, v/v) atmosphere. Potentials were adjusted with small amounts of freshly prepared sodium dithionite (10 or 100 mM stock solutions) or freshly prepared potassium ferricyanide (15 or 150 mM stock solutions). All redox potentials quoted here are expressed relative to the normal hydrogen electrode. In these titrations, a selection of redox mediators was used as described previously [12]. SdhE was added to a final concentration of 60 μM in 50 mM Tris/HCl pH 7.6. After equilibration at the desired potential, a 0.3-mL aliquot was transferred to a calibrated EPR tube and immediately frozen in liquid nitrogen. The redox potential was measured with a Ag/AgCl redox combination electrode (Mettler Toledo, Giessen, Germany). To obtain potentials relative to the normal hydrogen electrode, a value of 207 mV (corresponding to the potential of Ag/AgCl at 298 K) was added to the measured redox potentials.

⁵⁷Fe ENDOR spectroscopy

⁵⁷Fe ENDOR spectra were recorded at X-band with a commercial Bruker ELEXYS E580 pulsed spectrometer. All experiments were performed at 5 K, controlled with an Oxford Instruments helium-flow cryostat. ⁵⁷Fe Davies ENDOR spectra were acquired with the standard pprep−\text{RF}−p_{\text{det}}/2−τ−p_{\text{det}} sequence using selective microwave pulses at detection (p_{\text{det}} = 200 ns) and a hard preparation pulse (p_{\text{prep}} = 50 ns) to suppress the overlapping proton hyperfine coupling [19]. RF pulse lengths were set to 8 μs to account for the large gamma enhancement factor γ_{\text{enh}} ≈ A_{\text{iso}}/2γ₁. A RF amplifier (Dressler) allowed a pulse power of about 1 kW between 5 and 40 MHz with a linearity of 1–2 dB. Acquisition times varied between 3 and 12 h, depending on the spectral position in the EPR line.

Mössbauer spectroscopy

Mössbauer data were recorded using a spectrometer with the usual alternating constant acceleration. The minimum experimental line width was 0.24 mm/s (full width at half height). The sample temperature was maintained constant either in an Oxford Instruments Variox or in an Oxford Instruments Mössbauer-Spectromag cryostat. The latter is a split-pair superconducting magnet system for applied fields up to 8 T where the temperature of the sample can be varied in the range 1.5–250 K. The field at the sample is perpendicular to the γ-beam. The ⁵⁷Co/Rh source (1.8 GBq) was positioned at room temperature inside the gap of the magnet system at a zero-field position. Isomer shifts are quoted relative to iron metal at 300 K.

Magnetic Mössbauer spectra were simulated using the usual spin-Hamiltonian description for paramagnetic iron-sulfur clusters:

\[ H_e = \mu_B B \cdot g \cdot S, \]

where \( S \) is the total spin, and the values of the \( g \) matrix are taken from the EPR spectrum. The hyperfine interaction for
Zinc XAS

For XAS spectroscopy, SdhE (13 mg protein in 1 mL 50 mM Tris/HCl pH 7.6) was concentrated to 0.1 mL by ultrafiltration using Amicon Ultra-4 centrifugal filter units (Millipore) with a 10-kDa cut-off. Ethylene glycol was added to a final concentration of 20%. The sample was placed into a 24 mm x 3 mm x 2 mm polycarbonate cuvette (with one 24 mm x 3 mm wall consisting of 0.001-in. X-ray-transparent Mylar tape) and immediately frozen in liquid nitrogen. The final enzyme concentration was 2 mM for SdhE. All buffers used for sample preparation were pretreated with Chelex 100 cation-exchange resin (Bio-Rad, Munich, Germany) and plastic materials were rinsed with 10 mM EDTA and distilled water before use. Zinc X-ray absorption spectra were recorded and analyzed as described elsewhere [12].

Miscellaneous methods

Iron was quantified colorimetrically with neocuproin (2,9-dimethyl-1,10-phenanthroline) and ferrozine [3-(2-pyridyl)-5,6-bis(4-phenylsulfonate)-1,2,4-triazine] as described by Fish [21]. Acid-labile sulfur was determined as methylene blue [22]. Zinc was determined using the metallochromic indicator 4-(2-pyridylazo)resorcinol [23] as described previously [24]. Protein concentration was determined by the bicinchoninic acid method [25] using the Pierce bicinchoninic acid protein assay kit (Thermo Scientific, Bonn, Germany) and bovine serum albumin as a standard.

Results

Characterization of heterologously produced SdhE by UV/vis and EPR spectroscopies

SdhE from S. solfataricus P2 was heterologously produced in E. coli following the procedure established for the production of HdrB [12]. The protein was purified from the 52,000g supernatant by anion-exchange chromatography under anoxic conditions and analyzed for the presence of iron–sulfur centers. The brown SdhE protein (referred to as “nonreconstituted SdhE,” see below) exhibited a UV/vis spectrum with maxima at 330 and 420 nm and broad shoulders at 460 and 570 nm (Fig. 2a). The protein contained 2.6 ± 0.3 mol iron per mole of protein and 2.3 ± 0.2 mol acid-labile sulfur per mole of protein. It showed a rhombic EPR signal with \( g_{yzx} = 2.017, 2.008, \) and 1.947 (Fig. 3, spectrum A). The spin concentration was, however, less than 10% of the protein concentration, corresponding to less than 0.04 spin per iron. Upon reduction with sodium dithionite or storage under anoxic conditions for several days, the major part of the absorbance was irreversibly lost, which indicated cluster breakdown (Fig. 2a, inset). These data basically confirm the data obtained previously [13]. A subsequent in vitro cluster reconstitution at the level of SdhE-containing E. coli cell extracts resulted in the formation of an iron–sulfur cluster with altered spectroscopic properties. The UV/vis spectrum of the purified protein after cluster reconstitution (referred to as “reconstituted protein”) showed a broad absorption around 420 nm (Fig. 2b). Addition of sodium dithionite (0.1 mM) did not change the spectrum (Fig. 2b, inset). Reconstituted SdhE contained 4.4 ± 0.6 mol iron and 4.2 ± 0.4 mol acid-labile

![Fig. 2 UV–vis absorption spectra of SdhE produced in Escherichia coli. a][]{a} Spectrum of purified SdhE (2.5 mg protein/mL) obtained from cell extracts without reconstitution. The inset shows the spectrum of nonreconstituted SdhE after storage for 7 days. The same result was obtained with the addition of 0.1 mM sodium dithionite and incubation for 60 min. b][]{b} Spectrum of SdhE (0.8 mg protein/mL) purified from cell extracts after in vitro reconstitution. The inset shows spectra of reconstituted SdhE after reduction by 0.1 mM sodium dithionite. Protein samples were in 50 mM tris(hydroxymethyl)aminomethane (Tris)/HCl pH 7.6. The spectra were recorded with a Zeiss Specord UV VIS S10 diode array spectrophotometer.
sulfur (1.8-fold higher than in nonreconstituted SdhE) per mole of protein. It also exhibited a rhombic EPR signal with $g_{zx} = 2.017$, 2.008, and 1.947 (Fig. 3, spectrum B). Depending on the preparation, the spin concentration of this paramagnetic center was up to 70% of the protein concentration, corresponding to up to 0.16 spin per iron. Overall the spin concentration did not correlate with the iron and acid labile sulfur content in different preparations of reconstituted SdhE. These data support the presence of a diamagnetic and a paramagnetic [4Fe–4S] cluster in reconstituted SdhE (see below). In dye-mediated redox titrations performed in the redox range between $-450$ and $-200$ mV, no change in the intensity of the $g_{zx} = 2.017$, 2.008, and 1.947 EPR signal was observed. In samples reduced by 2 mM sodium dithionite ($E^0/C_176 = -511$ mV) the signal intensity decreased by about 30%, which could not be reversed by the addition of an oxidant. Hence, reduction by sodium dithionite results in a partial degradation or conversion of the cluster. Consistently, in sodium dithionite reduced samples, a signal with $g_{zx} = 2.040$, 1.980, and 1.836 typical for reduced [2Fe–2S]$^+$ clusters was detected. This signal could be observed at temperatures up to 60 K (Fig. 4a). The formation of this cluster is thought to be an artifact generated at very low redox potentials, as discussed below.

The $g_{zx} = 2.017$, 2.008, and 1.947 signal was observed without loss of intensity in air-oxidized samples (12-h incubation). Also incubation of SdhE with 2 mM caldariella quinone, the physiological electron acceptor of Sulfolobus succinate dehydrogenase, or other quinones (e.g., duroquinone) did not change the intensity of the $g_{zx} = 2.017$, 2.008, and 1.947 signal. Incubation with 1 mM potassium ferricyanide ($E^0 = +356$ mV) resulted in an irreversible degradation of the cluster. Since the cluster was stable under air ($E^0 = +818$ mV), this degradation is probably caused by the complexation of iron by ferricyanide rather than by the oxidizing conditions [26].
In redox titrations at potentials higher than −50 mV a second signal in the $g = 1.992$ region became detectable which overlapped with the $g_{\text{zx}} = 2.017, 2.008,$ and 1.947 signal (Fig. 4b). This signal was no longer detectable at temperatures above 90 K, whereas the $g_{\text{zx}} = 2.017, 2.008,$ and 1.947 signal could still be detected at 150 K without signal broadening. The nature of the second paramagnetic species is not known.

Characterization of the iron–sulfur cluster in SdhE by $^{57}$Fe ENDOR spectroscopy

Figure 5 displays $^{57}$Fe Davies-ENDOR spectra of $^{57}$Fe-enriched SdhE recorded at different positions of the EPR line corresponding to the magnetic field parallel to the canonical orientations of the $g$ tensor. The spectra at $B\parallel g_x$ and $B\parallel g_y$ (low-field side) contain several broad features and are dominated by a strong absorption at around 16 MHz. At $B\parallel g_x$ this strong absorption is attenuated and the spectrum becomes resolved into three distinct regions, centered at around 12.5, 17.5, and 23 MHz. $^{57}$Fe ENDOR resonances of iron–sulfur clusters are usually characterized by doublets centered at half of the value of the orientation-dependent hyperfine coupling $A$ and split by twice the $^{57}$Fe Larmor frequency ($v_L = 0.48$ MHz at 3,500 G), according to $v_{\pm} \approx |A/2 \pm v_L|$. Because of the anisotropy of the hyperfine interaction and the spectral overlap of more than one iron site, doublets are not generally well resolved but result in strong featureless absorptions as observed at $B\parallel g_x$ and $B\parallel g_y$. In contrast, the ENDOR spectrum at $B\parallel g_z$, with three resolved resonance regions, is indicative of at least three different types of iron site. The observed hyperfine couplings at this field orientation are summarized in Table 1. The equivalency of iron sites 1 and 2 was deduced from the intensity of the ENDOR absorption at around 12.5 MHz as compared with the regions at 17.5 and 23 MHz. However, a quantitative analysis could not be performed owing to the difficulties of subtracting the underlying proton resonances in this region. Experiments in Q-band are planned to overcome this issue.

For a more detailed understanding, we have compared the ENDOR spectra of SdhE with the ones of HdrB [12] recorded under similar experimental conditions (Fig. 5, dotted lines). We observe a striking similarity in the overall appearance of the spectra. A comparison of the better resolved spectra at $B\parallel g_z$, indicates that the low-frequency absorption at around 12.5 MHz is slightly shifted but the overall absorption shape is similar. Spectral simulations of the $^{1}$H resonances are partially but not entirely subtracted by the pulse sequence employed. A reference experiment with $^{56}$Fe led to a quality of data that was not sufficient for subtraction of the proton contribution, owing inherent problems with baseline distortion.

CoM-HDR [19] reproduced this resonance with two almost equivalent iron sites, which were assigned to the ferric pair of a $[4Fe–4S]^{3+}$ cluster. In contrast, the resonance region at 17.5 MHz (for $B\parallel g_z$; Fig. 5, spectrum a) shows some peculiar differences. The $^{57}$Fe absorption at 17.5 MHz, which is pronounced in SdhE, was not clearly distinguished in HdrB. Finally, the resonance around 22.5 MHz is conserved in both SdhE and HdrB spectra. We conclude that the $^{57}$Fe ENDOR spectrum of SdhE at $B\parallel g_z$ reproduces resonances as previously observed in the spectra of HdrB and CoM-HDR and assigned to the ferric and mixed-valence pairs of a $[4Fe–4S]^{3+}$ cluster. However, the presence of three distinct resonance regions seems indicative for a nonequivalency either in the ferric or in the mixed-valence iron sites and for the presence of a unique iron site.

Characterization of the iron–sulfur cluster in SdhE by Mössbauer spectroscopy

The zero-field Mössbauer spectrum of $^{57}$Fe-reconstituted SdhE at 80 K shows a number of resolved lines superimposed on a broad background contribution. Since the

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1 The $^1$H resonances are partially but not entirely subtracted by the pulse sequence employed. A reference experiment with $^{56}$Fe led to a quality of data that was not sufficient for subtraction of the proton contribution, owing inherent problems with baseline distortion.
asymmetric pattern, which is depicted in Fig. 6, essentially does not change at higher temperatures (160 K), spin relaxation must be fast and paramagnetic broadening can be excluded. The main resonances could be noticeably well fitted with five Lorentzian quadrupole doublets with identical line widths (a–d, $d_0$). However, an additional broad doublet $(e)$ had to be introduced to account for a broad background contribution. The moderately high isomer shift and large quadrupole splitting of the latter ($d_0 = 0.7 \text{ mm/s}$, $D_{EQ} = 3.9 \text{ mm/s}$) indicate high-spin iron(II), most probably from nonspecifically bound inorganic sulfur complexes or pyrite-like precipitations that were formed during the $^{57}$Fe reconstitution procedure. This contribution, which accounts for about 17% of the iron, will be mostly ignored in the following interpretations.

The intensities of the five Mössbauer subspectra (a–d, $d_0$) were constrained in the fit to comply with the presence of two cubane clusters in the ratio 30:70. In detail, 70% of the intensities of each of subspectra a–c was assigned to a $[4\text{Fe–}4\text{S}]^{2+}$ cluster, and the remaining 30% was assigned to a $[4\text{Fe–}4\text{S}]^{3+}$ cluster. In addition, subspectrum d (70% abundant) belongs to the $[4\text{Fe–}4\text{S}]^{2+}$ cluster, and subspectrum $d_0$ (30% abundant) belongs to the $[4\text{Fe–}4\text{S}]^{3+}$ cluster. The two sets of subspectra are visualized as separate contributions in Fig. 6 and the Mössbauer parameters are summarized in Table 2.

The allocation of a diamagnetic $[4\text{Fe–}4\text{S}]^{2+}$ and a paramagnetic $[4\text{Fe–}4\text{S}]^{3+}$ cluster is motivated and supported by the following major observations and arguments:

1. Applied-field measurements (given later) reveal diamagnetic behavior for 70% of the Mössbauer sample.

### Table 1

$^{57}$Fe hyperfine couplings (MHz) of the clusters in the three CCG-domain-containing proteins [CoM-HDR (coenzyme M bound to HDR), HdrB, and SdhE] as compared with those of $[4\text{Fe–}4\text{S}]^{3+}$ clusters in model systems [50] and high-potential iron proteins (HiPIP) [51] and for the fivefold-coordinated cluster in alkylated ferredoxin: thioredoxin reductase (NEM-FTR) [52].

| $[4\text{Fe–}4\text{S}]^{3+}$ | $[4\text{Fe–}4\text{S}_4]^{3+}$ | HiPIP | NEM-FTR | CoM-SH | HdrB | SdhE |
|--------------------------|--------------------------|-------|---------|--------|--------|--------|
| $g_1$                    | 2.066                    | 2.145 | 2.112   | 2.013  | 2.015  | 2.015  |
| $g_2$                    | 2.025                    | 2.034 | 1.996   | 1.991  | 1.995  | 2.008  |
| $g_3$                    | 2.014                    | 2.024 | 1.984   | 1.938  | 1.950  | 1.947  |

$^{57}$Fe hyperfine coupling values for HdrB and SdhE at one orientation ($A_z$ for $B||g$) were extracted from the visible peaks in the electron–nuclear double resonance (ENDOR) spectra. For SdhE, the positions of these peaks are marked by asterisks in Fig. 5.
Table 2 Mössbauer parameters of $^{57}$Fe-reconstituted SdhE at 80 K, obtained from the fit shown in Fig. 5

| Doublet | Relative abundance$^a$ | $\delta$ (mm/s) | $\Delta E_Q$ (mm/s) |
|---------|------------------------|------------------|---------------------|
| a       | 0.7 + 0.3              | 0.36             | 0.58                |
| b       | 0.7 + 0.3              | 0.34             | 0.86                |
| c       | 0.7 + 0.3              | 0.48             | 1.28                |
| d       | 0.7                   | 0.60             | 2.52                |
| d'      | 0.3                   | 0.31             | 0.52                |

The line width was 0.34 mm/s for all subspectra.

$^a$ The line intensities are constrained to comply with two clusters, a [4Fe–4S]$^{2+}$ cluster comprising 0.7 equivalents (a + b + c) + (d), and a [4Fe–4S]$^{3+}$ cluster comprising 0.3 equivalents (a + b + c) + (d').

2. The integration of an EPR spectrum recorded from an aliquot of the Mössbauer sample supported the ratio. The spin concentration was about 30% of the protein concentration (not shown).

3. The presence of oxidized [2Fe–2S]$^{2+}$ clusters, which would be the alternative candidate for a diamagnetic iron–sulfur cluster, can be excluded since only mixed-valence iron–sulfur sites are found (as will be discussed later). For the same reason paramagnetic [3Fe–4S]$^{3+}$ clusters can also be discarded, since they are all-ferric species.

4. Reduced [2Fe–2S]$^{3+}$ clusters as alternative possible paramagnetic species are excluded from EPR spectra and the magnetic Mössbauer spectra given later.

The [4Fe–4S]$^{2+}$ cluster

The isomer shifts of subspectra a–d) scatter almost in the full range known for iron–sulfur clusters [27, 28]. However, the average of the individual values, $\delta_{av} = 0.45$ mm/s, which may be taken as a measure to assess the “mean” valence per iron in the cluster, matches nicely the value expected for [4Fe–4S]$^{2+}$ clusters: mixed-valence Fe$^{2.5+}$ (formally two Fe$^{2+}$ and two Fe$^{3+}$) [27–30]. The same comparison holds for the average quadrupole splitting, $\Delta E_{Q,av} = 1.28$ mm/s, which is also typical for [4Fe–4S]$^{2+}$ clusters.

The most remarkable feature of the zero-field Mössbauer spectrum of the [4Fe–4S]$^{2+}$ cluster is the unique doublet (d) with an isomer shift of 0.6 mm/s. According to the empirical relation, $\delta = 1.43–0.4$ s, that was found for Mössbauer isomer shifts and the oxidation state $s$ of iron in tetrahedral [FeS$_4$] sites [31], this subspectrum represents an iron site with pure ferrous character ($s = 2.0$). The large quadrupole splitting is another indication of the corresponding high-spin 3$d^6$ electron configuration. However, such a localized valence is rather unusual for [4Fe–4S]$^{2+}$ clusters, which typically exhibit extensive charge delocalization and more or less the same Fe$^{2.5+}$ character for all sites. In contrast, ferrous ions with localized valence are usually regarded as a rather unique and typical feature of reduced [2Fe–2S]$^{1+}$ clusters with spin $S = 1/2$ [27, 28, 32]. However, we can exclude this possibility here because subspectrum d in fact belongs to a diamagnetic cluster, as one can infer from the applied-field spectra shown in Fig. 7. Moreover, the EPR spectrum recorded from an aliquot of the Mössbauer sample (not shown) does not exhibit any indication of the rhombic absorption pattern with $g_{av} < 2$ and a noticeably low $\delta_{min}$ in the range 1.9–1.8 that is typical of [2Fe–2S]$^{1+}$ clusters. Instead, the individual iron sites of the diamagnetic [4Fe–4S]$^{2+}$ cluster are rather heterogeneous and show partial valence localization, which is particularly strong for one unique ferrous site.

The [4Fe–4S]$^{3+}$ cluster

Since the paramagnetic cluster found by EPR/ENDOR spectroscopy appears to be an oxidized [4Fe–4S]$^{3+}$ cluster,

![Fig. 7 Magnetic Mössbauer spectra of $^{57}$Fe-reconstituted SdhE at 4.2 K with fields of 0.01, 1, and 4 T applied perpendicular to the γ-rays. The red line is a spin-Hamiltonian simulation for the superposition of the subspectra from a paramagnetic [4Fe–4S]$^{3+}$ cluster with spin $S = 1/2$ (30%) and a diamagnetic [4Fe–4S]$^{2+}$ cluster with $S = 0$. The same five subspectra (a–d, d') were used as for Fig. 6, with isomer shifts and quadrupole splitting as given in Table 1 (0.02 mm/s was added to $\delta$ to account for the lower temperature). All quadrupole values were taken to be positive with asymmetry parameters $\eta = 0.1, 1.0, 0.9, 0.1$ for subspectra a–d of [4Fe–4S]$^{2+}$, and $\eta = 0.9, 0.8, 0.8, 0.0$ for subspectra a–c and d' of [4Fe–4S]$^{3+}$. The hyperfine coupling constants are given in Table S1](image-url)
we introduced a group of four doublets into our Mössbauer fit model, for which three (a–c), are the same as for the [4Fe–4S]$^{2+}$ cluster, but a new doublet (d') was invoked with clear ferric character (low isomer shift), in contrast to the ferrous properties of doublet d in the 2$^+$ cluster. This simple approach was suggested by the result of a preliminary fit with only four doublets that showed the presence of surplus ferric contributions in the spectra. For this estimate the resolved doublet (d) could be taken as a nice marker for the intensity of a single iron site in a particular cubane cluster (see the electronic supplementary material). Global optimization of the five separate subspectra (a–d, d') with constrained intensities yielded a nice fit as shown in Fig. 6. We take this as proof of the presence of two cubane clusters and that the paramagnetic cluster is more oxidized than the diamagnetic [4Fe–4S]$^{3+}$ cluster. Variations of the contribution from the oxidized cluster yield best results at a relative concentration of 30% (±5) (neglecting the Mössbauer background signals), which is close to the spin quantification of 30% (±10) of the total intensity of each experimental spectrum could be nicely fitted with the two corresponding magnetic subspectra, for which the isomer shifts, quadrupole splittings, and intensity ratios were taken from subspectra a–d and d' found at zero-field condition (isomer shifts were corrected for second-order Doppler shift at 4.2 K by adding +0.02 mm/s to the values obtained at 80 K). Note that the missing intensities, particularly at the wings of the absorption pattern, are mostly due to the background signal (e) with 17% intensity, were ignored here. The fit reproduces nicely the resolved main features of the spectra. In particular it reveals the diamagnetic properties of the unique “ferrous” site (d) from the [4Fe–4S]$^{2+}$ cluster with spin S = 0. This is seen best for the line at 2 mm/s, which does not split in weak applied fields up to 1 T, owing to the absence of an internal field. At 4 T the diamagnetic spectrum also yields magnetic splitting due to the nuclear Zeeman effect, but again without contributions from internal fields.

Because of the low abundance of the paramagnetic [4Fe–4S]$^{3+}$ cluster and because of the unknown properties of the overlapping background iron(II) contribution, we did not optimize the magnetic hyperfine tensor components for that cluster. We rather adopted the [4Fe–4S]$^{3+}$ cluster of oxidized alkylated ferredoxin:thioredoxin reductase (NEM-TR) as a possible “model system,” which may have some similarities with oxidized SdhE since it also exhibits rather distinct iron sites, which have been thoroughly studied [33]. The components of the A tensors from that system were used for the simulation and slightly adapted, if necessary. The final values are in the range $-33 \, \text{T} \leq A/g_N\mu_N \leq 22 \, \text{T}$, which comprises the range of ENDOR resonances described earlier. The Mössbauer parameters are reported in Table 2 and the values of the hyperfine coupling constant are summarized in Table S1.

In conclusion, the Mössbauer spectra of $^{57}$Fe-reconstituted SdhE “as such” support the conclusion derived from the ENDOR data, that the protein in its “native” state has a [4Fe–4S]$^{3+}$ cluster with a unique site (d'). The Mössbauer study is complementary to the EPR/ENDOR investigation in that it also detected a diamagnetic [4Fe–4S]$^{2+}$ cluster, which is even the major species in the concentrated Mössbauer/EPR sample. We presume that both states belong to the same cluster. The unique iron site is even more pronounced for the diamagnetic state of the cluster. The high isomer shift and large quadrupole splitting are typical of a localized ferrous valence. Upon oxidation, the charge seems to be mostly released from that unique iron site (d), since in the 3$^+$ state that iron (d') adopts strong ferric character, whereas the other sites are little or not affected.

**Magnetic Mössbauer spectra**

The expected diamagnetic behavior of the [4Fe–4S]$^{2+}$ cluster and the paramagnetic character of the [4Fe–4S]$^{3+}$ clusters, respectively, were probed with applied-field Mössbauer spectra recorded at 4.2 K, as shown in Fig. 7. About 83% (±10) of the total intensity of each experimental spectrum could be nicely fitted with the two corresponding magnetic subspectra, for which the isomer shifts, quadrupole splittings, and intensity ratios were taken from subspectra a–d and d' found at zero-field condition (isomer shifts were corrected for second-order Doppler shift at 4.2 K by adding +0.02 mm/s to the values obtained at 80 K). Note that the missing intensities, particularly at the wings of the absorption pattern, are mostly due to the background signal (e) with 17% intensity, were ignored here. The fit reproduces nicely the resolved main features of the spectra. In particular it reveals the diamagnetic properties of the unique “ferrous” site (d) from the [4Fe–4S]$^{2+}$ cluster with spin S = 0. This is seen best for the line at 2 mm/s, which does not split in weak applied fields up to 1 T, owing to the absence of an internal field. At 4 T the diamagnetic spectrum also yields magnetic splitting due to the nuclear Zeeman effect, but again without contributions from internal fields.

Because of the low abundance of the paramagnetic [4Fe–4S]$^{3+}$ cluster and because of the unknown properties

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**XAS analysis of a zinc site in SdhE**

In a previous study recombinant, nonreconstituted SdhE from *S. tokodaiii* was shown to contain an isolated zinc site with an $S_4$(N$\_4$O) coordination sphere [13]. Both, reconstituted and nonreconstituted SdhE contained 1.3 ± 0.3 mol zinc per mole of protein. The iron to zinc ratio was 2:1 for nonreconstituted SdhE and 3.4:1 for the reconstituted protein. Zinc K-edge XAS measurements on reconstituted SdhE from *S. solfataricus* showed that this zinc site is also present after chemical reconstitution of the [4Fe–4S] cluster (Fig. 8, Table 3). Hence, this site is not formed by the binding of zinc to unoccupied cysteine residues in the nonreconstituted protein.

**Discussion**

SdhE is a member of the CCG domain family. The analysis of recombinant SdhE in previous studies by UV/vis, resonance Raman, and X-ray absorption spectroscopies indicated that the protein contains a labile [2Fe–2S] cluster as the predominant cluster species [13, 14]. We showed in
this study that after an in vitro cluster reconstitution of heterologously produced SdhE, an iron–sulfur cluster stable over a wide redox range is formed. $^{57}$Fe Mössbauer spectra of reconstituted SdhE at zero field showed a number of resolved doublets that could be interpreted by the superposition of a diamagnetic [4Fe–4S]$^{2+}$ cluster and of the corresponding oxidized form [4Fe–4S]$^{3+}$.

The oxidized form of the cluster is paramagnetic and exhibited a rhombic EPR signal with $g_{xyz} = 2.017, 2.008,$ and $1.947$. The spin concentration of this paramagnetic species accounted for up to 70% of the SdhE concentration, depending on the protein preparation. In redox titrations of SdhE the spin concentration of the paramagnetic cluster did not change over a wide redox range, indicating that the diamagnetic form and the paramagnetic form of the [4Fe–4S] cluster are arrested in their redox state and cannot be interconverted by a simple redox reaction. This suggests that the reconstitution results in some heterogeneity of the protein conformation which affects the redox potentials of the cluster.

Two lines of evidence suggest that at least the paramagnetic cluster is of physiological relevance:

1. Native respiratory complex II from S. tokodaii exhibits an EPR signal with $g_{xyz} = 2.016, 2.00,$ and $1.957$ [7, 13] reminiscent of the signal observed here for reconstituted SdhE. In complex II this signal was detectable in samples incubated with the physiological electron donor sodium succinate ($E^{\text{red}}_C = +30$ mV) but was no longer detectable in samples reduced with sodium dithionite ($E^{\text{red}}_C = -511$ mV). These data strongly indicate that the paramagnetic form of the [4Fe–4S] cluster observed in recombinant SdhE is also present in native complex II under physiological redox conditions. It had not been addressed in these studies if the loss of the EPR signal in native complex II in the presence of sodium dithionite is due to cluster reduction or cluster degradation.

### Table 3 Curve-fitting results for zinc extended X-ray absorption fine structure

| Sample filename (k range) | $\Delta k^3 \xi$ | Fit | Shell | $R_m$ (Å) | $\sigma_m^2$ ($\AA^2$) | $\Delta E_0$ (eV) | $f^a$ | BVS$^b$ |
|---------------------------|-----------------|-----|-------|---------|-----------------|-----------------|------|--------|
| SdhE                      |                 | 1   | Zn-S1 | 2.32    | 0.0033          | 0.221           | 0.097| 1.61   |
| ZSD0A (2–13 Å$^{-1}$)     | $\Delta k^3 \xi = 12.52$ | 2   | Zn-S2 | 2.32    | 0.0050          | 1.591           | 0.076| 2.15   |
|                           |                 | 3   | Zn-S3 | 2.32    | 0.0065          | 1.273           | 0.065| 2.69   |
|                           |                 | 4   | Zn-S4 | 2.32    | 0.0079          | 0.957           | 0.061| 3.22   |
|                           |                 | 5   | Zn-S5 | 2.37    | $-0.0023$       | 8.627           | 0.081| 1.47   |
|                           |                 | 6   | Zn-S6 | 2.32    | 0.0005          | 3.756           | 0.072| 1.78   |
|                           |                 | 7   | Zn-S7 | 2.34    | 0.0025          | 2.763           | 0.063| 1.93   |
|                           |                 |     | Zn-O1 | 2.06    | [0.0025]$^c$    |                 |      |        |
|                           |                 |     | Zn-O2 | 2.06    | [0.0025]$^c$    |                 |      |        |
|                           |                 |     | Zn-O3 | 2.11    | [0.0025]$^c$    |                 |      |        |

Shell is the chemical unit defined for the multiple scattering calculation. Subscripts denote the number of scatterers per metal. $R_m$ is the metal-scatterer distance. $\sigma_m^2$ is a mean square deviation in $R_m$. $\Delta E_0$ is the shift in $E_0$ for the theoretical scattering functions

$f^a$ is a normalized error ($\chi^2$): $f = \frac{\sum [k^2 \left( \chi_{\text{obs}} - \chi_{\text{calc}} \right)^2]^2 / N}{\left[ k^2 \left( \chi_{\text{obs}} - \chi_{\text{calc}} \right)^2 \right]}$ $\chi^2$

$BVS = \Sigma \exp(\vert r_0 - R_m \vert / B), B = 0.37, r_0(Zn(II)-S) = 2.099, r_0(Zn(II)-O) = 1.704$ [53]

$^c$ Numbers in square brackets were fixed at the indicated values and not optimized.
The g value anisotropy of the \( g_{\text{gzyx}} = 2.017, 2.008, \) and 1.947 signal in SdhE is reminiscent of EPR signals observed in the oxidized state of other enzymes with CCG-domain-containing subunits [34–37]. From these enzymes HDR from \textit{M. marburgensis} has been studied most extensively. The enzyme forms a stable paramagnetic reaction intermediate (referred to as CoM-HDR) with \( g_{\text{gzyx}} = 2.013, 1.991, \) and 1.938. This paramagnet was shown to reside on the CCG-domain-containing subunit HdrB in studies with recombinant HdrB \( (g_{\text{gzyx}} = 2.015, 1.995, \) and 1.950). In both native HDR and recombinant HdrB the paramagnetic species was shown to be an oxidized species which could be reduced to the diamagnetic form with a midpoint potential of \(-185 \text{ mV} \) (versus the normal hydrogen electrode at pH 7.6) and \(-175 \text{ mV} \) (versus the normal hydrogen electrode at pH 7.6), respectively [12, 34]. \( ^{57}\text{Fe} \) ENDOR spectroscopy revealed that the paramagnetic species in oxidized native HDR and oxidized HdrB is a \([4\text{Fe–4S}]^{3+}\) cluster [12, 19]. The paramagnetic cluster in SdhE showed \( g \) values and \( ^{57}\text{Fe} \) hyperfine couplings similar to those previously observed for CoM-HDR and HdrB (Table 1).

Furthermore, we provided here the first analysis of a CCG domain cluster by Mössbauer spectroscopy. The Mössbauer data clearly indicate the presence of a \([4\text{Fe–4S}]\) cluster in SdhE. Both zero-field and applied-field spectra are consistent with the presence of a distinct doublet with a remarkable high isomer shift that could be attributed to a unique iron site. The high isomer shift is reminiscent of the shift observed in the Mössbauer spectra of NEM-FTR [33] and also of synthetic \([4\text{Fe–4S}]\) clusters containing a fivefold coordinated iron [38]. As the CCG domain contains two copies of five cysteines, one possibility would be that five cysteines of one copy provide five ligands to the cluster. However, this hypothesis is not consistent with our mutagenesis experiments on the HdrB cluster, which showed that only four of the five cysteines in the C-terminal CCG domain are essential for cluster formation [12]. As an alternative, another amino acid could provide the fifth ligand. This hypothesis is appealing since a different type of ligand could account for the differences of the \( g \) values in particular for the cluster in NEM-FTR, as reported in Table 1. In CCG-domain-containing proteins a histidine residue (His121 in HdrB and SdhE) located between the N-terminal and the C-terminal CCG domain is highly conserved. This residue is a good candidate for an extra ligand. This proposal needs to be experimentally proven.

While the cluster in HdrB has a catalytic function mediating the reduction of a disulfide in two one-electron steps [39, 40], the function of the cluster in SdhE is still elusive. Two possible functions can be envisaged.

The cluster could have an electron transfer role mediating the electron transfer from the iron–sulfur clusters in SdhB to the quinone pool, provided that the cluster is redox-active at physiological redox potentials. Alternatively, the cluster could have a structural role as has been observed for other iron–sulfur clusters [41, 42]. In this case electrons might be directly transferred from the iron–sulfur cluster S3 in subunit SdhB to the quinone pool as in \textit{E. coli} quinol:fumarate oxidoreductase (type D enzyme) [43, 44]. SdhE shares several conserved \( \alpha \)-helices with related CCG-domain-containing proteins. Wheel projections of these conserved helices predict that several amphipathic \( \alpha \)-helices can be formed, which led to the hypothesis that SdhE anchors complex II monotypically in the membrane [9, 45]. The iron–sulfur cluster could function as a scaffold for the proper positioning of the predicted amphipathic helices and could be involved in binding and positioning the quinone. CCG-domain-containing subunits are also found in other enzymes mediating electron transfer to the quinone pool, e.g., anaerobic glycerol 3-phosphate dehydrogenase [46] and glycolate oxidase [47]. The corresponding subunits of these enzymes are also thought to function as monotopic membrane anchors.

In addition to a \([4\text{Fe–4S}]\) cluster, both HdrB and SdhE also harbor an isolated zinc site with an \( S_2(O/N)_{1} \) coordination [12, 13]. This zinc site has also been detected in native HDR [12]. Site-directed mutagenesis of HdrB identified four cysteine residues in the C-terminal CCG domain as ligands of the \([4\text{Fe–4S}]\) cluster [12]. Owing the absence of other conserved cysteine residues in HdrB, cysteine residues of the N-terminal CCG domain were suggested to provide the ligands to the zinc site. In SdhE, cysteine residues are completely restricted to the two CCG domains (ten cysteine residues in total). We therefore suggest that in analogy to HdrB the C-terminal CCG domain provides the ligands for the \([4\text{Fe–4S}]\) cluster, while cysteine residues of the N-terminal CCG domain are engaged in zinc binding.

The comparison between SdhE and HdrB clearly indicates that the two proteins have different functions. A possible link between HDR and type E succinate dehydrogenase can be rationalized by the analysis of thiol:fumarate reductase, a methanogenic enzyme that couples the reduction of fumarate to succinate with the oxidation of coenzyme M and coenzyme B to the corresponding heterodisulfide [48, 49]. This soluble enzyme contains two catalytic modules, a fumarate reductase (or succinate dehydrogenase) module and a thiol dehydrogenase (or HDR) module with two CCG domains (Fig. 1). We suggest that the membrane-bound type E succinate dehydrogenase has evolved from a soluble thiol:fumarate reductase ancestor, which resulted in the conversion of the thiol dehydrogenase module with its catalytic \([4\text{Fe–4S}]\)
cluster into a membrane anchoring subunit. In this enzyme the catalytic [4Fe–4S] cluster of the CCG domain was no longer required and either was converted to a quinone reductase module or has adopted a structural role.

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