Paramagnetic $^1$H NMR spectroscopy to investigate the catalytic mechanism of radical S-adenosylmethionine enzymes

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Abstract

Iron-sulfur clusters in radical S-adenosylmethionine (SAM) enzymes catalyze an astonishing array of complex and chemically challenging reactions across all domains of life. Here we showed that $^1$H NMR spectroscopy experiments tailored to reveal hyperfine-shifted signals of metal-ligands is a powerful tool to monitor the binding of SAM and of the octanoyl-peptide substrate to the two [4Fe-4S] clusters of human lipoyl synthase. The paramagnetically shifted signals of the iron-ligands were specifically assigned to each of the two bound [4Fe-4S] clusters, and then used to examine the interaction of SAM and substrate molecules with each of the two [4Fe-4S] clusters of human lipoyl synthase. $^1$H NMR spectroscopy can therefore contribute to the description of the catalytic mechanism of radical SAM enzymes.

Keywords

Lipoyl synthase; iron-sulfur proteins; enzyme mechanism; electron transfer; metallo enzyme.
Lipoyl synthase (LIAS in humans) is a member of the radical S-(5′-Adenosyl)-L-Methionine (SAM) superfamily of enzymes and uses two [4Fe-4S] clusters to catalyze the final step of the biosynthesis of the lipoyl cofactor [1-7]. The mechanism of lipoyl synthase consists of a two-step reaction where a protein-bound octanoyl chain is converted into lipoic acid by a consecutive insertion of two sulfur atoms at C6 and C8 positions of the octanoyl chain [8, 9]. The two [4Fe-4S] clusters are both involved in the catalytic mechanism [9, 10]. One of them (Fig. 1), typical of all radical SAM enzymes (hereafter named FeSRS), performs a reductive cleavage of a SAM molecule to obtain methionine and a 5′-deoxyadenosyl radical (5′-dA•) [1]. The 5′-dA• radical serves to generate a radical on the octanoyl chain. The other cluster (usually defined as auxiliary cluster and named FeSaux hereafter, Fig. 1) provides the inorganic sulfides to the formed octanoyl chain radical [5, 9-11]. The FeSRS cluster is bound to a CX₃CX₂ motif, i.e. three iron ions are covalently bound to three Cys residues of the motif (Cys 137, Cys 141 and Cys 144 in LIAS) and the fourth iron ion, termed as catalytic iron ion, is exposed for the binding of SAM (Fig. 1) [12, 13]. The FeSaux cluster is bound through a conserved CX₄CX₅C motif (Cys 106, Cys 111 and Cys 117 in LIAS) and a serine (Ser 345 in LIAS) (Fig. 1) [12, 13]. The complete turnover of lipoyl synthase requires two equivalents of SAM (one per sulfur insertion) and two sulfur atoms from the auxiliary cluster. Therefore, lipoyl synthase typically catalyzes no more than one turnover in in vitro reactions as a consequence of the FeSaux cluster disruption [1, 14]. Recently, it was shown that the Fe-S cluster carrier protein NfuA from Escherichia coli can regenerate the auxiliary cluster of E. coli lipoyl synthase (LipA) after each turnover. In such a way, although the consumption of the FeSaux cluster after each turnover, LipA can act catalytically upon the continuous supply of [4Fe-4S] clusters by NfuA [15]. On the other hand, this mechanism of action, i.e. the FeSaux cluster disruption, implies the release of free iron and inorganic sulfide ions and it would seem to present a problem for the cell considering the high iron and sulfide toxicity. Thus, it is reasonable that several important chemical aspects remain to be answered in order to have a complete description of the various steps of the enzymatic cycle [16]. The most enigmatic questions
concerns how the second sulfur atom is inserted in the substrate and the ultimate fate of the degraded auxiliary cluster.

![Figure 1. Structure of lipoic synthase from *Mycobacterium tuberculosis* in ribbon representation (PDB ID 5EXJ). The FeS<sub>RS</sub> and FeS<sub>aux</sub> clusters and the iron ligands, numbered according to LIAS sequence, are shown in CPK mode and sticks, respectively.](image)

This study reports for the first time that $^1$H paramagnetic NMR spectroscopy is a valuable tool investigating the binding of SAM and substrate molecules to the two clusters of LIAS, in such a way resulting instrumental to contribute answering the still open questions of the enzymatic mechanism. To achieve this, we have exploited the hyperfine-shifted $^1$H NMR signals of iron ligands of FeS<sub>RS</sub> and FeS<sub>aux</sub> clusters and follow their changes upon cluster interaction with SAM, with compounds mimicking SAM (S-(5’-Adenosyl)-L-Homocysteine, SAH) and the breakdown product of SAM (5’-deoxy-5’-MethylThioAdenosine, MTA), and with a synthetic
octanoyl-peptide substrate (Glu-Ser-Val-(N\textsuperscript{6}-octanoyl)Lys-Ala-Ala-Ser-Glu, named octanoyl-peptide hereafter) (Fig. S1).

Wild-type LIAS lacking the N-terminal mitochondrial targeting sequence was expressed and purified from *E. coli* cells (WT LIAS, hereafter). The protein was obtained highly pure and monomeric in solution (Fig. 2a) with a characteristic brown colour, displaying, in the UV-visible absorption spectrum, a weak shoulder at 330 nm and a broad peak at 400 nm, in addition to the peak at 280 nm due to aromatic amino acids (Fig. 2b). The peak at 400 nm is characteristic of [4Fe-4S]\textsuperscript{2+} clusters with a delocalized redox state, i.e. with two Fe\textsuperscript{2.5+}-Fe\textsuperscript{2.5+} pairs [17-19]. The UV-visible spectrum of WT LIAS does not significantly change upon its chemical reconstitution; consistently the signals in the paramagnetic \textsuperscript{1}H NMR spectra of WT LIAS remain unperturbed before and after chemical reconstitution (data not shown). Therefore, the chemically reconstituted procedure was avoided for all WT LIAS samples used in the following spectroscopic characterization.

Figure 2. Analytical gel filtration and UV-visible absorption spectra of LIAS. (a) Analytical gel filtration chromatogram (protein concentration 500 μM) in 50 mM potassium phosphate, pH 7.0, with 5 mM DTT. The protein eluted at a volume corresponding to a molecular mass of 41 kDa. Right inset: Standard calibration curve for the Superdex 200 10/300 increase column. Left inset: SDS-PAGE of the last purification step of WT LIAS, 1. proteins obtained after the first HiTrap chelating column; 2. protein marker in kDa ; 3. protein obtained after TEV protease cleavage and a second passage on HiTrap chelating column. (b)
UV-visible absorption spectra of WT LIAS (blue line), chemically reconstituted C137/C141/C144A LIAS variant (green line), as purified C106/C111/C117A LIAS variant (black line). The buffer was 50 mM potassium phosphate, pH 7.0, with 5 mM DTT, and the sample concentration was 25 μM. The spectra were recorded in 1 cm cuvette.

The paramagnetic ¹H NMR spectrum of WT LIAS (Fig. 3a) shows two intense (a and b) and one weak (c) hyperfine shifted signals in the 20-13 ppm spectral region. Their size and linewidths are typical of βCH₂ and αCH of cysteine/serine bound to a [4Fe-4S]²⁺ cluster [20-22]. The temperature dependence of hyperfine shifted NMR signals provides information on the electronic state of the paramagnetic center [23]. In the case of [4Fe-4S]²⁺ clusters, the antiferromagnetic couplings among the four equivalent iron ions (each of them being formally Fe².5⁺) give rise to a S=0 diamagnetic ground state; the paramagnetism arises from the excited levels of the energy diagram, that become more populated as temperature increases [24, 25]. The hyperfine shifts due to the contribution of the excited states therefore increase when increasing temperature, giving rise to the so-called antiCurie temperature dependence. Conversely, in the [4Fe-4S]⁺ clusters which contain two Fe².5⁺ ions and two Fe³⁺ ions, magnetic couplings produce a paramagnetic S=1/2 ground state. Here, paramagnetism decreases at increasing temperature and the chemical shifts follow the classical Curie law. As a consequence, the temperature dependence of ¹H NMR spectrum shown in Fig. 3d allowed us to obtain information about the oxidation state of the [4Fe-4S] clusters of LIAS. Signal a experiences an anti-Curie temperature dependence and the weak signal c exhibits a Curie temperature dependence (Fig. 3d). Temperature dependence of signal b suggests that at least two protons, having respectively Curie and anti-Curie temperature dependence, are overlapped in this signal (Fig. 3d). Longitudinal relaxation times (T₁) of signals a-d have been measured in WT LIAS in D₂O. T₁ values are in the 1-3 ms range (signal a, 1.2 ± 0.2 ms; signal b, 2.1 ms ± 0.4; signal c, 2.5 ± 0.2 ms; signal d 1.6 ± 0.6 ms). They are slightly smaller than
previous findings in electron transfer Fe-S proteins [26-28] and account for electronic correlation time of iron ions around $1 \times 10^{-11}$ s.

**Figure 3.** Paramagnetic $^1$H NMR spectra of WT LIAS and of the triple LIAS variants and temperature dependence of hyperfine-shifted signals of cluster ligands. $^1$H NMR spectra acquired at 400 MHz of: (a) WT LIAS in the 30-10 ppm spectral region and in the far-shifted 90-30 ppm region (inset), in degassed 50 mM phosphate buffer pH 7.0, and 5 mM DTT; (b) chemically reconstituted C137/C141/C144A LIAS variant in degassed 50 mM phosphate buffer pH 7.0, and 5 mM DTT; (c) as purified C106/C111/C117A LIAS variant in degassed 50 mM phosphate 100% D$_2$O buffer pH 7.0, and 5 mM DTT; (d) Temperature dependence of hyperfine-shifted signals of cysteine/serine ligands in WT LIAS (black dots) and triple C137/C141/C144A (red dots) and C106/C111/C117A (green dots) LIAS variants.

We used site-directed mutagenesis to obtain a cluster specific assignment of these NMR signals. In a LIAS variant, the three cysteine residues that binds the FeS$_{RS}$ cluster (i.e. Cys 137, Cys 141 and Cys 144) were mutated into alanines; in another LIAS variant, the same type of mutations were performed on the FeS$_{aux}$ cluster Cys ligands (i.e. Cys 106, Cys 111 and Cys 117). The triple C137/C141/C144A and C106/C111/C117A LIAS variants can thus
bind only the FeS$_{aux}$ or the FeS$_{RS}$ cluster, respectively. The UV-visible spectra of both variants are similar to that of WT LIAS, possessing the same distinctive features at 330 and 400 nm characteristic of [4Fe-4S]$^{2+}$ clusters and not displaying features that are indicative of [2Fe-2S]$^{2+}$ clusters (Fig. 2b). The paramagnetic $^1$H NMR spectrum of the C137/C141/C144A LIAS variant shows two signals with chemical shift values very similar to those of the signals $a$ and $b$ observed in WT LIAS (Fig. 3b), both having an anti-Curie temperature dependence (Fig. 3d). The chemical shifts and the temperature dependence of these two signals are typical of $\beta$CH$_2$ of Cys or Ser residues bound to a [4Fe-4S]$^{2+}$ cluster with an S=0 electronic ground state. As previously reported for [4Fe-4S] and [2Fe-2S] clusters [22, 29], a Cys cluster ligand mutation to Ser affects the overall electronic distribution within the cluster but does not change dramatically the features of the $^1$H NMR spectrum. Signals of serine $\beta$CH$_2$ will experience very similar hyperfine shift as cysteine $\beta$CH$_2$ signals. The shorter metal-to-proton distance of the oxygen-bound serine, compared to the sulfur-bound cysteine, may result in a faster nuclear relaxation of serine $\beta$CH$_2$; however, metal-to proton distances are affected also by Fe-S(O)-C-H dihedral angles and therefore we cannot discriminate between Cys and Ser ligands. However, comparing the paramagnetic $^1$H NMR spectrum of the C137/C141/C144A LIAS variant, containing exclusively the auxiliary cluster, with those of wild-type [4Fe-4S]$^{2+}$ HIPIP and a stable variant [4Fe-4S]$^{2+}$ HiPIP in which a cysteine ligand had been replaced by serine ligand [22, 30, 31], it results that the chemical shift values observed for the beta protons of the auxiliary cluster are more similar to those reported for the mutated [4Fe-4S]$^{2+}$ HIPIP with respect to those of wild-type [4Fe-4S]$^{2+}$ HIPIP. This comparison suggest that Ser 345 is a ligand in the auxiliary cluster of human LIAS, in agreement with what observed in the crystal structures of bacterial LIAS homologues, in which the same serine was found to bind the auxiliary cluster [12, 13]. Moreover, signal $a$ has a $T_1$ value smaller than what previously found for $\beta$CH$_2$ of cysteines bond to [4Fe-4S]$^{2+}$ cluster [26, 27], and this could be interpreted
as a further indirect evidence of serine coordination. In conclusion, all the NMR data on C137/C141/C144A LIAS variant allowed us to assign the $^1$H signals due to the $\beta$CH$_2$ of Cys/Ser residues of the FeS$_{aux}$ cluster in WT LIAS and to determine that the FeS$_{aux}$ cluster is in an oxidized [4Fe-4S]$^{2+}$ state ([4Fe-4S]$_{aux}^{2+}$, hereafter). The paramagnetic $^1$H NMR spectrum of the C106/C111/C117A LIAS variant contains two contiguous intense signals with very close chemical shifts that are similar to that of signal $b$ observed in WT LIAS spectrum (Fig. 3c). As a result, the paramagnetic $^1$H NMR spectra of both variants indicated that signal $b$ in WT LIAS spectrum arises from three resonances and that one of these protons is due to FeS$_{aux}$ cluster and the other two to the FeS$_{RS}$ cluster. Both WT and C106/C111/C117A LIAS when dissolved in 100% D$_2$O buffer, which determines the disappearance of exchangeable signals, showed the presence of a signal ($d$) (Fig. 3c), which is hidden in the spectrum acquired in H$_2$O buffer by the sharp NH signal at 12.9 ppm (Fig. 3a). The anti-Curie temperature dependence of this signal (Fig. 3d) and its chemical shift are typical of a $\beta$CH$_2$ of a cysteine residue bound to an oxidized [4Fe-4S]$^{2+}$ cluster. A weak shoulder signal with the same chemical shift and Curie temperature dependence as those of the weak signal $c$ observed in WT LIAS is also present in the paramagnetic $^1$H NMR spectrum of the C106/C111/C117A LIAS variant (Fig. 3c). The detection of this Curie temperature dependent signal in the $^1$H NMR spectra of both WT and C106/C111/C117A LIAS suggest that a small fraction of the FeS$_{RS}$ cluster is in the reduced state. By comparing the chemical shift of this signal with known paramagnetic $^1$H NMR spectra of small electron transfer reduced [4Fe-4S]$^+$ ferredoxins [32], it appears that signal $c$ may arise from a $\alpha$CH proton of a Cys residue bound to a reduced [4Fe-4S]$^+$ cluster. According to this interpretation, $\beta$CH$_2$ protons of cysteines bound to a [4Fe-4S]$^+$ cluster should be observed through a paramagnetic tailored experiment performed over a wide spectral window using fast repetition rates [33, 34]. Three very weak signals with Curie temperature dependence have been, indeed, detected, in the WT protein, in the 90-30 ppm
region (inset of Fig. 3a). Overall, the NMR data on the C106/C111/C117A LIAS variant allowed us to identify the $^1$H signals due to the $\beta$CH$_2$ protons of the Cys residues of the FeS$_{RS}$ cluster in WT LIAS, and indicated that the FeS$_{RS}$ cluster in WT LIAS is mainly present in an oxidized [4Fe-4S]$^{2+}$ state ([4Fe-4S]$_{RS}^{2+}$ hereafter), and only a small fraction of the FeS$_{RS}$ cluster is in a reduced [4Fe-4S]$^+$ state ([4Fe-4S]$_{RS}^+$ hereafter).

The NMR data described in the last paragraph on the variants were acquired i) on a C106/C111/C117A LIAS variant that was not chemically reconstituted since this variant is characterized by a low protein stability and largely precipitates upon chemical reconstitution, and ii) on a C137/C141/C144A LIAS variant that was, on the contrary, chemically reconstituted to form a [4Fe-4S]$^{2+}$ cluster, since the paramagnetic $^1$H NMR spectrum of the as purified C137/C141/C144A LIAS variant shows signals of $\beta$CH$_2$/aCH cysteine ligands typical of a mixture of [4Fe-4S]$^{2+}$ a [3Fe-4S]$^+$ clusters [35, 36] with a 30/70 intensity ratio for the two species (Fig. S2). Overall, the NMR spectra on the two variants showed that a structural cooperativity is present between the two clusters. Indeed, the absence of the FeS$_{RS}$ cluster exposes the FeS$_{aux}$ cluster to oxidative conversion from [4Fe-4S]$^{2+}$ to [3Fe-4S]$^+$, and when FeS$_{RS}$ cluster is the only cluster present in the enzyme, the protein stability is greatly reduced.
The cluster-specific assignment of the paramagnetic $^1$H NMR signals of WT LIAS allowed us to follow the interactions of SAM and substrate molecules with the two [4Fe-4S] clusters of WT LIAS. We have first investigated the interaction of WT LIAS with SAM (Fig. S1). Stepwise additions of SAM to WT LIAS (up to two equivalents) give rise to a new set of intense peaks in the 22-13 ppm region (labelled with symbols $\alpha$-$\delta$ in Fig. 4a), at decrement of signals $b$ and $c$, which contain resonances from the FeS$_{RS}$ cluster ligands, while signal $a$ of the FeS$_{aux}$ cluster remains essentially unaffected by the SAM-LIAS interaction.

Figure 4. Paramagnetic $^1$H NMR spectra of WT LIAS in the presence of SAM and the octanoyl-peptide substrate. $^1$H NMR spectra acquired at 400 MHz of: (a) WT LIAS in the
absence (black) and in the presence of one (red) and two (blue) equivalents of SAM; (b) WT LIAS in the absence (black) and in the presence of one equivalent of octanoyl-peptide substrate (violet); (c) WT LIAS in the presence of one equivalents of SAM (red), and one equivalent of SAM and one equivalent of octanoyl-peptide substrate (green). Sample conditions are degassed 50 mM phosphate buffer pH 7.0, and 5 mM DTT.

All the new $\alpha$-$\delta$ signals have chemical shift values and anti-Curie temperature dependences consistent with a new species containing a $[4\text{Fe-4S}]^{2+}$ cluster (Fig. S3). All these data indicated the formation of a SAM-LIAS adduct in which SAM closely interacts with the $[4\text{Fe-4S}]^{2+}$ cluster, while it is not interacting with FeS$_{\text{aux}}$ cluster. This is in agreement with previous ENDOR studies showing that SAM binds to radical SAM Fe-S cluster via coordination of the amino and carboxylate of SAM to the catalytic iron ion of the $[4\text{Fe-4S}]^{2+}$ cluster [37, 38], and with the crystal structure of LipA complexed with the SAM-mimic compound SAH (Fig. S1), which showed that SAH is located close to the FeS$_{\text{RS}}$ cluster while being far from the FeS$_{\text{aux}}$ cluster [13]. Consistent with this analysis, the same spectral changes were observed when monitoring the interaction between WT LIAS and the SAM-mimic compound SAH, i.e. signals with chemical shifts similar to those of the $\alpha$-$\delta$ signals, all having anti-Curie temperature dependence, were observed (data not shown). Moreover, the stepwise disappearance of signal $c$ in Fig. 4a indicate that the small fraction of reduced $[4\text{Fe-4S}]^{+}$ cluster gets oxidized upon addition of SAM, as expected by the $[4\text{Fe-4S}]^{+}$-induced reductive cleavage of SAM. No signals of $\beta$CH$_2$ of cysteines bound to the small fraction of reduced $[4\text{Fe-4S}]^{+}$ cluster were indeed observed in the 70-30 ppm region in the final mixture, in agreement with the FeS$_{\text{RS}}$ cluster oxidation.

In the interaction between WT LIAS and SAM, two signals $z$ and $z'$ at 26.9 and 23.5 ppm, having lower intensity than $\alpha$-$\delta$ signals, appear by SAM additions. The chemical shift values and the anti-Curie temperature dependence of these two signals (Fig. S3) are consistent with
the formation of a [4Fe-4S]$^{2+}$ cluster species. The observed oxidation of the small fraction of [4Fe-4S]$_{RS}$$^+$ cluster induced by SAM addition produces the formation of methionine and of the 5'-dA• radical. This radical is a transient and highly reactive intermediate and was only very recently observed at low temperature using a novel approach involving cryogenic photoinduced electron transfer [38-40]. It is thus impossible that it would survive in these experiments long enough to produce interactions observable by NMR. Therefore, 5'-dAdo• is reasonably quenched upon production to generate 5'-deoxyadenosine (5'-dA). Bearing in mind that 5'-dA is expected to go close the FeS$_{aux}$ cluster in order to form the radical on the octanoyl chain of the substrate, we can suggest that the $z$ and $z'$ signals originate from the $\beta$CH$_2$ of the cysteines bound to the 5'-dA-interacting FeS$_{aux}$ cluster species. According to this interpretation, the low intensity of the $z$ and $z'$ signals with respect to the $\alpha$-$\delta$ signals is in agreement with the low percentage of the reduced [4Fe-4S]$_{RS}$$^+$ cluster present in the protein that can promote SAM cleavage and thus form the 5'-dA-interacting species. To validate this model, the interaction between WT LIAS and a compound mimicking the 5'-dA (MTA, Fig. S1) was investigated. Unlike what we observed following the additions of SAM, an excess of MTA (up to 2.5 equivalents) was required to observe some spectral changes in WT LIAS. The formation of new signals with chemical shifts similar to those of $\alpha$-$\delta$ signals was, however, not observed upon MTA additions (Fig. S4), thus indicating that MTA does not interact with FeS$_{RS}$ cluster. However, two new weak signals at 26.8 and 23.7 ppm, with very similar chemical shift to the $z$ and $z'$ signals observed in the SAM-WT LIAS complex, appeared (Fig. S4). These new signals have chemical shift values and anti-Curie temperature dependences consistent with a [4Fe-4S]$^{2+}$ state, as the two signals $z$ and $z'$ have (Fig. S3). Considering that MTA is proximal to one iron ion of the FeS$_{aux}$ cluster in the LipA crystal structure complexed with MTA [13], we conclude that these two signals originate from the presence of a species formed by the [4Fe-4S]$_{aux}$$^{2+}$ cluster interacting with MTA in the 1:2.5 WT LIAS-MTA mixture. This observation confirms the interpretation described above, i.e. 5'-dA and
methionine are formed by SAM cleavage by the small fraction of reduced FeS\textsubscript{RS} cluster present in WT LIAS and the formed 5’-dA interacts with the FeS\textsubscript{aux} cluster. The interaction between WT LIAS and the octanoyl-peptide substrate mimicking the amino acid sequence of the glycine cleavage system H protein was followed by acquiring paramagnetic \textsuperscript{1}H NMR spectra on a 1:1 mixture of WT LIAS and octanoyl-peptide, and on a 1:1:1 mixture of WT LIAS, octanoyl-peptide and SAM (sequential or simultaneous addition of SAM and octanoyl-peptide produce the same final \textsuperscript{1}H NMR spectrum). The paramagnetic \textsuperscript{1}H NMR spectrum of the 1:1 mixture of WT LIAS and octanoyl-peptide is fully superimposable with that of WT LIAS, indicating that the octanoyl-peptide does not interact with the protein in the absence of SAM (Fig. 4b). On the contrary, the paramagnetic \textsuperscript{1}H NMR spectrum of the 1:1:1 mixture of WT LIAS, octanoyl-peptide and SAM shows several spectral changes. Specifically, two new intense signals at 28.8 and 24.1 ppm appear, signal \textit{b} increases while signal \textit{a} decreases in intensity (Fig. 4c). The two new signals have chemical shift values and anti-Curie temperature dependences consistent with the presence of [4Fe-4S]\textsuperscript{2+} clusters in the mixture (Fig. S3). These spectral changes occur on signals of the FeS\textsubscript{aux} cluster cysteines. Indeed, signal \textit{a} is exclusively due to \(\beta\text{CH}_2\) of cysteine/serine of the FeS\textsubscript{aux} cluster, signal \textit{b} includes \(\beta\text{CH}_2\) of a cysteine/serine of the FeS\textsubscript{aux} cluster, and the signals at 28.8 ppm and 24.1 ppm have chemical shift values similar to those of \(\zeta\) and \(\zeta’\) signals, which characterize the FeS\textsubscript{aux} cluster interactions, as discussed above. Therefore, the observed spectral changes monitor the interaction between the octanoyl-peptide and the FeS\textsubscript{aux} cluster. On the contrary, the signals \(\alpha-\delta\) assigned to the SAM-bound FeS\textsubscript{RS} cluster remains essentially unperturbed (Fig. 4c), indicating that SAM remains bound to the cluster. Overall, the SAM:peptide:WT LIAS interaction studies indicate that, only once SAM is bound to the protein, the octanoyl-peptide substrate can recognize its binding site. This is in agreement with what found in the radical SAM enzyme biotin synthase, where dethiobiotin substrate binding occurs only in the presence of SAM and the substrate binding is highly cooperative [41].
In conclusion, the data here presented showed that $^1$H NMR spectroscopy tailored to reveal hyperfine-shifted signals of metal-ligands [42] is a powerful tool to investigate the sequential binding of SAM and of the octanoyl-peptide substrate to the [4Fe-4S] clusters of human lipoyl synthase. We can specifically monitor the interaction between SAM and the FeS$_{RS}$ cluster, the redox changes of the FeS$_{RS}$ cluster upon SAM interaction, follow the reductive cleavage of SAM, and monitor the substrate binding at the FeS$_{aux}$ cluster. These information can be exploited to characterize the catalytic mechanism of the radical SAM superfamily of enzymes. In particular, we believe that, while X-ray, EPR and ENDOR spectroscopies offer a more detailed but also more static view by trapping the intermediates of the catalytic mechanism and freezing transient interactions, paramagnetic NMR provides a picture at room temperature of the kinetics of substrate uptake and its transformation, as well as the dynamics of site specific interactions of SAM and the substrate occurring at both clusters, thus being essential to obtain a comprehensive view of the catalytic mechanism.
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Competing Interests.

The authors declare no competing financial interests.

Author Contributions


Appendix A. Supplementary data

Supplementary information includes Materials and Methods and Supplementary Figures S1-S4.
References


