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Designed metalloprotein stabilizes a semiquinone radical

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Abstract

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Enzymes use binding energy to stabilize their substrates in high-energy states that are otherwise inaccessible at ambient temperature. Here we show that a *de novo* designed Zn(II) metalloprotein stabilizes a chemically reactive organic radical that is otherwise unstable in aqueous media. The protein binds tightly to and stabilizes the radical semiquinone form of 3,5-di-*tert*-butylcatechol. Solution NMR spectroscopy in conjunction with molecular dynamics simulations show that the substrate binds in the active site pocket where it is stabilized by metal–ligand interactions as well as by burial of its hydrophobic groups. Spectrochemical redox titrations show that the protein stabilized the semiquinone by reducing the electrochemical midpoint potential for its formation via the one-electron oxidation of the catechol by approximately 400 mV (9 kcal mol⁻¹). Therefore, the inherent chemical properties of the radical were changed drastically by harnessing its binding energy to the metalloprotein. This model sets the basis for designed enzymes with radical cofactors to tackle challenging chemistry.

How to dial functions in a protein is the key question in protein engineering. *De novo* protein design has proved to be a powerful tool to pinpoint the structural features that determine function within proteins—a minimized protein scaffold is systematically varied and tested to engineer new functions^{1–11}. One of the key questions in protein design is to understand how a protein's environment can direct the properties of the bound cofactor or substrate. In previous studies, optimization of the electrostatic and hydrophobic interactions guided the tuning of cofactors like haem and flavins in affinity, reduction potentials and O2 binding^{12–14}. A series of designed three-helix coiled coils with mononuclear Zn(n) to mimic carbonic anhydrase were shown to activate water for ester hydrolysis and CO₂ hydration^{15–17}. An artificial metallo- β -lactamase was designed to self-assemble into a

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Competing financial interests

The authors declare no competing financial interests.

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Author contributions

G.U. and W.F.D. conceived and designed the research. G.U. and W.F.D. wrote the manuscript. G.U. prepared the samples and reagents and performed the research. G.U. and G.T.G. designed and conducted the spectrochemical redox titration experiments. T.L. designed and performed the QM/MM molecular modelling. Y.W. recorded and analysed NMR data. T.L. and Y.W.

tetramer and use catalytic Zn(n) sites to hydrolyse β -lactams. The enzyme was functional in *E. coli*, where it hydrolysed the β -lactam antibiotic, ampicillin, which resulted in cell survival¹⁸. Our group studied designed Due Ferri (DF) proteins¹⁹ as models for natural non-haem di-iron proteins, and showed that subtle modifications to the first- and second-coordinating spheres of the site were able to direct the O₂-dependent chemistry²⁰. Recently, protein design was used to trap the unstable coplanar conformation of a *p*-biphenylalanine residue²¹. Here we extend the use of rational protein design to stabilize a chemically unstable species—an organic radical.

Enzymes efficiently use radicals, an unstable and generally highly reactive species, in challenging chemical transformations. These radicals are not destructive like the reactive oxygen species because they are 'controlled'; that is, (1) generated during catalytic turnover and quenched at the end, and (2) contained in the enzymatic environment to prevent any undesired off-pathway reactivity. However, the mechanism by which enzymes stabilize these otherwise kinetically and thermodynamically unstable species remains poorly understood. Protein engineering has been used to test principles of radical stabilization through the de novo design of 'maquette' proteins with covalently attached radical-forming amino acids Trp[•] and Tyr[•] or mercaptophenol derivatives^{22–25}. Although it has been possible to stabilize Tyr[•] kinetically with half-lives up to six seconds²⁶, they were thermodynamically destabilized by approximately 100 mV relative to the corresponding small molecule in aqueous solution 24,27 . Here we focus on the thermodynamic and kinetic stabilization of ortho-semiquinones, which are radical intermediates in essential redox processes and catalysis in nature²⁸⁻³¹. The 3,5-di-*tert*-butyl-semiquinone radical anion (SQ[•]), a smallmolecule analogue of native o-semiquinone radicals, is the one-electron oxidized intermediate in the redox triad 3,5-di-tert-butylcatechol/semiquinone/o-benzoquinone (QH₂/SQ[•]/Q) (Fig. 1a). Although SQ[•] is naturally less stable than QH₂ and Q, its thermodynamic potential can be shifted. In aprotic organic solvents, coordination to cations^{32,33} and transition-metal ions^{34–36} stabilizes the SQ[•]. In organic solvents the presence of Zn(n) ([$Zn(H_2O)_6$]²⁺) shifts the one-electron reduction potential of Q, $E_{O/SO^{\bullet}}$, by ~220 mV compared with the free SQ[•] in solution, which makes the Zn(n)-SQ[•] complex more stable by 5 kcal mol⁻¹ (ref. 34). In aqueous buffers only a two-electron reduction of Q to QH₂ is observed, and SQ[•] is undetectable under steady-state conditions³⁷.

We previously used QH_2 as a substrate for a di-Fe(m) bound DF protein variant, DF3³⁸. In the catalytic cycle, QH_2 is first oxidized to Q by di-Fe(m). The dimetal centre is then reoxidized by ambient O₂ to di-Fe(m) to complete the catalytic turnover. As the di-Fe(m) centre catalyses two-electron chemistry, the substrate was converted from QH_2 into Q, and no SQ[•] intermediate was observed. We rationalized that a redox-inert transition metal should be used to observe and stabilize SQ[•] instead. Here we used the single-stranded form of DF-type proteins, DFsc, in which we exchanged the Fe(m) with Zn(n). The DFsc variant used in this work, 2A3H-DFsc (referred to as DFsc) binds two Zn(n) to form [DFsc-Zn(n)₂], a wellstructured four-helix bundle (PDB 2LFD)²⁰.

We hypothesized that inherent chemical properties of SQ[•] could be used to distinguish it from QH₂ and Q, and guide its binding to [DFsc-Zn(π)₂]; the SQ[•] radical is the predominant form of the compound at neutral pH (p $K_a = 6$)³⁹, and is an excellent chelator of Zn(π)

(Supplementary Table 1). In contrast, Q is a poor chelator, and is observed to dissociate away from Zn(π) when Zn(π)–Q is electrochemically generated from Zn(π)–SQ[•] (ref. 35). QH₂ mono- and dianions are minor species in solution at neutral pH (p $K_a = 10$)³⁹, which means that binding to Zn(π) is also unlikely to result from the thermodynamic cost of deprotonation. Moreover, SQ[•] is stabilized in apolar solvents relative to water, and therefore binding to the hydrophobic cleft of DFsc should stabilize the SQ[•] by removing the radical from bulk water.

In this study, we show that $[DFsc-Zn(n)_2]$ strongly stabilizes SQ[•] over the otherwise more stable QH₂ and Q forms. Optical and magnetic spectroscopy, along with spectrochemical redox titrations, demonstrate that the binding of SQ[•] to DFsc effectively pulls the equilibrium towards SQ[•] given a mixture of 1:1 Q:QH₂. By examining derivatives of QH₂ we show that tight binding and radical stabilization requires the presence of the hydrophobic *t*-butyl groups, which are partially buried on interaction with the protein. Molecular dynamics (MD) simulations were used to gain further insight into the structural characteristics of the [DFsc-Zn(n)₂]–SQ[•] complex. Taken together, these results established how binding energy can be harnessed to stabilize an otherwise inaccessible radical.

Results and discussion

Optical and magnetic spectroscopy methods were used to confirm SQ[•] binding to DFsc and define the new characteristics of the radical in the protein environment. When [DFsc-Zn(n)₂] was added to an equimolar mixture of Q and QH₂, the equilibrium shifted to favour the corresponding SQ[•] (Fig. 1b), as evidenced by a large decrease in the absorption band of Q ($\lambda_{max} = 415 \text{ nm}$) and the appearance of a new broad band that spanned 740–850 nm (Fig. 2a), typical of a Zn(n)-bound SQ[•] radical³⁵. An isosbestic point at 492 nm suggests the lack of additional intermediates. A strong band at the position of the SQ[•] was also observed in the circular dichroism spectrum (Supplementary Fig. 2), which suggests that the achiral SQ[•] is bound to DFsc in a unique asymmetric conformation. Taken together, optical spectroscopy indicates that the *in situ* generated SQ[•] is in complex with the metal-bound DFsc, and yields the [DFsc-Zn(n)₂]–SQ[•] moiety. The SQ[•] is most probably bound to the Zn(n), as the absorption features closely match that of small molecule Zn(n)–SQ[•] complexes (Supplementary Table 1).

Factors that direct SQ[•] binding were evaluated by using derivatives of QH₂ or Q, which were examined for SQ[•] formation. *o*-Quinones/catechols with electron-releasing or electron-withdrawing substituents, such as -OMe, $-NO_2$ and -H, do not form SQ[•] in the presence of [DFsc-Zn(π)₂] (Supplementary Fig. 3). Therefore, it is unlikely that the *t*-butyl groups play an electronic role in binding. The only variant to form SQ[•] was 4-*t*-butylcatechol, with a lower yield than its di-*t*-butyl analogue, which highlights the importance of the hydrophobic *t*-butyl groups in the tight binding and, thus, stability of SQ[•] in the context of the protein.

Room-temperature electron paramagnetic resonance (EPR) spectra confirmed the formation of the paramagnetic SQ[•] radical in complex with [DFsc-Zn(π)₂]. The spectrum of [DFsc-Zn(π)₂]–SQ[•] is consistent with the presence of an organic radical (g = 2.003). The signal is broadened (peak-to-peak line-width, 8 Gauss) and lacks hyperfine features, which suggests

the immobilization of the SQ[•] radical. Overlaid spectra of SQ[•] generated in the presence of $[DFsc-Zn(\pi)_2]$, apo DFsc or Zn(π) are shown in Fig. 2b. Spin quantification shows a yield of $72 \pm 7\%$ radical formation in the presence of $[DFsc-Zn(\pi)_2]$ (with respect to the protein). The apo DFsc control does not show any trace of a radical. The Zn(π)-only control showed a low yield of radical formation (2% yield). Although it is expected that Zn(π) would partially stabilize SQ[•] (refs 34,40), the radical experiences a markedly different environment in the protein, as evidenced by the line-shape differences of $[DFsc-Zn(\pi)_2]$ –SQ[•], Zn(π)-only and SQ[•]-only spectra (Fig. 2b and Supplementary Fig. 6).

The midpoint reduction potential of [DFsc-Zn(π)₂]–SQ[•] was determined by a series of redox titrations with dithionite in the presence of a redox indicator dye (potassium indigo tetrasulfonate (ITS)) under anaerobic conditions at neutral pH (Supplementary Fig. 4). Relative populations of [DFsc-Zn(π)₂]–SQ[•] ($\lambda_{max} = 740$ nm) and oxidized ITS ($\lambda_{max} = 594$ nm, $E_{mid,7} = -46$ mV)⁴¹ under steady-state conditions were determined, and fit in the Nernst equation to yield the midpoint potential $E_{mid,7} = -21$ mV versus NHE for the reduction of [DFsc-Zn(π)₂]–SQ[•] at neutral pH, a value that is ~400 mV less than the reported reduction potential of free SQ[•] in aqueous solution³⁹. This finding indicates that [DFsc-Zn(π)₂]–SQ[•] is strongly stabilized in the protein environment (by 9 kcal mol⁻¹).

Solution NMR spectroscopy confirmed the structure and paramagnetic nature of the [DFsc- $Zn(u)_2$]–SQ[•] complex. The bound organic radical caused considerable paramagnetic relaxation enhancement (PRE) of the backbone amides, observed as a significant decrease in the corresponding peak intensity⁴². To determine which backbone resonances are affected by the SQ[•] radical anion, ¹⁵N heteronuclear single quantum coherence (HSQC) spectra (Fig. 2c) were recorded before and after the addition of the substrate to reach a 74% conversion to [DFsc-Zn(u)_2]–SQ[•]. As PRE that resulted from a radical with an isotropic *g* factor is not expected to yield pseudocontact shifts or residual dipolar coupling, reassignment of the residues in [DFsc-Zn(u)_2]–SQ[•] was not necessary^{42,43}. Residues that experienced a significant decrease in peak intensity were then mapped onto the solution NMR structure of [DFsc-Zn(u)_2] (Fig. 3). The highest peak-intensity decrease that was detected was found in residues proximal to the active site, consistent with the SQ[•] binding at the active site.

To interpret our results further, we used MD to gain insights into the structural properties of the [DFsc-Zn(π)₂]–SQ[•] complex. First, a metadynamics simulation was used to sample possible conformations of the SQ[•] when interacting with the di-Zn(π) site. The two collective variables used defined the distance and angle between the centre of mass of the SQ[•] oxygen atoms and the Zn(π) ions. An ensemble of 20 different interacting conformers (3.0 Å distance cutoff) were then used to seed individual 50 ns MD simulations, summing up to a total of 1 μ s. The SQ[•] binding was characterized by an enlargement of the helix 1 and 2 interface, which allowed the SQ[•] to interact directly with the Zn(π) cations (Fig. 4 and Supplementary Fig. 5). A total of 10,000 snapshots were then clustered to yield three different conformations (root mean square deviation of the centroids was less than 1.5 Å over the Zn(π)–SQ[•] site). Each centroid structure was further optimized using a Gaussian09 ONIOM QM/MM⁴⁴ (quantum mechanics/molecular mechanics) method and all converged to a single geometry, in which the semiquinone was bound to the only coordinatively unsaturated Zn(π) as a bidentate ligand.

The resulting structure is consistent with a SQ[•]-bound conformation. A notable feature is the sequestration of the *t*-butyl groups in a very hydrophobic pocket adjacent to the di- $Zn(\pi)$ site surrounded by the apolar side chains of A10, G14, I17, A43, G47, V50 and Y51 (Fig. 4c). Favourable interactions with the protein cleft would contribute to the binding of SQ[•], an observation also validated by the lack of SQ[•] formation in derivatives missing the *t*-butyl groups. A major difference of the SQ[•]-bound DFsc model is the change in the coordination number of the active site Zn(I). The di-Zn(I) are five- and six-coordinate in the starting structure, $[DFsc-Zn(\pi)_2]$ (ref. 20), and both become six-coordinate in the SQ[•]-bound state. The only His ligand (H107) to the pentavalent $Zn(\pi)$ in [DFsc- $Zn(\pi)_2$] dissociates to create a tetravalent $Zn(\pi)$, which then acquires SQ[•] to become six-coordinate. This finding is consistent with the expectation that SQ[•] binding would require a coordinatively unsaturated metal site, and provides an indication that a His rotation could be responsible in forming the tetravalent $Zn(\pi)$ intermediate. It is interesting that in the structure of an alternative oxidase with a related di-iron active site, proximal His ligands show variable coordination to iron depending on substrate and inhibitor binding (3VVA, PDB http://www.pdb.org/pdb/search/ structidSearch.do?structureId=3VVA)⁴⁵.

Based on the computational model, we hypothesized that mutating H107 could leave the $Zn(\pi)$ four-coordinate in the [DFsc-Zn(π)₂] state and so be more available for the bidentate SQ[•] binding. Variants of DFsc in which H107 was mutated to the non-coordinating residues Ala and Asn were made and characterized. Spectral titrations indicated that H107A-DFsc and H107N-DFsc variants bind approximately 2.3 equiv. Zn(π) per protein (versus the theoretical value of 2 equiv. (Supplementary Fig. 10)). Interestingly, both variants form SQ[•] complexes more rapidly than the wild-type DFsc (Supplementary Fig. 8), which confirms the hypothesis that Zn(π) loses a ligand to become four coordinate on SQ[•] binding. The variants also form a 4-*t*-butylcatechol-derived SQ[•] complex, reminiscent of the wild-type DFsc (Supplementary Fig. 9). These findings are consistent with our prediction that H107 is not necessary for protein–SQ[•] complex formation.

In conclusion, we used a designed metalloprotein to stabilize successfully the SQ[•] radical anion using binding energy to pull the equilibrium towards the otherwise unstable SQ[•] state at room temperature in aqueous medium. Through optical and magnetic spectroscopic characterization, we demonstrate that SQ[•] is bound to the metalloprotein, most probably anchored to Zn(π) as a bidentate ligand, and is encased within the protein when bulk solvent is removed. Structural analogues of SQ[•] provide further insight that one of the crucial driving forces is favourable hydrophobic interactions with the cleft of the protein. Computational modelling of the bound structure allowed us to rationalize our results further, which are consistent with the requirement for hydrophobic interactions initiated by the *t*-butyl groups and Zn(π)-binding in enhancing the binding energy to drive the equilibrium towards the otherwise unstable SQ[•]. We assessed that the protein–SQ[•] complex is stabilized by 10⁶–10⁷ (9 kcal mol⁻¹) in comparison with the free radical SQ[•] in neutral solution. Thus, these studies lead to a deeper understanding of how proteins stabilize radical species.

Methods

Ultraviolet/visible (UV/vis) spectroscopy

DFsc was dissolved in 50 mM MOPS (pH 7.0) and 100 mM NaCl to $50-100 \mu$ M concentration. The protein was reconstituted with Zn(II) by adding ZnSO₄ stock in milliQ water to the protein in a 2:1 Zn(II):protein ratio and incubated at room temperature for one hour. After incubation, the [DFsc-Zn(II)₂] solution was transferred to a quartz cuvette (1 cm pathlength, Starna Cells) and the initial spectra recorded with a Cary 300 Bio UV/vis spectrophotometer. Substrates, premixed in a separate vessel (Q:QH₂ in a 1:1 ratio, dissolved in dimethylformamide (DMF)), were added to the protein solution, and then quickly mixed by pipetting. Derivatives were screened under identical conditions, with 4-*t*-butylcatechol, 4-nitrocatechol, 3-methoxycatechol and catechol added to the protein solution in ten molar equivalents, and incubated overnight to air oxidize.

EPR spectroscopy

Measurements were performed with a Bruker EMX EPR spectrometer (Bruker Instruments). All the measurements were performed at room temperature (298 K). The spectra were recorded at a frequency of 9.83 GHz, with a microwave power of 25 mW, modulation amplitude of 3 Gauss and modulation frequency of 100 kHz. Each spectrum was collected as an average of three scans, sweeping 100 Gauss. The protein sample contained 50 μ M [DFsc-Zn(π)₂] in 50 mM MOPS (pH 8.0), 100 mM NaCl. To the protein solution, Q was added to yield 1 mM (stock solution in DMF), followed by the addition of NADH (stock solution in milliQ) to yield 500 μ M. A 25 μ l aliquot of this mixture was transferred to a 25 μ l micropipette (Drummond, Wiretrol) and sealed with clay at one end.

NMR spectroscopy

The DFsc was labelled with ¹⁵N by bacterial expression in M9 minimal medium supplemented with ¹⁵NH₄Cl (Sigma-Aldrich). Two-dimensional (2D) ¹⁵N-HSOC spectra were collected at 25 °C on a Bruker Avance-I 800 MHz spectrometer with a 5 mm x, y,zshielded pulse-field gradient triple resonance probe. The protein (640 μ M) was reconstituted with 2.5 molar equivalents of ZnSO₄ in 100 mM MOPS (pH 7.51), 100 mM NaCl; 5% (v/v) D₂O was added before data collection. To the protein solution, 213 μ M of *N*-acetyl-¹⁵N,¹³Cglycine (synthesized following the literature protocol⁴⁶) was added as the internal concentration standard. After the $[DFsc-Zn(n)_2]$ spectrum was recorded, 5 equiv. of the Q (400 mM stock prepared in 1:1 DMF:MeOH mixture) and 2.5 equiv. of NADH (500 mM stock prepared in milliQ) were added to the sample. UV/vis spectroscopy of the sample indicated that this resulted in a 74% conversion to the [DFsc-Zn(π)₂]-SQ[•] complex. 2D ¹⁵N-HSQC spectra were recorded along $t_1(^{15}N)$ and $t_2(^{1}H)$ with, respectively, 128 and 2,048 complex points, and $t_{1,\max}({}^{15}N) = 30$ ms and $t_{2,\max}({}^{1}H) = 91$ ms. The ¹H chemical shift was referenced to the water line at 4.74 ppm and ¹⁵N chemical shifts were referenced indirectly via gyromagnetic ratios. The ¹⁵N carrier frequency was set at 119 ppm. NMR data were processed, analysed and visualized using NMRPipe⁴⁷ and Sparky⁴⁸. Prior to Fourier transformation, time-domain data were performed by linear prediction once, multiplied by the esine square bell window functions shifted by 75° and zero-filled once.

Equilibrium spectrochemical redox titrations

DFsc (100 μ M) was reconstituted with 200 μ M Zn(π) in 50 mM MOPS (pH 7.0) and 100 mM NaCl, and the SQ[•] complex was formed by adding a premixed mixture of 1:1 Q:QH₂ to yield 200 μ M each in the final concentration. After overnight incubation with the Q:QH₂ mixture (at ambient temperature and atmosphere), the solution was spun down and sterile filtered. To this solution, a redox indicator dye (dye, ITS, with midpoint reduction potential at neutral pH, $E_{m7} = -46 \text{ mV}$ versus NHE)⁴¹ was added in a 2 μ M concentration. The sample was then transferred to an anaerobic quartz chamber and degassed by sequential vacuum pumping and purging with $N_2(g)$. The quartz chamber was fitted with a 250 µl Hamilton Gastight syringe containing degassed 1.6 mM sodium dithionite solution. After an initial spectrum collection, a 5 µl aliquot of the dithionite solution was introduced into the 700 µl protein solution (equivalent to 10 µM dithionite per injection), mixed and left to equilibrate for 60 minutes, at which point a spectrum of the final equilibrated state was collected. Dithionite was added in increments of 5 µl until the SQ[•] signal no longer decreased. Absorbances at 596 and 740 nm were used to follow the oxidized dye and SQ[•] concentrations, respectively. Concentrations of the oxidized dye and SQ[•] were set to 100% in the absence of dithionite (black spectrum in Supplementary Fig. 4). Populations of oxidized dye and SQ[•] were thus quantified as a ratio for each spectrum collected after equilibrium was reached following each dithionite aliquot addition. Under equilibrium conditions, a common solution potential is reached⁴⁹, and we can assume that the E_{ref} equals E_{SO} , as shown in equation (1). The ratio of oxidized-to-reduced dye measured from each spectrum and the known E_{ref} for the dye were used to calculate the E_m for the semiquinone in the SQ[•]/QH₂ couple (where QH₂ is 3,5-di-*t*-butylcatechol (DTBC)).

$$E_{\rm m}^{\rm ref} - \frac{\rm RT}{nF} \ln \frac{\rm [Dye_{\rm red}]}{\rm [Dye_{\rm ox}]} = E_{\rm m}^{\rm SQ} - \frac{\rm RT}{nF} \ln \frac{\rm [DTBC]}{\rm [SQ]} \quad (1)$$

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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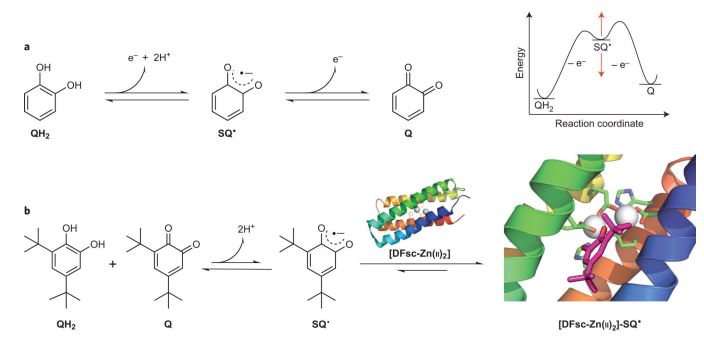


Figure 1. Semiquinone is unstable as a free radical, but its stability can be tailored in a protein environment

a, Scheme of a canonical equilibrium between QH_2 , the anion radical SQ^{\bullet} and Q. The SQ^{\bullet} anion radical is an intermediate between the fully reduced QH_2 and the fully oxidized Q. The corresponding energy versus reaction coordinate plot is representative of the expected differences in thermodynamic potential for each discrete oxidation state. The red arrows indicate that the environment of SQ^{\bullet} can change the SQ^{\bullet} stability by shifting its thermodynamic potential. **b**, General scheme for reconstituting DFsc with a radical. SQ^{\bullet} is generated *in situ* via comproportionation of QH_2 and Q and consumed by disproportionation as a free radical. In the presence of $Zn(\pi)$ -bound DFsc ([DFsc- $Zn(\pi)_2$]), the equilibrium is shifted towards the otherwise unstable SQ^{\bullet} , which complexes with the protein to form [DFsc- $Zn(\pi)_2$]– SQ^{\bullet} .

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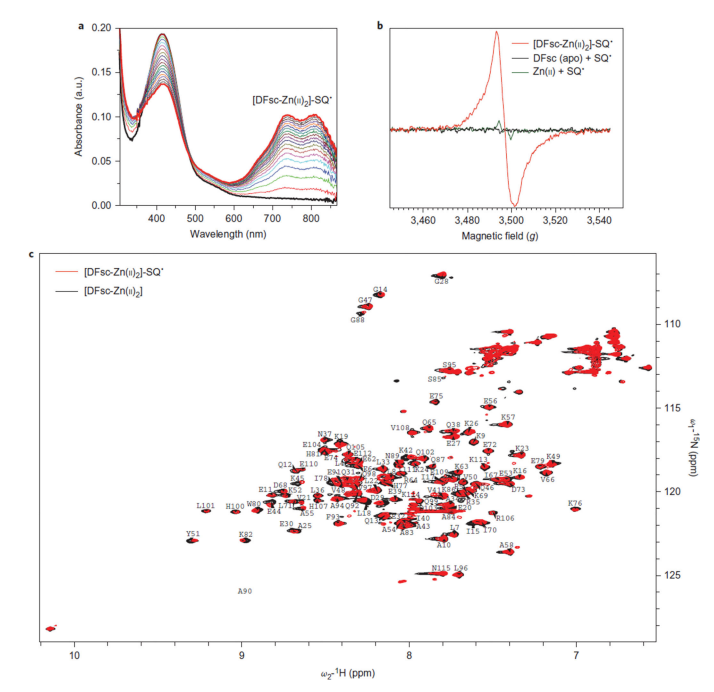


Figure 2. Observation of SQ[•] in complex with the *de novo* metalloprotein $[DFsc-Zn(\pi)_2]$ by optical and magnetic spectroscopy

a, UV/vis absorption spectra were recorded starting from the time of QH_2 and Q mixing to generate SQ^{\bullet} *in situ* (black line), and at ten minute intervals up to 180 minutes (red line) (a.u., arbitrary units). **b**, EPR spectra of [DFsc-Zn(π)₂], apo DFsc and Zn(π) in the presence of SQ[•] generated *in situ*. **c**, 2D HSQC spectra of [DFsc-Zn(π)₂] (black) and after 74% of all the protein was converted into [DFsc-Zn(π)₂]–SQ[•] (red). Backbone assignments are labelled and are in good agreement with previous studies²⁰. All amide resonances are visible at this contour level except for A90.

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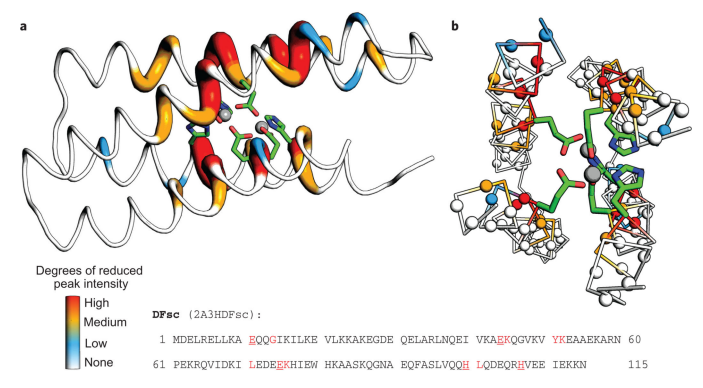


Figure 3. Analysis of results extracted from the $[DFsc-Zn(n)_2]-SQ^{\bullet}$ HSQC spectra colourmapped on the $[DFsc-Zn(n)_2]$ structure (PDB 2LFD), with the relative degrees of peak intensities compared

a,b, The highest degree of reduced peak intensity corresponds to the highest PRE. Colourmapped structures are from the side (**a**), and top-down views, with the backbone amide nitrogen atoms displayed as spheres and the loop regions omitted for clarity (**b**). For both panels, $Zn(\pi)$ is shown as grey spheres and $Zn(\pi)$ -coordinating residues as sticks. Residues that experience the highest PRE are coloured red on the structural models and below in the corresponding sequence. Residues that coordinate to $Zn(\pi)$ are underlined in the sequence. All molecular graphics were generated with PyMOL⁵⁰.

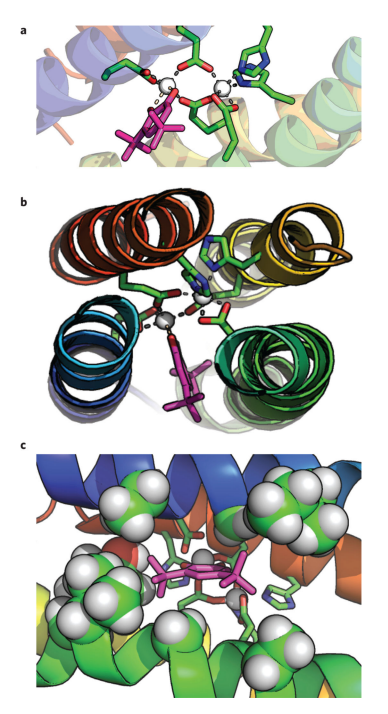


Figure 4. QM/MM-optimized model of [DFsc-Zn(n)2]–SQ[•]

a, View of the DFsc active site showing the first coordination shell of the di-Zn(π) (white spheres), with the SQ[•] coloured magenta. Dashed lines indicate bonds. **b**, Top-down view of the [DFsc-Zn(π)₂]–SQ[•] complex showing the enlargement of helix 1 (blue) and helix 2 (green) to accommodate SQ[•] binding. **c**, View of the hydrophobic residues that line the helix 1 and 2 interface (shown as spheres: A10, G14, I17, A43, G47, V50, Y51), which interact with the *t*-butyl groups of SQ[•] (in magenta).