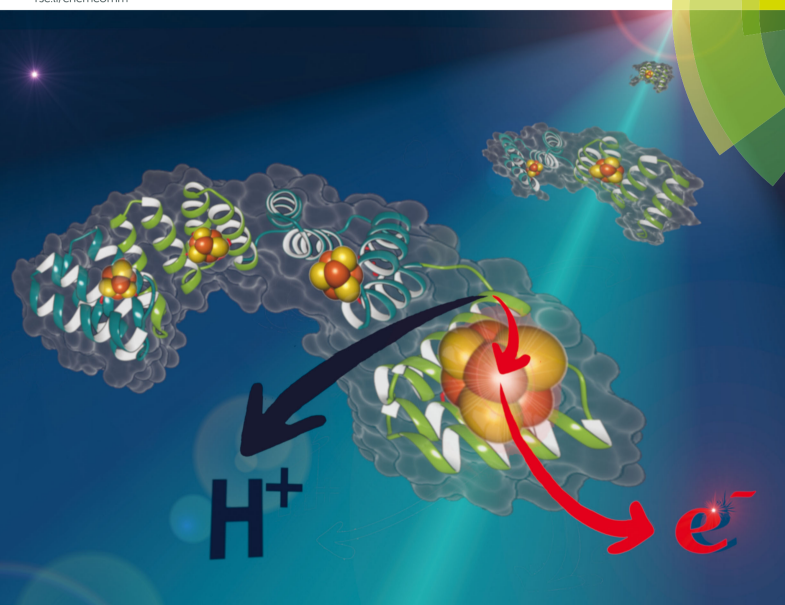
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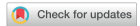
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Repeat proteins as versatile scaffolds for arrays of redox-active FeS clusters†

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Arrays of one, two and four electron-transfer active [4Fe-4S] clusters were constructed on modular tetratricopeptide repeat protein scaffolds, with the number of clusters determined solely by the size of the scaffold. The constructs show reversible redox activity and transient charge stabilization necessary to facilitate charge transfer.

Molecular materials for solar energy harvesting, molecular electronics and nanoscale devices require systems capable of facilitating long-range electron transfer (ET) with high quantum yield.^{1,2} As biological ET becomes better understood, natureinspired ET systems are emerging as alternatives to purely synthetic designs. In nature, medium and long distance electron transfer is often performed by arrays of cofactors held at specific distances within complex proteins.³⁻⁵ Fe-S clusters are some of the most common cofactors utilized for this purpose. Cubanelike [4Fe-4S] clusters are the most abundant type of Fe-S clusters found in protein structures, involved in many natural ET processes (e.g. photosynthesis, respiration).^{3,5–8} A common coordination motif utilizes four conserved cysteines coordinated to the cluster and ensconced in the hydrophobic core of the protein.9-15 Most synthetic models aimed towards engineering of functional redox enzymes have been designed to bind a single electronically isolated cluster in the protein scaffold or in the proximity of a catalytically active cofactor. 9-11 Recently, first examples of two [4Fe-4S] clusters incorporated at design-determined distances in the hydrophobic core of a three-helix bundle have been reported. 12,13 However, designing multi-center Fe-S proteins remains a challenge.

Modular assembly of multi-center ET-active systems based on repeat proteins was identified as a viable strategy. The structural simplicity and intrinsic modularity of the repeat proteins allow to use them as simple units for bottom-up fabrication, where each repeat unit can be used as a building block with individually engineered properties and functionalities. 16,17 Consensus tetratricopeptide repeat (CTPR) protein, composed of helix-turn-helix motifs that connect in a sequence to form a right-handed superhelical structure, 18 possesses the robustness and stability needed to support supramolecular assemblies^{19–21} or act as support structures in solution and solid state. 22-24 In the current work, a four-sulfur coordination site was 'grafted' onto CTPR protein scaffold, forming arrays of functional ET-active [Fe-S] clusters with precisely defined distance and orientation. The system establishes a foundation for the modular design of long-range ET conduits. To the best of our knowledge, this is the first example of a single scaffold used to assemble a controlled number of clusters.

Based on the crystal structure of tryptophanyl-tRNA synthethase (PDB ID: 2G36) and on a previous design of a [4Fe-4S] clustercoordinating peptide, 12,13 a four-cysteine [4Fe-4S] coordination site was modeled at the interface of two adjacent CTPR repeats (PDB: 2HYZ), on the concave surface of CTPR superhelix. Of the several possible positions for the cysteines, the design that supported the correct distances and rotamers for the four coordinating cysteines, Y5C and N9C in the first repeat and E2C and N6C in the second repeat, was selected. A [4Fe-4S] cluster was then manually docked into the designed binding pocket using PyMOL software and the resulting model was subjected to three rounds of energy minimization using GROMACS, with iterations of manual modeling. The side chain conformations and backbone geometry of the designed cysteines were compatible with the TPR helical fold and with the cluster coordination distances observed in natural coordination sites. The two-repeat cluster-forming unit was sequentially aligned with consecutive sections of an eight-repeat model to provide models of two- and four-cluster assemblies. The optimized models show no global structural deviations from the crystal structure of CTPR protein. Based on the models, the clusters in the multi-cluster constructs are about 16 Å apart (Fig. 1).

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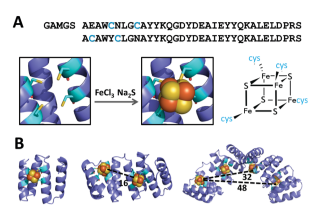


Fig. 1 Design of CTPR proteins for the incorporation of [4Fe-4S] clusters. (A) Top: Sequence of modified CTPR2 cluster binding unit; residues mutated to cysteines are highlighted in blue: Y5C and N9C in the first repeat and E2C and N6C in the second repeat. Bottom: Structural model of the designed 4-Cys site (left) and the [4Fe-4S] cluster (right). (B) Models of CTPR2-[4Fe-4S], CTPR4-2[4Fe-4S], and CTPR8-4[4Fe-4S], containing 1, 2 and 4 [4Fe-4S] clusters, respectively. Distances listed in angstroms.

Cubane-type [4Fe–4S] clusters within the proteins were formed using an established *in situ* synthetic procedure from iron(III) chloride and sodium sulfide under anaerobic conditions.²⁵ The UV-vis spectra of the resulting CTPR2-[4Fe–4S], CTPR4-2[4Fe–4S] and CTPR8-4[4Fe–4S] constructs (Fig. S1, ESI†) show broad absorption peaks at 345, 430 and 630 nm, characteristic of charge transfer excitations from sulfur to iron in [4Fe–4S]²⁺ clusters,²⁶ but redshifted compared with inorganic [4Fe–4S] clusters due to the hydrophobic environment of the protein.^{10,26,27} The absorption at 430 nm disappeared upon reduction of the clusters to [4Fe–4S] state with sodium dithionite, as expected for cuboidal [4Fe–4S] clusters.^{28–30}

Iron content (measured using the ferrozine method³¹) in the protein complexes (determined using Bradford assay32,33) was in agreement with a single [4Fe-4S] cluster per binding cavity (Table 1): one cluster in CTPR2 scaffold, two clusters in CTPR4 and four clusters in CTPR8. Circular dichroism (CD) analysis showed the CTPR proteins retained α-helical structure in all three constructs, indicating that the incorporation of the [4Fe-4S] clusters did not affect the structure of the protein scaffolds (Fig. S2, ESI†). To the contrary, thermal denaturation analysis showed that the clusters increased global stability of the proteins, with the effect most pronounced for the 2-repeat protein, which cannot benefit from the stabilizing effect of the extended superhelical structure due to its size (Fig. S3, ESI†). The $T_{\rm m}$ values for CTPR2, CTPR4 and CTPR8 were 55, 68, and 75 °C, respectively, while the respective $T_{
m m}$ values for CTPR2-[4Fe-4S], CTPR4-2[4Fe-4S] and CTPR8-4[4Fe-4S] were 65, 73

Table 1 Quantification of iron content in designed proteins using ferrozine method

Construct	Fe atoms per protein chain
CTPR2-[4Fe-4S] CTPR4-2[4Fe-4S] CTPR8-4[4Fe-4S]	$egin{array}{l} 4.60 \pm 0.38 \ 7.62 \pm 0.22 \ 16.44 \pm 1.33 \end{array}$

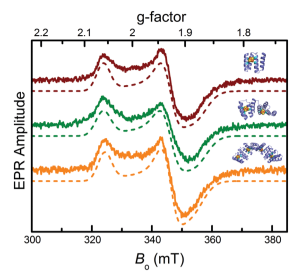


Fig. 2 EPR characterization of the CTPR-bound [4Fe-4S] clusters. From top to bottom: EPR spectra of sodium dithionite-reduced CTPR2-[4Fe-4S] (red), CTPR4-2[4Fe-4S] (green) and CTPR8-4[4Fe-4S] (orange) constructs (solid lines) and numerically simulated EPR spectra (dotted lines). Simulation parameters provided in ESI. \dagger

and 75 $^{\circ}$ C. This stabilizing effect, arising from a rigid [4Fe–4S] cluster holding protein helical fragments in place, is in line with previous results. ^{12,13}

Electron paramagnetic resonance (EPR) spectroscopy confirmed the iron–sulfur clusters initially formed in an EPR-silent [4Fe–4S]²⁺ resting state. The lack of signal further confirmed cluster identity, ruling out other possible clusters (*e.g.* [3Fe–4S]) that are EPR active in the resting state. Reduction with sodium dithionite produced the EPR active [4Fe–4S]¹⁺ state: all three constructs displayed nearly identical spectra with principal *g*-values of 1.88, 1.93, and 2.06 (Fig. 2), typical of [4Fe–4S]¹⁺ clusters.³⁴ Features indicative of exchange and/or magnetic dipole interactions between the clusters, observed with natural ferredoxins containing two [4Fe–4S]¹⁺ clusters at 10–15 Å from each other,³⁵ were not detected for CTPR4-2[4Fe–4S] and CTPR8-4[4Fe–4S] constructs, suggesting lack of sufficiently strong interactions between the clusters.

Square wave voltammetry was used to assess the redox potential of the clusters. Electrochemically quasi-reversible signals were observed for all three constructs, with redox potentials of ca. -0.21 V, -0.24 V and -0.24 V vs. SCE for CTPR2-[4Fe-4S], CTPR4-2[4Fe-4S], and CTPR8-4[4Fe-4S], respectively (Fig. 3 and Table S1, ESI†), and peak to peak separation of about 250 mV, indicative of moderate electron transfer rate between the electrode and proteins.36 These values fall within the range expected for the relatively solvent exposed low-potential [4Fe-4S] clusters. 26,27,37 The similar behavior of the three constructs suggests the clusters react mostly independently, with little interaction between neighboring clusters within the larger assemblies. At constant protein concentration, the current was proportional to the number of clusters in the constructs, as expected (Fig. S4, ESI†). However, the larger constructs showed increasingly more negative redox potentials, despite identical environment around the clusters. This trend could be attributed to the difficulty of successive cluster reduction after a

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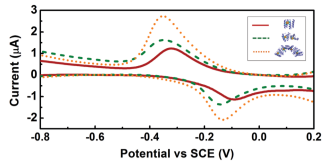


Fig. 3 Square wave voltammetry of the CTPR-[4Fe-4S] complexes. Voltammograms of CTPR2-[4Fe-4S], CTPR4-2[4Fe-4S] and CTPR8-4[4Fe-4S] are shown in red, green and orange, respectively.

first cluster in a construct has been reduced, thus suggesting at least some degree of interaction between the clusters. Structural factors were then considered as a reason for the lack of charge transfer between the clusters. Transient absorption (TA) spectroscopy was thus employed to investigate charge transfer pathways of the clusters.

Laser pulses of ~ 100 fs at 400 nm were used to promote ligand to metal charge transfer (LMCT) transitions from sulfur to iron atoms. ³⁸ Global analysis of the transient absorption data gave three evolution-associated-difference spectra (EADS). ³⁹ EADS from all three proteins have similar features and the TA was satisfactorily fitted with almost identical lifetimes (Fig. 4A and Fig. S5, ESI†), indicating at least 3 transient states (species) involved in the photo induced charge transfer process, but no direct electron transfer between the [4Fe–4S] clusters in the protein. However, the EADS showed that the 400 nm laser pump induces an internal reduction of the [4Fe–4S] cluster, and the

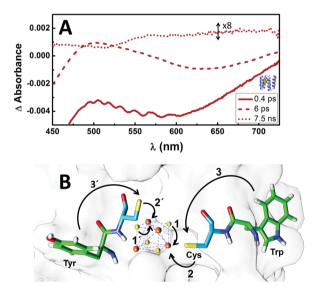


Fig. 4 (A) EADS for the CTPR2-[4Fe-4S] from the global fit of TA data with three lifetimes: 0.4 ps (straight line), 6 ps (dashed line) and 7.5 ns (dotted line). The 7.5 ns EADS is magnified by a factor of 8 for better visibility. (B) Proposed electron transfer pathways. Arrows show the direction of electron transfer. Iron is in orange, sulfur is yellow, cysteines (Cys) are light blue, tyrosine (Tyr) and tryptophan (Trp) are green.

surrounding protein is capable of stabilizing that state via long-range redox reaction to its outside, creating a long-lived (about 7 ns) charge-separated state. As such, the EADS corresponding to the species with 0.4 ps lifetime shows broad ground state bleaching (GSB) with maxima below 460 nm and around 600 nm (Fig. 4A and Fig. S5, ESI \dagger), characteristic of the S(3p) \rightarrow Fe(3d) LMCT excited state³⁸ that, due to the short lifetime, may be attributed to the transient internal electron transfer (ET) from a bridging sulfide (ET pathway 1, Fig. 4B). Similarly, the 6 ps EADS show GSB maxima below 460 nm and around 650 nm. In this case, due to the longer lifetime and the red-shifted GSB, the most probable path involves a cysteinyl sulfur transferring an electron to the iron in the [4Fe-4S] cluster (ET pathway 2, Fig. 4B), thus generating an electron hole in the cysteine. Interestingly, this state decays in part by forming a transient state with 7.5 ns lifetime (7.5 ns EADS) and characteristic bleaching of tyrosyl or tryptophanyl radical around 510 nm.40 This suggests the electron hole on the cysteine is most likely quenched by proton coupled electron transfer (PCET) promoted by the linked tryptophan where the electron acceptor is the cysteine hole and the proton acceptor is possibly water (pathway 3, Fig. 4B).41 Cysteine-linked tyrosine might be involved in this PCET as well (pathway 3'), but shorter distance (fewer connecting bonds between chromophores) and bigger driving force for the electron transfer would probably favor the tryptophan.

This study has demonstrated the potential of CTPR proteins as a scaffold for [4Fe-4S] clusters, enabling to coordinate a desired number of redox active clusters in a single assembly, with the cluster count controlled solely through the size of the protein. The repeat nature of the protein also imposed a consistent distance of about 1.6 nm between the clusters in multi-cluster assemblies. The designed proteins coordinate the clusters with expected stoichiometry, with one cluster per engineered binding site, and with spectral characteristics similar to those observed in natural ferredoxins. The CTPR-[4Fe-4S] constructs remain well folded, with the characteristic helical structure of the TPR domain. The EPR spectra of the reduced protein-cluster complexes correspond to the [4Fe-4S]¹⁺ state. Square wave voltammetry confirmed that the [4Fe-4S] clusters coordinated by the designed proteins are redox active and have redox potentials within the range of low-potential protein-bound clusters. Transient absorption spectroscopy showed that clusters form long-lived charge-separated states stabilized by the protein matrix, as observed in proteins involved in long range energy transfer. Direct interaction between the [4Fe-4S] clusters was not observed, but instead the excitation pathway was traced to the surface of the protein. Thus, the lack of cross-talk between the clusters is attributable to the structural factors of the system, in particular the large distance (>15 Å) between the clusters, the exposure of the clusters to the environment and the lack of driving force (e.g. difference in potentials) for inter-cluster communication, making the charge transfer to the environment (water) more favorable. Nevertheless, the reactivity of the clusters shows the potential of CTPR-[4Fe-4S] assemblies as a simplified platform for the studies of long-range ET, establishing the foundation for the modular design of long-range ET conduits.

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Overall, this communication serves to further demonstrate the versatility of CTPR proteins as a platform for designing functional materials or assemblies with order maintained at a molecular level. CTPR proteins have proven to be robust scaffolds for complex multifunctional systems with diverse functional elements. 22,23 The stabilization of redox active clusters brings the possibility of combining photo- and redox-active elements for full-fledged energy transfer conduits. As such, CTPR proteins show virtually unmatched potential as a protein scaffold for nanotechnological systems, including potential artificial photosystems.

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Conflicts of interest

There are no conflicts to declare.

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