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Elucidating the molecular function of human BOLA2 in

GRX3-dependent anamorsin maturation pathway

Lucia Banci^{1,2,§}, Francesca Camponeschi^{1,2}, Simone Ciofi-Baffoni^{1,2,§}, Riccardo Muzzioli^{1,2}

 ¹Magnetic Resonance Center CERM, University of Florence, Via Luigi Sacconi 6, 50019, Sesto Fiorentino, Florence, Italy
 ²Department of Chemistry, University of Florence, Via della Lastruccia 3, 50019 Sesto Fiorentino, Florence, Italy

[§] To whom correspondence should be addressed:

Lucia Banci, E-mail: banci@cerm.unifi.it, Phone: +39-055-4574273, Fax: +39-055-4574923 Simone Ciofi-Baffoni, E-mail: ciofi@cerm.unifi.it, Phone: +39-055-4574192; Fax: +39-055-4574923

Abstract

In eukaryotes, the interaction between members of the monothiol glutaredoxin family and members of the BolA-like protein family has been involved in iron metabolism. To investigate the still unknown functional role of the interaction between human glutaredoxin-3 (GRX3) and its protein partner BOLA2, we characterized at the atomic level the interaction of apo BOLA2 with the apo and holo states of GRX3, and studied the role of BOLA2 in the GRX3-dependent anamorsin maturation pathway. From these studies, it emerged that apo GRX3 and apo BOLA2 form a heterotrimeric complex, composed by two BOLA2 molecules and one GRX3 molecule. This complex is able to bind two [2Fe-2S]²⁺ clusters, each being bridged between a BOLA2 molecule and a monothiol glutaredoxin domain of GRX3, and to transfer both [2Fe-2S]²⁺ clusters to apo anamorsin producing its mature holo state. Collectively, the data suggest that the heterotrimeric complex can work in the cytosol as a [2Fe-2S]²⁺ cluster transfer component in Fe/S protein maturation pathways.

Introduction

In eukaryotes, the biosynthetic pathways responsible for the maturation of Fe/S proteins is a complex, multi-step process involving more than 30 different proteins.¹ As a result, atomic level studies are vital to fully unravel the molecular function of each single player in the Fe/S protein maturation process. In this work we characterized at the atomic level the interaction between human glutaredoxin-3 (GRX3) and its protein partner BOLA2, and identified a possible role for BOLA2 in the GRX3-dependent anamorsin maturation pathway.

The functional role of eukaryotic monothiol glutaredoxins (Grx) has been largely studied in S. cerevisiae, which encodes three CysGlyPheSer(CGPS)-type monothiol Grx homologues, Grx3, Grx4, and Grx5.^{2,3,4} Grx5 is a single domain protein, located in the mitochondrial matrix, which binds a [2Fe-2S] cluster ligated by two protein molecules, each providing a cysteine ligand, and by two glutathione molecules, each providing a further cysteine ligand.^{5,6} Grx5 has been shown to participate in the mitochondrial iron-sulfur (Fe/S) cluster (ISC) assembly machinery, acting, in its holo-homodimeric state, as a donator of a [2Fe-2S] cluster to partner apo-proteins.^{7,8} Grx3 and Grx4 are located in the cytoplasm and each consists of a N-terminal thioredoxin (Trx) domain, with no Trx-related activity,^{9,10} and of a glutaredoxin domain capable of binding a [2Fe-2S] cluster through protein dimerization and glutathione binding,¹¹ in the same way as Grx5 does. Grx3 and Grx4 have been linked to iron regulation through their binding to the transcriptional activator Aft1/2, which regulates iron uptake in yeast.¹²⁻¹⁴ Specifically, to perform this regulatory function, Grx3 (or Grx4) needs to be complexed with the BolA-like protein Fra2, forming a [2Fe-2S]-bridged heterodimeric complex,^{11,15} which, at variance with the holo-homodimeric species, can transfer the cluster to the iron-responsive transcription factor Aft2.¹⁶ Based on this information on yeast, holoheterodimeric Grx/BolA complexes have been generally linked to iron homeostasis regulation, while holo-homodimeric Grx complexes have been related to Fe/S protein biogenesis.¹⁷ Recently, it has been also shown that double depletion of yeast Grx3/4

specifically impaired all iron-requiring reactions in the cytosol, in mitochondria, and in the nucleus, including the synthesis of Fe/S clusters, of heme, and of di-iron centers.⁹ These data suggest that holo-homodimeric cytoplasmatic Grxs species can also function in intracellular iron trafficking.

In humans, two monothiol Grxs are present, GRX3 and GRX5.^{2,18} GRX5 is located in mitochondria essentially performing the same function as the yeast Grx5 homolog, i.e. it acts as a [2Fe-2S] cluster transfer protein in the ISC machinery.^{19,20} GRX3 consists of three domains: one N-terminal Trx domain and two Grx domains, each able to bind a glutathionecoordinated [2Fe-2S] cluster via protein dimerization.^{21,22} GRX3 is located in the cytosol and, similarly to what found for yeast Grx3/Grx4, the molecular function of its holo-homodimeric state ([2Fe-2S]₂ GRX3₂) has been associated to intracellular iron trafficking, being responsible for iron redistribution to virtually all iron-binding proteins or iron-dependent pathways in the cell.²³ Recently, we proposed that human GRX3 is an active component of the cytosolic Fe/S cluster assembly (CIA) machinery, as its homodimeric [2Fe-2S]₂ GRX3₂ form is able to mature the cytosolic [2Fe-2S]-binding protein anamorsin, a component of the CIA machinery.²⁴ The CIA machinery is responsible for the maturation of all cytosolic and nuclear Fe/S proteins, which perform key functions in metabolic catalysis, iron regulation, protein translation, DNA synthesis, and DNA repair.^{25,26} Therefore, the cellular function of GRX3 in maturing the CIA component anamorsin indirectly affects several crucial life processes.

Similarly to yeast, GRX3 can form a holo-hetero complex with a BolA-like protein, the cytosolic BOLA2.²¹ However, while the yeast Grx3/4 forms a [2Fe-2S]-bridged heterodimer with a Fra2 molecule, the human GRX3 forms a [2Fe-2S]-bridged heterotrimer with two BOLA2 molecules ([2Fe-2S]₂ GRX3-BOLA2₂). Since there is no human ortholog of the transcription factor Aft1, being indeed cellular iron homeostasis in humans regulated at the RNA-level by the Iron Regulatory Proteins 1 and 2,²⁷ [2Fe-2S]₂ GRX3-BOLA2₂

heterotrimers cannot be involved in the same [2Fe-2S] cluster transfer pathway found in yeast for regulating iron homeostasis. Although other functions of the [2Fe-2S]₂ GRX3-BOLA2₂ heterotrimers in human cells might be possible, it seems conceivable that this complex takes a role in iron metabolism given that: i) [2Fe-2S]-bridged Grx3-BolA2 interaction is conserved from *S. cerevisiae* to humans,¹⁷ ii) human GRX3 binds iron *in vivo*,²² and iii) human GRX3 partially rescues the growth defects and iron accumulation in some *S. cerevisiae grx3*\Delta*grx4*\Delta mutants, suggesting that human GRX3 partially can complement the functions of yeast Grx3 and Grx4 in iron homeostasis.²⁸

In this work we have characterized *in vitro* the interaction of apo BOLA2 with the apo and holo states of GRX3, and investigated the role of BOLA2 in the GRX3-dependent anamorsin maturation pathway.

Experimental Section

Protein production

The cDNA coding for human BOLA2 (UniProtKB/Swiss-Prot: Q9H3K6) was acquired from Life technologies. BOLA2 gene was amplified by PCR and subsequently inserted into the pET21a vector using the restriction enzymes NdeI and BanHI. BL21(DE3)gold competent *E. coli* cells (Stratagene, La Jolla, CA) were transformed with the obtained plasmid, and cells were grown in LB or minimal media (with (¹⁵NH4)₂SO4 and/or [¹³C]-glucose) containing 1 mM ampicillin at 37 °C under vigorous shaking up to a cell OD₆₀₀ of 0.6. Protein expression was induced by adding 0.5 mM IPTG and cells were grown for 4 hours at 25 °C. Cells were harvested by centrifugation at 7500 x g and resuspended in lysis buffer (25 mM MES pH 6 containing 0.01 mg/ml DNAase, 0.01 mg/ml lysozyme, 1 mM MgSO4, 0.5 mM EDTA and 5 mM DTT). Cell disruption was performed on ice by sonication and the soluble extract, obtained by ultracentrifugation at 40000 x g, was loaded on HiTrap SP FF column (GE Healthcare) and BOLA2 protein was eluted with 25 mM MES pH 6, 1 M NaCl and 5 mM

DTT. The protein was then concentrated with Amicon Ultra-15 Centrifugal Filter Units with a MWCO of 3 kDa (Millipore) and the buffer exchanged by PD10 desalting column in 50 mM phosphate buffer pH 7, 5 mM DTT and 5 mM GSH.

Various constructs of human GRX3 (full-length protein, Trx and GRX3(GrxA/B) constructs in their apo and/or [2Fe-2S]-bound forms) were obtained following previously reported procedures.²⁴ [2Fe-2S]₂ GRX3₂ was obtained by chemical reconstitution following a previously reported procedure.²⁴

Biochemical and UV/vis, CD, EPR spectroscopic methods

The aggregation state of isolated apo and holo proteins and of protein mixtures was analyzed using analytical gel filtration on Superdex 75 HR 10/30 and Superdex 200 10/300 increase columns (Amersham Bioscience), calibrated with gel filtration marker calibration kit, 6500-66000 Da (Sigma Aldrich), to obtain the apparent molecular masses of the detected species. Purified samples in degassed phosphate buffer 50 mM pH 7, 5 mM DTT, 5 mM GSH were loaded on the column pre-equilibrated with degassed 50 mM phosphate buffer pH 7, 5 mM GSH, 5 mM DTT. Elution profiles were recorded at 280 nm with a flow rate of 0.5 ml/min for Superdex 75 HR 10/30 column and 0.75 ml/min for Superdex 200 10/300 increase column.

UV/vis and CD spectra were anaerobically acquired on a Cary 50 Eclipse spectrophotometer and JASCO J-810 spectropolarimeter, respectively, in degassed 50 mM phosphate buffer pH 7, 5 mM GSH and 5 mM DTT.

EPR spectra of the 1:4 [2Fe-2S]₂ GRX3₂-apo BOLA2 mixture and of the chemically reconstituted [2Fe-2S]₂ GRX3-BOLA2₂ heterotrimeric complex were recorded after the anaerobic reduction of the cluster by stoichiometric addition of sodium dithionite and immediate freezing of the protein solution in liquid nitrogen. EPR spectra were acquired in degassed 50 mM phosphate buffer pH 7, 5 mM GSH, 5 mM DTT and 10% glycerol at 45 K, using a Bruker Elexsys E500 spectrometer working at a microwave frequency of ca. 9.45

GHz, equipped with a SHQ cavity and a continuous flow He cryostat (ESR900, Oxford instruments) for temperature control. Acquisition parameters were as following: microwave frequency, 9.640928 GHz; microwave power, 5 mW; modulation frequency, 100 KHz; modulation amplitude, 2.0 G; acquisition time constant, 163.84 ms; number of points 1024; number of scans 8; field range 500-6000 G or 2300-4300 G.

The iron and inorganic sulfur content and the protein concentration were estimated following standard chemical assays as already reported.²⁹

NMR spectroscopy

Standard ¹H-detected triple-resonance NMR experiments for backbone resonance assignment were recorded on 1 mM ¹³C, ¹⁵N labeled samples (apo forms of GRX3(GrxA/B) and BOLA2) in degassed 50 mM phosphate buffer pH 7, 5 mM DTT and 5 mM GSH at 298 K, using Bruker AVANCE 500 MHz and 700 MHz spectrometers. ¹⁵N heteronuclear relaxation experiments were performed on ¹⁵N-labeled apo BOLA2 and on ¹⁵N-labeled apo GRX3(GrxA/B) in the presence and in the absence of two equivalents of unlabeled apo BOLA2, in degassed 50 mM phosphate buffer pH 7, 5 mM DTT and 5 mM GSH at 298 K, to measure ¹⁵N backbone longitudinal (R₁) and transverse (R₂) relaxation rates and heteronuclear ¹⁵N{¹H} NOEs. All NMR data were processed using the Topspin software package and were analyzed with the program CARA.

An apparent dissociation constant (K_d) for the interaction between apo GRX3(GrxA/B) and apo BOLA2 proteins, in degassed 50 mM phosphate buffer pH 7, 5 mM DTT at 298 K, was obtained by plotting the average chemical shift differences δ_{av} (i.e. (((Δ H)² + (Δ N/5)²)/2)^{1/2}, where Δ H and Δ N are chemical shift differences for ¹H and ¹⁵N, respectively) of five well resolved backbone NH signals of ¹⁵N labeled GRX3(GrxA/B) and of six well resolved backbone NH signals of ¹⁵N labeled BOLA2, as a function of BOLA2 and GRX3(GrxA/B) concentrations, respectively. These data were then fitted to a simple two-component model and averaged to obtain the final K_d value.^{30,31}

Protein-protein interaction and cluster transfer

Apo GRX3 (or apo GRX3(GrxA/B))-apo BOLA2 interaction was investigated by ¹H-¹⁵N HSQC NMR spectra, titrating ¹⁵N labeled apo forms of GRX3 or GRX3(GrxA/B) with unlabeled apo BOLA2, and ¹⁵N labeled apo BOLA2 with unlabeled apo forms of GRX3 or GRX3(GrxA/B), in degassed 50 mM phosphate buffer pH 7, 5 mM GSH and 5 mM DTT containing 10% (v/v) D₂O at 298K. Spectral changes were monitored after the addition of increasing amounts of the unlabeled partner. NMR data were analysed with CARA program and plotted following standard procedures.

To follow changes in cluster coordination upon interaction between [2Fe-2S]₂ GRX3₂ and apo BOLA2, [2Fe-2S]₂ GRX3₂ was incubated under anaerobic conditions with increasing concentrations of apo BOLA2up to a 1:4 protein ratio, in degassed 50 mM phosphate buffer pH 7, 5 mM GSH and 5 mM DTT. UV/vis, EPR and CD spectra were then recorded as described above, and compared with those collected on a [2Fe-2S]₂ GRX3-BOLA2₂ complex, which was obtained by chemically reconstituting a 1:2.5 apo GRX3-apo BOLA2 mixture following a previously reported procedure.²⁴ Protein-protein interaction between [2Fe-2S]₂ GRX3₂ and apo BOLA2 was investigated by ¹H-¹⁵N HSQC NMR spectra performed on ¹⁵N labeled [2Fe-2S]₂ GRX3₂ titrated with unlabeled apo BOLA2, and on ¹⁵N labeled apo BOLA2 titrated with unlabeled [2Fe-2S]₂ GRX3₂. Spectral changes were monitored and analysed after the addition of increasing amounts of the unlabeled partner.

The chemically reconstituted [2Fe-2S]₂ GRX3-BOLA2₂ complex was titrated under anaerobic conditions with increasing concentrations of apo anamorsin up to a 1:1 protein ratio in degassed 50 mM phosphate buffer pH 7, 5 mM GSH and 5 mM DTT. Cluster transfer and the protein-protein interaction were followed by UV/vis and CD spectroscopy and analytical gel filtration chromatography, respectively.

Results

Interaction of apo GRX3 with apo BOLA2

BOLA2 Full-length protein was overexpressed in E. coli cells and purified in its apo form. Analytical gel filtration ^{15}N NMR relaxation and data (Supporting Information Fig. **S1**), which provided an overall reorientational correlation time of 6.88 ± 0.23 ns. showed that the purified apo protein is monomeric. Purified apo BOLA2 was unable to bind Fe/S clusters when a chemical reconstitution approach²⁹ was applied.

To investigate the interaction between the apo forms of GRX3 and BOLA2, chemical shift changes of backbone NHs were followed by ¹H-¹⁵N HSQC NMR experiments upon titration of ¹⁵N-labeled apo GRX3 with unlabeled apo BOLA2, and of ¹⁵N-labeled apo BOLA2 with unlabeled apo GRX3, in the presence of 5



Figure 1. Apo GRX3 interacts with apo BOLA2. (A) Overlay of ¹H-¹⁵N HSQC spectra of ¹⁵N labeled apo GRX3 (black), of ¹⁵N labeled Trx domain of GRX3 (green) and of a 1:2 mixture of ¹⁵N labeled apo GRX3 and unlabeled apo BOLA2 (red). The chemical shift changes of two backbone NHs belonging to GrxA and GrxB domains, K192 and D286, respectively, are reported in the inset for ¹⁵N labeled apo GRX3 (black), for 1:1 (blue) and 1:2 (red) mixtures of ¹⁵N labeled apo GRX3 and unlabelled apo BOLA2. (**B**) Overlay of ¹H-¹⁵N HSQC spectra of ¹⁵N labeled apo BOLA2 (black) and of a 1:2 mixture of unlabeled apo GRX3 and ¹⁵N labeled apo BOLA2 (red).

mM GSH and 5 mM DTT. Chemical shift variations, which were in a fast/intermediate exchange regime on the NMR time scale, were observed in the ¹H-¹⁵N HSQC maps of GRX3 for the backbone NHs of both Grx domains, when increasing amounts of unlabeled BOLA2

were added up to a 1:2 GRX3-BOLA2 molar ratio, while no significant effects were observed on the backbone NHs of the Trx domain of GRX3 (**Fig. 1A**). These chemical shift changes occur simultaneously on both Grx domains, thus indicating that BOLA2 does not have a preferential interaction towards one of the two Grx domains (**Fig. 1A**, inset). Similarly, chemical shift variations in a fast/intermediate exchange regime on the NMR time scale were observed in the ¹H-¹⁵N HSQC maps of BOLA2 when increasing amounts of unlabeled GRX3 were added up to a 1:2 GRX3-BOLA2 molar ratio (**Fig. 1B**).

The interaction between apo BOLA2 and apo GRX3 was also followed by analytical gel filtration chromatography, performed on protein mixtures with different apo GRX3 and apo BOLA2 ratios (**Fig. 2A**). The formation of two peaks was observed by adding apo BOLA2 to apo GRX3 up to a 1:2 GRX3-BOLA2 ratio: i) a peak with an apparent molecular mass of 70.4 kDa, predominant at 1:1 protein ratio; ii) a peak with an apparent molecular mass of 74.3 kDa, predominant at 1:2 protein ratio (**Fig. 2A**).



Figure 2. Apo GRX3 and apo BOLA2 form a heterotrimeric 1:2 complex. (A) Analytical gel filtration (Superdex 200 10/300 increase column) chromatograms of BOLA2 and GRX3 proteins in their apo states and of their protein mixtures. Black line: apo GRX3; red line: apo BOLA2; green line: 1:0.5 mixture of apo GRX3 and apo BOLA2; blue line: 1:1 mixture of apo GRX3 and apo BOLA2; cyan line: 1:1.5 mixture of apo GRX3 and apo BOLA2; magenta line: 1:2 mixture of apo GRX3 and apo BOLA2. The apparent molecular masses are reported at the top of each chromatographic peak. (B) Analytical gel filtration (Superdex 75 HR 10/30 column) chromatograms of isolated BOLA2, GRX3(GrxA/B) and GRX3 proteins in their apo states and of their protein mixtures. Black line: apo BOLA2; red line: apo GRX3; green line: 1:2.5 mixture of apo GRX3(GrxA/B) and apo BOLA2; blue: 1:2.5 mixture of apo GRX3 and apo BOLA2. The apparent molecular masses are reported at the top of each chromatographic peak.

These results indicated that apo GRX3 forms a heterodimeric complex with apo BOLA2 at the 1:1 protein ratio (the 70.4 kDa value matches indeed with the sum of the apparent molecular masses of apo GRX3 and apo BOLA2), which then evolves, upon addition of a further equivalent of apo BOLA2, to form a species which has an apparent molecular mass (74.3 kDa) intermediate between that of the heterodimeric complex (70.4 kDa) and that of an apo heterotrimeric complex formed by two BOLA2 molecules and one GRX3 molecule (82 kDa, obtained by the sum of the apparent molecular masses of apo GRX3 and two molecules of apo BOLA2). The presence of this peak with an intermediate apparent molecular mass can be a consequence of the applied gel filtration conditions, which determined, indeed, the presence of a fast exchange equilibrium between the two heterodimeric and heterotrimeric complexes. Therefore, we performed the analytical gel filtration chromatography using a different column, i.e. Superdex 75 HR 10/30, and it resulted that apo GRX3 protein interacts with apo BOLA2 forming a peak with an apparent molecular mass of 76.8 kDa predominant at 1:2.5 GRX3-BOLA2 protein ratio (Fig. 2B). This value essentially corresponds to the sum of the apparent molecular masses of apo GRX3 (52.7 kDa) and two molecules of apo BOLA2 (11.1 kDa) (Fig. 2B), thus definitively indicating the formation of the apo GRX3-BOLA22 heterotrimeric complex. Since the Trx domain is not involved in the interaction between apo BOLA2 and apo GRX3, we produced a two-domain construct containing the GrxA and GrxB domains only, (GRX3(GrxA/B)), to map, by solution NMR, the residues involved in the formation of the heterotrimeric complex. With this shorter construct, the chemical shift data analysis was indeed simplified by the reduction of NMR signal overlap and broadening due to the reduced protein size. We titrated ¹⁵N-labeled BOLA2 with unlabeled GRX3(GrxA/B) and ¹⁵N-labeled GRX3(GrxA/B) with unlabeled BOLA2 and followed the chemical shift changes of backbone NHs via ¹H-¹⁵N HSQC experiments. Chemical shift variations are in a fast/intermediate exchange regime on the NMR time scale in both titrations, as it occurs for the apo GRX3-apo BOLA2 interaction. By fitting the chemical shifts to a simple twocomponent model,^{30,31} an apparent dissociation constant of $25 \pm 15 \mu M$ was obtained (Fig. 3A and 3B insets).



Figure 3. GrxA and GrxB domains interact with apo BOLA2. (A) Overlay of ¹H-¹⁵N spectra of ¹⁵N labeled apo HSOC GRX3(GrxA/B) (black) and of a 1:2 mixture of ¹⁵N labeled apo GRX3(GrxA/B) and unlabeled apo BOLA2 (red). In the inset, a zoom of ¹H-¹⁵N HSQC spectra on the K192 and D286 backbone NH signals is reported for 15N labeled apo GRX3(GrxA/B) (black), and for 1:0.5 (blue), 1:1 (green), 1:1.5 (violet) and 1:2 ^{15}N labeled mixtures of (red) ano GRX3(GrxA/B) and unlabeled apo BOLA2. The inset also shows the δ_{av} changes measured for T199 (red circles) and Q202 (blue triangles) in the GrxA domain of GRX3(GrxA/B), and of K245 (green squares) and F263 (magenta circles) in the GrxB domain of GRX3(GrxA/B) as a function of apo BOLA2 concentration. Solid lines show the fitting curves. (B) Overlay of ¹H-¹⁵N HSQC spectra of ¹⁵N labeled apo BOLA2 (black) and of a 2:1 mixture of ¹⁵N labeled apo BOLA2 and unlabeled apo GRX3(GrxA/B) (red). In the inset, a zoom of ¹H-¹⁵N HSQC spectra on the Q51 backbone NH signal is reported for ¹⁵N labeled apo BOLA2 (black), and for the 0.1:1 (blue), 0.15:1 (green), 0.25:1 (violet), 0.5:1 (dark green), 0.75:1 (grey), 1:1 (brown) and 2:1 (red) mixtures of unlabeled apo GRX3(GrxA/B) and ¹⁵N labeled apo BOLA2. The inset also shows the δ_{av} changes measured for the L50, O51. V56 and N57 residues of BOLA2 as a function of apo BOLA2 concentration. Solid lines show the fitting curves.

The ¹H-¹⁵N HSQC signals of the final protein mixture, i.e. 1:2 ¹⁵N-labeled apo GRX3(GrxA/B)-apo BOLA2, well superimposed with the corresponding signals of a 1:2 apo ¹⁵N-labeled GRX3-apo BOLA2 mixture, indicating that the same complex involving the same interacting region was present in solution in the two mixtures, and confirming that Trx domain has no specific role in the protein-protein recognition between apo GRX3 and apo BOLA2 (**Supporting Information Fig. S2**). Comparing the ¹H-¹⁵N HSQC map of the final apo GRX3(GrxA/B)-apo ¹⁵N-labeled BOLA2 mixture with that of ¹⁵N-labeled BOLA2 (**Fig.**

3B), fifteen signals of BOLA2 experienced significant chemical shift variations and two NH signals broadened beyond detection (**Fig. 4A**).



Figure 4. Mapping the interaction surfaces between BOLA2 and each GrxA and GrxB domain in the apo heterotrimeric complex. The left panels show backbone weighted average chemical shift differences δ_{av} for BOLA2 residues, observed upon addition of unlabeled apo GRX3(GrxA/B) to ¹⁵Nlabeled apo BOLA2 up to a 2:1 BOLA2-GRX3(GrxA/B) ratio (A), and for GRX3(GrxA/B) residues observed upon addition of 2 eq. of unlabeled apo BOLA2 to ¹⁵N-labeled apo GRX3(GrxA/B) (B). A threshold of 0.15 ppm (mean value of δ_{av} plus 1 σ , black dashed line) for BOLA2 and of 0.10 ppm for GRX3(GrxA/B) were used to identify significant chemical shift differences. Cyan bars identify residues whose backbone NHs broaden beyond detection upon complex formation. The δ_{av} value for Pro residues was arbitrarily set to zero. The right panels show backbone NHs experiencing significant chemical shift differences in the ¹H-¹⁵N HSQC spectra upon the formation of the heterotrimeric apo GRX3(GrxA/B)-BOLA2₂ complex mapped as blue and cyan spheres on a structural model of apo BOLA2 (A) and on the crystal structure of GrxA (PDB ID 3ZYW) and GrxB (PDB ID 2YAN) domains of apo GRX3 (B). Blue spheres identify NHs showing significant chemical shift variations, and cyan spheres identify backbone NHs broaden beyond detection. Side-chains of His, Cys and Pro residues are in green, yellow and orange, respectively.

In the reverse titration, when detecting ¹⁵N labeled GRX3(GrxA/B) (**Fig. 3B**), fifteen NH signals broadened beyond detection and twenty-one NH signals experienced chemical shift variations in the ¹H-¹⁵N HSQC spectrum of the final 1:2 GRX3(GrxA/B)-BOLA2 mixture compared with that of ¹⁵N-labeled GRX3(GrxA/B) (**Fig. 4B**).

Once the affected residues are mapped on the structural model of apo BOLA2 (calculated with MODELLER 9.15 software using the solution structure of apo BOLA2 from *Mus musculus*³² as template with a 87% sequence identity) and on the crystal structures of the apo form of the single GrxA (PDB ID 3ZYW) and GrxB (PDB ID 2YAN) domains of GRX3, well-defined interacting regions were identified on both proteins. On BOLA2 side, it comprises helix α 3 and the following strand β 3, containing His 68, a ligand of the [2Fe-2S]²⁺ cluster²¹ (**Fig. 4A**). On each GrxA and GrxB domain, the interacting surface involves helices α 2 and α 3, strand β 4 and the surrounding loops, and comprises the Cys ligand of the [2Fe-2S]²⁺ bound cluster²¹ (**Fig. 4B**).

¹⁵N backbone NMR relaxation experiments, performed on ¹⁵N-labeled apo GRX3(GrxA/B) in absence or in presence of two equivalents of apo BOLA2, showed an increase in the molecular tumbling value (τ_m) from 12.0 ± 1.2 ns (consistent with a monomeric state of GRX3(GrxA/B) in solution) to 27.7 ± 4.5 ns. This increase is consistent with the formation of a 1:2 GRX3(GrxA/B)-BOLA2 protein complex, being the τ_m value of apo BOLA2 6.4 ± 1.0 ns (consistent with a monomeric protein state in solution). This result is in agreement with the analytical gel filtration data collected with a Superdex 75 HR 10/30 column (**Fig. 2B**), which showed the formation of a peak with an apparent molecular mass of 49.9 kDa in the chromatogram of the protein mixture, which essentially corresponds to the sum of the apparent molecular masses of apo GRX3(GrxA/B) (29.6 kDa) and two molecules of apo BOLA2 (11.1 kDa).

Interaction of [2Fe-2S]₂ GRX3₂ with apo BOLA2

The interaction of apo BOLA2 with the holo-dimeric form of GRX3, i.e. $[2Fe-2S]_2$ GRX3₂, was characterized by UV/vis, CD, EPR, NMR and analytical gel filtration. Upon stepwise additions of apo BOLA2 to $[2Fe-2S]_2$ GRX3₂, the absorbance peaks typical of the oxidized $[2Fe-2S]^{2+}$ clusters bound to GRX3 disappeared in the UV/vis and CD spectra and new absorbance peaks appeared. The final spectra (**Fig. 5A** and **5B**) at a 1:4 [2Fe-2S]₂ GRX3₂-BOLA2 ratio are very similar to that of the holo complex obtained by chemically reconstituting apo GRX3-BOLA2₂ with two $[2Fe-2S]^{2+}$ clusters ([2Fe-2S]₂ GRX3-BOLA2₂, hereafter), as assessed by iron, and acid-labile sulfide chemical analysis (Fe and acid-labile S values, reported as mol Fe or S per mol of trimer, are 3.8 ± 0.1 and 3.9 ± 0.1 , respectively).



Figure 5. Formation of the [2Fe-2S] GRX3-BOLA2₂ complex from the interaction between [2Fe-2S]₂ GRX3₂ and apo BOLA2. (A) UV/vis and (B) CD spectra of [2Fe-2S]₂ GRX3₂ (black line), of the 1:4 [2Fe-2S]₂ GRX3₂-apo BOLA2 mixture (blue line), and of chemically reconstituted [2Fe-2S]₂ GRX3-BOLA2₂ (green line). (C) Analytical gel filtration (Superdex 75 HR 10/30 column) chromatograms of BOLA2 (magenta) and GRX3 (cyan) proteins in their apo states, of [2Fe-2S]₂ GRX3₂ complex (black) and of a 1:4 [2Fe-2S]₂ GRX3₂-apo BOLA2 mixture (green). The apparent molecular masses are reported at the top of each chromatographic peak. (D) In the upper panels, the overlay of ¹H-¹⁵N HSQC spectral regions of ¹⁵N labeled [2Fe-2S]₂ GRX3₂ (black), and of 1:2 (red) and 1:4 (blue) ¹⁵N labeled [2Fe-2S]₂ GRX3₂ unlabeled apo BOLA2 (black), and of 2:1 (red) and 4:1 (blue) ¹⁵N labeled apo BOLA2-unlabeled [2Fe-2S]₂ GRX3₂ mixtures.

The 1:4 protein-protein ratio is needed to fully saturate the four Grx domains present in the dimeric [2Fe-2S]₂ GRX3₂ complex with BOLA2 molecules. The final mixture containing [2Fe-2S]₂ GRX3₂ and apo BOLA2 at a 1:4 molar ratio was EPR silent, as expected for the presence of a S=0 ground state of oxidized [2Fe-2S]²⁺ clusters. Upon chemical reduction with sodium dithionite, the EPR signal of a reduced [2Fe-2S]⁺ protein-bound cluster is observed with g values of 2.01, 1.91, and ~1.87 (g_{av} ~1.93), which are the same as those observed on the chemically reconstituted heterotrimeric [2Fe-2S]₂ GRX3-BOLA2₂ complex upon sodium dithionite reduction (Supporting Information Fig. S3). Analytical gel filtration chromatography, performed with a Superdex 75 HR 10/30 column on the same final mixture, showed the presence of a main peak with an apparent molecular mass of 73.2 kDa, which is very close to the sum of the apparent molecular masses of apo GRX3 (52.7 kDa) and two molecules of apo BOLA2 (11.1 kDa) (Fig. 5C). These results indicate that [2Fe-2S]₂ GRX3₂ interacts with apo BOLA2 to form, in the final protein mixture, a GRX3-BOLA22 heterotrimeric complex that binds [2Fe-2S]²⁺ cluster(s). The EPR, UV/vis, CD and analytical gel filtration data collected on this final mixture mirror the data previously obtained on a [2Fe-2S]₂ GRX3(GrxA/B)-BOLA2₂ complex by co-expression of human BOLA2 and GRX3(GrxA/B) in *E. coli* cells,²¹ thus indicating that the same type of cluster is bound to the two heterotrimeric complexes following different holo protein production approaches, i.e. chemical reconstitution vs. in vitro and in cell Fe-S metallation, as well as different protein constructs, i.e. full-length protein vs. a GrxA/GrxB domain construct.

Holo complex formation between [2Fe-2S]₂ GRX3₂ and apo BOLA2 was also followed by NMR, performing ¹H-¹⁵N HSQC spectra on ¹⁵N-labeled [2Fe-2S]₂ GRX3₂ titrated with unlabeled apo BOLA2 up to a protein-protein ratio of 1:4, and, viceversa, on ¹⁵N labeled BOLA2 titrated with unlabeled [2Fe-2S]₂ GRX3₂ up to a 4:1 protein-protein ratio. In the final mixture of the first titration, the NH signals of GRX3 affected by BOLA2 additions belong to residues of both Grx domains, while those of the Trx domain remain unperturbed. In the ¹H-

¹⁵N HSQC NMR spectra acquired throughout the titration steps (Fig. 5D), a decrease in the intensity of some backbone NH signals of [2Fe-2S]₂ GRX3₂ (located more than 10 Å away from the paramagnetic [2Fe-2S]²⁺ cluster and therefore only affected by BOLA2-GRX3 interaction) was observed, with the concomitant appearance of new backbone NH signals. This indicates the formation of a new species in solution in slow exchange, on the NMR time scale, with the [2Fe-2S]₂ GRX3₂ species (Fig. 5D). This species has chemical shifts slightly different from those of the heterotrimeric complex formed by two apo BOLA2 molecules and an apo GRX3 molecule. In the final mixture, the NH signals of the [2Fe-2S]₂ GRX3₂ species fully disappeared in favor of the new NH signals, indicating the complete consumption of [2Fe-2S]₂ GRX3₂, which therefore completely evolves to the new species (Fig. 5D). In the reverse titration, upon addition of unlabeled [2Fe-2S]₂ GRX3₂, some backbone NHs of ¹⁵N labeled BOLA2 broaden beyond detection and others display chemical shift changes in a slow exchange regime on the NMR time scale (Fig. 5D). The residues undergoing such spectral changes are essentially the same as those involved in the apo/apo interaction, indicating that the interaction region is the same in both apo and holo GRX3-BOLA2 interactions. Overall, the spectroscopic and gel filtration data indicate: i) a complete conversion of the homodimeric [2Fe-2S]₂ GRX3₂ into a GRX3-BOLA2₂ heterotrimeric complex that contains [2Fe-2S]²⁺ cluster(s); ii) similarly to what we found for the apo/apo heterotrimeric complex, the GrxA and GrxB domains of GRX3 interact with two BOLA2 molecules, while Trx domain is not involved in such interaction.

Two possible scenarios can occur at the 1:4 [2Fe-2S]₂ GRX3₂-BOLA2 ratio, i.e. the formation of a heterotrimeric GRX3-BOLA2₂ complex containing two [2Fe-2S]²⁺ clusters together with the formation of a heterotrimeric apo GRX3-apo BOLA2₂ complex, or the formation of two heterotrimeric GRX3-BOLA2₂ complexes each containing only one [2Fe-2S]²⁺ cluster. The two scenarios result from two possible occurring molecular mechanisms, **A** and **B** shown in **Scheme 1**.



Scheme 1. Possible solution equilibria upon reaction of $[2Fe-2S]_2 GRX3_2$ with apo BOLA2 up to a 1:4 ratio. (A) Two molecules of apo BOLA2 (in blue) interact with a $[2Fe-2S]_2 GRX3_2$ molecule (GRX3 molecules are in red) to form a heterotrimeric GRX3-BOLA2₂ complex that binds two $[2Fe-2S]^{2+}$ clusters and an apo GRX3 molecule. The further addition of two equivalents of apo BOLA2 determines the formation of an apo heterotrimeric GRX3-BOLA2₂ complex; (B) two distinct heterodimeric GRX3-BOLA2 complexes, containing one $[2Fe-2S]^{2+}$ cluster each, are first formed. Then, the further addition of two equivalents of apo BOLA2 determines the formation of two distinct heterotrimeric GRX3-BOLA2₂ complexes still containing one $[2Fe-2S]^{2+}$ cluster each.

The key experimental evidence, that would allow us to provide information on whether a favorite mechanism between **A** and **B** is operative, is the detection of the formation of the apo free GRX3 form, which is observed indeed in the **A** mechanism only (**Scheme 1**). Analytical gel filtration were therefore performed on [2Fe-2S]₂ GRX3₂-apo BOLA2 protein mixtures at different protein:protein ratios, using a Superdex 75 HR 10/30 column to investigate the possible presence of free apo GRX3. These gel filtration analysis showed: i) the presence of a main peak with an apparent molecular mass which increases from 68.0 kDa to 73.2 kDa upon addition of apo BOLA2, ii) the presence, only for [2Fe-2S]₂ GRX3₂-apo BOLA2 ratios greater than 1:4, of a peak with the apparent molecular mass of apo GRX3 at any investigated protein:protein ratio (**Fig. 6A**).



Figure 6. The formation of apo GRX3 is not observed upon the interaction of [2Fe-2S]₂ GRX3₂ with apo BOLA2. Analytical gel filtration chromatograms of (**A**) BOLA2 (black) and GRX3 (red) proteins in their apo states, of [2Fe-2S]₂ GRX3₂ (blue) and of a 1:2 (cyan), 1:4 (magenta), 1:6 (green) [2Fe-2S]₂ GRX3₂-apo BOLA2 mixtures (Superdex 75 HR 10/30 column), and of (**B**) BOLA2 (black) and GRX3 (red) proteins in their apo states, of [2Fe-2S]₂ GRX3₂ (blue) and of a 1:0.25 (orange), 1:0.5 (light gray), 1:1 (light magenta) [2Fe-2S]₂ GRX3₂-apo BOLA2 mixtures (Superdex 200 10/300 increase column). The apparent molecular masses are reported at the top of each chromatographic peak.

The analytical gel filtration chromatography was also performed at higher resolution with a Superdex 200 10/300 increase column on protein mixtures at sub-stoichiometric [2Fe-2S]₂ GRX3₂-apo BOLA2 ratios of 1:0.25, 1:0.5 and 1:1 (**Fig. 6B**). It was observed the concurrent formation of i) the 1:1 heterodimeric complex, corresponding to the peak with an apparent molecular mass of 71.3 kDa, which is indeed very close to the sum of the apparent molecular masses of one molecule of apo BOLA2 and one molecule of apo GRX3 (i.e. 70.4 kDa), and ii) of the heterotrimeric complex, corresponding to the peak with an apparent molecular mass of 87.4 kDa, which is indeed close to the sum of the apparent molecular mass of 87.4 kDa, which is indeed close to the sum of the apparent molecular mass of 87.4 kDa, which is indeed close to the sum of the apparent molecular mass of 87.4 kDa, which is indeed close to the sum of the apparent molecular mass apparent molecules and one molecule of apo GRX3 (i.e. 82.0 kDa). The peak of apo GRX3, which is always present at a low percentage in the chemically reconstituted [2Fe-2S]₂ GRX3₂ sample, does not increase in its intensity upon BOLA2 additions (**Fig. 6B**), as expected if the **A** mechanism was the favored process. From these gel filtration data we therefore conclude that the **B** mechanism is the preferential occurring process.

[2Fe-2S]₂ GRX3-BOLA2₂ complex transfers [2Fe-2S]²⁺ clusters to apo anamorsin

Cluster transfer from [2Fe-2S]₂ GRX3-BOLA2₂ to apo anamorsin was followed by UV/vis, CD spectroscopy and analytical gel filtration. Once the chemically reconstituted [2Fe-2S]₂ GRX3-BOLA2₂ heterotrimeric complex was titrated with apo anamorsin up to a 1:1 proteinprotein ratio, in the UV/vis and CD spectra the absorbance peaks typical of [2Fe-2S]₂ GRX3-BOLA2₂ disappeared and the absorbance peaks typical of the [2Fe-2S]²⁺ cluster-bound form of anamorsin ([2Fe-2S] anamorsin)^{29,33} appeared (**Fig. 7A** and **7B**).



Figure 7. [2Fe-2S]₂ GRX3-BOLA2₂ transfers its clusters to apo anamorsin upon complex formation. (A) UV/vis spectra of [2Fe-2S]₂ GRX3-BOLA2₂ (black line) titrated with 0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.8 equivalents of apo anamorsin (black dashed lines). The arrow indicates the direction of the change in intensity and wavelength of the absorbance peaks with the increase of apo anamorsin concentration. The UV/vis spectra of the final 1:1 protein mixture and of [2Fe-2S] anamorsin are shown as red and green lines, respectively. (B) CD spectra acquired to follow cluster transfer from [2Fe-2S]₂ GRX3-BOLA2₂ to apo anamorsin. Black line: chemically reconstituted [2Fe-2S]₂ GRX3-BOLA2₂; green line: [2Fe-2S] anamorsin; red line: 1:1 mixture of [2Fe-2S]₂ GRX3-BOLA2₂ and apo anamorsin. (C) Analytical gel filtration (75 HR 10/30 column) chromatograms of BOLA2 (magenta) and anamorsin (blue) proteins in their apo states, of [2Fe-2S]₂ GRX3-BOLA2₂ (black) and of a 1:1 mixture between [2Fe-2S]₂ GRX3-BOLA2₂ and apo anamorsin (red). The apparent molecular masses are reported at the top of each chromatographic peak.

The CD spectrum of the final 1:1 mixture essentially corresponds to that of [2Fe-2S] anamorsin, being clearly distinguishable from that of [2Fe-2S]² GRX3-BOLA2². Analytical gel filtration chromatography performed with Superdex 75 HR 10/30 column on the final 1:1 mixture showed the presence of a main peak with an apparent molecular mass of 128.2 kDa, which essentially corresponds to the sum of the apparent molecular masses of [2Fe-2S]² GRX3-BOLA2² (73.2 kDa) and apo anamorsin (52.0 kDa) (**Fig. 7C**). Overall, the data showed that: i) both [2Fe-2S]²⁺ clusters are transferred from [2Fe-2S]² GRX3-BOLA2² to apo anamorsin; ii) the final product is a complex formed by an apo GRX3 molecule, two apo BOLA2 molecules and a [2Fe-2S] anamorsin molecule (GRX3-BOLA2²-[2Fe-2S] anamorsin, hereafter).

Moreover, by gel filtrating a 1:1 mixture of apo GRX3(GrxA/B)-BOLA22 and [2Fe-2S] anamorsin, a peak corresponding to a ternary complex formed by GRX3(GrxA/B), BOLA2 and anamorsin is not observed (Supporting Information Fig. S4), at variance to what detected in the 1:1 GRX3-BOLA2₂-[2Fe-2S] anamorsin mixture, where the corresponding ternary complex is formed (Fig. 7C). This demonstrates that, in the final GRX3-BOLA22-[2Fe-2S] anamorsin complex, the interaction between apo GRX3-BOLA22 and [2Fe-2S] anamorsin depends on the presence of the Trx domain of GRX3. We had previously shown that the Trx domain of GRX3 interacts with the N-terminal domain of anamorsin mediating the cluster transfer between the two proteins.²⁴ Here we show that the Trx domain, in addition of being essential for the complex formation between anamorsin and GRX3-BOLA22 in the GRX3-BOLA22-[2Fe-2S] anamorsin complex, maintains the same conformational freedom as it has in the homodimeric [2Fe-2S]₂ GRX3₂ complex. Therefore, we propose that the cluster transfer mechanism previously reported for [2Fe-2S]₂ GRX3²⁴ is also operative in the [2Fe-2S]₂ GRX3-BOLA2₂-apo anamorsin interaction, i.e. the protein recognition, specifically occurring between the N-terminal domains of the two proteins, plays a role in the cluster transfer process.

Discussion

BolA-like proteins have recently emerged as novel players in iron metabolism, being involved, on the basis of genetic and biochemical studies, in intracellular iron regulation pathways and in the maturation of Fe/S cluster-containing proteins.^{3,17,34} Specifically, a role for the cytosolic *S. cerevisiae* BolA2 (usually named Fra2) in iron regulation^{14,16,35} and for the mitochondrial human BOLA3 in the production of the lipoate-containing 2-oxoacid dehydrogenases and in the assembly of the respiratory chain complexes³⁶ have been established. However, there are no studies exploring analogous functions for BolA homologues in *S. cerevisiae* and humans. In particular, since the Fra2-dependent, iron signalling pathway described in *S. cerevisiae* is not conserved in humans,²⁷ a homologous regulatory function of BOLA2 in human cells can be excluded.

In this work we have investigated by *in vitro* studies the role of the GRX3-BOLA2 interaction in cytoplasmic Fe/S protein biogenesis, i.e. in the CIA pathway, being inspired i) by the general accepted view that BOLA2 protein is involved in human iron metabolism once interacting with its protein partner GRX3;^{17,37} ii) by knowing that human BOLA2 forms *in vitro* a [2Fe-2S]-bridged hetero complex with each monothiol Grx domain of GRX3;²¹ and iii) by our recent finding that the holo form of GRX3 is capable to mature the cytosolic Fe/S protein anamorsin, thus proposing GRX3 as a component of the CIA machinery.²⁴

Our data showed that apo BOLA2 and the Grx domains of apo GRX3 significantly and specifically interact forming an apo heterotrimeric complex composed by a GRX3 molecule and two BOLA2 molecules. This complex is able to bind two [2Fe-2S]²⁺ clusters upon chemical reconstitution. The same cluster content was achieved in the heterotrimeric complex obtained by co-expressing in *E. coli* cells human BOLA2 and a construct of GRX3 containing only the two Grx domains.²¹ A heterotrimeric GRX3-BOLA2₂ complex was also obtained by mixing *in vitro* [2Fe-2S]₂ GRX3₂ with apo BOLA2. However, in this case the formed heterotrimeric GRX3-BOLA2₂ complex contains only one [2Fe-2S]²⁺ cluster per complex.

These data indicate that a heterotrimeric GRX3-BOLA2₂ complex containing two [2Fe-2S]²⁺ clusters can be formed in one-step process only by inserting two [2Fe-2S]²⁺ clusters in an apo heterotrimeric GRX3-BOLA2₂ complex. Therefore, it is reasonable to suggest that the apo heterotrimeric GRX3-BOLA2₂ complex, similarly to apo GRX3 alone,²⁴ might be a physiologically relevant species that receives two [2Fe-2S]²⁺ clusters in the cytoplasm from a still unknown protein partner. In support of this, our data showed that, similarly to what observed for the holo homodimeric form of GRX3 containing two [2Fe-2S]²⁺ clusters,²⁴ the heterotrimeric complex containing two [2Fe-2S]²⁺ clusters matures the CIA machinery component anamorsin by transferring both clusters to the CIAPIN1 domain of apo anamorsin to generate [2Fe-2S] anamorsin. Both complexes, i.e. the homodimeric GRX3 and the heterotrimeric GRX3-BOLA2₂ complexes containing two [2Fe-2S]²⁺ clusters, can thus be active players, at the cellular level, for maturing anamorsin. The present data also suggest that the mechanism of cluster transfer relying on the interaction between the N-terminal domains of anamorsin and GRX3²⁴ is common to both cluster transfer processes, i.e. from [2Fe-2S]₂ GRX3₂ to apo anamorsin and from [2Fe-2S]₂ GRX3-BOLA2₂ to apo anamorsin.

Now, the arising question is whether and how the cells can select the homodimeric [2Fe-2S]² GRX3² *vs.* the heterotrimeric [2Fe-2S]² GRX3-BOLA2² complex to mature anamorsin. Assuming that the interaction between apo BOLA2 and apo GRX3 is physiologically relevant, the relative cellular levels of GRX3 and BOLA2, which can be regulated by cellular conditions (i.e. aerobic *vs.* anaerobic cellular growth, oxidative stress etc.), should select which of the two complexes mature anamorsin. The sensitivity of Fe/S protein biogenesis pathways towards oxygen and/or oxidative stress has been observed for members of both bacterial and eukaryotic Fe/S cluster assembly machineries.³⁸⁻⁴² Recently, two new CIA yeast proteins were found to be specifically involved in the maturation of the ribosome-associated ABC protein Rli1 and this function is fundamental under oxidative stress cellular conditions, as in the absence of oxygen these factors can be overcome to some extent without losing cell

viability.⁴³ It might be possible therefore that the homodimeric [2Fe-2S]₂ GRX3₂ complex is operative in normal cellular conditions, while the heterotrimeric [2Fe-2S]₂ GRX3-BOLA2₂ complex works under oxidative stress. Consistently with this possible model, bacterial BolA proteins have been observed to be specifically required under aerobic and oxidative stress conditions. For instance, in *E. coli* BolA protein is upregulated under oxidative stress conditions⁴⁴ and in bacterial operons, BolA tends to occur not only with a monothiol glutaredoxin, but also with proteins involved in defense against oxidative stress.⁴⁵ Also in eukaryotic genomes, the presence of BOLAs strongly correlates with an aerobic metabolism.⁴⁶ Last but not least, [2Fe-2S]₂ GRX3-BOLA2₂ complex is stable in air, while the homodimeric [2Fe-2S]₂ GRX3₂ is oxidatively labile and is gradually degraded in air over a period of ~1 h.²¹ Thus, BOLA2 binding to GRX3 stabilizes [2Fe-2S]²⁺ Fra2-Grx3/4 complexes.¹¹

Conclusion

In conclusion, we propose that the GRX3-BOLA2 interaction might have a role in the CIA pathway. GRX3 and BOLA2 form indeed an apo complex composed by two molecules of BOLA2 bound to a molecule of GRX3. This species, upon binding two [2Fe-2S]²⁺ clusters, is able to transfer both of them to anamorsin, thus maturing it in one-step process, similarly to what we already showed for the holo homodimeric GRX3 form.²⁴ The [2Fe-2S]²⁺ cluster bound forms of GRX3 and GRX3-BOLA22 might thus act as [2Fe-2S]²⁺ cluster transfer components in the cytosol, in a similar way as monothiol glutaredoxins do in mitochondria.^{7,19,47,48}

Supporting Information

Four figures reporting: ¹⁵N backbone amide relaxation parameters of apo BOLA2; an overlay of ¹H-¹⁵N HSQC spectra of different constructs of apo GRX3 in the presence or absence of apo BOLA2; EPR spectra of [2Fe-2S]₂ GRX3-BOLA2₂ complexes obtained by chemical reconstitution and by mixing [2Fe-2S]₂ GRX3₂ with apo BOLA2; gel filtration of a 1:1 mixture of apo GRX3(GrxA/B)-BOLA2₂ and [2Fe-2S] anamorsin.

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Competing financial interests

The authors declare no competing financial interest.

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TOC graphic



Supplementary Information

Elucidating the molecular function of human BOLA2 in GRX3-dependent anamorsin maturation pathway

Lucia Banci^{1,2}, Francesca Camponeschi^{1,2}, Simone Ciofi-Baffoni^{1,2}, Riccardo Muzzioli^{1,2}

¹Magnetic Resonance Center CERM, University of Florence, Via Luigi Sacconi 6, 50019, Sesto Fiorentino, Florence, Italy

²Department of Chemistry, University of Florence, Via della Lastruccia 3, 50019 Sesto Fiorentino, Florence, Italy



Figure S1. ¹⁵N backbone amide relaxation parameters of apo BOLA2. ¹⁵N R₁, R₂, and ¹⁵N{¹H} NOE values versus residue number of apo BOLA2 obtained at 600 MHz and 298 K. Missing bars indicate Pro residues and the not observable backbone NHs of Met 1, Glu 2, Leu 49.



Figure S2. Apo GRX3 interacts with apo BOLA2 as the apo GRX3(GrxA/B) construct does. Overlay of ¹H-¹⁵N HSQC spectra of a 1:2 mixture between ¹⁵N labeled apo GRX3 and unlabelled apo BOLA2 (black), of a 1:2 mixture between ¹⁵N labeled apo GRX3(GrxA/B) and unlabelled apo BOLA2 (red), and of ¹⁵N labeled Trx domain of GRX3 (green).



Figure S3. EPR spectra monitoring the cluster properties upon GRX3 and BOLA2 interaction. EPR spectra at 45 K obtained mixing [2Fe-2S]₂ GRX3₂ and apo BOLA2 in a 1:4 ratio (black) and of the chemically reconstituted heterotrimeric apo GRX3/BOLA2₂ complex (red) after anaerobic reduction with stoichiometric sodium dithionite and rapid freezing in 50 mM phosphate, 5 mM GSH, 5 mM DTT buffer at pH 7, 10% glycerol. The g-values are indicated.



Figure S4. The Trx domain of GRX3 is essential for complex formation between GRX3-BOLA2₂ and anamorsin. Analytical gel filtration (Superdex 200 10/300 increase column) chromatograms of apo BOLA2 (red) and [2Fe-2S] anamorsin (green), of apo GRX3(GrxA/B) (black) and of a 1:1 mixture between apo GRX3(GrxA/B)-BOLA2₂ and [2Fe-2S] anamorsin (blue). The apparent molecular masses are reported at the top of each chromatographic peak.