Review Article

Computational approaches for de novo design and redesign of metal-binding sites on proteins

Gunseli Bayram Akcapinar and Osman Ugur Sezerman
Department of Statistics and Medical Informatics, School of Medicine, Acibadem University, Istanbul, Turkey

Correspondence: Osman Ugur Sezerman (Ugur.Sezerman@acibadem.edu.tr)

Metal ions play pivotal roles in protein structure, function and stability. The functional and structural diversity of proteins in nature expanded with the incorporation of metal ions or clusters in proteins. Approximately one-third of these proteins in the databases contain metal ions. Many biological and chemical processes in nature involve metal ion-binding proteins, aka metalloproteins. Many cellular reactions that underpin life require metalloproteins. Most of the remarkable, complex chemical transformations are catalysed by metalloenzymes. Realization of the importance of metal-binding sites in a variety of cellular events led to the advancement of various computational methods for their prediction and characterization. Furthermore, as structural and functional knowledgebase about metalloproteins is expanding with advances in computational and experimental fields, the focus of the research is now shifting towards de novo design and redesign of metalloproteins to extend nature's own diversity beyond its limits. In this review, we will focus on the computational toolbox for prediction of metal ion-binding sites, de novo metalloprotein design and redesign. We will also give examples of tailor-made artificial metalloproteins designed with the computational toolbox.

Introduction

Function of a protein and its interactions with other molecules such as proteins, nucleic acids are determined by protein's 3D structure. There are over 124000 protein structures deposited in the Research Collaboratory for Structural Bioinformatics (RCSB) Protein Databank (PDB) by June 2016 [1]. In 2012, Yu and colleagues estimated that approximately one-third of the proteins deposited in the PDB database contained metal ions essential for function [2]. In cells, 30–40% of the proteins depend upon at least one metal ion to carry out their biological functions [3,4]. It has been well known that regions or residues of proteins that interact with the metal ligands are very well conserved both in sequence and in structure [5,6].

In many biological systems, proteins that require a metal ion to carry out their physiological function, aka metalloproteins, are very widespread and they perform a variety of functions as storage and transport proteins, enzymes, regulators of gene expression and signal transduction cascade proteins [7-10]. Metalloproteins have been in the focus of the biological research for many years. This intense focus could be traced back to the early X-ray crystallography studies that led to the discovery of presence of a metat atom (iron) along with 3D structure of myoglobin [11,12] and haemoglobin proteins [13].

Metal ions can be crucial for protein structure and function. Proteins are involved in a variety of cellular events and catalytic reactions. Chemical nature of the side chains of amino acids that made up proteins provides them with limited proportion of the chemical functionality seen in nature. On the other hand, association of proteins with cofactors such as small organic molecules, single metal ions or clusters with
metal and non-metal atoms granted these proteins with a diversity of functions [12]. Metal ions are involved in nucleophlic catalysis events, in induction of the conformational changes in proteins, in electron transport, in folding and stabilization of protein structures [14]. Metalloproteins play crucial roles in development and progress of a number of diseases including brain diseases such as prion, amyotrophic lateral sclerosis (ALS), Parkinson’s and Alzheimer’s where an effective treatment and cure is still missing [15]. They are also implicated in apoptosis and aging [16,17]. In plants and other microorganisms, they improve metal-adsorption capacities [18]. Well-characterized plant metalloproteins such as metallothioneins and phytochelatins are involved in the uptake of the essential micronutrients required for plant metabolism and detoxification of heavy metal ions [19]. Many metalloproteins in plants are involved in the electron transport chain of the photosynthetic machinery [20,21]. Furthermore, metal ions play a crucial role in protein folding and stability [22-24]. Realization of the importance and involvement of metal-binding sites in a diversity of cellular processes and functions, led to the advancement of various computational tools and algorithms for prediction, identification and engineering of these sites. These advances present us new means to understand the biological function of a protein; to decipher the underlying mechanisms of protein folding and stability and thus improve protein function and stability. An active research area was already established with studies involving de novo design, redesign, prediction, optimization and stabilization of protein structure. It has been almost 30 years since the introduction of a helical protein designed from the first principles [25]. In this era, focus is now shifting towards introduction of novel functions to proteins; protein stabilization by de novo protein design or redesign; use of metal cofactors. Subsequently, this approach extends the nature's repository of protein structure and function beyond its natural limits.

In this review, we will focus on the computational tools for the prediction of metal-binding sites on a protein and emphasize the computational tools that are used in de novo design and redesign of metal ion-binding sites. We will also highlight some noteworthy artificial metalloproteins that were designed using the developed computational tools.

### Computational tools for de novo design and redesign of metalloproteins

Unprecedented increase in the number of protein sequences and structures deposited in the public databases due to advances in genome-sequencing technologies and experimental methods for structure determination, poses a great challenge for researchers in terms of prediction of the biological functions of the deposited proteins. Since metal ions play crucial roles in many biological processes, their presence in protein structures reveals essential information about protein's inherent function(s). Owing to the fact that determination of structural and functional features of a protein such as metal ion-binding sites using experimental methods is still challenging because of the problems related to cost, time and automation of processes, there is an increasing demand for prediction of those features using computational methods. Use of computational methods for the prediction of metal ion-binding sites not only contributes to expansion of the existing knowledgebase, but also aids de novo metalloprotein design and redesign by providing necessary structural information about the metal co-ordination environment. This information is used in the design of tailored and stable metalloproteins with improved functions. Therefore, prosperity of a metalloprotein design largely depends on the quantity and quality of available information on metal ion-binding sites, protein scaffolds and physiochemical rules that direct the folding of a polypeptide into a functional protein [26,27].

### Computational prediction of metal ion-binding sites

Metal ion-binding sites in proteins exhibit a wide range of diversity. In some proteins, backbone oxygen and nitrogen atoms are involved in metal ion binding, whereas in some proteins side chain oxygen, nitrogen and sulfur atoms are involved. Metal ions were also found to be selective in their binding to their respective ligands. In a 2012 paper, Yu and colleagues used 1109 metal ion-binding polypeptides and predicted the metal ion-binding sites and verified that metal ions preferentially bind to certain residues on the protein [2]. For Ca^{2+}, favoured amino acids were predicted to be D, E, N and G; for Cu^{+} H; for Mg^{2+} D and E; for Fe^{3+} H, D, E, C and Y. Residues that reside partially within 3.5 Å of the metal ion were considered as metal ion-binding residues. As the metal ion-binding sites diversify, correct identification of the ligands in metal ion-binding sites with experimental methods becomes more problematic without prior knowledge of the identity of the bound metal ion since current high-throughput methods based on X-ray absorption spectroscopy can only identify its presence but cannot identify the residues involved in metal ion binding [28,29]. Besides, most of the computational tools for metal ion-binding site prediction use the information mostly derived from crystals of the metal ion-bound form of the proteins, holo (metal ion-bound) forms whereas a majority of the recently resolved structures are from the apo (metal ion-free) forms [30]. Moreover, metal ion binding is a dynamic
process that often results in structural rearrangements of the residues in binding pockets [31,32]. Additionally, in some cases, a metal ion is buried in a prosthetic group, tightly bound, specific non-polypeptide unit required for the biological function of some proteins, such as haem that consists of a central Fe (II/III) atom in a protoporphyrin ring [13] as in haemoglobin structure (Figure 1). Consequently, prediction and identification of metal ion-binding sites pose a much more complex problem that cannot be solved solely based on simple geometrical criteria or sequence information. Therefore, different computational methods and algorithms have been introduced for the prediction of metal ion-binding sites and the residues involved in this process. Moreover, prediction of most probable sites of metal ion-binding motifs in a newly identified protein would facilitate redesign of that site to better accommodate a metal to modify its structural and functional properties. After prediction and comparison with known motifs, this knowledge would enable establishment of a metal ion-binding site library that could be utilized for protein engineering purposes.

Most computational methods rely on the sequence information and they target the identification of metal-binding motif or motifs in well characterized metal ion-binding proteins. These motifs are then used to search for similar patterns in newly identified proteins. Machine learning methods have also been introduced to the problem only recently [7,33-37]. Programs such as MetalPredator [38], MetalDetector v2.0 [39], SeqCHED Server [34], ZincFinder [35], SVMProt [36,37] employ sequence based approaches for the prediction of the metal ion-binding sites. For instance MetalDetector v2.0 [39] uses cysteine and histidine residues and therefore focuses only on transition metals, haem and Fe/S clusters for the prediction, by predicting the metal-bonding state of these residues and number of bound metal ions. On the other hand, MetalPredator [38] utilizes two libraries of Hidden Markov Model profiles representative of Pfam domains and structural motifs that bind Fe/S clusters. Some prediction software such as FINDSITE-metal [40], mFASD [41], TEMSP [42], FEATURE metal scanning [43-45] utilize structural information for prediction of the metal ion-binding sites. A few of them such as MetSite [46], 3DLigandSite [47], Fragment Transformation Method [2] combine information from both protein sequence and structure. Use of ab initio methods as in IonCom [48] server has also been recently realized. Several helper tools such as MetalS2 [49], MetalS3 [50], FindGeo [51], CheckMyMetal (CMM) [52] were developed for pairwise structural alignment, database mining of metal-binding sites, determination of metal co-ordination geometry, validation of metal-binding sites in PDB structures respectively. Software developed for the prediction of metal ion-binding sites, their rationale and web links are summarized in Table 1. Although these servers provide a large selection of different algorithms for the prediction, some of them are only limited to the prediction of a single metal (i.e. MetalPredator and MetalDetectorv2.0 only predict Fe–S cluster-binding sites whereas FEATURE metal scanning tool, TEMSP, ZincFinder predict Zn$^{2+}$-binding sites, the rest of the servers provide a diverse range of metals that are associated with metalloproteins). Databases such as MetalPDB [4], MetalMine [53], Metal Interactions in Protein Structures (MIPS) [54], MESPEUS.10 [55], COMe [56], MetLigDB [57] as listed in Table 2, encompass information on both sequence motifs that define metal ion-binding sites and structure of metal ions and their corresponding residues that are deposited in various sequence and structural databases.

Figure 1. Structure of chain C of cross-linked carbonmonoxy haemoglobin A from Homo sapiens (left) Haem-coordinating residues are shown on the right. Haem group (red) with bound Fe (yellow) and carbonmonoxy (black). PDB ID: 1SDK.
Table 1 Software designed for metal ion-binding site prediction

<table>
<thead>
<tr>
<th>Software</th>
<th>Short Description</th>
<th>Ref.</th>
<th>Link</th>
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<tr>
<td>FINDSITE-metal</td>
<td>Sequence and structure based</td>
<td>[40]</td>
<td><a href="http://cssb.biology.gatech.edu/findsite-metal">http://cssb.biology.gatech.edu/findsite-metal</a></td>
</tr>
<tr>
<td>MetalPredator</td>
<td>Sequence based</td>
<td>[38]</td>
<td><a href="http://metalweb.cerm.unifi.it/tools/metalpredator/">http://metalweb.cerm.unifi.it/tools/metalpredator/</a></td>
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<tr>
<td>MetalDetector v2.0</td>
<td>Sequence based</td>
<td>[39]</td>
<td><a href="http://metaldetector.dsi.unifi.it/v2.0/">http://metaldetector.dsi.unifi.it/v2.0/</a></td>
</tr>
<tr>
<td>SeqCHED Server</td>
<td>Sequence based</td>
<td>[34]</td>
<td><a href="http://login1.weizmann.ac.il/~ronenle/Web/SeqCHED/">http://login1.weizmann.ac.il/~ronenle/Web/SeqCHED/</a></td>
</tr>
<tr>
<td>MetSite</td>
<td>Sequence and structure based</td>
<td>[46]</td>
<td><a href="http://bioinf.cs.ucl.ac.uk/structure/">http://bioinf.cs.ucl.ac.uk/structure/</a></td>
</tr>
<tr>
<td>mFASD</td>
<td>Structure based</td>
<td>[41]</td>
<td>source code: <a href="http://staff.ustc.edu.cn/~liangzhi/mfasd/">http://staff.ustc.edu.cn/~liangzhi/mfasd/</a></td>
</tr>
<tr>
<td>SVMProt</td>
<td>Sequence based</td>
<td>[37,36]</td>
<td><a href="http://obkd2.nus.edu.sg/cgi-bin/svm-prot/svmprot.cgi">http://obkd2.nus.edu.sg/cgi-bin/svm-prot/svmprot.cgi</a></td>
</tr>
<tr>
<td>3DLigandSite</td>
<td>Sequence and structure based</td>
<td>[47]</td>
<td><a href="http://www.sbg.bio.ic.ac.uk/3dligandsite/">http://www.sbg.bio.ic.ac.uk/3dligandsite/</a></td>
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Table 2 List of databases for metalloproteins, metal ion-binding site motifs and structural information on metal ion-binding sites

<table>
<thead>
<tr>
<th>Databases</th>
<th>Reference/Link</th>
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<tr>
<td>MetLigDB</td>
<td>[57] <a href="http://silver.sejong.ac.kr/MetLigDB">http://silver.sejong.ac.kr/MetLigDB</a></td>
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Computational design tools, strategies for de novo design and redesign of metalloproteins

Many of the biological and chemical reactions that establish the foundations of life such as water oxidation, carbon dioxide reduction, nitrogen fixation, photosynthesis require involvement of metalloproteins or metalloenzymes [58]. Chemical transformations achieved by metalloproteins are diverse [59]. Despite the diversity of these reactions, we are still far from completely understanding the principles that govern these metalloproteins when performing their functions. Much effort has been put to fully mimic or exploit these processes. As de novo protein design and redesign methods have been advancing for the past decades, application of these methods to artificial metalloprotein and metalloenzyme design and redesign is inevitable. Previous research has created fully functional de novo designed metalloproteins that exhibit activity for a variety of reactions ranging from ester and organophosphate hydrolysis [60,61] to nitric oxide reduction [60,61].

In metalloproteins, metal ions either interact with amino acid side chains or accompanying residues. Most of the metal-binding sites are known to be promiscuous and can accommodate non-native metals with similar properties [62]. Redesign of metalloproteins usually involves modification of a well-characterized and well-known native protein through engineering or introduction of metal-binding ligands to protein structure to accommodate non-native metals. For de novo design of metalloproteins, protein structures should be designed from scratch starting from the amino acid sequences. Residues for metal ion binding should also be introduced to the structure [63]. Both methods require correctly folded and functional final protein structures with introduced metal ions or cofactors. De novo design approach is more complex since physiochemical rules that drive the protein folding [64] have not yet been deciphered completely. Moreover, “inverse folding problem” adds another layer to this complexity. De novo protein design uses the “inverse folding problem”, which states that different combinations of amino acid sequences can fold into the same 3D structure and resulting proteins can perform the same function [65,66].
In protein design, we start with a rigid or flexible backbone structure and try to determine a sequence that would fold into this structure. As a variety of sequences fold into the same structure, there is degeneracy. Therefore, the accuracy and availability of the template structures for protein design has a huge affect in the success of the final design.

The current collection of metalloproteins is largely coming from nature’s own repository. This repository is now being diversified by de novo design and redesign approaches with the aid of the knowledgebase of metal ion-binding sites. A typical workflow for this procedure is shown in Figure 2. Successful application of de novo design and redesign approaches create a diversity of advanced proteins with either new or modified functions. Additionally, through the application of these methods, an excellent platform is formed to study structure–function relationship and protein folding problem further.

Construction of artificial or redesigned metalloproteins requires detailed information on the nature of the metal ion-binding motifs, how the metal ions are co-ordinated (ligand–metal interactions and binding pocket geometry, geometrical arrangements, redox states) and finally a detailed information on the structure–function of the scaffold protein that would accommodate the metal ion. This, in turn, requires the use of state of the art computational tools for design and experimental techniques for analysis and characterization. A variety of deterministic, stochastic and probabilistic methods have been applied for de novo protein design [67]. With the recent advances in the computational
field (high performance computing, better algorithms, faster processors), backbone flexibility is also incorporated into the calculations [68].

A compelling number of programs such as Protein WISDOM [68], HostDesigner [69], METALSEARCH [70], CORE [71], RosettaDesign [72], EvoDesign [73], RosettaBackrub [74], IPRO (OptGraft procedure) [75], PyRosetta [76] has been in use for de novo protein design (Table 3). These software provide in silico workbenches for the design of novel, stable and functional proteins. Most of these programs use force fields to represent molecular interactions and to search for sequences with lowest energy in the target fold since the minimum free energy conformation of a protein is considered as the native fold of the protein.

Although all of them have been used for de novo design of metalloproteins, only three of them, HostDesigner, METALSEARCH and IPRO (OptGraft) are specific for de novo design or redesign of metalloproteins. HostDesigner is a combinatorial chemistry-based program that uses two complementary algorithms LINKER and OVERLAY for creation of a number of candidate structures and to discover metal ion receptors in these structures [69]. Two user-defined complex fragments are connected with linking fragments deposited in the library and structures are built by LINKER. Next, a single user-defined complex structure is built by OVERLAY through the superimposition of the linking fragments on to this structure.

METALSEARCH, being one of the earliest design programs, creates lists of four residues that might form tetrahedral metal ion-binding sites if amino acids in the native protein were replaced by cysteine or histidine residues. Program also provides dihedral angles of the amino acids and the co-ordinates of the predicted metal ion as output [70]. OptGraft is a computational program that is present in IPRO suite of programs and it is specifically designed for transferring a binding site on to an existing protein scaffold. For this purpose, possible binding pocket placement combinations are exhaustively explored using mixed-integer linear optimization and a ranked list of possible designs that fit the geometric criteria and orientation of the native binding pocket is generated. Moreover, this procedure also assesses the impact of the new pocket on the protein structure and if there is a potential distortion in the structure, small mutations in the neighbouring residues are introduced to counteract their probable distortional effects [75].

Applications
Computational design of novel redox centres
METALSEARCH was used for de novo design of a rubredoxin-like Fe site [77], for the introduction of zinc-binding sites to the designed four helix bundle protein α4 and to the Ig-binding B1 domain of the Streptococcal protein G [78, 79]. It has been long known that iron ion in rubredoxin proteins is co-ordinated by four cysteine residues. Farinas et al. [77] designed a redox centre that mimics rubredoxin’s into the Ig-binding B1 domain of the Streptococcal protein G by taking into account the backbone movements while maintaining the structural integrity and stability of the B1 protein. For this purpose, they first used METALSEARCH to identify potential metal ion-binding sites that are capable of tetrahedral co-ordination in the B1 protein structure. Contrary to previous work of the same group [78, 79], they made an attempt to use a more realistic backbone flexibility description in the computational part. As an input to the METALSEARCH, rather than using the averaged NMR structure, researchers selected 6 randomly calculated NMR structures out of 60 that reflected a more realistic backbone flexibility. Co(II) and Cd(II) bound designed variants’ stability and integrity were confirmed with spectroscopic methods. Fe(II) complex of the designed
variant was produced and analysis and characterization of the rubredoxin centre revealed that the mutant rubredoxin centre mimicked oxidized rubredoxin. Nanda et al. [80] also managed to design a redox active minimal β protein scaffold (RM1) that exhibit rubredoxin activity through formation of a stable, redox-active 4-Cys thiolate Fe(II/III) site. The designed complex was further shown to be stable through repeated cycles of oxidation and reduction even in the presence of an oxygen containing environment.

In another design approach, computationally designed metalloprotein using an in-house developed CORE software [71] that functions as an artificial redox centre was produced with the ability to mimic photosynthesis [81]. An antiparallel four helix bundle containing two helix–loop–helix peptides connected through a disulfide bridge was used as the scaffold. This scaffold is functionalized through histidine-coordinated Ruthenium(II) Bipyridine (Ru(bpy)2) and haem cofactors by engineering of the binding sites. The exterior hydrophilicity of the bundle was provided by the salt bridges formed primarily between glutamic acid and lysine residues, whereas identity of the hydrophobic core residues inside the helices were determined using the CORE software. An exemplary structure of Ru(bpy)2(mbpy)-modified bovine adrenodoxin protein was shown in Figure 3 to show the metal co-ordination. The backbone structure that was used as an input to the software contains alanine in all positions with the exception of the exterior hydrophilic residues and cofactor-coordinating residues. Software predicts the optimum hydrophobic core residues based on the protein thermal stability and co-operativity. Consequently, the resulting protein mimics photosynthesis following photo-excitation participating in multiple oxidation and reduction cycles with exogenous electron acceptors and donors.

**Computational design of artificial metalloproteins and metalloenzymes**

Computational protein design software Rosetta Design, is a highly versatile tool that can be exploited for analysis of protein–metal interactions in the design of artificial metalloproteins. Recently, this software was used to improve the catalytic performance of an artificially designed metalloenzyme and to design a promiscuous metalloprotein where an incorporated unnatural amino acid (2,2’-bipyridin-5yl) alanine (Bpy–Ala) residue co-ordinates the metal ion with octahedral co-ordination geometry [80,81]. Heinisch et al. [82] adopted a computational design approach for the genetic optimization of a first-generation artificial metalloenzyme using this software. Previous research produced an assembly of Noyori-type pianostool complex [(5-Cp*)Ir(pico)Cl] in a native human carbonic anhydrase II enzyme that is capable of asymmetric transfer hydrogenation of a salsolidine precursor (a cyclic imine) with moderate enantioselectivity [83] and combinations of these first-generation artificial metalloenzymes with naturally occurring enzymes were also successfully utilized in redox cascade reactions [84]. The crystal structure of the complex revealed that the metal-coordinating moiety was not fully occupied, which led to a modest activity and selectivity. Therefore, RosettaDesign [72] was used to improve cofactor-binding sites and the second co-ordination sphere around cofactor...
in the first generation artificial metalloenzymes. The software produced four scaffold variants that have mutations for protein backbone stabilization and that enable hydrophobic cofactor burial. The produced designs not only exhibited increased affinity towards the cofactor but also increased activity, turnover number and enantioselectivity with respect to the first-generation artificial metalloenzyme. Mills et al. [85] employed a similar design approach with the use of RosettaMatch algorithm [86] and RosettaDesign [72] for the co-ordination of metal ions by a metalloprotein through the use of unnatural amino acids. For this purpose, an active site capable of catalysing oxidative ring opening of catechol substrates was designed. Four theoretical enzyme active sites, also referred as “theozymes” were constructed including unnatural amino acid Bpy–Ala, catechol containing dopamine molecule, a tyrosine and a histidine ligand for metal co-ordination. RosettaMatch was adopted for the identification of a set of backbone positions in native scaffolds that could accommodate theozyme geometrical restraints. RosettaDesign was then employed to introduce additional stabilizing interactions for the Bpy–Ala complex to the RosettaMatch output. An additional filtering step was applied to find the optimum structures. Although 13 genes that encode artificial metalloproteins are designed and synthesized, only five of the designed proteins yielded soluble, full length proteins in the presence of the unnatural amino acid. These proteins were shown to be promiscuous and bind to divalent cations including Co²⁺, Zn²⁺, Fe²⁺ and Ni²⁺. Moreover, structural analysis via X-ray crystallography revealed that the designed proteins exhibit only slight deviations from the designed models.

Additionally, the OptGraft procedure in IPRO software suite was used by Fazelinia et al. [75] to graft a calcium-binding pocket of thermitase protein to the first domain of CD2 protein (a rat cell adhesion protein that does not bind calcium). Minimally perturbed structures were created by identification of suitable locations for the grafting using OptGraft procedure. Candidate residues that would ensure the desired spatial restraints for the binding pocket were determined using CHARMM energy calculations. Designs with higher stability were further selected. Novel CD2 variants were shown to exhibit higher affinities for terbium and selective binding to calcium.

Researchers also developed in-house programs for the de novo design of metal ion-binding sites. Zhu et al. [87] managed to incorporate computationally engineered zinc-binding sites to de novo designed DS119 protein with βαβ fold with 1:1 stoichiometry using an in-house developed program. Likewise, Yeung et al. [88] used myoglobin as scaffold to introduce nitric oxide reductase activity by grafting three histidines and one glutamate residues that would accommodate a haem and non-haem Fe₆ centre. They followed a molecular dynamics (MD)-based modelling approach for the in silico introduction of mutations to myoglobin scaffold and to evaluate the ability of the designed metal ion-binding site to bind an iron ion. To achieve this three in silico mutations (L29H, F43H and V68E) were introduced into sperm whale myoglobin structure using the VMD software suite. Additionally, molecular dynamics simulations (NVT ensemble, 310 K, 10000 steps, 1 fs/step) with NAMD software were performed to estimate the ability of the mutant protein to bind Fe atom through the engineered metal ion-binding site. The triple mutant of the sperm whale myoglobin was constructed and mutant protein was expressed in Escherichia coli. The mutant protein was further purified and characterized. Finally, the tailor-made protein was shown to exhibit nitric oxide reductase activity.

Computational design of metal-templated interfaces

In 2010, Salgado et al. used Metal Templated Interface Redesign (MeTIR) approach to gain insights about the role played by metal co-ordination via metal co-ordinated nucleation process in the formation of early protein folds and complexes throughout evolution [89]. For this purpose, they have redesigned a monomeric protein, namely cytochrome bc₅₆₂ with non-self-associating surface and haem groups in its structure to obtain a variant that exhibits Zn co-ordinated self-association properties. A second redesign step was introduced to Zn co-ordinated D2-symmetrical tetramer to engineer the complex further into a complex that exhibits self-assembly in the presence or absence of metals. Mutations were introduced only into the residues on the surface of cytochrome bc₅₆₂ to allow easy tracing of the structure with crystallographic methods and to preserve the overall fold. Residues inside the protein (with low solvent accessible surface are (SASA) values), residues contacting heme groups or Zn atoms, residues that were involved in side chain–main chain H-bonding interactions remained and they were not selected for mutations. A variant of RosettaDesign algorithm was employed for the rotamer optimization of the selected residues to find minimal number of mutations that make a maximum impact towards the self-assembly of the monomeric protein. The energy and packing scores obtained, ranked and position of candidate residues with respect to the undesirable residues were evaluated to find optimal residues at the interface for redesign.

Knowledge-driven metalloprotein designs

In some cases, computational programs were not used for the design. Instead, designs were based on previous information derived from sequence, structure and function of proteins, previous literary knowledgebase and intuition.
of the researchers [90–92]. In most of the cases, previously designed protein scaffolds are modified to engineer the binding pockets [63, 88, 91, 93–96]. In one recent case, noble metals were introduced to iron-binding sites of the haem proteins such as myoglobin and the designed protein was subjected to directed evolution [97]. First, apo-form of haem proteins were expressed in *E. coli* using minimal media devoid of Fe for the minimization of haem biosynthesis, at low temperature to alleviate the instability of the apo-form. Myoglobin from *Physeter macrocephalus* and cytochrome P450 BM3h from *Bacillus megaterium* were found to be overexpressed with or without mOCR stability tags in relatively high yields. These haem lacking apo-forms were purified and holo-forms of the proteins were reconstituted with noble metal-porphyrin IX cofactors. Next, eight myoglobin variants with mutations in the axial ligand position (His31) were expressed in apo-form and then reconstituted with various noble metal-porphyrin IX cofactors. Consequently, the resulting proteins had the capability to catalyse the functionalization of C—H bonds to form C—C bonds by carbene insertion and add carbenes to both β-substituted vinylarenes and unactivated aliphatic α-olefins.

Another approach involves the use of biotin–streptavidin (Sav) system to anchor metal complexes to a specific location in a protein [59, 98–100]. This methodology was applied to artificial cupredoxin synthesis recently [59]. A systematic modulation of the primary and secondary co-ordination spheres of artificial copper proteins was performed. Biotinylated copper complexes within a cysteine containing variant of Sav have exhibited properties similar to native cupredoxins. Fine tuning of the position of the Cu centres inside the modified Sav was achieved by changing length of the linker between biotin and the copper complex. A similar approach was employed by Heinisch et al. [82] and Quinto et al. [98] in the design of an artificial metalloenzyme that exhibits activity as an NAD(P)H dependent transfer hydrogenase recently. To this end, a biotinylated iridium cofactor was introduced to variants of Sav scaffold that resulted in the successful activity of the artificial metalloenzymes.

**Conclusions and future perspectives**

Metalloproteins advanced through millions of years of protein evolution. Considering this huge evolutionary history, we are still far from perfecting our *de novo* design attempts. Computational design offers an alternative but complementary route to directed evolution experiments for the introduction of novel functionalities. Learning from nature’s toolbox and rules that drive protein evolution, we can improve our designs [95]. Considerable efforts have been put for the design and production of new artificial metalloproteins [85]. However, there are still challenges such as achieving accuracy at the atomic level since flexible polar amino acid side chains form metal-binding sites through alternative ways. Computational design principles can be used to design a library of variants that could co-ordinate a metal cofactor. Additionally, it can be used to reduce number of the variants for experimental characterization [82]. Initially, the metalloprotein design was focused on the design and formation of mono-metal centres on the designed proteins. This horizon was expanded by the introduction of di-metal centres [91, 101] and current research is focused on building multi-metal cluster containing artificial metalloproteins [58] and specifically metalloenzymes since most remarkable and complex chemical transformations are achieved through the use of multi-metal centres [59]. As these horizons expand, design of different scaffolds from first principles that would host different metal centres is also needed. Moreover, directed evolution should complement *de novo* protein design efforts to produce artificial metalloproteins with improved properties. Although an increasing number of graphical user interfaces for protein design and metal ion-binding site prediction are being developed, use of computational programs for the design of artificial metalloproteins does not follow the same pace. Therefore, there is an emerging need to develop user-friendly tools specific for metalloprotein design and engineering.

**Competing interests**
The authors declare that there are no competing interests associated with the manuscript.

**Abbreviations**

BIM, bioinorganic motif; bpy, bipyridine; Bpy–Ala, (2,2′-bipyridin-5yl) alanine; CHARMM, Chemistry at Harvard Macromolecular Mechanics; CMM, Check My Metal; fs, femtosecond; MIPS, metal interactions in protein structures; PDB, Protein Databank; Ru(bpy)2, ruthenium(II) bpy; Sav, Streptavidin; SASA, Solvent Accessible Surface Area.

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