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Authors: Vincent Louis Pecoraro; Leela Ruckthong, Ph.D.; Jeanne A. Stuckey, Ph.D.

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# How Outer Coordination Sphere Modifications Can Impact Metal Structures in Proteins: A Crystallographic Evaluation 

Leela Ruckthong ${ }^{1,2}$, Jeanne A. Stuckey ${ }^{3,4}$ and Vincent L. Pecoraro ${ }^{* *}$<br>${ }^{1}$ Department of Chemistry, University of Michigan, Ann Arbor, Michigan 48109, USA<br>${ }^{2}$ Department of Chemistry, Faculty of Science, King Mongkut's University of Technology, Thonburi (KMUTT), Bang Mod, Thung Khru, Bangkok, 10140, Thailand<br>${ }^{3}$ Life Sciences Institute, University of Michigan, Ann Arbor, Michigan 48109, USA<br>${ }^{4}$ Department of Biological Chemistry, University of Michigan, Ann Arbor, Michigan 48109, USA<br>*e-mail: vlpec@umich.edu


#### Abstract

A challenging objective of de Novo metalloprotein design is to control of the outer coordination spheres of an active site to fine tune metal properties. The well-defined three stranded coiled coils, TRI and CoilSer peptides, are used to address this question. Substitution of Cys for Leu yields a thiophilic site within the core. Metals such as $\mathrm{Hg}(\mathrm{II}), \mathrm{Pb}(\mathrm{II})$ and As (III) result in trigonal planar or trigonal pyramidal geometries; however, spectroscopic studies showed $\mathrm{Cd}(\mathrm{II})$ formed $3-$, 4 - or 5 -coordinate $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{OH}_{2}\right) \times$ (where $x=0-2$ ) when the outer coordination spheres were perturbed. Unfortunately, there has been little crystallographic examination of these proteins to explain the observations. Herein, we compare the high-resolution x-ray structures of apo- and mercurated proteins to explain the modifications that lead to metal coordination number and geometry variation. It reveals that Ala substitution for Leu opens a cavity above the Cys site allowing for water excess, facilitating $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{OH}_{2}\right)$. Replacement of Cys by Pen restricts thiol rotation, causing a shift in the metal binding plane that displaces water, forming $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$. D-Leu, above the Cys site, reorients the side chain towards the Cys layer diminishing the space for water accommodation yielding $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$, while D-Leu below opens more space, allowing for equal $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{OH}_{2}\right)$ and $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{OH}_{2}\right)_{2}$. These studies provide insights on how to control desired metal geometries in metalloproteins using coded and non-coded amino acids.


Keywords: De novo protein engineering, D-amino acids, Non-natural amino acids, Metalloprotein Engineering

## Introduction

We have employed de novo designed proteins containing thiol residues to chelate metals in geometries that are relevant for understanding heavy metal sequestration in sulfur rich sites of human chaperones and metalloregulator proteins. ${ }^{1-10}$ Using the TRI-family peptides (sequences given in Table 1) we have established a well-defined scaffold using three-stranded coiled coil (3SCC) forming peptides that can evaluate metal binding within a hydrophobic core (Figure 1). These peptides are based on a heptad repeat approach in which hydrophobic leucine (Leu) residues at $\boldsymbol{a}$ and $\boldsymbol{d}$ positions generate the helical core and salt bridge interactions between $\boldsymbol{e}$ and $\boldsymbol{g}$ residues on the helical interface stabilize the aggregation state and a parallel orientation of helices. ${ }^{10-12}$ The substitution of Leu with cysteine (Cys) in one of the hydrophobic a or d positions generates a layer of three Cys residues forming a tri-sulfur chelating site. Previous reports have shown that the cysteine side chains in these apo-proteins are preorganized for binding metals into trigonal pyramidal geometries (i.e. $\mathrm{Pb}(\mathrm{II}) \mathrm{S}_{3}$ and $\mathrm{As}(\mathrm{III}) \mathrm{S}_{3}$ ), but are simply predisposed for encapsulating metals that are trigonal planar or pseudo-tetrahedral. ${ }^{13}$ In the preorganized systems, the ligands in the unbound state, which are directed toward the N -termini and helical core, remain almost in the same position upon metal
complexation. This is mainly because trigonal pyramidal geometry does not require the metal to bind in the same plane as the Cys sulfur atoms, but rather it may achieve the necessary bond lengths and angles when it is situated below the plane of coordinating atoms. However, predisposition of Cys occurs when the metal binding side-chains must rotate away from the helical core toward the helical interface, in order to increase space for metal binding within ( $\mathrm{Hg}(\mathrm{II})$ ) or close to ( $\mathrm{Zn}(\mathrm{II})$ ) the Cys plane. Since $\mathrm{Cd}(\mathrm{II})$ in a $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ environment most likely binds into a geometry similar to trigonal planar $\mathrm{Hg}(\mathrm{II})$ rather than trigonal pyramidal $\mathrm{Pb}(\mathrm{II})^{14,15}$, it is likely that Cys residues are predisposed rather than pre-organized toward trigonal planar $\mathrm{Cd}(\mathrm{II})$ sites in 3SCCs.

We have paid specific attention to $\mathrm{Cd}(\mathrm{II})$ binding to the TRIfamily peptides to understand coordination number control in $\alpha$ helical systems. ${ }^{1-10}{ }^{113} \mathrm{Cd}$ NMR, ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC, X-ray Absorption and UV-vis spectroscopies demonstrated that the incorporation of $\mathrm{Cd}(\mathrm{II})$ to the (TRIL16C) ${ }_{3}$ peptide generated a mixture of trigonal planar $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ and pseudo-tetrahedral $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right) .{ }^{6}$, ${ }_{8,16-23}$ Unlike $\mathrm{Hg}(I)$, which prefers linear or trigonal planar structures, $\mathrm{Cd}(\mathrm{II})$ easily accepts 4-coordination when an exogenous ligand is available. The evidence for the formation of a $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ clearly implies that Leu residues in the twelfth position provide a certain amount of space that allows water

Table 1. Peptide sequences

| Peptides |  | abcdefg | $\begin{gathered} \text { abbcdef g } \\ 9 \end{gathered}$ | $\begin{aligned} & \text { abbcdefg } \\ & 16 \quad 19 \\ & \hline \end{aligned}$ | $\begin{aligned} & \text { abbcdefg } \\ & 23 \end{aligned}$ | $\begin{aligned} & \text { abcdefg } \\ & 30 \end{aligned}$ | PDB ID |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| TRI | Ac-G | LKALEEK | LKALEEK | LKALEEK | LKALEEK | G-NH2 |  |
| TRIL16C | Ac-G | LKALEEK | LKALEEK | CKALEEK | LKALEEK | G-NH2 |  |
| TRIL12AL16C | Ac-G | LKALEEK | LKAAEEK | CKALEEK | LKALEEK | G- $\mathrm{NH}_{2}$ |  |
| TRIL12DLL16C | Ac-G | LKALEEK | LKA ${ }_{\text {d }}$ LEEK | CKALEEK | LKALEEK | G-NH2 |  |
| TRIL2WL16CL19 ${ }_{\text {D }}$ | Ac-G | WKALEEK | LKALEEK | $\underline{\underline{C}} K A_{\text {d }}$ LEEEK | LKALEEK | G-NH2 |  |
| CoilSer (CS) | Ac-E | WEALEKK | LAALESK | LQALEKK | LEALEHG | $-\mathrm{NH}_{2}$ |  |
| CSL16C | Ac-E | WEALEKK | LAALESK | CQALEKK | LEALEHG | $-\mathrm{NH}_{2}$ | $5 \mathrm{K92}{ }^{13}$ |
| CSL9PenL23H | Ac-E | WEALEKK | PenAALESK | LQALEKK | HEALEHG | $-\mathrm{NH}_{2}$ | $3 P B J^{28}$ |
| GRAND-CoilSer (GRAND-CS) | Ac-E | WEALEKK | LAALESK | LQALEKK | LQALEKK | LEALEHG - $\mathrm{NH}_{2}$ |  |
| GRAND-CSL16CL30H | Ac-E | WEALEKK | LAALESK | CQALEKK | LQALEKK | HEALEHG- $\mathrm{NH}_{2}$ | $5 \mathrm{KB0}{ }^{13}, 5 \mathrm{~KB} 1^{13}$ |
| GRAND-CSL12AL16C | Ac-E | WEALEKK | LAAAESK | CQALEKK | LQALEKK | LEALEHG- $\mathrm{NH}_{2}$ | $5 \mathrm{KB2}{ }^{13}$, 6EGO* |
| GRAND-CSL12DLL16C | Ac-E | WEALEKK | LAA ${ }_{\text {d }}$ LESK | CGALEKK | LQALEKK | LEALEHG - $\mathrm{NH}_{2}$ | 6EGL* |
| GRAND-CSL16CL19DL | Ac-E | WEALEKK | LAALESK | $\underline{C}$ QA $_{\text {d }}$ LEKK | LQALEKK | LEALEHG - $\mathrm{NH}_{2}$ | 6EGM ${ }^{*}$, 6EGN* |
| GRAND-CSL12DLL16CL19 ${ }_{\text {D }}$ L | Ac-E | WEALEKK | LAA ${ }_{\underline{D}} \underline{L} E S K$ | $\underline{\underline{C}} \mathrm{QA}_{\underline{D}} \underline{\underline{L}}$ EKK | LQALEKK | LEALEHG - $\mathrm{NH}_{2}$ |  |

Bold and underlined residues indicate substitutions.
N -and C-termini are capped by Ac and $\mathrm{NH}_{2}$ groups, respectively.

* This work
access above the metal site, allowing the $\mathrm{Cd}(\mathrm{II})$ site to have bound water $60 \%$ of the time. However, when the sterics were altered by replacing Leu with alanine (Ala), TRIL12AL16C gave $100 \% \mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right) .{ }^{16,17}$ An exclusive trigonal planar $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ could be formed with two strategies. First, a more sterically demanding analogue of Cys ( $\beta$-dimethyl cysteine also called penicillamine (Pen)) was incorporated in lieu of Leu in the sixteenth position. ${ }^{12,16}$ The $\operatorname{Cd}\left(\right.$ II)(TRIL16Pen) ${ }_{3}$ formed $100 \%$ $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$. Second, when the chirality of Leu in the twelfth position is inverted to D-Leu (TRIL12DLL16C) ${ }^{24}$ the branched side chain was proposed to reorient toward the C-termini of the 3SCC to block the space above the metal site. As predicted, a $100 \% \operatorname{Cd}(I I) S_{3}$ was achieved. Based on this observation, the alternate configuration of D-Leu has been varied in the outer coordination spheres around the metal center to investigate how the coordination numbers of $\mathrm{Cd}(\mathrm{II})$ can be controlled. Based on ${ }^{113} \mathrm{Cd}$ NMR and ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC, the replacement of L-Leu by D-Leu at the nineteenth position (TRIL2WL16CL19DL) led to $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ and $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}$ in a percentage ratio of $50: 50 .{ }^{25}$ The evidence of this new $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}$ species suggested that D-Leu potentially opens space below the metal site, thus the (TRIL2WL16CL19DL) ${ }_{3}$ contains two possible cavities both above and below the metal site at the same time. The incorporation of two D-Leu simultaneously above and below the metal site in the GRAND-CSL12dLL16CL19DL design, reduced the amount of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}$ by $20 \%$, while the $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ species increases to $70 \%$. ${ }^{25}$ D-Leu obviously shows potential to engineer the steric environments that affects the availability of space around the metal center, which consequently controls water access around the metal site. Despite the success in ${ }^{113} \mathrm{Cd}$ NMR and ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC characterizations of this $\mathrm{Cd}(\mathrm{II})$-bound peptide series, structural details of such modifications have not yet been revealed.

We have employed 3SCC CoilSer (CS) and GRAND-CoilSer (GRAND-CS) (Table 1) to act as crystallographic analogues in structural studies of these designs. ${ }^{11-13,21,22}$ Both peptides, differ by length, and contain a histidine (His) at the $\boldsymbol{f}$ position of the last heptad. This His located on the helical interface is critical for crystallization as it ligates to a $\mathrm{Zn}(\mathrm{II})$ ion along with glutamates from other trimers. The external $\mathrm{Zn}(\mathrm{II})$ sites facilitate the 3D
packing of trimers in lattice form. Spectroscopic studies have shown that the substitution of Leu with Cys in these
crystallographic analogues results in identical heavy metal binding properties as the TRI-family peptides. ${ }^{22}$ A number of X ray crystallographic structures in both apo- and metallated forms of these peptides have been reported. ${ }^{11-13,23-24}$ Unfortunately, even though great effort has been spent on optimizing the crystal growth conditions of the designs, to date no crystal structures of CS or GRAND-CS constructs have been isolated with bound $\operatorname{Cd}(I I)$. We believe that $\operatorname{Cd}(I I)$ was unable to bind to the protein under the crystal growth conditions due to the high affinity of this metal ion toward the oxygen-containing precipitants that are present (e.g. polyethylene glycol, glycerol and ethoxyethanol). Usually these materials are at concentrations much higher than the protein, so it is likely that the $\mathrm{Cd}(\mathrm{II})$-thiolate center cannot compete successfully for the metal in the presence of these concentrated oxygen ligands. Thus, direct observation of the $\mathrm{Cd}(\mathrm{II})$ within these 3SCCs under crystalization conditions has been unsuccessful.

To still be able to gain insight for the impact of modifying outer coordination sphere hydrophobic residues, we have utilized $\mathrm{Hg}(\mathrm{II})$ to represent a trigonal planar $\mathrm{Cd}(\mathrm{II})$. The $\mathrm{Hg}(\mathrm{II})-\mathrm{S}$ bond distance in $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$crystal structure (PDB ID: 5KB1) was reported to be $2.38 \AA^{13}$, which is in good agreement with an X-ray absorption result for $\mathrm{Hg}(\mathrm{II})(\text { TRIL16C })_{3}{ }^{-}\left(2.43 \AA\right.$ A). ${ }^{7}$ At the same time, the EXAFS results for the $\mathrm{Cd}(\mathrm{II})-\mathrm{S}$ bond distance for the trigonal planar $\mathrm{Cd}(\mathrm{II})$ (TRIL16Pen) ${ }_{3}{ }^{-}$is $2.46 \AA,{ }^{29}$ which leads one to predict the trigonal planar structures of $\mathrm{Hg}(\mathrm{II}) \mathrm{S}_{3}$ and $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ are similar. Thus, regardless of the metal size difference, the crystallographic $\mathrm{Hg}(\mathrm{II}) \mathrm{S}_{3}$ structures could be used to explain general characteristics of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$. In this study, we have also achieved a variety of crystal structures based on the sequences designed for $\mathrm{Cd}(\mathrm{II})$ studies. We have obtained the $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL} 12 \mathrm{AL} 16 \mathrm{C})_{3}{ }^{\text {² }}$, representing the TRIL12AL16C environment, to analyze the effect of Ala (twelfth position) above the metal site in comparison with the 12Leu packing of the known $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL3OH })_{3}{ }^{+13}$. The analysis will explain why $\mathrm{Cd}(\mathrm{II})(\mathrm{TRIL} 16 \mathrm{C})_{3}$ can form a mixture of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ and $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ centers, whereas $\mathrm{Cd}(\mathrm{II})(\mathrm{TRIL} 12 \mathrm{~A} 16 \mathrm{C})_{3}{ }^{-}$ results in a $100 \% \mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3} \mathrm{O}$. Moreover, the exclusion of water from the $\mathrm{Pen}_{3}$ site has been investigated using a combination
between $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}$and $[\mathrm{Hg}(\mathrm{II})]_{s}[\mathrm{Zn}(\mathrm{II})$ $\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}\right.$

$)]_{N}(\text { CSL9PenL23H })_{3}{ }^{{ }^{+}}$. We have crystallized apo-(GRANDCSL12dLL16C) $)_{3}$ and apo-(GRAND-CSL16CL19DL) $)_{3}$ to

Figure 1. General overview of GRAND-CS structure that contains a $\mathrm{Hg}(\mathrm{II}) \mathrm{S}_{3}$ binding site at the sixteenth position. Helical core residues are shown as sticks. Leucine residues in the $12^{\text {th }}$ and $19^{\text {th }}$ positions are shown in pink and orange, respectively. Cys residues in the $16^{\text {th }}$ position are colored green. $\mathrm{Hg}(\mathrm{II})$ is shown as a blue sphere. The Leu layer at the $12^{\text {th }}$ position and the interlayer between the $12^{\text {th }}$ and $16^{\text {th }}$ positions are defined as 'above' the metal binding site. The Leu layer at the $19^{\text {th }}$ position and the interlayer between the $16^{\text {th }}$ and $19^{\text {th }}$ positions are defined as 'below' the metal binding site.
investigate the steric interference caused by D-Leu. Additionally, the metalated $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 19 \mathrm{DL})_{3}{ }^{\text {a }}$ structure is used to examine cavities around the metal site. This structural analysis explains how steric engineering can be applied to vary Cd (II) geometries from 3-, 4- to 5 -coordinate around the metal site of 3SCCs. This knowledge is useful for biophysical applications when one would want to design a desired metal site in a protein to control coordination number or provide access for solvent or substrate in catalytic reactions.

## Results and Discussion

The objective of these studies is to understand how outer coordination sphere residues influence coordination number on metals bound within the hydrophobic region of a 3SCC structure. These studies are not only interesting from a theoretical viewpoint, as they suggest strategies to control metal ion coordination number or substrate access to a metallo-catalytic
center in designed proteins, but also to elucidate factors that may define the stability of metal binding to native 3SCC regions as found in the ORF1p protein of the LINE-1 human retrotransposon, which also contains layers of cysteine thiolates within the hydrophobic core of a 3SCC domain. ${ }^{30-32}$

In this section, we will address how changing the steric factors of side chain residues located toward the N -terminus (above the metal site), the C-terminus (below the metal site) or on the ligands themselves (Pen) influence the structure of the metal binding site. In general, while crystals using the parent TRI peptides can form, they diffract poorly as they are not ordered in one dimension. To resolve this problem, CS peptides (either CS or GRAND-CS which is one heptad longer leading to a more stable scaffold) have been examined. In all cases, the metal binding behaviors between TRI and CS derivatives are identical. While it is preferable to have completed these studies using the relevant $\mathrm{Cd}(\mathrm{II})$, crystals of CS derivatives with this ion have not been forthcoming. Therefore, we will use $\mathrm{Hg}(\mathrm{II})$ as an analogue of $\mathrm{Cd}(\mathrm{II})$ binding in trigonal planar and trigonal bipyramidal binding environments. This substitution is reasonable given that previous EXAFS analysis has shown that trigonal planar Cd(II) complex in these peptides have $\mathrm{Cd}(\mathrm{II})$-S distances of $2.46 \AA^{19,20}$, while the $\mathrm{Hg}(\mathrm{II})$ structure exhibited $\mathrm{Hg}(\mathrm{II})-\mathrm{S}$ of $2.38 \AA .{ }^{13}$

The parent peptide CSL16C binds $\operatorname{Cd}($ II $)$ with a 60:40 mixture of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ and $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$. We will first discuss, why this ratio occurs when L-leucine residues are located above and below the sulfur metal binding plane. We will then explain how replacing L-Leu with L-Ala (GRAND-CSL12AL16C) provides an environment that allows isolation of a $100 \% \mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$. Following this discussion we will explain how two alternative methods, replacing cysteine by Pen (CSL16Pen) or altering the chirality of the Leu above the cysteine layer (GRANDCSL12pLL16C), constrict the metal environment to give exclusively $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$. Finally, we will show how alteration of chirality below the sulfur plane allows greater access to solvent, ultimately leading to a structure that has a significant degree of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}$ species. These studies demonstrate how altering chirality around the metal binding site can enhance or diminish solvent access, depending on the placement of the substitution.

## Allowing 4-coordinate Cd(II) by Removal of Steric Bulk: Leu to Ala Mutation

Modifications of residues in outer coordination spheres play an important role in modulating solvent access to the metalbinding site, as well as metal site hydration and metal ion coordination. ${ }^{33-36}$ Unlike Leu, Ala contains a single methyl group ( $\mathrm{C}_{\beta}$ carbon) attached to the $\alpha$-carbon. Alber et al. had shown that conversion of Leu to Ala allowed for the addition of four waters into the cavity generated in 4 -helix bundles by removing the leucine isopropyl groups. ${ }^{36}$ Though the 3SCC is a narrower construct, it might be expected to behave similarly. Lee et al. substituted Ala for the more bulky Leu at the twelfth position to provide a water pocket above the metal site in (TRIL12AL16C) $3^{16,17}$ The design resulted in an exclusive 100\% $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right) .^{16,17}$ While a structure of $\mathrm{Cd}(\mathrm{II})(\mathrm{CSL} 12 \mathrm{AL} 16 \mathrm{C})_{3}{ }^{-}$ has not been obtained, structural understanding of the design
can still be achieved using the related $\mathrm{Hg}(\mathrm{II})(G R A N D-$ CSL12A16C) $3^{-}$.

To see the effect of 12Ala compared to 12Leu, the $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND-CSL12A16C) $3^{-}$structure is overlaid onto the known $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$(Figure 2). In both cases, $\mathrm{Hg}(\mathrm{II})$ is found as a trigonal planar structure. Previous work has shown the GRAND-CS peptides are predisposed to bind trigonal planar or pseudo-tetrahedral metals ${ }^{13}$, meaning that a large rotation of the thiol from the apoprotein is required upon metal complexation. In trigonal planar structures, $\mathrm{Hg}(\mathrm{II})$ induces $\sim 100^{\circ}$ of apo-Cys rotation from a position pointing upward toward the N -termini to being directed downward toward the C-terminal end. This rotamer reorientation expands the hydrophobic cavity above the sulfur plane sufficiently to accommodate a water molecule as seen in $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+} .{ }^{13}$ The three Cys residues in $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12A16C) })^{-}$are symmetric due to crystallographic requirements of the R32 space group. Each Cys contains two rotamers (Figure S1) in which only the major conformer is suitable to bind the metal with an orientation toward the helical interface $\left(\chi_{1}=-150.92^{\circ}\right)$. This $\chi_{1}$ value is close to the $-150.35^{\circ}$ observed in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}\left(\right.$ GRAND-CSL16CL30H) $3^{+}$ indicating that the orientations of the bound Cys ligands in both structures are similar (Figure 2c). Such arrangements make the metal pocket sizes comparable ( $\mathrm{S}_{\gamma}-\mathrm{S}_{\gamma}$ separation of $4.24 \AA$ for $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND-CSL12A16C) $3^{-}$and 4.08 A for $\left.\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}\right)$. The $\mathrm{Hg}(\mathrm{II})$ ion in $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12A16C) })^{-}$is situated at a distance of $0.26 \AA$ below the 16Cys plane with an averaged $\mathrm{Hg}(\mathrm{II})-\mathrm{S}$ distance of $2.44 \AA$ and average $\mathrm{S}-\mathrm{Hg}(\mathrm{II})-\mathrm{S}$ angle of $118.21^{\circ}$ (Table 3). These values correspond closely to those observed in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$. Taken together these parameters confirm that both designs show essentially identical first coordination environments for $\mathrm{Hg}(\mathrm{II}) \mathrm{S}_{3}$. Moreover, the apoprotein is also predisposed for Hg (II) binding in $\mathrm{Hg}(\mathrm{II})$ (GRAND$\left.\mathrm{CSL}^{2 A 16 C}\right)_{3}{ }^{-}$(Figure S2). These indicate that a trigonal planar $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ is not restricted from forming in the 12 Leu peptide. Therefore, we can expect that the change in the $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ to $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ ratio is not a consequence of the first coordination sphere, but rather depends on factors associated with the outer coordination spheres that surround the metal pocket. Figure 3 emphasizes the steric hindrance generated from the aliphatic isobutyl side chain of Leu compared to the methyl group of Ala. It is obvious that Ala generates a hole above the metal site confirming the proposed impact of the modification. As a consequence, the larger space in $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND-CSL12A16C) $3^{-}$allows for up to four waters to access the metal binding site (Figure 2b). This observation is consistent with Alber's previous study ${ }^{31}$, providing a convenient explanation for the shift in coordination mode to fully $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$. In contrast, in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}($ GRANDCSL16CL30H) $)^{+}$, only a single, unbound water that sits on the threefold axis directly above the metal at a $2.79 \AA$ distance is observed (Figure 2a). In $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND- CSL12A16C) $3^{-}$, one of the waters behaves in the same way as the observed water in the 12Leu-structure. Indeed, it is again located

on a threefold axis at a non-bonding distance of $3.55 \AA$ from Hg (II) (Figure 2b). Moreover, the other three waters are threefold related, but located close to the helical interface between two neighbouring strands with a distance of $4.34 \AA$ from the $\mathrm{Hg}(\mathrm{II})$ center. These waters form a hydrogen bonding network and are separated by a distance of $2.78 \AA$ from the central water. Each solvent is found within the same plane (with respect to the N-termini). Such distances of $\mathrm{Hg}(\mathrm{II})$ to water are too long to be a $\mathrm{Hg}(\mathrm{II})-\mathrm{O}$ bond (predicted to be $\sim 2.20 \AA$ ), therefore, all of the waters found within the cavity are considered to be uncoordinated and stabilized through H bonding interactions
Figure 2. Comparison of the $\mathrm{Hg}(\mathrm{II}) \mathrm{S}_{3}$ sites and the amount of observed waters above the metal site between the $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{N}(\mathrm{GRAND}-$ CSL16CL30H) ${ }_{3}{ }^{+}$(PDB code: $\left.5 \mathrm{KB1}\right)^{13}$ and $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12AL16C) })_{3}{ }^{-}$. Left to right columns demonstrating top-down and side-on views of the Hg (II) binding sites in the sixteenth position. In a) $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}($ GRAND CSL16CL30H) ${ }_{3}{ }^{+}$, b) $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12AL16C) })^{-}$and c) an overlay between the two structures. Main chain atoms of $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}($ GRAND $\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}$and $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12AL16C) })_{3}{ }^{-}$are green and pink, respectively. 16Cys, 12Leu and 12Ala side-chains are shown as sticks (sulfurs=yellow). $\mathrm{Hg}(\mathrm{II})$ atoms in the $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$ and $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL} 12 \mathrm{AL} 16 \mathrm{C})_{3}{ }^{-}$are present as blue and gray spheres, respectively. The observed waters in $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$ and $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL} 12 \mathrm{AL} 16 \mathrm{C})_{3}{ }^{-}$are shown as small red and cyan spheres.
between each other and the backbone of peptides. Another compelling point to support the large size of the cavity formed
with 12Ala is the observation of a longer $\mathrm{Hg}(\mathrm{II})$ to the central water distance in $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL} 12 \mathrm{~A} 16 \mathrm{C})_{3}{ }^{-}$( 3.55 A ) than in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}(2.79 \AA$ Å). This increase in $\mathrm{Hg}(\mathrm{II})-\mathrm{O}$ separation clearly demonstrates that more space is available in the 12Ala-containing structure for the water to move upward because it can form H -bonding to the additional three waters that also occupy in the cavity. The different number of water molecules between the two structures can explain the different degrees of solvation of $\mathrm{Cd}(\mathrm{II})$ between TRIL12A16C and TRIL16C designs. This observation proves that a cavity for solvent exists and it may allow for some water access when $\mathrm{Cd}(\mathrm{II})$ is bound to the metal site. Furthermore, EXAFS data indicate that


Figure 3. Packing of residues (shown as spheres) in the twelfth position above the metal site representing less hydrophobic character of Ala in $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL12AL16C})_{3}{ }^{-}$compared to Leu in $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}($ GRAND CSL16CL30H) ${ }_{3}{ }^{+}$(PDB code: 5KB1) ${ }^{13}$. From top down view of the N -termini, a) 12Leu residues in the $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL3OH })_{3}{ }^{+}$, and b) 12Ala residues in the $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12AL16C) })_{3}$. In c) an overlay between a) and b). Main chain atoms of $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$and $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL12AL16C})_{3}$ are shown in green and pink, respectively. Cys residues are shown as sticks (sulfurs=yellow). $\mathrm{Hg}(\mathrm{II})$ atoms in the $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$and $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12AL16C) })_{3}$ are present as blue and grey spheres, respectively. The observed waters in the $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}\left(\right.$ GRAND-CSL16CL30H) ${ }_{3}{ }^{+}$and $\mathrm{Hg}(\mathrm{II})($ GRAND CSL12AL16C) $)^{-}$are shown as small red and cyan spheres
a $\mathrm{Cd}(\mathrm{II})-\mathrm{O}$ bond in a $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ structure is $2.35 \AA,{ }^{19}$ which would appear achievable based on the $\mathrm{Hg}(\mathrm{II})$ to water separation in the crystal structures. However, one must remember that Hg (II) forms a trigonal planar structure whereas Cd(II) would have a 4-coordinate pseudo-tetrahedral polyhedron. This means that the $\mathrm{Cd}(\mathrm{II})$ would need to be displaced above the three sulfur atom plane toward the solvent ligand. As the spectroscopic data indicated that only $60 \%$ of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ is present in $\mathrm{Cd}(\mathrm{II})(\mathrm{TRIL} 16 \mathrm{C})_{3}{ }^{-7}$, this suggests that in a four coordinate structure the cavity may not be capable of stabilizing water well in the hydrophobic core. This is likely due to a combination of steric clashes between the bound water and the isobutyl side chains of leucine and the lack of additional hydrogen bonding atoms in close proximity that would stabilize the coordinated water. In the TRIL12AL16C design, this steric restriction is no longer operative, even for a four coordinate complex, and multiple waters that may H -bond to the bound water are present. Thus, $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ is the only species that forms in Cd(II)(TRIL12AL16C) $3^{\circ}$.

This structural analysis explains the factors that control the binding of water to $\mathrm{Cd}(\mathrm{II})$ in TRIL16C and TRIL12AL16C type peptides; however, equally interesting are the constraints that exclude water from the $\mathrm{Cd}(\mathrm{II})$ coordination environment to yield $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ systems in other designed peptides.

## Enforcing 3-coordinate Cd(II) Through Steric Interference of the Metal Binding Ligand: Cys to Pen Mutation

Lee et al. demonstrated that the incorporation of the Pen ligand in lieu of Cys at the sixteenth position (TRIL16Pen) led to $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ coordination as confirmed by ${ }^{113} \mathrm{Cd}$ NMR and ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC spectroscopies. ${ }^{16}$ The spectroscopic evidence has been confirmed by structural analysis. Upon metal binding, the methyl groups prevent the thiol side chains from rotating downward toward the C-terminal end as observed for the L-cys derivative. Thus, the ligands stay in roughly the same position as in the apo protein indicating that Pen is highly preorganized for metal binding. The consequence of this modification is that the sulfur plane cannot shift toward the C-termini and must remain close to the Leu layer above the metal site (as compared to the Cys derivative). In this situation, the space above the Pen layer becomes insufficient for water accommodation. Consequently, the formation of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ is favorable in $\mathrm{Cd}(\mathrm{II})(\mathrm{CSL} 16 \mathrm{Pen})_{3}{ }^{\circ}$.

Three possible explanations are considered for the perturbation of metal coordination environments by penicillamine. First, Pen ligands could have positioned their $\gamma$-methyl groups toward the space above the metal plane resulting in a smaller cavity above the site that excludes solvent access. Second, Pen might have undergone conformational changes upon metal complexation that excluded the water. Third, Pen ligands could have perturbed the primary coordination sphere of the metal in a specific way that encouraged a $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ structure.

The first hypothesis was refuted by analysis of the aligned apo-(CSL16Pen) $)_{3}\left(\mathrm{PDB}\right.$ ID: 3H5F) ${ }^{12}$ and apo-(CSL16C) ${ }_{3}$ (PDB ID: 5K92) ${ }^{13}$ structures. The helical backbones of the two structures are well-overlaid (RMSD $=0.17$ ). The incorporation of Pen does not perturb the helical framework (Supporting Information Figure S3, a). Only the major conformers (with $95 \%$ occupancy) are considered to be oriented for metal binding as the thiols are directed into the helical core (Supporting Information Figure S4, b) which resembles the major Cys residues in apo-(CSL16C) $)_{3}$, where the $\mathrm{S}_{\gamma}$ atoms point at the central core and toward the N -termini (Supporting Information Figure S3, b and c). The thiol pocket in the apo-(CSL16Pen) ${ }_{3}$ (average $\mathrm{S}_{\gamma}-\mathrm{S}_{\gamma}$ separation of $3.71 \AA$ ) is slightly larger than in the apo-(CSL16C) $)_{3}(3.32 \AA)$. While the $S_{\gamma}$ atoms of Pen in apo(CSL16Pen) $_{3}$ are oriented toward the interior of the coiled coil, the $\gamma$-methyl groups are pointing to the exterior. The fact that the Pen rotamers are almost at full occupancy ( $95 \%$ ) suggests that these side-chain conformations are geometrically preferred when Pen is placed at the sixteenth position; however, the similar orientations of $\gamma$-thiols observed in the Cys structure exhibits only $70 \%$ occupancy, implying that there is more free rotation of $\gamma$-thiols in Cys rather than Pen. The
rigidity of these $\mathrm{S}_{\gamma}$ angles in apo-(CSL16Pen) $)_{3}$ likely results from the restricted thiol rotation around $C_{\beta}$ atom due to the steric constraint imposed by the two $\gamma$-methyl groups of Pen. Based on these observations, the $\gamma$-methyl groups of Pen, which are oriented toward the helical interface, are not positioned to block the space above the metal site (between the 12Leu and 16Pen layers) that potentially excludes water from binding to $\mathrm{Cd}(\mathrm{II})$.


Figure 4. Comparison of the trigonal planar structures of $\mathrm{Hg}(\mathrm{II})(\mathrm{S}-\mathrm{Cys})_{3}{ }^{-}$ from $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}$(PDB code: 5KB1) ${ }^{13}$ and $\mathrm{Hg}(\mathrm{II})(\mathrm{S}-\mathrm{Pen})_{3}{ }^{-}$from the $[\mathrm{Hg}(\mathrm{II})]_{\mathrm{S}}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right)\right]_{\mathrm{N}}(\mathrm{CSL9PenL23H})_{3}{ }^{\mathrm{n+}}$ (PDB code: 3PBJ) ${ }^{28}$. a) Top-down from the N -termini and b) side-on views of the overlay. Main chain atoms of $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$are colored in green and $[\mathrm{Hg}(\mathrm{II})]_{S}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right)\right]_{N}(\mathrm{CSL9PenL23H})_{3}{ }^{\mathrm{n+}}$ in cyan. Cys, Pen and Leu (above the metal site) are shown as sticks (sulfurs=yellow). $\mathrm{Hg}(\mathrm{II})$ atoms in the $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}$and $[\mathrm{Hg}(\mathrm{II})]_{s}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right)\right]_{N}(\mathrm{CSL9PenL23H})_{3}{ }^{\mathrm{n+}}$ are labelled as blue and grey spheres, respectively.

The second hypothesis can be dismissed by comparing the previously published structure of $[\mathrm{Hg}(\mathrm{II})]_{\mathrm{s}}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right.\right.$ $)]_{N}\left(\text { CSL9PenL23H) } 3^{\text {n+ }} \text { with that of the apo-(CSL16Pen) }\right)_{3}$ to reveal the behavior of trigonal planar $\quad \mathrm{Hg}(\mathrm{II})$ binding to Pen-ligand in an a site. The $[\mathrm{Hg}(\mathrm{II})]_{\mathrm{s}}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right.\right.$ $)]_{N}(\mathrm{CSL} 9 \mathrm{PenL} 23 \mathrm{H})_{3}{ }^{\mathrm{n}+}$ contains a trigonal planar $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3 .}{ }^{28}$ The bound Pen ligands (Supporting Information Figure S5, b) direct their $S_{\gamma}$ atoms toward the interior core while positioning the $\gamma$-methyl groups out toward the helical interface. This $\mathrm{S}_{\gamma}$ configuration is similar to that observed in the apo-(CSL16Pen) $3_{3}$ (Supporting Information Figure S5, a). ${ }^{12}$ The invariance of the thiol layer between the non-metallated and metallated proteins can be underscored by their very tiny torsion angle shift (Table 1), that results in almost equal $S_{\gamma}-S_{\gamma}$ separations between the two structures ( 3.71 versus $3.84 \AA$ for apo-(Pen) ${ }_{3}$ and $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$, respectively) (Supporting Information Figure S5, c). The highly similar sulfur planes emphasize that apo-Pen ligands exhibit a high degree of preorganization for metal binding, which could be due to the rigidity caused by the bulky $\gamma$ methyl substitution that prevents the $\gamma$-thiol from moving freely through space. Therefore, the second hypothesis that a conformational change of the Pen side chain is responsible for the presence of $100 \% \mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ structure appears to be incorrect.

The third hypothesis suggested a change in the first coordination sphere orientation that is imposed by the remote
methyl groups. This can be assessed by comparing the coordination spheres of the $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$ and $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\mathrm{Cys}}\right)_{3}$ in $\left[\mathrm{Hg}(\mathrm{II})_{S}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right)\right]_{N}(\mathrm{CSL9PenL23H})_{3}{ }^{{ }^{n+}}\right.$ and $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$(Figure 4). We observe that the thiol orientations of the bound-Cys and bound-Pen point
in completely opposite directions. While the Pen ligands are
oriented to the interior of the 3SCC, the Cys residues instead are directed further out toward the helical interface as confirmed by the different $\chi_{1}$ torsion angles of the two ligands $\left[-49.85^{\circ}\right.$ observed in $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$ and $-150.35^{\circ}$ in $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\mathrm{Cys}}\right)_{3}$ ] (Figure 4a). The average $\mathrm{S}_{\gamma}-\mathrm{S}_{\gamma}$ separation in Pen is subsequently shorter than Cys by $0.24 \AA$, verifying that Cys can make a larger triangular metal plane compared to Pen. This observation indicates that the $\gamma$-methyl groups of Pen inhibit the expansion of the three atom sulfur plane to the requisite distances that are optimal for a trigonal $\mathrm{Hg}(\mathrm{II})$ species. The predisposed apo-Cys peptide reorients the $\gamma$-thiols downward toward the C-termini and facing out toward the helical interface in order to accept Hg (II) into the metal plane. ${ }^{13}$ The Hg (II)-Cys plane is shifted $\sim 1.30 \AA$ down toward the C-termini with respect to the apo-Cys structure ${ }^{13}$, whereas due to the preorganization of Pen ligands the sulfur plane is essentially unaltered on $\mathrm{Hg}(\mathrm{II})$ binding. Figure 4b illustrates the difference between the metallated planes in these proteins. The $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$ is positioned about $1.80 \AA$ more toward the N - termini relative to $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\mathrm{Cys}}\right)_{3}$. Clearly, the immobility of the Pen side chains requires the metal to bind in a more compressed trigonal plane that is located further toward the $N$-termini. Since the packing of the 12Leu layers remain unchanged between the $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\mathrm{Cys}}\right)_{3}$ and $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$ while the metallated-Pen plane is higher than in the bound-Cys form, there is a less space available for a fourth ligand in the Pen structure. Supporting Information Figure $\mathbf{S 6}$ (top panels) shows that the packing of the Leu residues above the $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$ site is, in fact, slightly tighter than in the $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\mathrm{Cys}}\right)_{3}$. As a consequence, the Leu layer is at a distance of $4.86 \AA$ from the bound-16Pen plane, whereas the related distance determined from the $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\mathrm{Cys}}\right)_{3}$ lengthens to $5.92 \AA$ (Figure 5). The impact of this difference in interlayer spacing on water access

Table 2: Crystallographic parameters observed from the crystal structures ${ }^{\text {a }}$

| Peptides | $\begin{aligned} & \text { apo-(CSL16C) } \\ & \text { (PDB code: } 5 \mathrm{~K} 92)^{14} \end{aligned}$ | $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}$ (GRANDCSL16CL3OH) ${ }_{3}{ }^{+}$ (PDB code: 5KB1) | apo-(CSL16Pen) ${ }_{3}$ (PDB code: 3H5F) ${ }^{12}$ |  | Hg(II)(GRANDCSL12A16C) ${ }_{3}$ <br> (PDB code: 6EGO) | apo-(GRAND- <br> CSL12ㄷLL16C) ${ }_{3}$ <br> (PDB code: 6EGL) | apo-(GRANDCSL16CL19ㄴㄴ) (PDB code: 6EGM) | Hg(II)(GRANDCSL16CL19。L) ${ }_{3}$ <br> (PDB code: 6EGN) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 16Cys rotamers $\chi_{1}$ (interior rotamers ${ }^{\text {b }}$ $\mathrm{S}_{y}-\mathrm{S}_{y}$ distance $(\hat{A})^{)^{c}}$ $\chi_{1}$ (exterior rotamers) $\mathrm{S}_{y}-\mathrm{S}_{y}$ distance $(\tilde{A})^{d}$ | $-66.24^{\circ}$ (average) 3.32 (average) $-176.47^{\circ}$ (average) $^{h}$ 5.33 (average) | $\begin{gathered} -150.35^{\circ} \\ 4.08 \\ -169.58^{\circ} \\ 5.06 \end{gathered}$ | $-49.85^{\circ}$ (average) <br> 3.71(average) <br> $72.99^{\circ}$ (average) <br> 8.45(average) | $-50.23^{\circ}$ (average) 3.84 (average) | $\begin{gathered} -150.92^{\circ} \\ 4.24 \\ -56.73^{\circ} \\ 3.24 \end{gathered}$ | $\begin{gathered} -68.57^{\circ} \\ 3.22 \\ -174.79^{\circ} \\ 5.93 \end{gathered}$ | $\begin{gathered} -61.13^{\circ} \\ 3.31 \\ - \end{gathered}$ | $\begin{gathered} -153.11^{\circ} \text { (average) } \\ 4.19(\text { average) } \\ -28.85^{\circ},-94.80^{\circ 1} \\ 3.66(\text { average }) \end{gathered}$ |
| Metal site <br> M-S bond length ( $A \AA$ ) <br> S-M-S angle(average) <br> Distance of metal from to the bound Cys plane $(\AA \AA)^{e}$ | - - - | $\begin{gathered} 2.38, \mathrm{Hg}(11)-\mathrm{S} \\ 118.50^{\circ} \\ -0.3 \end{gathered}$ | - <br> - <br> - | $2.23, \mathrm{Hg}(\mathrm{II})-\mathrm{S}($ average $)$ $\begin{gathered} 119.90^{\circ} \\ -0.06 \end{gathered}$ | $\begin{gathered} \text { 2.44, } \mathrm{Hg}(11)-\mathrm{S} \\ 118.21^{9} \\ -0.26 \end{gathered}$ | - - - | - - - | 2.44, $\mathrm{Hg}(\mathrm{II})-\mathrm{S}$ (average) $\begin{gathered} 119.69^{\circ} \\ -0.12 \end{gathered}$ |
| Leu rotamers above the metal site <br> Interior $\mathrm{C}_{5}$ separation $(\dot{A})^{1}$ <br> Exterior $\mathrm{C}_{5}$ separation $(\hat{A})^{9}$ Distance of the layer from the interior sulfur plane (Å) | $\begin{gathered} \hline \text { (12L-Leu) } \\ 4.40 \\ 6.73 \\ 4.92 \end{gathered}$ | $\begin{gathered} \hline \text { (12L-Leu) } \\ \\ 3.89 \\ 6.13 \\ 5.92 \end{gathered}$ | $\begin{gathered} \hline \text { (12L-Leu) } \\ 4.94 \\ 6.74 \\ 4.95 \end{gathered}$ | $\begin{gathered} \hline \text { (5L-Leu) } \\ 3.60 \\ 5.84 \\ 4.86 \end{gathered}$ | $:$ | $\begin{gathered} \hline \text { (12D-Leu) } \\ 3.93 \\ 4.53 \\ 2.32 \end{gathered}$ | $\begin{gathered} \hline \text { (12L-Leu) } \\ 4.04 \\ 6.45 \\ 4.72 \end{gathered}$ | $\begin{gathered} \hline \text { (12L-Leu) } \\ 4.06 \\ 6.39 \\ \sim 6.17 \end{gathered}$ |


| Leu rotamers below the metal site | (19L-Leu) | (19L-Leu) | (19L-Leu) | (12L-Leu) | (19L-Leu) | (19L-Leu) | (19D-Leu) | (19D-Leu) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\frac{\text { Interior } \mathrm{C}_{\bar{\delta}}}{}$ Separation (A) | 4.64 | 6.17 | 5.86 | 5.28 | 6.47 | 5.30 | 5.17 | 8.09(average) |
| Exterior $\mathrm{C}_{5}$ separation (A) | 7.25 | 9.08 | 8.66 | 8.45 | 9.30 | 8.22 | 8.96 | 11.49(average) |
| Distance of the layer from the interior sulfur plane (A) | 4.41 | 3.30 | 4.52 | 4.60 | 3.23 | 4.04 | 5.18 | 3.76 |

${ }^{\text {a P Peptides that were crystallized in R32 space group have crystallographically imposed } 3 \text {-fold symmetry along the } z \text { axis that runs through the center of the three }}$ helices of the 3SCC. The consequence of symmetry is that structures in R32 will have a single reported value for the following crystallographic parameters (x1 dihedral angles, atomic distances, and M -S distances), while average values are usually given for the structurecrystallizing in C 2 in which the three helices are independent. ${ }^{\mathrm{b}} \chi_{1}$ of Cys residue is determined from the dihedral angle of $\mathrm{N}-\mathrm{C}_{\alpha}-\mathrm{C}_{\beta}-\mathrm{S}_{\gamma},{ }^{c}$ The distance determined between $\mathrm{S}_{\gamma}$ atoms of the interior Cys conformers., ${ }^{d}$ The distance determined between $S_{\gamma}$ atoms of the exterior Cys conformers., ${ }^{e}$ Plus sign (+) indicates the metal is situated above the bound Cys plane. Minus ( - ) indicates the metal is situated below the bound Cys plane., Interior $\mathrm{C}_{\delta}$ separation define the average distance between the interior $\mathrm{C}_{\delta}$ atoms of Leu residues of all chains. ${ }^{9}$ Exterior $\mathrm{C}_{\delta}$ separation define the average distance between the exterior $\mathrm{C}_{\delta}$ atoms of Leu residues of all chains., ${ }^{\text {h }}$ Average X 1 dihedral angle determined from minor Cys conformers observed from two of the chains. ${ }^{i}$ Average Sy -Sy separation determined from the two minor Cys conformations found on two chains and the third Cys (major) from the remaining chain., ${ }^{j} \times 1$ dihedral angle determined from minor Cys conformers observed from two chains.
is supported by the absence of a water molecule in Enforcing 3-coordinate Cd(II) By Modification of Outer the area above the Penligand binding
site in the

$[\mathrm{Hg}(\mathrm{II})]_{S}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}^{-}\right)\right]_{N}(\mathrm{CSL9PenL23H})_{3}{ }^{\mathrm{n}} .{ }^{28} \quad$ Moreover, the aligned binding sites (Supporting Information Figure S6, bottom panel) also show that the coordinated $\mathrm{Hg}(\mathrm{II})$ in $\mathrm{Hg}(\mathrm{II})\left(\mathrm{S}_{\text {Pen }}\right)_{3}$ occupies the space where the water was previously observed in $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$. It is obvious that Pen cannot generate enough space to accommodate a polar molecule within the hydrophobic core above the metal site. Therefore, these observations support the third formulated hypothesis.

The restriction of the metal binding $\mathrm{S}_{3}$ plane in the presence of Pen ligands likely translates to the $\mathrm{Cd}(\mathrm{II})$ proteins. The shift of the metal plane toward the Leu residues and concurrent induction of a tighter hydrophobic packing above the metal site would act in concert to generate less space for solvents. This reasoning supports a model for complete water exclusion (full $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ formation) in $\mathrm{Cd}(\mathrm{II})(\text { TRIL16Pen })_{3}$. This hypothesis has been confirmed by the corresponding ${ }^{113} \mathrm{Cd}$ NMR and ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC results. ${ }^{16}$

## Coordination Sphere Chirality: L-Leu to D-Leu Mutation on N -terminal Side

GRAND-CSL12DLL16C serves as a crystallographic analog for TRIL12DLL16C (See Supplementary Discussion). To illustrate the effect of alternate chirality on the internal hydrophobic residues in the outer coordination spheres, the apo-(GRAND-CSL12dLL16C) ${ }_{3}$ is overlaid onto the known apo-
$(\mathrm{CSL16C})_{3}($ Supporting information Figure S7, a). This figure compares a parent peptide that contains solely L -amino acids in the sequence to one with a single D-Leu substitution. Both of the peptides fold into parallel 3SCCs as predicted. Although they are different in length by one heptad, the $\alpha$-helical backbones of the two structures are extremely similar ( $\mathrm{RMSD}=0.36$ ). Intriguingly, there are no kinks in the helical backbones observed in apo-(GRAND-CSL12LL16C) ${ }_{3}$ suggesting that the incorporation of a D-Leu does not disturb the coiled coil secondary structure. In the sixteenth position, the $\mathrm{C}_{\beta}$ carbons of the Cys residues of the apo-(GRAND- CSL12DLL16C) $)_{3}$ point toward the N-termini of

the helices and the $\mathrm{S}_{\gamma}$ atoms adopt two conformations (Supporting information Figure S8). The major Cys rotamers have the thiols positioned toward the metal binding core of the peptide, exhibiting a similar range of torsion angles with the apo-$(\mathrm{CSL16C}))_{3}\left(-68.57^{\circ}\right.$ versus $\left.-66.24^{\circ}\right)$ (Supporting information Figure S8, b). The $\mathrm{S}_{\gamma}-\mathrm{S}_{\gamma}$ distances are comparable between both structures; $3.22 \AA$ in apo-(GRAND-CSL12DLL16C) ${ }_{3}$ and $3.32 \AA$ (average) in apo-(CSL16C) ${ }_{3}$ (Supporting information Figure S9, a and b). The minor Cys orientations of apo-(GRAND-CSL12DLL16C) $)_{3}$ point their thiol groups to the outer interface, subsequently causing a long $\mathrm{S}_{\gamma}-\mathrm{S}_{\gamma}$ separation ( $5.93 \AA$ ) between minor Cys conformers, which are not suitable for metal binding (Supporting information Figure S8, c). According to this first structural analysis of the layer at the sixteenth position, it appears that the apo-structures of (GRAND-CSL12DLL16C) ${ }_{3}$ and (CSL16C) $)_{3}$ present a relatively similar metal binding environment. The effect of D-Leu above the metal site is pronounced at the twelfth position. The packing of 12D-Leu residues in apo-(GRAND-CSL12DLL16C) $)_{3}$ and 12L-Leu in apo-(CSL16C) $)_{3}$ are
Figure 5. Interlayer spaces around the thiolate site of designed peptides. a) apo-(CSL16C) $)_{3}$ (PDB code: 5 K 92$)^{13}$, b) apo-(CSL16Pen)3 (PDB code: 3H5F) ${ }^{12}$, c) apo-(GRAND-CSL16CL19DL) ${ }_{3}$, d) apo-(GRAND-CSL12 ${ }^{\left.\text {LLL16C })_{3}, ~ e\right) ~}$ $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}$(PDB code: $\left.5 \mathrm{KB1}\right)^{13}$, f) $[\mathrm{Hg}(\mathrm{II})]_{S}\left[\mathrm{Zn}(\mathrm{II})\left(\mathrm{H}_{2} \mathrm{O} / \mathrm{OH}-\right)\right]_{N}(\mathrm{CSL9PenL23H})_{3}{ }^{\mathrm{nt}}$ (PDB code: 3PBJ) ${ }^{28}$, g) Hg(II)(GRANDCSL16CL19DL) $)^{\circ}$ and h) $\mathrm{Hg}(\mathrm{II})(G R A N D-C S L 12 A L 16 C)_{3}$. Main chain atoms are shown as ribbon diagrams. Residue side chains are present as sticks. D-Leu in c), d) and g ) are colored in red. $\mathrm{Hg}(\mathrm{II})$ atoms and observed waters are shown as big and small spheres, respectively.

Figure 6. Effect of D-Leu in the twelfth position above the metal site (sixteenth) in the 3SCC environment. a) Side-on and b) top-down views of the overlay between apo-(GRAND-CSL12DLL16C) $)_{3}$ (red) and apo-(CSL16C) ${ }_{3}$ (PDB code: 5K92, orange) ${ }^{13}$ structures showing the difference in $\mathrm{C}_{\beta}$ carbon positions between D-Leu (red sticks) and L-Leu (orange sticks). From top down view of the N -termini, c) and d) representing the packing comparison between D-Leu and L-Leu residues (shown as spheres) in apo-(GRANDCSL12DLL16C) ${ }_{3}$ and apo-(CSL16C) $)_{3}$, respectively.
compared in Figure 6. It is obvious that 12D-Leu residues are more tightly packed than seen for 12L-Leu, causing greater steric
hindrance above the metal binding layer. This perturbation occurs because the D-configuration repositions the $C_{\beta}$ atoms from directing toward the N -termini (in L-Leu) to the C -termini direction (Figure 6, a and b). This $\mathrm{C}_{\beta}$ deviation twists the positions of $\delta$-methyl groups $\left(\mathrm{C}_{\delta 1}, \mathrm{C}_{\delta 2}\right)$ toward the center of the coiled coil. In the apo-(CSL16C) $)_{3}$ structure, only one of the two $\delta$ methyl atoms of each L-Leu residue is pointed toward the center, while the other points to the helical interface, thus opening up more space above the metal binding site and potentially making it less well-packed compared to 12D-Leu (Figure 6, cand d). This D-Leu effect shortens the separation between 12D-Leu and 16Cys layer to $2.32 \AA$ as compared to the $4.92 \AA$ observed in the apo-(CSL16C) $)_{3}$ (Figure 5). The differential orientations of leucine layer in the outer coordination spheres, therefore, could represent an important effect of the amino acid side chains chirality on metal structures and binding mode preferences in the metallated-forms.

One may compare the known $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{N}$ (GRANDCSL16CL30H) ${ }^{+}$to apo-(GRAND-CSL12DLL16C) $)_{3}$ to obtain insight on Cd (II) complexation. As expected, the GRANDCSL16CL30H is predisposed toward $\mathrm{Hg}(\mathrm{II})$-binding as described above for CSL16C (Supporting information Figure S10). ${ }^{13}$ The metal induces significant rotation of the interior Cys
conformations by moving the thiols downward and to the side. This shift orients the cysteine sulfur atoms more toward the helical interface leading to an expansion of $S \gamma-S \gamma$ separations from 3.22 to $4.08 \AA$. The $\chi 1$ dihedral angle changes from $-66.24^{\circ}$ (average) in the apo-structure to $-150.35^{\circ}$ (average). Unsurprisingly, the orientation of 12D-Leu in apo-(GRANDCSL12 ${ }_{D}$ LL16C) $)_{3}$ differs from the 12L-Leu in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$. First, both $\delta$-methyl groups of each D-Leu residue, point toward the core of the helices, whereas in $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$only one of the $\mathrm{C}_{\delta}$ atoms $\left(\mathrm{C}_{\delta 1}\right)$ of each 12L-Leu is in the core while the other is facing out toward the helical interface as shown in
Supporting Information Figure S11, a and b. Moreover, the analysis of the aligned structures demonstrates that the $\mathrm{C}_{\beta}$ atoms of D-Leu are drastically different in position from the Lchirality, causing the $\mathrm{C}_{\delta 2}$ atoms in the D-Leu layer to tuck toward the center and move closer to the observed water in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$. This causes the 12DLeu layer ( $\mathrm{C}_{82}$ plane) to move closer to where the water would reside (only $1.30 \AA$ distance), while the interior $\mathrm{C}_{\delta}$ plane $\left(\mathrm{C}_{\delta 1}\right)$ of $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$is at a distance of $3.80 \AA$ from the water (Supporting Information Figure S11, c). It is assumed that if the (GRAND-CSL12 $\left.{ }_{D} L L 16 C\right)_{3}$ were to bind a metal, the shift of the sulfur plane toward the binding would likely cause the layers (12D-Leu versus 16Cys) above the metal site to be separated by approximately $4.30 \AA$, while in the actual $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$structure the related distance determined from $12 \mathrm{~L}-\mathrm{Leu}$ is $5.92 \AA$. This strongly emphasizes that the D-Leu layer above the metal site is tightlypacked suggesting that the water should no longer exist within this tiny space. Therefore, steric encumbrance appears to be the basis for water exclusion in TRIL12DL16C design.

## Increasing the Coordination Number of Cd(II) By Modification of Outer Coordination Sphere Chirality: L-Leu to D-Leu Mutation on C-terminal Side

The combination of ${ }^{13} \mathrm{Cd}$ NMR, ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC and EXAFS spectroscopies confirmed that the TRIL2WL16CL19pL peptide binds $\mathrm{Cd}(\mathrm{II})$ with a higher coordination number than found for TRIL12DLL16C with two species appearing in equal quantities as $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ and $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2} .^{25}$ Structural analysis of the apo-peptide is completed to evaluate hydrophobic packing in the absence of the metal site. The overlay of apo-(GRANDCSL16CL19pL) ${ }_{3}$ with apo-(CSL16C) $)_{3}$ illustrates that the $\alpha$-helical backbones are well-aligned with no kinks observed in the D-Leu region (Supporting information Figure S7, b). The Cys residues of apo-(GRAND-CSL16CL19DL) ${ }_{3}$ display a single rotamer pointing toward the core of the structure resembling the major conformer of apo-(CSL16C) ${ }_{3}$ and apo-(GRANDCSL12DLL16C) ${ }_{3}$ (Supporting information Figure S9). This can be confirmed by their close values in side-chain torsion angles and $\mathrm{S}_{\gamma}-\mathrm{S}_{\gamma}$ separations (Table 2). The similarity in Cys layers reveals that
the D-Leu does not affect the first coordination sphere ligands in the non-metallated form, regardless of the position where it is placed (12th or 19th position) in the outer coordination spheres. The effect of 19D-Leu is determined by overlaying the apo-(GRAND-CSL16CL19pL) ${ }_{3}$ onto the apo-(CSL16C) ${ }_{3}$ structure. Both the 19D-Leu and the 19L-Leu side chains appear to direct the $\delta$-methyl groups out toward the helical interface; however, the reorientation of the $\mathrm{C}_{\beta}$ atoms with the 19D-Leu in the apo-(GRANDCSL16CL19DL) ${ }_{3}$ causes both of the $\delta$-methylene groups to move further to the outer face than the 19L-Leu in apo-


Figure 7. Effect of D-Leu in the nineteenth position below the metal site (sixteenth) in the 3SCC environment. Top panels: Overlays between apo-(GRAND-CSL16CL19DL) $)_{3}$ (blue) and apo-(CSL16C) $)_{3}\left(\right.$ PDB code: $5 \mathrm{~K}^{2} 2^{13}$, orange) structures demonstrate the deviation of a) $\mathrm{C}_{8}$ and b ) $\mathrm{C}_{\beta}$ positions of D-Leu (blue sticks) from L-Leu (orange sticks). Bottom panels: The packing in the nineteenth position of c) D-Leu (blue spheres) in apo-(GRAND-CSL16CL19 $\left.{ }^{\text {D L }}\right)_{3}$ and d) L-Leu (orange spheres) in apo-(CSL16C) . In e) an overlay between e) and d).
$(\text { CSL16C) })_{3}$ (Figure 7, top panel). Moreover, the hydrophobic pocket below the metal site made by 19D-Leu appears to be bigger than 19L-Leu (Figure 7, bottom panel). Thus, the altered chirality of D-Leu can remove the steric constraints when it is placed below the metal site by rearranging the bulky $\delta$ methyl groups away from the center of the coiled coil. Consequently, it generates more open space with the less-wellpacked hydrophobes, which is believed to allow for better water access below the binding site. This conclusion is consistent with formation of the $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ corresponding to the ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC angular frequency characteristics of $0.316 \mathrm{rad} / \mathrm{ns}$. ${ }^{25}$ As shown in the L16C variant, the cavity that lies between the 12L-Leu and 16 Cys layers can bind $\mathrm{Cd}(\mathrm{II})$ from the top. Due to the limitation of spectroscopic techniques, these two conformations of $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ products (either with the water ligand positioned above or below) cannot be distinguished using the $0.316 \mathrm{rad} / \mathrm{ns}$ ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC angular frequency value. In order to assess this model, the $12 \mathrm{~L}-$ Leu packing in apo-(GRAND-CSL16CL19DL) ${ }_{3}$ is further investigated. If this hypothesis is true, the 12L-Leu in apo-(GRAND-CSL16CL19DL) ${ }_{3}$ should show sufficient space for water, comparable to the 12L-Leu layer in apo-(CSL16C) ${ }_{3}$. The overlay structures of both apo-peptides (Figure 7) reveals that all
the $\mathrm{C}_{\beta}$ carbons in 12L-Leu are directed toward the N -termini due to their L-configuration. The 12L-Leu sidechains in apo-
(GRAND-CSL16CL19DL) ${ }_{3}$ point one $\delta$-methyl group toward the core, while the other is oriented more toward the helical interface. A similar observation is noted for the apo-(CSL16C) ${ }_{3}$ parent
peptide. The packing in both structures look very similar even though the layer in apo-(GRAND-CSL16CL19DL) $)_{3}$ is slightly tighter-packed and the cavity is smaller (Supporting Information Figure S12). However, both structures generate a larger space above the metal site when compared to the smaller pocket made by the 12D-Leu in apo-(GRAND-CSL12dLL16C) ${ }_{3}$ (Figure 5). These crystal structures show that the cavity generated by the 12L-Leu in the apo-(GRAND-CSL16CL19DL) ${ }_{3}$ is large enough to house a water ligand above the metal site that can allow for $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ formation. Apparently, there are two spaces available in apo-(GRAND-CSL16CL19DL) $)_{3}$ for water access: one above that is likely partially occupied and a larger cavity below the
by the 12D-Leu in apo-(GRAND-CSL12dLL16C) $3_{3}$ (Figure 5). These crystal structures show that the cavity generated by the 12 L -Leu in the apo-(GRAND-CSL16CL19 $\left.{ }_{\mathrm{D}}\right)_{3}$ is large enough to house a water ligand above the metal site that can allow for $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ formation. Apparently, there are two spaces available in apo-(GRAND-CSL16CL19 $\left.{ }_{\mathrm{D}} \mathrm{L}\right)_{3}$ for water access: one above that is likely partially occupied and a larger cavity below the metal site that could be fully occupied by solvent. Therefore, the observation of the $0.316 \mathrm{rad} / \mathrm{ns}$ angular frequency from ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC could represent both 4-coordinate $\mathrm{Cd}(\mathrm{II})$ conformations where one has water bound on top with respect to the metal binding plane and the other has water bound below.

Apart from the $0.316 \mathrm{rad} / \mathrm{ns}$ angular frequency species observed in ${ }^{111 \mathrm{~m}} \mathrm{Cd}$ PAC, another $50 \%$ of the products from TRIL2WL16CL19D was reported to exhibit a 0.159 rad/ns


Figure 8. Side-on view of the metallated 3SCCs representing the existence of waters around the 16Cys coordinate site. In a) Hg(II)(GRAND$\left.\mathrm{CSL} 16 \mathrm{CL} 19_{\mathrm{D}} \mathrm{L}\right)_{3}{ }^{3}$, b) $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{N}(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}(\mathrm{PDB} \text { code: } 5 \mathrm{~KB} 1)^{13}$ and c ) an overlay between a) and b). Waters in Hg(II)(GRAND-CSL16CL19DL) ${ }^{-}$ and $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 30 \mathrm{H})_{3}{ }^{+}$are shown as yellow and red small spheres, respectively. $\mathrm{Hg}(\mathrm{II})$ ions in $\mathrm{Hg}(\mathrm{II})\left(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 19_{\mathrm{D}}\right)_{3}{ }^{-}$and $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$are represent in cyan and blue spheres, respectively. 12L-Leu, 16Cys (sulfurs=yellow) and 19D-Leu residues are shown as sticks with color corresponding to the ribbon diagram of each structure.
frequency that is uncommon in these designed peptide systems. ${ }^{25}$ However, this lower frequency is consistent with a 5-
coordinate $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}$ complex. Since the angular frequency is closer to zero, the NQI around the metal site is relatively symmetrical (where the prefect tetrahedral geometry ideally
shows $\omega_{0}$ of $\left.0 \mathrm{rad} / \mathrm{ns}\right){ }^{37}$ Therefore, this $0.159 \mathrm{rad} / \mathrm{ns}$ value suggests an axially symmetric trigonal bipyramidal structure. As shown by the crystallography described above, the existence of two cavities above and below the metal site in the apo-(GRAND-CSL16CL19 $\left.{ }^{\text {L }}\right)_{3}$ supports this possibility as space is available for waters to form a $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}$ structure.

Excitingly, additional evidence supporting the possibility of simultaneous water access in this peptide has been obtained through the crystal structure of $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND-CSL16CL19DL) $3_{3}$. Though the metal center is not $\mathrm{Cd}(\mathrm{II})$-bound, the mercurated binding site reveals some interesting aspects related to the previous predictions. Five water molecules are observed within both cavities around the metal layer of $\mathrm{Hg}(\mathrm{II})(G R A N D-$ CSL16CL19DL) $)^{-}$(Figure 8a). The first water is situated above the metal site, at a $2.76 \AA$ distance from the $\mathrm{Hg}(\mathrm{II})$ center. This value is close to the previously observed water found in $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}(2.79 \AA) .{ }^{13}$ It is stabilized by H -bonding interactions with Cys ligands and the second water which is located close to the helical interface between two helical chains of the 3SCC. The second water is $3.11 \AA$ from the first water and $4.25 \AA$ from the $\mathrm{Hg}(\mathrm{II})$ center. The reason this second water is observed in this $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL16CL19DL) })_{3}$ structure, but not in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$, is probably because $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND-CSL16CL19DL) ${ }^{-}$was crystalized in $\mathrm{P} 2{ }_{1} 2{ }_{1} 2_{1}$ space group which does not impose three fold symmetry on the helices. However, the R32 space group for $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$is tightly packed and the three-fold crystallographic symmetry constraints may exclude any waters which are not aligned on the three fold axis, thus resulting in the appearance of there being only one axial water positioned above the $\mathrm{Hg}(\mathrm{II})$ (Figure 8b). The third, fourth and fifth waters
are positioned within the nineteenth D-Leu cavity below the Cys plane with separations of $3.71 \AA, 6.03$ and $6.06 \AA$ away from $\mathrm{Hg}(\mathrm{II})$, respectively. The third water is situated near one of the 3SCC helices above the $\mathrm{C}_{\delta 2}$ plane of the 19D-Leu, lying close to the helical interface and showing strong H -bonding interactions with the thiol and carbonyl oxygen of the 16Cys residue of the corresponding helix. Moreover, it is at a distance of $3.24 \AA$ and 4.28 $\AA$ from the fourth and fifth waters, respectively that are situated toward the C-termini. The fourth water, is oriented more to the center of the helix, on the same plane as $\mathrm{C}_{\delta 1}$ atom of the19D-Leu. The fifth water is H-bonded with the carbonyl oxygen of one of the 19D-Leu residues. These observations strongly suggests that once the metal is bound in the Cys plane in $\mathrm{Hg}(\mathrm{II})\left(\text { GRAND-CSL16CL19 }{ }_{\mathrm{D}}\right)_{3}{ }^{-}$, the pocket made by the 19DLeu is sufficiently large as to accommodate more than just one water molecule.

To analyze the impact of D-Leu on hydrophobic packing in the metallated structures further, the $\mathrm{Hg}(\mathrm{II})$ (GRANDCSL16CL19DL) ${ }_{3}$ is aligned to the previously published $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$. Figure 8c represents an excellent overlay of the bound $S_{\gamma}$ conformers between the two structures as confirmed from their similarity in $\chi_{1}$ (Table 3). The $\mathrm{Hg}(\mathrm{II})$ ions of the two Hg (II)-structures are in the same plane (Figure 8c). The averaged $\mathrm{Hg}(\mathrm{II})-\mathrm{S}$ distance in $\mathrm{Hg}(\mathrm{II})(G R A N D-$

CSL16CL19DL) ${ }_{3}{ }^{-}$is $2.43 \AA$, which is close to the distances of trigonal planar $\mathrm{Hg}(\mathrm{II})$
structures in $\mathrm{Hg}(\mathrm{II})_{S} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$and reported small molecule complexes. ${ }^{14,15,38}$ As previously noted, all the waters observed in the outer coordination spheres of $\mathrm{Hg}(\mathrm{II})$-bound structures are non-bonded and are believed to have H -bond interactions to the bound Cys ligands, which helps compensate the negative charge of the metallated site. Figure 5 ( $\mathbf{e}$ and $\mathbf{g}$ ) confirms that the interlayer space above the metal site in $\mathrm{Hg}(\mathrm{II})\left(\text { GRAND-CSL16CL19 } \mathrm{DL}^{2}\right)_{3}$ is slightly bigger than $\mathrm{Hg}(\mathrm{II})_{\mathrm{S}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$due to the presence of the second water above the metal containing layer that is located
near the helical interface between two of the helical chains. The


Figure 9. Packing comparison of hydrophobic residues around the metal site between $\mathrm{Hg}(\mathrm{II})-(\text { GRAND-CSL16CL19 } \mathrm{L})_{3}{ }^{-}$and $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}($ GRAND CSL16CL30H) $3^{+}$(PDB code: 5KB1) ${ }^{13}$. Packing of residues in the nineteenth position below the metal site, a) 19D-Leu of $\mathrm{Hg}(\mathrm{II})\left(\right.$ GRAND-CSL16CL19pL) ${ }_{3}$, b) $19 \mathrm{~L}-\mathrm{Leu}$ of $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$and c$)$ an overlay between a) and b), which emphasizes the similarity of the bound $\mathrm{S}_{\mathrm{Y}}$ conformers (sticks) in the sixteenth position from the top-down view ( Hg (II) ions and observed waters of both structures are omitted for clarity.) Main chain atoms are represent as ribbon diagrams, 16Cys as sticks, D-Leu and L-Leu as spheres.
axial water observed directly on top of the Hg (II) atom in $\mathrm{Hg}(\mathrm{II})\left(\mathrm{GRAND}-\mathrm{CSL} 16 \mathrm{CL} 19_{\mathrm{D}} \mathrm{L}\right)_{3}{ }^{-}$is within $0.10 \AA$ of the position of water observed in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL3OH })_{3}{ }^{+}$ (Figure 8c). While the hydrophobicity of the layer at the twelfth positions are slightly different, both metallated-structures easily accommodate a water molecule axially above the $\mathrm{Hg}(\mathrm{II})$. Moreover, due to the low symmetry requirement of $\mathrm{P} 2_{1} 2_{1} 2_{1}$ space group, the cavity above the metal site in $\mathrm{Hg}(\mathrm{II})$ (GRANDCSL16CL19DL) ${ }_{3}{ }^{-}$is also amenable for a second water to H -bond with the centrally axial water. These structural comparisons confirm that the space in the layer at the 12th position is suitable for water access, demonstrating that the $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}\left(\mathrm{H}_{2} \mathrm{O}\right)$ species with a water ligand oriented toward the N -terminus in TRIL2WL16CL19DL is reasonable. The significant change in leucine orientations in the layer at the nineteenth position strongly suggests that the 19L-Leu layer in $\mathrm{Hg}(\mathrm{II})_{\mathrm{s}} \mathrm{Zn}(\mathrm{II})_{\mathrm{N}}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$packs tighter than the 19D-Leu in $\mathrm{Hg}(\mathrm{II})\left(\text { GRAND-CSL16CL19 } \mathrm{D}^{2}\right)_{3}{ }^{-}$(Figure 9). It is obvious that the $\mathrm{C}_{\beta}$ deviation of D -Leu forces the whole side chain to be directed more toward the helical interface, generating a larger interlayer space below the metal site compared to the $\mathrm{Hg}(\mathrm{II})_{s} \mathrm{Zn}(\mathrm{II})_{N}(\text { GRAND-CSL16CL30H })_{3}{ }^{+}$. The alignment of both structures demonstrates that the tighter 19LLeu packing in the metallated-L16C peptide would cause drastic steric clashes if waters were to be present as similar to the $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL16CL19 } \mathrm{L})_{3}{ }^{\circ}$. This not only explains the
reasons why the altered D-Leu side chain removes the steric hindrance below the metal binding site, allowing for more water access in TRIL2WL16CL19DL, but also hints to why there is no observation of water below the metal binding site that could bind to $C d$ (II) when the 19L-Leu configuration is placed in the TRIL16C peptide.

## Conclusion

One of the most challenging aspects of de Novo metalloprotein design is developing strategies to control coordination geometry within a protein environment. To achieve this objective, one must not only understand the positioning of first coordination sphere ligands, but also comprehend what features of the outer coordination spheres are necessary to obtain a desired geometry. We have used correlated X-ray crystallographic structures of Hg (II)-peptide complexes to evaluate how steric changes in either the first or second coordination sphere influence the coordination numbers of Cd (II) complexes in 3SCC environments. In some ways, the results are surprising in that they illustrate that a well-reasoned modification may achieve the desired structural result, but for reasons that might not have originally been predicted. An example of this is the ability to control water access to the metal. Clearly, sterics of second coordination sphere side chains influence the available space around the metal site. The predisposition of Cys upon trigonal planar binding allows for an expansion of the interlayer space between the Leu layer (above) and the metal site. The presence of a water molecule in such a cavity of the metallated structure strongly suggests that $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3} \mathrm{O}$ formation is possible when $\mathrm{Cd}(\mathrm{II})$ is bound to the L16C peptide. However, the fourth ligand is only available at a significant price due to the strong hydrophobicity of Leu residues above the metal site. As a consequence, a mixture of 3 - and 4 -coordinate $\mathrm{Cd}(\mathrm{II})$ is formed. The shift to $100 \% 3$-coordinate $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$ can be achieved by reducing the space for water above the metal site. This can be done in two ways. Exploiting the chirality of D-Leu, one can reorient, or "lower", the hydrophobic side chain toward the metal binding plane while keeping the $\operatorname{Cd}(\mathrm{II})$ at the relatively same position within the helical scaffold. As predicted, structural analysis confirms that D-Leu side chains are reoriented toward the C-termini of the structure, causing steric interference above the metal site. However, the second approach, use of Pen ligand to perturb the first coordination sphere ligand, achieves the same objective by an unpredicted structural change. The bulky Pen restricts thiol rotation, causing a shift in the metal plane towards the Leu layer above the site, thus "raising" the metal binding sulfur layer and the $\mathrm{Cd}(\mathrm{II})$ towards the N -termini, which blocks water access. Therefore in the D-Leu case, the roof above the metal site is lowered, whereas in the second substitution with Pen, the floor containing the metal is raised. Both effects diminish space for solvation of the $\mathrm{Cd}(\mathrm{II})$ center, generating a pure $\mathrm{Cd}(\mathrm{II}) \mathrm{S}_{3}$.

In contrast, when the size of the leucine side chain is diminished with alanine, a larger space is generated, which allows for four waters to occupy the newly formed cavity.

Moreover, the structural analysis confirms that the position of DLeu in the outer coordination spheres generates a different steric effect on the metal site, as this lowers the bulky isopropyl groups of L-leucine away from the metal center. The consequence of this change is a trigonal bipyramidal Cd (II) because two cavities are simultaneously available above and below the binding site. These studies provide insights on how to control desired metal geometries in proteins, which are potentially useful for broader applications in future metalloprotein designs.

## Experimental Section

Materials: Fmoc-protected amino acids and the MBHA rink amide resin were purchased from Novabiochem; $N$ hydroxybenzotriazole (HOBt) and 2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU) were bought from Anaspec Inc.; diisopropy- lethylamine (DIEA), acetic anhydride, and pyridine were purchased from Aldrich; piperidine was supplied by Sigma; and $N$-methylpyrrolidinone (NMP) and $N, N$-dimethylformamide (DMF) were obtained from Fisher Scientific.

Peptide synthesis and purification: All peptide variants were synthesized on an Applied Biosystems 433A automated peptide synthesizer with Fmoc-protected amino acids using the standard Fmoc protocol (Applied Biosystems). ${ }^{39}$ The C-terminus of the peptides was amidated on the solid support MBHA rink amide resin ( 0.25 mmole scale) with HBTU/HOBt/DIEPA coupling methods. The N-terminus was acetylated with a solution of $4 \%(\mathrm{v} / \mathrm{v})$ acetic anhydride, $4.3 \%$ ( $\mathrm{v} / \mathrm{v}$ ) pyridine, and 91.7\% $N, N$-dimethylformamide (DMF). The peptides were cleaved from the resin using a cleavage mixture of $90 \%$ trifluoroacetic acid (TFA), 5\% anisole, 3\% thioanisole, and 2\% ethanedithiol for 3.5 hours. The cleaved peptide solution was filtered and evaporated under a dry $\mathrm{N}_{2}$-flow until a glassy film appeared on the surface. Cold diethyl ether was then added to the thin film to obtain a precipitated white crude peptide. This crude was re-dissolved in $\mathrm{ddH}_{2} \mathrm{O}$ and lyophilized to get a fluffy white powder, which was subsequently dissolved in $10 \%$ acetic acid. The peptide was purified by reversed phase HPLC on a Waters 600 Semiprep HPLC peptide C-18 using a linear gradient of $0.1 \%$ TFA in water to $0.1 \%$ TFA in 9:1 $\mathrm{CH}_{3} \mathrm{CN} / \mathrm{H}_{2} \mathrm{O}$ program over 30 mins (flow rate $10 \mathrm{~mL} / \mathrm{min}$ ). The purified peptides were identified by electrospray mass spectrometry. Concentration of peptide stock solutions was determined by quantitation of the cysteine thiols using Ellman's test, which uses dithionitrobenzoate (DTNB) as an indicator. ${ }^{40}$

Crystallizations: All peptides were crystallized by sitting drop vapor diffusion experiments at $20^{\circ} \mathrm{C}$ with drops containing equal volumes of peptide $(0.75 \mu \mathrm{~L})$ and precipitant $(0.75 \mu \mathrm{~L})$ solutions. The $\mathrm{Hg}(\mathrm{II})(\mathrm{GRAND}-\mathrm{CSL} 12 \mathrm{~A} 16 \mathrm{C})_{3}{ }^{-}$crystals were prepared from a peptide solution $(20 \mathrm{mg} / \mathrm{mL}$ GRANDCSL12A16C, 0.92 eq of $\mathrm{HgCl}_{2}$ per 3SCC peptide, 15 mM $\mathrm{Zn}(\mathrm{OAc})_{2}$ and 0.5 mM Tris buffer pH 8.5 ) and a well solution (0.1 M MES pH 6.5 and $25 \%$ (w/v) PEG-1000). The apo-GRAND-CSL12DLL16C was grown from $20 \mathrm{mg} / \mathrm{mL}$ GRAND-

CSL12dLL16C, $15 \mathrm{mM} \mathrm{Zn}(\mathrm{OAc})_{2}$ and 0.5 mM Tris buffer pH 8.5 . The precipitant solution contains $40 \%$ (v/v) PEG-400, sodium acetate buffer pH 4.5 at a final well solution pH 5.4 . The apo-GRAND-CSL16CL19 ${ }_{D}$ L solution was prepared from 20 $\mathrm{mg} / \mathrm{mL}$ peptide, $15 \mathrm{mM} \mathrm{Zn}(\mathrm{OAc})_{2}$ and 0.5 mM Tris buffer pH 8.5. The well solution contains $25 \%(\mathrm{v} / \mathrm{v})$ PEG-2000 MME and 0.1 M MES pH 6.5. The crystals of $\mathrm{Hg}(\mathrm{II})$ GRAND-CSL16CL19DL were crystallized from a peptide solution $(20 \mathrm{mg} / \mathrm{mL}$ GRANDCSL16CL19ㄴ, 0.92 eq of $\mathrm{HgCl}_{2}$ per 3SCC peptide, 15 mM $\mathrm{Zn}(\mathrm{OAc})_{2}$ and 0.5 mM Tris buffer pH 8.5$)$ against well solution (0.2 M Lithium acetate and 20\% (v/v) PEG-3350). Crystals were cryoprotected in a mother liquor containing 20\% glycerol prior to supercooling in liquid $\mathrm{N}_{2}$ for data collection.

Data collections and refinements: Data were collected at the Advanced Photon Source of the Argonne National Laboratory on the LS-CAT Beamline 21-ID-F, equipped with a Mar 225 CCD detector, respectively. All data were collected with a $1^{\circ}$ oscillation then processed and scaled with HKL2000. ${ }^{41}$ All structures presented were solved by molecular replacement using Molrep ${ }^{42}$ in the CCP4 suite of programs ${ }^{43-46}$, then underwent iterative rounds of electron density fitting and refining in Coot ${ }^{42}$ and Buster 2.11.2 program ${ }^{47}$, respectively. The X-ray crystal structures of well-folded, three-stranded parallel coiled coil peptides of apo-(GRAND-CSL12DLL16C) $3_{3}$, apo-(GRANDCSL16CL19DL) $)_{3}$ and $\mathrm{Hg}(\mathrm{II})(G R A N D-C S L 16 C L 19 \mathrm{~L})_{3}{ }^{-}$were determined to $1.34 \AA, 1.83 \AA$ and $1.93 \AA$ resolution, respectively. The crystallographic data for the crystal structures is shown in Table 3. The apo-(GRAND-CSL12DLL16C) ${ }_{3}$ crystallized in the space group R32, contains one single strand of peptide per asymmetric unit with a Matthew's coefficient of 2.38 corresponding to $47.67 \%$ solvent content. The three stranded coiled coil is obtained by the combination of three adjacent symmetric units that are crystallographic imposed by the threefold axis. The structure was solved using a previously published method. ${ }^{48}$ The structure was refined to $1.42 \AA$ ( $R_{\text {working }}=19.6 \%$, $R_{\text {free }}=20.3$ \%).

Sharing similar lattice packing of the R32 space group, the refined apo-(GRAND-CSL12DLL16C) was subsequently employed to be a search model for apo-(GRAND-

CSL16CL19DL) by mutating the 12 D-Leu to L-Leu. 19 D-Leu was replaced after the first round of refinement. The solvent content per asymmetric unit of this structure is $48.60 \%$. The structure was refined to $1.83 \AA$ ( $R_{\text {working }}=20.0 \%, R_{\text {free }}=20.6 \%$ ). The helix of $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12A16C) })_{3}$ was solved by a GRAND-CSL12A16C model from the previously published $\mathrm{Zn}(\mathrm{II})\left(\right.$ GRAND-CSL12A16C) $3^{\circ}$. The structure was refined to 1.93 $\AA\left(R_{\text {working }}=23.14 \%, R_{\text {free }}=25.15 \%\right)$. The Hg(II)(GRANDCSL16CL19DL) ${ }_{3}$, assigned to space group $\mathrm{P} 2_{1} 2_{1} 2_{1}$, was solved using AutoSol Wizard in Phenix. ${ }^{49-51}$ To solve the structure, the anomalous difference of heavy atoms, Hg (II) and Zn (II), was determined to generate the experimental phases. The obtained solution reveals a possible three-stranded coiled coil packing per asymmetric unit, yet the third strand was broken in the middle, missing the residues 19Leu, 20Glu, 21Lys and 22Lys. By using the $2 F_{o}-F_{c}$ electron density as a guide, all missing residues were built back into the chain to generate the final starting model which consequently served as the search model in MolRep(reference). D-Leu at the nineteenth position was replaced with L-Leu after one round of refinement according to the difference density shown in the $F_{0}-F_{c}$ map. The Matthew's coefficient is 4.68 corresponding to $73.74 \%$ solvent. The structure was refined to $1.84 \AA\left(R_{\text {working }}=21.1 \%, R_{\text {free }}=22.6 \%\right)$. The validity of the models were verified using the MolProbity software. ${ }^{52}$ All non-glycine residues of these structures fall in the preferred right handed $\alpha$-helical region of the Ramanchandran plot. Every side chain is present in the preferred rotameric conformation.

Table 3: Data collection and refinement statistics of the crystal structures

| Peptides | apo-(GRANDCSL12DLL16C) ${ }_{3}$ PDB code: 6EGL | apo-(GRAND- <br> CSL16CL19DL) ${ }_{3}$ <br> PDB code: 6EGM | $\mathrm{Hg}(I)$ (GRANDCSL12AL16C) ${ }_{3}$ PDB code: 6EGO | Hg(II) (GRANDCSL16CL19 $\left.{ }^{\text {L }}\right)_{3}{ }^{\text {² }}$ PDB code: 6EGN |
| :---: | :---: | :---: | :---: | :---: |
| Data collection |  |  |  |  |
| Cell dimensions |  |  |  |  |
|  |  |  |  |  |
| a, b, c (Å) | 38.213, 38.213, 140.655 | 37.898, 37.898, 140.667 | 38.186,38.186, 142.385 |  |
| $\alpha, \beta, \gamma\left({ }^{\circ}\right)$ | 90.00, 118.78, 90.00 | 90.00, 90.00, 120.00 | 90.00, 90.00, 120.00 | 2.636, $80.508,88.730$ |
|  |  |  |  | 90.00, 90.00, 90.00 |
| Wavelength ( $\AA$ ) | 0.97872 | 0.97872 | 0.98756 | 0.98756 |
| Resolution $(\AA)^{1}$ | 1.42(1.42-1.40) | 1.83 (1.87-1.83) | 1.84 (1.87-1.84) | 1.84 (1.87-1.84) |
| $\mathrm{R}_{\text {sym }}(\%)^{2}$ | 5.6 (43.4) | 9.4 (48.3) | 6.9 (54.4) | 12.9 (60.8) |


| <l/ $/>^{3}$ | >50 (2) | >50 (2) | > 50 (2) | >50 (2) |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Completeness (\%) ${ }^{4}$ | 99.3 (100) | 99.4 (100) | 98.6 (100) | 97.6 (99.6) |  |
| Redundancy | 5.6 (5.5) | 35.6 (39.8) | 15.8 (12.4) | 8.3 (7.6) |  |
| Refinement |  |  |  |  |  |
| Resolution ( A ) | 1.42 | 1.83 | 1.92 | 1.84 |  |
| R-Factor (\%) ${ }^{5}$ | 19.6 | 20.0 | 23.1 | 21.1 |  |
| $\mathrm{R}_{\text {free }}(\%)^{6}$ | 20.3 | 20.6 | 25.1 | 22.6 |  |
| Protein atoms | 302 | 273 | 318 | 870 |  |
| Metal ions | 1 Zn (II) | 1 Zn (II) | $\begin{aligned} & 1 / 3 \mathrm{Hg}(\mathrm{II}) \\ & 1 \mathrm{Zn}(\mathrm{II}) \text { on surface } \end{aligned}$ | $\begin{aligned} & 1 \mathrm{Hg}(\mathrm{II}) \\ & 3 \mathrm{Zn}(\mathrm{II}) \end{aligned}$ |  |
| Water Molecules | 52 | 44 | 29 | 189 |  |
| Unique Reflections | 8093 | 2584 | 3266 | 20219 | $\square$ |
| R.m.s.d. ${ }^{7}$ |  |  |  |  |  |
| Bonds | 0.01 | 0.01 | 0.06 | 0.01 |  |
| Angles | 1.15 | 1.01 | 0.685 | 1.08 |  |
| MoIProbity Score ${ }^{8}$ | 1.11 | 0.50 | 1.25 | 1.45 |  |
| Clash Score ${ }^{8}$ | 3.17 | 0.00 | 3.09 | 4.20 |  |

[^0]
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Technology, Thonburi, Thailand for funding. Use of the Advanced Photon Source, an Office of Science User Facility operated for the U.S. Department of Energy (DOE) Office of Science by Argonne National Laboratory, was supported by the U.S. DOE under contract no. DE-AC02-06CH11357. Use of the LS-CAT Sector 21 was supported by the Michigan Economic Development Corporation and the Michigan Technology TriCorridor (Grant 085P1000817).

## Accession Codes

apo-(GRAND-CSL12DLL16C) ${ }_{3}$, PDB ID: 6EGL apo-(GRAND-CSL16CL19DL) 3 , PDB ID: 6EGM $\mathrm{Hg}(\mathrm{II})\left(G R A N D-C S L 16 \mathrm{CL} 19_{\mathrm{D}} \mathrm{L}\right)_{3}{ }^{\circ}$, PDB ID: 6EGN $\mathrm{Hg}(\mathrm{II})(\text { GRAND-CSL12AL16C) })_{3}{ }^{\circ}$, PDB ID: 6EGO

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## TOC Figure





[^0]:    ${ }^{1}$ Statistics for highest resolution bin of reflections in parentheses
    ${ }^{2} R_{\text {sym }}=\Sigma_{h} \Sigma_{j}\left|I_{\mathrm{hj}}-\ln _{\mathrm{h}}>\right| / \Sigma_{\mathrm{h}} \Sigma_{\mathrm{j}} \mathrm{l}_{\mathrm{h}}$, where $I_{\mathrm{hj}}$ is the intensity of observation j of reflection h and $<\mathrm{l}_{\mathrm{h}}>$ is the mean intensity for multiply recorded reflections
    ${ }^{3}$ Intensity signal-to-noise ratio
    ${ }^{4}$ Completeness of the unique diffraction data
    ${ }^{5} \mathrm{R}$-factor $=\Sigma_{\mathrm{h}}| | \mathrm{F}_{0}\left|-\mathrm{IF}_{\mathrm{c}} \| / \Sigma_{\mathrm{h}}\right| \mathrm{F}_{0} \mid$, where $\mathrm{F}_{0}$ and $\mathrm{F}_{\mathrm{c}}$ are the observed and calculated structure factor amplitudes for reflection.
    ${ }^{6}$ Rfree is calculated against a $10 \%$ random sampling of the reflections that were removed before structure refinement.
    ${ }^{7}$ Root mean square deviation of bond lengths and bond angles
    ${ }^{8}$ Reference 52

