Review

Designed Metal-ATCUN Derivatives: Redox- and Non-redox-Based Applications Relevant for Chemistry, Biology, and Medicine

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SUMMARY

The designed “ATCUN” motif (amino-terminal copper and nickel binding site) is a replica of naturally occurring ATCUN site found in many proteins/peptides, and an attractive platform for multiple applications, which include nucleases, proteases, spectroscopic probes, imaging, and small molecule activation. ATCUN motifs are engineered at periphery by conjugation to recombinant proteins, peptides, fluorophores, or recognition domains through chemically or genetically, fulfilling the needs of various biological relevance and a wide range of practical usages. This chemistry has witnessed significant growth over the last few decades and several interesting ATCUN derivatives have been described. The redox role of the ATCUN moieties is also an important aspect to be considered. The redox potential of designed M-ATCUN derivatives is modulated by judicious choice of amino acid (including stereochemistry, charge, and position) that ultimately leads to the catalytic efficiency. In this context, a wide range of M-ATCUN derivatives have been designed purposefully for various redox- and non-redox-based applications, including spectroscopic probes, target-based catalytic metalloenzymes, inhibition of amyloid-β toxicity, and telomere shortening, enzyme inactivation, biomolecules stitching or modification, next-generation antibiotic, and small molecule activation.

INTRODUCTION

Over the last few decades, the research on designed metal-ATCUN derivatives (ATCUN; amino terminal copper and nickel binding motif possessing H2N–X-X-His sequence; X = any amino acid) has greatly progressed due to the wide range of applications envisaged, relevant for biology and chemistry, which include DNA (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018), RNA (Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Ross et al., 2017), proteins (Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Pinkham et al., 2018a, 2018b), sugar (Yu and Cowan, 2017a, 2017b), and lipids cleavage (Alexander et al., 2017), antitumor (Kimoto et al., 1983), and antimicrobial activity (Alexander et al., 2018, 2019), inhibition of enzyme activity (Gokhale et al., 2008), telomere shortening (Yu et al., 2015), aggregation of amyloid-β peptides (Zhang et al., 2016), nitration of tyrosine (Maiti et al., 2019), protein-protein cross-linking (Horowitz et al., 2012; Brown et al., 1998), water oxidation (Deng et al., 2018), hydrogen evolution (Kandemir et al., 2016), nitrite to NH3 (Guo et al., 2018), useful in spectroscopic probes (Donaldson et al., 2001; Maiti et al., 2017a, 2017b; Deng et al., 2019a, 2019b; Torrado et al., 1998), and imaging (Miyamoto et al., 2016).

The storyline started in 1960s, when ATCUN motif was first found in human serum albumin (HSA) (Harford and Sarkar, 1997; Laussac and Sarkar, 1984). Since then, it was found in a large number of other naturally occurring proteins/peptides, including BSA (Harford and Sarkar, 1997; Laussac and Sarkar, 1984), hepcidin (Fleming and Sly, 2001), neurokinins C and K (Harford and Sarkar, 1995), human sperm protamine P2a (Mackay et al., 1986), and histatins (Conklin et al., 2017). During this period, the understanding of structure and function of this metal binding ATCUN motif in proteins and peptides was intensively investigated (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Alexander et al., 2018, 2019). The structure of ATCUN motif in albumin is well characterized, and its primary function is assigned to the Cu-transport in blood (Harford and Sarkar, 1997; Laussac and Sarkar, 1984). Therefore, the design of ATCUN motif is of interest for
mimicking the structure and function of naturally occurring proteins and peptides, as well as for other new performances. Many efforts are being made in designing ATCUN motifs with amino acid variants, as well as its conjugation with protein, peptide, or fluorophore for fulfilling the needs of various practical and biological relevant usages (Gonzalez et al., 2018; Pinkham et al., 2018a, 2018b; Donaldson et al., 2001; Maiti et al., 2017a; Deng et al., 2019a, 2019b; Torrado et al., 1998; Miyamoto et al., 2016). The designed parameters include stereochemistry, position, and charge of amino acid, modulating the redox potential of the metal center in M-ATCUN derivatives (M = CuII, NiII, and CoII), leading to catalytic efficiency (Jin and Cowan, 2005, 2007).

In this context, a wide range of applications of M-ATCUN derivatives can be classified into two classes: redox- and non-redox-based applications. However, Cu-ATCUN and Ni-ATCUN derivatives (little information has been so far reported on Co-ATCUN) are mostly introduced for cleavage or modification of biomolecules (such as DNA, RNA, proteins, and amino acids) that are catalyzed by a redox-dependent mechanism (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018). The cleavage of biomolecules by M-ATCUN derivatives is not a random process. This chemistry is designed intentionally, where catalytic center (M-ATCUN) is coupled with the target domain to yield a cocktail complex that selectively binds and oxidatively modifies the therapeutically relevant biomolecules (Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Yu et al., 2015; Joyner and Cowan, 2011). In addition, M-ATCUN derivatives also perform a variety of small molecule activation processes (Deng et al., 2018; Kandemir et al., 2016; Guo et al., 2018).

On the other hand, ATCUN motifs stabilized in a particular oxidation state of metal ion, specially CuII-ATCUN derivatives, are considered as a useful spectroscopic probes such as paramagnetic nuclear magnetic resonance (NMR) (Maiti et al., 2017b), sensor (Torrado et al., 1998), and imaging (Miymoto et al., 2016). Due to high stability of CuII-ATCUN complex, the ATCUN motif can be used as a Cu-chelation therapy, such as inhibition of aggregation of amyloid-ß peptides (Zhang et al., 2016). The combination of ATCUN motif with selected recombinant proteins, peptides, or fluorophores are also used in order to understand the structure and function of biomolecules (Donaldson et al., 2001; Maiti et al., 2017b; Deng et al., 2019a, 2019b; Torrado et al., 1998; Miymoto et al., 2016).

Over the decades, designed M-ATCUN derivatives rely on various redox- and non-redox-based applications, especially spectroscopic probes, target-based catalytic metallodrugs, inhibition of amyloid-ß toxicity and telomerase activity, biomolecules stitching or modification, next-generation antibiotic and small molecules activation. With such a wide range of applications, it is clear that interest in the ATCUN motif is likely to expand in various field in the coming years. Review highlights on the rational design, redox chemistry, and applications of M-ATCUN derivatives in different fields, and viewpoint on this emerging field.

**BIOLOGICAL ROLE OF M-ATCUN MOTIF IN PROTEINS/PEPTIDES**

The metal-binding ATCUN motif is present in many natural proteins and peptides, which plays a fundamental role in metal homeostasis under physiological conditions (Harford and Sarkar, 1997; Laussac and Sarkar, 1984; Fleming and Sly, 2001). HSA is one of the most important copper transport proteins in blood, known as labile Cu pool in extracellular space, and ensures copper homeostasis in human body by utilizing the ATCUN tag (Harford and Sarkar, 1997; Laussac and Sarkar, 1984). Otherwise, dysregulation of Cu homeostasis leads to several number of human diseases, which are WD, cancer, diabetes, and Alzheimer’s disease (AD) (Lowe et al., 2017; Maiti and Moura, 2020). Histamin is one class of natural host defense peptide, possessing ATCUN site that involves in antimicrobial activity (Conklin et al., 2017). Hepcidin-25, an iron-regulatory hormone containing ATCUN site, is responsible for iron homeostasis in mammals (Fleming and Sly, 2001). Another class of peptides with ATCUN site, neuromedin C, a neurotransmitter, acts as a growth factor for some tumors and also plays a key role in metal homeostasis in the central nervous system (Harford and Sarkar, 1995). Therefore, these findings provide inspiration to design synthetic ATCUN derivatives with useful metal-binding properties as well as reactivity.

**DESIGN OF ATCUN DERIVATIVES**

Artificial ATCUN motifs have been devoted considerable attention ever since its discovery in HSA. Decades of investigations have been put forward several interesting ATCUN derivatives, which have been designed purposefully in order to understand the stability, redox chemistry, efficacy, and selectivity. The basic design strategies of ATCUN derivatives are introduction of variable amino acid (X) at first and
second positions in NH$_2$-X-X-His sequence and/or conjugation of ATCUN motif with protein, peptide, or fluorophore (Figure 1). The initial design of ATCUN motif was interested from NH$_2$-terminal protein sequences of albumins (Harford and Sarkar, 1997; Sankararamakrishnan et al., 2005; Lau et al., 1974), and antimicrobial peptides (AMPs) (Alexander et al., 2018, 2019). Variable side chains of amino acids surrounding His residue in ATCUN site significantly influence the activity and metal binding affinity, but they do not directly involve in metal coordination. This design is of interest, specifically, for catalytic activities like DNA, RNA, protein cleavage/modification as described below (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018).

A series of ATCUN motifs are synthesized by introduction of various amino acid residues at first and second positions of ATCUN, such as case-I: simplest amino acid (Gly), case-II: hydrophobic and aromatic amino acids (Phe and Tyr); case-III: hydrophilic and neutral amino acids (such as; Asn and Thr), case-IV hydrophilic and negatively charged amino acid (Val, Asp), case-V: hydrophilic and positively charged amino acids (Arg and Lys) and case-VI: L-to D-amino acid (Jin and Cowan, 2005, 2007; Miyamoto et al., 2016; Kozłowski et al., 1999; Sóvágo and Osza, 2006; Sóvágo et al., 2016; Bal et al., 1996). Generally, these simple and small peptides based ATCUN derivatives are synthesized by a conventional solid-phase peptide synthesis method employing the Fmoc strategy (Neupane et al., 2013).

Dervan et al. first designed ATCUN motif possessing GGH sequence that was the simplest replica of naturally occurring ATCUN motif in proteins or peptides (Mack and Dervan, 1990, 1992; Mack et al., 1988). In this regard, varieties of ATCUN derivatives have been designed. For example, the positively charged amino acid, like lysine (K) and arginine (R) at positions 1 and 2 in ATCUN motif enhances the Cu$^{II}$ binding affinity, whereas negatively charged amino acid, like aspartic acid (D), glutamic acid (E) suppress the Cu$^{II}$ binding.
The above designed ATCUN derivatives are also conjugated with small organic molecules at positions 1, 2, 3 and NH$_2$- (in N-terminus site) with or without a linker moiety. This designed has been, particularly, developed as a sensor (Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Torrado et al., 1998; Wendee and Kulak, 2015; Libardo et al., 2014). Torrado et al. have designed three ATCUN-organic derivatives where organic molecule, S-(dimethylamino)naphthalene-1-sulfonamide (Dns), as a fluorophore has been incorporated into NH$_2$-GGH at NH$_2$- site with a linker (-CH$_2$-, -C$_2$H$_4$- and -C$_3$H$_6$-) (Torrado et al., 1998). Several ATCUN-dns derivatives are reported where Dns group is attached to the vicinity of the ATCUN motif at any amino acid residue (at position, 1, 2, or 3) in ATCUN (Zheng et al., 2002; Choi et al., 2015).

Another strategy is the conjugation of ATCUN motif with protein and/or peptide that is generally synthesized genetically. Usually, a DNA fragment-encoded ATCUN coupled with protein or peptide sequence is amplified by polymerization chain reaction. Afterward, it is ligated into the vector for ATCUN-protein or -peptide expression. ATCUN-fusion proteins or peptides are generally obtained from heterologous overexpression in E. coli (Donaldson et al., 2001; Choi et al., 2015; Pauluta et al., 2007). Mack et al. successfully introduced artificial ATCUN tag into NH$_2$-terminus of the DNA binding Hin recombinase protein genetically (Mack and Dervan, 1990). Based on this concept, several hybrid ATCUN-protein (or peptides) derivatives are reported, which include multi-domain human ubiquitin protein (Donaldson et al., 2001), orange protein (ORP) (Marit et al., 2017b), green fluorescent protein (GFP) (Choi et al., 2015), and AMPs (Alexander et al., 2018). In addition, the redox catalytic M-ATCUN domain is linked with a gene-specific recognition domain (peptide or substrate) that selectively targets the disease-associated gene, like DNA, RNA, and protein, leading to oxidative damage/modification.

Over the last few decades, the designed strategies have been implicated as a potential catalytic metallo-drug toward nucleases and proteins inactivation as described below. The stability of M-ATCUN derivatives, in vivo, is also a key factor for clinical diagnosis because it must reach to target site with its active form (Mjos and Orvig, 2014). Overall, design of ATCUN derivative has been developed for wide applicability either as spectroscopic probe or as catalytic machine for nucleases, modification of biomolecules, and small molecule activation.

**INTERACTION OF ATCUN MOTIF WITH METAL IONS**

The ATCUN motif, a promising metal binding scaffold, binds with metal ions in a specific manner, defining four nitrogens (terminal amine, two intervening peptide bond nitrogen atoms and the histidine N3 nitrogen) coordinated square planar complexes. It is common to know that ATCUN motif interacts mainly Cu$^\text{II}$ and Ni$^\text{II}$ ions. In fact, it can interact with many metal ions, and this interaction depends on the metal ions, deprotonation of the amide groups, and coordination of histidine nitrogen (Burger, 1990). Deprotonation of amide nitrogen in aqueous medium requires strongly basic conditions (pK$_\text{b}$ = 15) in the absence of metal ion. But the process is significantly enhanced in the presence of metal ions, when the imidazole ring of the histidine residue is available to act as an anchor for the metal ion (Bortolus et al., 2010; Sundberg and Martin, 1974). Only few metal ions are able to induce the deprotonation of CONH groups, forming M-ATCUN species at physiological pH. Among them, Cu$^\text{II}$ and Ni$^\text{II}$-ATCUN are formed at physiological pH, but Co$^\text{II}$-ATCUN is formed at slightly higher pH (~9) (Neupane et al., 2013; Hawkins and Martin, 1983). ATCUN motifs have been also shown to bind other metal ions, which are Au$^\text{III}$, Pt$^\text{IV}$, and Pd$^\text{II}$ at very low pH (~1–2) to form square planar complexes as Ni$^\text{II}$, Cu-ATCUN complexes (Best et al., 1997; Kirvan and Margerum, 1985; Lihi et al., 2017). These findings provide a useful information for the synthesis of new M-ATCUN derivatives.
Since HSA is known as the first source of ATCUN motif and participates in the Cu^{II} transport in blood (Harrow and Sarkar, 1997; Laussac and Sarkar, 1984; Carter and Ho, 1994), the understanding of metal-albumin interaction is necessary to be considered in this respect. HSA possesses two specific Cu^{II} binding sites, ATCUN site, and the multimetal binding site (MBS), in which Cu^{II} binds strongly with ATCUN site over MBS (Bal et al., 2013; Al-Harthi et al., 2019). ATCUN motif in albumin not only interacts Cu^{II} or Ni^{II}, but also interacts other metal ions, which are Zn^{II}, V^{IV}, and Cd^{II} (Bal et al., 2013). Based on competition studies, V^{IV}O prefers ATCUN site, while Zn^{II} or Cd^{II} prefers MBS (Al-Harthi et al., 2019).

The stability constant of Cu-ATCUN in HSA and other proteins/peptides is in range, 12 < log $K_{7.4}$ < 15, depending on surrounding amino acid residues of ATCUN (Gonzalez et al., 2018; Rozga et al., 2007). In addition, ATCUN motif shows higher selectivity for Cu^{II} binding over other physiological cations because Cu^{II} may has strong lewis acid character and prefers square-planar geometry. In general, the binding affinity of the ATCUN motif for Cu^{II} is higher (about six order of magnitude) than for Ni^{II}. The binding affinity of some synthetic as well as naturally occurring ATCUNs are tabulated in Table 1 (Gonzalez et al., 2018; Rozga et al., 2007).

### REDOX CHEMISTRY OF M-ATCUN DERIVATIVES

The redox chemistry of M-ATCUN derivatives is a useful index due to its wide variety of catalytic activities. So, intense interests in electrochemical studies on M-ATCUN derivatives are described in literature (Gonzalez et al., 2018; Neupane et al., 2013; Mital et al., 2015; Hureau et al., 2011; Wiloch et al., 2017). The redox potential of M^{n+/n} in M-ATCUN derivatives highly depends on the nature of amino acid at periphery of ATCUN motif and geometry of metal-complexes. For instance, copper exhibits mainly two accessible redox couples, Cu^{II}/Cu^{I} and Cu^{III}/Cu^{II} in various catalytic activities. The Cu^{II}/Cu^{I} is more available in biological system over Cu^{III}/Cu^{II} redox couple (Solomon et al., 2014). The inherent problem between Cu^{I} and Cu^{II} is in coordination chemistry (Solomon et al., 2014; Maiti et al., 2018). For instance, Cu^{II} (Cu^{III} also) adopts a square planar geometry, but Cu^{I} cannot adopt the same geometry as Cu^{II} and it prefers tetrahedral or tri-, bi-coordinated geometry (Solomon et al., 2014; Maiti et al., 2018). Thus, Cu^{II}/Cu^{I} redox couple is highly geometric reorganization, resulting less efficient in redox chemistry, whereas the Cu^{III}/Cu^{II} redox couple involves low reorganization energy, resulting more efficient in redox chemistry. In Cu-ATCUN derivatives, the key point is the Cu^{III}/Cu^{II} redox couple, where Cu^{I} cannot bind at the same coordination site as Cu^{II}.

### Table 1. Stability Constant/Binding Affinity of Cu-ATCUN from Variable Sources

| Natural Protein/Peptide and Synthetic ATCUN Motif | Cu^{II} Binding Sequence | Stability Constant Log $K_{7.4}$ | Binding Affinity (K in M$^{-1}$) |
|-----------------------------------------------|--------------------------|---------------------------------|--------------------------------|
| HAS (Gonzalez et al., 2018; Rozga et al., 2007) | DAH                      | 12.0                            | 1.0 x 10^{12}                  |
| BSA (Rozga et al., 2007)                      | DTH                      | 12.0                            | 1.0 x 10^{12}                  |
| Hepcidin N-term (Plonka and Bal, 2017)        | DTH                      | 14.7                            | 5.0 x 10^{14}                  |
| des-angiotensinogen N-term (Sokolowska et al., 2002) | VIH                  | 13.0                            | 1.0 x 10^{13}                  |
| Human copper transporter (hCTR1-14aa) (Bosak et al., 2018) | MDH                  | 11.0                            | 1.0 x 10^{11}                  |
| Aβ_{42} (Mital et al., 2015)                  | RFH                      | 13.5                            | 3.2 x 10^{13}                  |
| HP2 N-term (Bal et al., 1997)                 | RTH                      | 14.5                            | 3.2 x 10^{14}                  |
| Endostain N-term (Kolozsi et al., 2009)       | HSH                      | 14.5                            | 3.2 x 10^{14}                  |
| GGH-COOH (Hay et al., 1993)                   | GGH                      | 12.4                            | 2.5 x 10^{12}                  |
| DAHK-NH$_2$ (Rozga et al., 2007)              | DAH                      | 13.8                            | 6.3 x 10^{13}                  |
| YH-COOH (Miyamoto et al., 2016)               | YH                       | 14.4                            | 2.5 x 10^{14}                  |
| MDH-NH$_2$ (Bosak et al., 2018)               | MDH                      | 13.1                            | 1.3 x 10^{13}                  |
| MNH-NH$_2$ (Bosak et al., 2018)               | MNH                      | 14.5                            | 3.2 x 10^{14}                  |
The Ni-ATCUN analogs, generally, involve Ni$^{II}$/Ni$^{III}$ redox couple in their potential catalytic applications. The transformation from Ni$^{II}$ (sq. planar) to Ni$^{III}$ (sq. pyramid) involves geometry organization accompanying large gain in ligand field stabilization energy with respect to Cu$^{II}$/Cu$^{III}$ redox couple in Cu-ATCUN system (Bossu et al., 1977). The other analog, Co-ATCUN derivative, generally, shows Co$^{II}$/Co$^{III}$ redox couple in their redox chemistry without geometry organization (both are sq. planar geometry). Such electrochemical studies are performed on variety of designed linear and cyclic ATCUN motifs (Jin and Cowan, 2005, 2007; Neupane et al., 2013, 2014; Mital et al., 2015; Hureau et al., 2011). The redox chemistry of M-ATCUN derivative is influenced by several factors, such as geometry, stability of M$^{III}$ center, variable amino acid, positioning of amino acid in ATCUN motif, and stereochemical orientation of amino acid residues relative to the M-ATCUN equatorial plane.

In this regard, Cowan et al. extensively studied on electrochemistry of a series of designed linear M-ATCUN derivatives (Table 2) (Jin and Cowan, 2007). Both Cu- and Ni-ATCUN derivatives show decrease in their redox potential with an increasing number of Lys residues in motif that influence the enhancement of $\sigma$-donor character relative to that of Gly on the stabilization of the M$^{III}$ center (Jin and Cowan, 2007). The redox potential is also influenced by the position of Lys in ATCUN derivatives. At first position, Lys residue has been greatly extended to stabilize the Cu$^{III}$ oxidation state than second position. The Lys residue at fourth position in Cu-GGHK, does not influence the Cu$^{III}$/Cu$^{II}$ redox potential as seen in Cu-KGH derivative because backbone amide of fourth position Lys does not directly bind to Cu center. In Ni-KGHK derivative, the side chain amine groups of Lys residues (at second and fourth positions) interact at the axial site of Ni$^{III}$ center, leading to more stabilization of Ni$^{III}$. Similarly, incorporation of two Lys at first and second positions in Ni-KKH sequence, the Ni$^{III}$ is effectively stabilized relative to the first or second position alone (Bossu et al., 1977; Murray and Margerum, 1982). So, stabilization of Cu$^{III}$ and Ni$^{III}$ in ATCUN derivatives depends not only on electrostatic interaction but also on the spatial orientation of the amino acid residues. In addition, the redox potential of Cu$^{III}$/Cu$^{II}$-GGH-COOH is increased to ~34 mV upon amidation at remote C-terminal COOH. Similar trend is observed in redox potential of Ni$^{III}$/Ni$^{II}$-GGH-CONH$_2$ (~80 mV with respect to free carboxylate in GGH-COOH) (Jin and Cowan, 2007). Further study with other derivative, KGHK-COOH, upon amidation of the more remote C-terminus, the redox potential is noticeable increased to ~0.9 mV for Cu$^{III}$/Cu$^{II}$-KGHK and ~20 mV for Ni$^{III}$/Ni$^{II}$-KGHK. The switching from L- to D-Lys, the Cu$^{III}$ is more stabilized, due to specific spatial orientation of D-Lys that leads to more negative redox potential corresponding to L-Lys isomers of Cu-ATCUN derivatives (Table 2). This redox effect is also observed in D-isomer, Ni$^{III}$/kkH, but it is significantly lower extent to redox potential (Jin and Cowan, 2007).

Dervan and coworkers first showed that designed GGHG-Hin-recombinase derivative performed DNA cleavage oxidatively (Mack et al., 1988). Therefore, Cu-GGHG derivative is of considerable interest for the study of redox chemistry (McDonald et al., 1997; Green et al., 2004). The Cu$^{II}$/GGHG-COOH shows

| Cu-ATCUN$^a$ | Ni-ATCUN$^a$ | Cu-ATCUN$^a$ | Ni-ATCUN$^a$ |
|----------------|----------------|----------------|----------------|
| Cu$^{II}$-GGH-CONH$_2$ | 1068 (00) | Ni$^{II}$-GGH-CONH$_2$ | 1087 (00) |
| Cu$^{II}$-GKH-CONH$_2$ | 1057 (−11) | Ni$^{II}$-GKH-CONH$_2$ | 1097 (10) |
| Cu$^{II}$-KGH-CONH$_2$ | 1051 (−17) | Ni$^{II}$-KGH-CONH$_2$ | 1099 (10) |
| Cu$^{II}$-KKH-CONH$_2$ | 1017 (−51) | Ni$^{II}$-KKH-CONH$_2$ | 1077 (−10) |
| Cu$^{II}$-KGHK-CONH$_2$ | 1049 (−19) | Ni$^{II}$-KGHK-CONH$_2$ | 1067 (20) |
| Cu$^{II}$-GkkH-CONH$_2$ | 1032 (−36) | Ni$^{II}$-GkkH-CONH$_2$ | 1157 (18) |
| Cu$^{II}$-GkkH-CONH$_2$ | 1033 (−35) | Ni$^{II}$-GkkH-CONH$_2$ | 1067 (20) |
| Cu$^{II}$-GkkH-CONH$_2$ | 993 (−69) | Ni$^{II}$-GkkH-CONH$_2$ | 1047 (−40) |
| Cu$^{II}$-GGH-COOH | 1034 (−34) | Ni$^{II}$-GGH-COOH | 1007 (−80) |

*CV is measured in 25 mM phosphate buffer, pH 7.4 and 0.1 M KCl. (Lower case letters represent D-amino acid).

The Ni-ATCUN analogs, generally, involve Ni$^{II}$/Ni$^{III}$ redox couple in their potential catalytic applications. The transformation from Ni$^{II}$ (Sq. planar) to Ni$^{III}$ (sq. pyramid) involves geometry organization accompanying large gain in ligand field stabilization energy with respect to Cu$^{II}$/Cu$^{III}$ redox couple in Cu-ATCUN system (Bossu et al., 1977). The other analog, Co-ATCUN derivative, generally, shows Co$^{II}$/Co$^{III}$ redox couple in their redox chemistry without geometry organization (both are sq. planar geometry). Such electrochemical studies are performed on variety of designed linear and cyclic ATCUN motifs (Jin and Cowan, 2005, 2007; Neupane et al., 2013, 2014; Mital et al., 2015; Hureau et al., 2011). The redox chemistry of M-ATCUN derivative is influenced by several factors, such as geometry, stability of M$^{III}$ center, variable amino acid, positioning of amino acid in ATCUN motif, and stereochemical orientation of amino acid residues relative to the M-ATCUN equatorial plane.

The redox potential of Cu$^{II}$/Cu$^{III}$-GGH-COOH is increased to ~34 mV upon amidation at remote C-terminal COOH. Similar trend is observed in redox potential of Ni$^{III}$/Ni$^{II}$-GGH-CONH$_2$ (~80 mV with respect to free carboxylate in GGH-COOH) (Jin and Cowan, 2007). Further study with other derivative, KGHK-COOH, upon amidation of the more remote C-terminus, the redox potential is noticeable increased to ~0.9 mV for Cu$^{III}$/Cu$^{II}$-KGHK and ~20 mV for Ni$^{III}$/Ni$^{II}$-KGHK. The switching from L- to D-Lys, the Cu$^{III}$ is more stabilized, due to specific spatial orientation of D-Lys that leads to more negative redox potential corresponding to L-Lys isomers of Cu-ATCUN derivatives (Table 2). This redox effect is also observed in D-isomer, Ni$^{III}$/kkH, but it is significantly lower extent to redox potential (Jin and Cowan, 2007).
the redox potential at 978 mV vs. NHE, which is very close to CuII-GGH-COOH system (~1003 mV vs. NHE) (McDonald et al., 1997). Another important naturally occurring ATCUN sequence, DAHK is found in the blood plasma, HSA (Hureau et al., 2011; Sendzik et al., 2017). The CV trace of designed CuI-DAHK derivative shows a reversible peak at $E_{1/2} = 0.77$ V (vs. AgCl/Ag) corresponding to the CuII/CuI redox couple, but no reduction redox couple, CuII/CuI is observed (Hureau et al., 2011; Guillereau et al., 2007). Therefore, the reduction of CuII to CuI in model peptide, CuI-DAHK is difficult by ascorbate, but it is relatively easier in HSA (Hureau et al., 2011). This unusual redox behavior of Cu-DAHK in serum albumin is assigned to the presence of two close proximity His residues near-by ATCUN motif that facilitates the reduction of CuII to CuI by ascorbate and stabilize it after reduction via His-CuI-His coordination (Haas et al., 2011; Wezynfeld et al., 2016; Hureau, 2012). The bis-His motif has higher magnitude for CuI binding over ATCUN motif (Zheng et al., 2002; Plonka and Bal, 2017). Therefore, ATCUN motif and near by bis-His motif both significantly play the redox interconversion between CuI and CuII (Haas et al., 2011; Wezynfeld et al., 2016; Hureau, 2012).

The redox chemistry is also examined in a series of designed cyclic and linear M-ATCUN derivatives (Neupane et al., 2013). Joshua Kritzer et al. reported a nice work on redox chemistry of cyclic and linear ATCUN derivatives, including effects of macrocyclization (Neupane et al., 2013). The copper bound cyclic- and linear-ATCUN derivatives show quasi-reversible peak with almost the same reduction potential value of CuIII/CuI redox couple (740–810 mV vs. SCE) and no reduction couple, CuII/CuI is observed. In addition, the cyclic Cu-ATCUN derivatives enhance the CuIII/CuII redox cycling (facilitates electron transfer) than linear Cu-ATCUN derivatives (Neupane et al., 2013).

Like Cu- and Ni-ATCUN, Co-ATCUN analogs, including Co-GGH and Co-KGHK derivatives show the redox interplay mainly between CoII and CoI, with relatively low redox potentials values at −119 and −228 mV vs. NHE, respectively (Joyner and Cowan, 2011). This result also supports that upon inclusion of positive charge amino acid (Lys) in ATCUN site, the redox potential is decreased.

**PRODUCTION OF HYDROXYL RADICAL BY M-ATCUN DERIVATIVES**

The understanding of mechanism and efficiency of hydroxyl radical formation by M-ATCUN are an important task because the oxidative cleavage/modification of biomolecules are induced by the hydroxyl radical (•OH) (see below). The formation of •OH is directly associated with redox potential of M-ATCUN derivatives (Jin and Cowan, 2007; Gokhale et al., 2008; Joyner et al., 2011; Fang et al., 2004). Therefore, the •OH production ability of varieties Cu-ATCUN derivatives is examined in presence of H2O2 (oxidant) and ascorbate (reductant) (Jin and Cowan, 2007; Neupane et al., 2013; Joyner and Cowan, 2011). Several experimental observations conclude that the production of •OH by Cu-ATCUN is more efficient in presence of H2O2/ascorbate system with respect to H2O2 or ascorbate alone (Jin and Cowan, 2007; Neupane et al., 2013; Libardo et al., 2014; Joyner et al., 2011; Santoro et al., 2018). For instance, a series of tri-peptide Cu-ATCUN derivatives (DAH, DMB, DSH, DTH, EAH, GGH, GKH, GNH, LKH, NGH, RTH, SMH, and VIH) are designed for testing the •OH production ability in presence of H2O2 and ascorbate. Among them, CuII-GGH shows the highest rate ($52.54 \pm 0.76 \mu M$ min$^{-1}$), whereas EAH shows the lowest rates (1.14 ± 0.04 \mu M min$^{-1}$) of •OH production in presence of H2O2/ascorbate (Asc$^-$) system (Libardo et al., 2014). The tri-peptide system is expanded to tetra-peptide, such as CuII-KGHK that has been also examined for the production of •OH in presence of Asc$^-$/H2O2 and the production rate of it is compared with CuII-GGH. Both Cu-peptides derivatives show the higher rate of ascorbate consumption in presence of H2O2 (turn over number >100 for GGH and 44 ± 7 for KGHK) than absence of H2O2 (turn over number 17 ± 2 for GGH and 11 ± 2 for KGHK) (Jin and Cowan, 2007; Joyner and Cowan, 2011).

Above results conclude that redox dependent mechanism of HO• formation by Cu-ATCUN derivatives is involved in the variable oxidation states (I, II, and III) of copper. At neutral pH, the reduction potential of H2O2/HO•, and ascorbyl radical/ascorbate are 380 mV (vs. NHE), and −66 mV (vs. NHE), respectively (Wood 1988; Borsook and Keighley, 1933). Therefore, H2O2/ascorbate system plays the redox chemistry with M-chelates complexes within +380 to −66 mV reduction potential window. In presence of ascorbate, the generation of •OH by CuII-ATCUN derivatives is proposed through a CuIII/CuI reduction couple, but the high reduction potential value of Cu-GGH (1038 mV vs. NHE) or Cu-KGHK (1058 mV vs. NHE) is incompatible with reduction of CuII-ATCUN derivatives by ascorbate (Jin and Cowan, 2007; Santoro et al., 2018). In this case, the key point is geometry reorganization in CuII/CuI redox couple, where CuI could not bind at the same coordination site as CuII. So, CuI may be a free state or weakly coordinated ATCUN motif, and thereby
Cu likely binds with biomolecules in vivo. This hypothesis is tested on CuII-KGHK, CuII-DAHK, or CuII-FRHD derivatives in presence of CuI chelating agent, bathocuproinedisulfonate (BCS). Upon addition of ascorbate or H2O2, the formation of CuI-BCS2 complex is increased and consequently the production of •OH is decreased or completely seized (Santoro et al., 2018). This observation indicates that the formation of CuI is directly connected with the formation of •OH. Still, it has a space to improve the redox mechanism.

Similarly, Cu-analogs, including Ni-ATCUN and Co-ATCUN derivatives, are often studied for the production of •OH. Ni-ATCUN shows mainly one accessible redox couple, NiIII/NiII, which significantly generates •OH in presence of H2O2/ascorbate system. Both Ni-GGH and Ni-KGHK derivatives show the higher rate of ascorbate consumption in presence of H2O2 (turn over number 40 ± 10 for GGH and 30 ± 8 for KGHK) than absence of H2O2 (turn over number 9 ± 2 for GGH and 9 ± 2 for KGHK) (Jin and Cowan, 2007; Joyner and Cowan, 2011). Unlike Ni- and Cu-ATCUN derivatives, both Co-GGH and Co-KGHK derivatives show the same turn over number (>100) of ascorbate consumption, even in absence of H2O2, due to lower redox potential of CoIII/CoII couple that facilitates •OH production (Joyner and Cowan, 2011).

APPLICATIONS OF M-ATCUN DERIVATIVES

In this context, a wide range of applications of M-ATCUN derivatives can be classified into two classes, – (1) redox-based and (2) non-redox-based applications (Figure 2). M-ATCUN, especially, Cu-ATCUN shows a redox silent component in presence of biological reductant, ascorbate due to its high redox potential (ranging from 0.87 to 1.07 V vs. NHE)2 and inherent geometry problem between CuII and CuI, it make a redox silent component in presence of biological reductant, ascorbate. Interestingly, Cu-ATCUN derivatives have shown to ability the production of HO• under H2O2/AscH•. Therefore, Cu-ATCUN derivatives can be utilized for various applications into two ways: presence of H2O2/AscH• (redox active) and absence of H2O2/AscH• (redox inactive). Based on literature survey, Cu/Ni-ATCUN derivatives are largely exploited toward redox-based applications, whereas Co-ATCUN derivatives are limited. Same redox couple of copper is more intensely used as a sensor probe in fluorescence turn-on/off switch. As a result, copper has received a considerable amount of attention in this field. In this review, we focus on Cu-, Ni-, and Co-based designed metal-ATCUN derivatives and their applications toward non-redox- and redox-based.

Non-Redox-Based Applications

The transition metal ions are extensively used for redox chemistry in a variety of chemical and biological transformations, but their specific stable oxidation state has been also used as a potential spectroscopic probe due to its inherent spectroscopic properties. Metal ions possessing observable spectroscopic features are used as probes in specific techniques including UV-vis, NMR, electron paramagnetic resonance (EPR), Mössbauer, fluorescence quencher and extended X-ray absorption fine structure in many metallo-proteins (Maiti et al., 2017a; Hagen 2006; Holm et al., 1996). For example, CuII is a paramagnetic species whereas CuI is a diamagnetic species. Based on this spectroscopic advantage, CuII-ATCUN state has been studied extensively as a paramagnetic probes and sensors (Donaldson et al., 2001; Maiti et al., 2017b; Deng et al., 2019a, 2019b; Torrado et al., 1998). Significantly, this review is mainly focused on stable CuII-ATCUN derivatives as spectroscopic probe and positron emission tomography (PET) imaging (Miyamoto et al., 2016). In addition, the ATCUN motif acts as a Cu-chelator for amyloid-β (non-redox-based application) (Zhang et al., 2016).

Spectroscopic Probes

Spectroscopic probes have been extensively investigated and widely used in many fields due to their powerful capability of enlightening and elucidating structural and/or functional features of small/macromolecules. Several spectroscopic tools are available in literature. However, only a few specific, efficient, and less toxic designed small metallo-peptide-based probes are available in literature. Nature provides a small ATCUN motif that represents a promising spectroscopic toolbox due to its small size, causing minimal perturbation in protein structure and function. The utility of ATCUN as a spectroscopic probe is studied by artificially introducing it into protein or peptide. This motif binds, specifically, paramagnetic transition metal ions like CuII ion to originate a CuII-ATCUN derivative that is useful for paramagnetic NMR study (Donaldson et al., 2001; Mal et al., 2002; Gaponenko et al., 2000; Brath et al., 2015). This NMR study can provide valuable information on long-distance interactions between metal ions and surrounding groups. This probe is also widely used as a universal fluorescence quencher that is highly suitable for various pharmaceutical and biomedical applications as discussed below.
Paramagnetic NMR Reporter. Paramagnetic-NMR has been acknowledged as a powerful tool for the study of biomolecules when the first time this tool successfully solved the solution structure of metalloproteins (Banci et al., 1994). Paramagnetic-NMR is well established (Clore and Iwahara, 2009), and its probe is widely used to obtain valuable information about the structure and function of biomolecules (Otting 2010). Those proteins in which paramagnetic transition metal ions are already present or the one in which diamagnetic transition metal ions are replaced, can be used as a paramagnetic probe for protein structural studies, including protein-protein interaction or protein-nucleic acid interfaces in folding and unfolding proteins (Cerofolini et al., 2018; Piccioli and Turano, 2015; Balayssac et al., 2008). For other proteins, paramagnetic metal binding scaffold, which is absent, can be introduced either by genetic engineering (Barthelmes et al., 2011) or chemically covalent attachment with wanted proteins or peptides. The short, three-residues Cu II binding ATCUN motif acts as a paramagnetic NMR probe where Cu II ion shows slow electron spin relaxation that causes broadening of proton NMR resonances (Clore and Iwahara, 1999; Viles et al., 1999; Kalverda et al., 1996). Donaldson et al. first successfully genetically introduced artificial ATCUN tag into multi-domain human ubiquitin protein (Phillips et al., 2001; Huang et al., 1999). Cu II-ATCUN-ubiquitin, a paramagnetic probe, is a suitable candidate for extracting long-range distance restraints from NMR study, which is useful for structure refinement (Donaldson et al., 2001). In this regard, Mal et al. had also designed a Cu II-ATCUN-protein model system (Maiti et al., 2017b), where Cu II-ATCUN domain acts as an NMR-probe for understanding the interaction mechanism between protein calmodulin (CaM) and its target serine/threonine protein kinases (CaM kinases). It is a very useful NMR technique. Therefore, it is rapidly growing in a wide array of intermolecular interactions of biomolecules. Yu et al. employed Cu-GCH as a paramagnetic NMR probe that characterized protein-protein interactions (Yu et al., 2009). In 2017, Maiti et al. also reported a paramagnetic NMR probe, Cu II-ATCUN motif that was inserted into N-terminus of ORP for understanding the molybdenum/copper heterometallic cluster assembly in protein pocket (Maiti et al., 2017b). The 1H-NMR study clearly indicates that His 53 is affected by Cu II-ATCUN probe, even though, it is out of paramagnetic region (the distance between His 53 and His 3 is ~27.9 Å). It is only possible when two molecules interact with each other in head-to-tail fashion (Figure 3). This Cu II-ATCUN-ORP derivative provides a model of intermolecular protein-protein interaction (Maiti et al., 2017a, 2017b).

These paramagnetic studies are expected to get more benefit from the diversity of ATCUN-conjugated protein model system, and undoubtedly it is a valuable addition into the NMR toolbox for the characterizing of macromolecular structure-function relation.

Fluorescence Sensors. Fluorescence assay is one of the most important optical analytical tools and has a wide range of applications in the fields of chemistry, biology, clinical diagnosis, food industry, pharmaceutical chemistry, and environmental science because of its high sensitivity, rapid response, and quantitative yield (Fabbrizzi et al., 1996; Krämer 1998). Fluorescence probes are constructed by combination of two
essential components, fluorophore (chromophore), and the quencher (sensor). Fortunately, naturally occurring Cu II binding ATCUN motif acts as a fluorescence quencher due to its paramagnetic character. For example, ATCUN motif in BSA selectively binds Cu II that significantly quenches the fluorescence emission (Tian et al., 2004; Kruck et al., 1976) by protein containing tryptophan residues (typical fluorescence signature of a protein) (Beechem and Brand, 1985). Therefore, this advantage of naturally occurring Cu-ATCUN motif has been employed as a chemo-sensors for detection and screening of certain cellular metal ions, catalytic redox reaction of DNA/RNA cleavage, protease, and bio-imaging.

Several fluorescence probes are designed where fluorophore, 5-(dimethylamino)naphthalene-1-sulfonamide (Dns) is attached to the vicinity of the receptor (Cu II-ATCUN motif) for detection of Cu ions as well as catalytic activity of Cu-ATCUN motif (Zheng et al., 2002). By applying this strategy, Torrado et al. first designed a series of ATCUN motifs (NH₂-G(Xn-Dns)GHG), where amino acid at first position of ATCUN motif was conjugated by dns with spacer (X = -CH₂, and n = 1, 2 and 3). The spacer makes a distance between the fluorophore (Dns) and receptor (Cu II), where shorter spacer quenches fluorescence signal more efficiently (Torrado et al., 1998). Torrado et al. first successfully applied this method in order to understand the fate of copper in catalytic DNA/RNA cleavage (Torrado et al., 1998). To expand the utility of Dns fluorophore, Zhen et al. developed a fluorescent chemosensor (Dns-NH₂-GGHG), where dansyl fluorophore was attached into N-terminal amino group, resulting Dns directly participating in the binding with Cu II.
This model significantly quenches the emission of fluorophore compared to side branch labeling method (Zheng et al., 2002). Young et al. also compared Cu\(^{1+}\) binding affinity in H(K\(^{+}\))HH with other three short peptides (HP(K\(^{+}\))DH, Ac-DH(K\(^{+}\))HD, and HP(K\(^{+}\))DHDH) by fluorescence quenching study (Young et al., 2015). The fluorescent-labeled (Dns) ATCUN probe is also used on CuO-NPs-based colorimetric immunoassay in clinical diagnosis for the detection of prostate-specific antigen (Deng et al., 2019a, 2019b).

This chemo-sensor has been also utilized toward catalytic activity of Cu-ATCUN motif. In this regard, Choi et al. designed a new fluorescence probe, where the fluorophore, GFP was added next to GGH sequence by genetically (Choi et al., 2015). Upon addition of Cu\(^{1+}\), the fluorescence of GFP is quenched by about 85%. In 2015, Wende et al. redesigned a series of ATCUN conjugated fluorophore (R-NH\(_2\)-Dap-β-Ala-His-Ser-Ser-CO\(_2\)H; R = fluorophore = Rhodamine B, dansyl chloride and fluorescein isothiocyanate) derivatives that significantly cleaved DNA through oxidative pathway. The fate of metal ion (Cu\(^{1+}\)), during the redox process, was monitored by fluorescence studies (Wendea and Kulak, 2015).

Protease activity is also detected by fluorescence assay. In this assay, generally, the substrate peptide for protease is attached with a pair of fluorophore-quencher system. Deng et al. (Deng et al., 2018) reported three fluorophore labeled peptides including EVNLDAHFWADK-Dns, and DAHFWADK-Dns (or SGHDEVDK-Dns) as proteases model for β-secretase (as known as precursor of amyloid) (Haass and Selko, 2007) and caspase-3 (central mediator of cell apoptosis) (Thornberry et al., 1997; Boeneman et al., 2009), respectively. In this proteases assay, initially, almost no change in the fluorescence signal intensity is observed even upon addition of Cu\(^{1+}\) in EVNLDAHFWADK-Dns but after cleaving in-between L and D amino acid residues in EVNLDAHFWADK-Dns by β-secretase (Folk and Franz, 2010), Cu\(^{1+}\) binds ATCUN motif (DAH) in fluorophore-labeled fragment (DAHFWADK-Dns), that significantly reduces the fluorescence intensity (Figure 4). This result shows the inhibition of β-secretase activity. As a result, β-secretase can be used as a therapeutic target for AD (see next section). The other proteases model (caspase-3), SGHDEVDK-Dns with Cu\(^{1+}\) exhibits poor fluorescence signal due to the presence of ATCUN motif (SGH) that strongly binds Cu\(^{1+}\) as a quencher. After proteases by caspase-3, the fluorophore-labeled fragment (K-Dns) is separated from Cu\(^{1+}\)-ATCUN and shows good fluorescence signal. This result may be potentially useful as fluorescence imaging for detection of cell apoptosis.

In the living system, copper-homeostasis is a complex process. Prior to entry into cell, Cu is present mainly as Cu\(^{1+}\) state and ensures copper homeostasis under physiological conditions by one of the most important proteins, albumin, which represents as labile Cu pool in human blood (Harford and Sarkar, 1997; Laussac and Sarkar, 1984). This labile copper pool has clinically significance as potential markers of copper-related pathologies (Lowe et al., 2017; Maiti and Moura, 2020). Therefore, measurement of this labile copper pool in biological samples is of considerable interest by fluorescence study. Quite recently, Falcone et al. (Falcone et al., 2020) have reported a turn-off luminescent sensor for Cu\(^{1+}\), which combines ATCUN motif with long-lifetime luminophore, lanthanide (i.e. Tb\(^{3+}\)), enabling Cu\(^{1+}\) detection in biological-like media.

**Inhibition of Amyloid-β Toxicity**

Above section, Figure 4 shows that the model peptide is cleaved to generate ATCUN fragments by β-secretase. It is well known that the formation of amyloid β-peptide (Aβ) by β-secretase is considered as a key step for the AD (Deshpande et al., 2006). The extracellularly released Aβ has different isoforms, which include Aβ\(_{1-40}\), Aβ\(_{1-42}\), and N-truncated Aβ\(_{4-42}\) members. Among them, Aβ\(_{4-42}\) isoforms of Aβ family is high abundance in brains of healthy and AD human, representing a dominant species in AD (Portelius et al., 2010). Other than, Aβ\(_{4-42}\) has strong ability to bind Cu\(^{1+}\), that not only induces Aβ aggregation into cytotoxicity elements but also generates reactive oxygen species (ROS), causing AD (Faller 2009; Jha et al., 2017; Young et al., 2014; Chegnon et al., 2017).

Mital et al. reported that a model peptide, Aβ\(_{1-42}\) binds Cu\(^{1+}\) with binding affinity, \(K_{D_{A}}\) of 3.4 x 10\(^{13}\) M\(^{-1}\), which is stronger than Aβ\(_{1-40}\) (~10\(^{10}\) M\(^{-1}\)) (Mital et al., 2015) but almost same in HSA (~10\(^{13}\) M\(^{-1}\)) (Bosak-Ahmad et al., 2020; Pushie et al., 2019). The electrochemical response of Cu\(^{1+}\)-Aβ\(_{1-42}\) shows a main irreversible peak at 1.04 V (vs. NHE) that is assigned to Cu\(^{2+}\)/Cu\(^{1+}\) redox couple. An additional peak at lower redox potential (~0.04 V vs. NHE) is assigned to the Cu\(^{3+}\)/Cu\(^{2+}\) redox couple that can be mediated by reducing agent like ascorbate (Mital et al., 2015). Note that Aβ\(_{4-42}\) peptide has two copper binding sites,
N-terminal ATCUN site binds Cu II strongly, suggesting redox silent site, and (2) bis-His motif (at 13th and 14th positions), binds Cu II relatively low affinity, suggesting redox active site, that facilitates the reduction of Cu II to Cu I. Upon addition of biological reductant i.e. ascorbate to Cu II in bis-His is reduced to yield His-Cu I-His complex, while the remaining Cu II in ATCUN site is unaffected (Figure 5) (Pushie et al., 2019). Nature designs such type of ATCUN site in Aβ sequence because it may trap Cu II under oxidative stress conditions, but in excess, it crosses the ability to bind Cu II, and thereby develops Aβ toxicity. Therefore, designed ATCUN motifs could be used as Cu-chelators for treatment of AD (Folk and Franz, 2010; Robert et al., 2015). For instance, a designed ATCUN motif, such as DAHK (replica of HSA) has stronger binding affinity of K7.4/C24 for Cu II and shows redox silence against ascorbate (Gonzalez et al., 2018; Rozga et al., 2007). Upon chelation of Cu II by DAHK from Aβ, the reduction of Cu II is inhibited by ascorbate, and thereby retards ROS production.

In addition, the hydrophobic part of Aβ sequence, Aβ16-20 (KLVFF) has strong tendency for Aβ aggregation (Goyal et al., 2017). Therefore, Aβ16-20 represents a promising drug target for the development of the peptide based inhibitor. For instance, designed KLVFF peptide acts as an Aβ inhibitor and has good outcome (Goyal et al., 2017). Several experimental observations conclude that a hybrid complex, Cu-ATCUN-inhibitor strongly suppresses the aggregation of Aβ, as well as ROS formation compared to Cu-ATCUN or inhibitor alone (Zhang et al., 2016; Jensen et al., 2012). In this regard, ATCUN motif is conjugated with peptide inhibitor to generate a bifunctional peptide that has ability to disaggregate Aβ and also to chelate Cu II ions. Faller et al. designed a bifunctional peptide by combination of DAHK with inhibitor (Aβ12-20 or Aβ13-20), that reduces the HO• formation in cell culture (Jensen et al., 2012). Similarly, Yuan et al. designed a bifunctional peptide GGHRYYAFFK, which significantly suppressed the cytotoxicity of the Cu-Aβ complex (Zhang et al., 2016) (Figure 5). Based on this concept, Meng et al. have also reported, recently, a cocktail peptide, RTHLVFFARK, that retards the aggregation of Aβ, and thereby suppresses the cytotoxicity of Cu-Aβ40 in cell culture (Meng et al., 2018). These findings may guide in the development of peptide-based inhibitors for treatment of AD.

**PET Imaging**

The radioactive ⁶⁴Cu II-ATCUN derivatives are thought to be a potential PET candidate for hypoxia (Wadas et al., 2010; Xie et al., 2016; Walke and Ruthstein, 2019). Hypoxia (or oxygen deficiency) is a hallmark of tumor-specific microenvironment (Brown and Wilson 2004). Therefore, PET-imaging of hypoxia is a very
important technique to visualize the target molecule in vivo. For clinical diagnosis, the stability of metallo-peptide in blood plasma and transportation of metallo-peptide to the target site are key factors. Miyamoto et al., designed a series of $^{64}$Cu–ATCUN–octreotide derivatives (ATCUN = YYH, VVH, NNH, TTH, GGH, and DDH), where ATCUN motif is conjugated with tumor-targeting peptide, octreotide (Oct), and tested the stability of these derivatives in blood plasma for medical applications (Figure 6) (Miyamoto et al., 2016). Among them, $^{64}$Cu-YYH–Oct and $^{64}$Cu-VVH-Oct are highly stable in blood plasma (85.8% and 79.1%, respectively, after incubation of 2 hr). The octreotide, analog of somatostatin, has high affinity for somatostatin receptor, which is highly expressed in various tumor cells, mostly neuroendocrine tumor (Martino et al., 2010). Therefore, octreotide is used as a specific molecular target for imaging therapy (Sun and Coy, 2011). Among them, the $^{64}$Cu–YYH–Oct derivative shows the highest stability in blood plasma and reaches to target tumor cell in tumor-bearing mouse model (Miyamoto et al., 2016). Therefore, this cocktail complex, $^{64}$Cu–ATCUN–octreotide derivative is a potential candidate for PET-imaging. Therefore, more research is needed to design a more stable $^{64}$Cu-ATCUN derivative that will be exploited more in clinical practice for hypoxia imaging in future.

**Redox-Based Applications**

Oxidation-reduction processes are considered as a heart of chemical and biological reactions. Nature uses proteins as scaffolds for building in metal cofactors, in order to control life at a minimal energetic cost. Chemists have learnt how to adopt naturally occurring scaffolds as catalysts and also explore for new performances. Beside the spectroscopic probe (above discussion), the designed metal-ATCUN derivatives, especially Cu-, Ni- and Co-ATCUN serve as a suitable catalytic metal-center for various catalytic activities, including DNA, RNA cleavage, protein inactivation (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018), small molecule activation (multi-electrons, multi-protons reactions) (Deng et al., 2018; Kandemir et al., 2016; Guo et al., 2018), as well as modification of biomolecules (protein-protein interaction, nitration of tyrosine) (Maiti et al., 2019; Horowitz et al., 2012; Brown et al., 1998). The catalytic metal center in M-ATCUN derivative works in two ways. One, the catalytic metal center generates ROS, which are mainly focused toward the cleavage or modification of biomolecules. Second, the same catalytic metal center activates the small molecules through the same redox cycle.
DNA Cleavage

Metallo-drug is of significant interest in the development of nuclease (DNA and RNA cleavage) related human diseases at the genetic level, especially, cancer, human immunodeficiency virus (HIV), and hepatitis C virus (HCV) (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Mjos and Orvig, 2014). In 1983, Pauling et al. remarkably investigated (Kimoto et al., 1983) the antitumor activity of the Cu II-GGH (Cu-ATCUN) complex against Ehrlich ascites tumor cells. That investigation raised the interest in metal-ATCUN motifs toward DNA cleavage (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018; Cowan 2001; Detmer et al., 1996; Patwardhana and Cowan, 2001; Pamatong et al., 1996).

Generally, in presence of AsCH⁺/H₂O₂, metal-ATCUN derivatives actively generate ROS such as hydroxyl radical (•OH) that facilitates DNA cleavage. Under physiological conditions, •OH abstracts hydrogen atom from any carbon atoms (C₁, C₂, C₃, C₄, or C₅) of deoxyribose rings to initialize strand-breaks of DNA (Pogożelski and Tullius, 1998) leading to yield variety of cleavage products (Burrows and Muller, 1998) (Figure 7).

In practical applications, the selectivity and efficacy of DNA strand scission are modulated by proper design of ATCUN motif. The nuclease activity of M-ATCUN derivatives could be influenced by some key factors (Figure 8), including overall higher positive charge on M-ATCUN, inclusion of positively charged amino acids (such as Arg or Lys in ATCUN motif), planarity of catalytic site, stereochemical/geometrical orientation (D/L configuration), position and size of amino acid (Jiang et al., 2007). To date, the most common target sites of DNA are the minor and major groove and G-quadruplex (guanine-rich nucleic acid sequences of telomere DNA) (Figure 8)(Jiang et al., 2007).

To meet this DNA cleavage efficiency, varieties of M-ATCUN-derivatives have been developed. In this simple modular design of M-ATCUN, M⁺ ion is in a square planar configuration having two accessible binding faces that promote effective DNA cleavage (Mack and Dervan, 1990, 1992; Jin et al., 2007; Mack et al., 1988; Mahmoudi and Sarkar, 1999; Harford et al., 1996). Binding affinity of M-ATCUN toward DNA highly depends on side chain of amino acid at the periphery of ATCUN motif. When periphery functional groups of entirely designed M-ATCUN derivatives are the same molecular recognition of DNA (e.g. guanidinium, amine, and amide moieties), those M-ATCUN candidates selectively bind and effectively cleave the DNA strands (Harford and Sarkar, 1997; Mack and Dervan, 1992; Jin and Cowan, 2005, 2007; Agbale et al., 2016; Yu and Cowan, 2017a, 2017b; Gonzalez et al., 2018).

Historically, the Ni⁺-GGH derivative was first studied as a potential nuclease candidate (Mack and Dervan, 1992; Liang et al., 1998). Alternatively, nuclease activity was also being tested by Cu⁺-ATCUN derivatives. The intensive work was carried out on DNA cleavage by Cu/Ni-ATCUN derivatives. Long et al. have investigated extensively the nuclease activity by Ni-ATCUN derivatives. These derivatives preferential cleave the minor groove of DNA (at AT-rich regions) but catalytic efficacy depends on the stereochemistry, charge, and position of amino acids in ATCUN motif (Mack and Dervan, 1992; Liang et al., 1998). DNA cleavage chemistry by Cu/Ni-ATCUN derivatives has been also well documented by Cowan et al. (Jin and Cowan, 2005). For instance, the reactivity rate of Cu-KGHK derivative toward oxidative DNA cleavage is higher than the simple Cu-GGH derivative (Jin and Cowan, 2005). The other metal-derivatives, Ni⁺/Cu⁺-KKH
and Ni$^{II}$/Cu$^{II}$-KGHK have been also studied where former M-derivative exhibits slightly higher cleavage rate than other (Jin and Cowan, 2005). This result indicates that the placement of Lys residue at first position in ATCUN has greater impact on cleavage efficiency than second position. It has been also accounted that the first position Lys (Arg analog) in KGH sequence is directed toward minor groove of DNA to generate iso-helical metallopeptide, which facilitates to yield stable DNA-ATCUN adduct, while the second position Lys is directed away from the minor groove of DNA. The slightly greater activity is simply accounted for by the strong electrostatic interaction between DNA (negative charge) and Lys or Arg (positive charge amino acid). In general, both Cu- and Ni-ATCUN derivatives show increase in DNA cleavage activity with increasing number of positive charge amino acid, such as Lys and Arg residues in ATUCN motif, that corroborate with redox potential (Jin and Cowan, 2005, 2007). Therefore, DNA cleavage activity does not depend only on positive charge of side chain amino acid residues but also important for proper orientation between Cu-ATCUN motif and DNA (Jin and Cowan, 2007).

The DNA cleavage chemistry is not a random process where square planar geometry of M-ATCUN interacts at minor groove and abstracts mainly C4'-H from deoxy-ribose ring (Jin and Cowan, 2007; Pushie et al., 2019). The abstraction of C4'-H is facilitated by the formation of transient H bonds between square-planar “edges” of M-RGH (or Lys analog) derivative (H-bond donors, N-terminal N-H protons, imidazole pyrrole N-H, and Arg side-chain) and minor groove of AT-rich regions of DNA (H-bond acceptors; O2 of T and N3 of A) (Figure 9). The DNA cleavage is also highly influenced by the stereochemistry of amino acids (Fang et al., 2004, 2006; Nagane et al., 2001). Switching from L- to D-amino acids at positions 1 and 2 in ATCUN motif, the D-ATCUN derivative improves the nuclease activity but shows lesser selectivity. In comparison with L-ATCUN, the D-ATCUN derivative is sterically lesser hinder complex that allows M-ATCUN deeper insertion into minor groove of DNA, resulting increase the DNA cleavage efficiency (Fang et al., 2006).

Interestingly, Ni-ATCUN has ability to produce another reactive species, oxy-sulfur radical through the same redox cycling between Ni$^{II}$ and Ni$^{III}$ in presence of sulfite/oxygen, that also significantly damages the biomolecules including DNA, RNA, and proteins (Neta and Huie, 1985; Brandt and van Eldik, 1995).

Figure 7. Schematic Representation of M-ATCUN Induces Oxidatively DNA Strand Scission by Sugar-Hydrogen Abstraction to Yield C1'-, or C2'-, or C3'-, or C4'-, or C5'-, Deoxyribosyl Radical.
(modified from Pogozelski and Tullius, 1998). B = Nucleobase.
For instance, Ni-KGH shows the single- and double-stranded DNA damage in presence of Na$_2$SO$_3$ and O$_2$ (Muller et al., 1997).

Cleavage of Telomeric DNA. DNA has another important therapeutic target site, G-quadruplex (G4) that is secondary structure of guanine (G)-rich sequence (TTAGGG) in telomeric DNA (Zahler et al., 1991). This G4-tetrad sequence is not read by the RNA template of telomerase complex (Xie et al., 2016), resulting downregulation of the telomerase activity in normal cells but up-regulation in tumor cells in human (>85%) (Kim et al., 1994; Sfeir et al., 2009). Therefore, G-quadruplex telomeric DNA plays an important role in cancer biology, suggesting as potential anticancer targets (Neidle 2017). To date, a large number of drugs have been published, but these are generally non-selective G4s cleavage drugs (Nadai et al., 2018). Alternative drugs are required for selective G4s cleavage. To achieve this goal, Cowan et al. (Yu et al., 2015) designed a Cu-GGHK-Acr derivative, where Acr (3,6,9-trisubstituted acridine) was a recognition domain of G-quadruplex (Figure 10B) that facilitated to bring the position of catalytic center, Cu-GGHK in close proximity to G4-tetrad through the π-π interaction between acridine ring and G4s surface (Yu et al., 2015; Campbell et al., 2008). So, this derivative selectivity cleaves the DNA sequence of G-quadruplex model (5′-FAM-dATT(TTAGGG)) at A$_1$-G$_2$ and T$_6$-A$_7$ nucleotides sites, resulting inhibition of cancer cell division (Figure 10A) (Yu and Cowan, 2017a, 2017b; Yu et al., 2015). Cu-GGHK-Acr derivative has been also tested on cancer cell (MCF7 and HuH7) and shows significant anticancer activity against both cell lines.

It is expanded to design a series of G-quadruplex telomeric DNA cleavage models that show selective and rapid telomere reduction in cancer cell (Yu et al., 2019). In this assay, three different Cu-ATCUN derivatives (Cu-GGH, Cu-GDH, and Cu-DGH) are selected, and these are conjugated with naphthalene diimide (ND) derivative (as a G-quadruplex recognition moiety) through aspartate side chain (D) or C-terminal carboxylate of ATCUN motif (Figure 10C). Like acridine, ND also interacts with G-tetrad through π-π interaction that facilitates to bring the DNA cleavage moiety, Cu-ATCUN in close proximity to this G-tetrad. In addition, binding affinity of ND derivatives toward G-tetrad is higher (at least two-fold) than acridine isomer, resulting significant inhibition of cancer cell growth (Cuenca et al., 2008). Both Cu-GDH-(p/m)ND derivatives exhibit the highest level of telomere reduction in cancer cells. These (Cu-ATCUN)$_2$-ND derivatives have two Cu-catalytic sites that have different binding patterns on G-tetrad. One Cu-center (ND1) is close to A$_1$-A$_2$, T$_6$-A$_7$, and G$_{20}$-G$_{22}$, whereas the second Cu-center (ND2) is close to G$_{14}$, and G$_{15}$ in G4-tetrad, resulting selective cleavage.
the nucleotide bases (Figure 10A). All (Cu-ATCUN)$_2$DN derivatives are also tested on human cancer cells (MCF7). Among them, (Cu-GDH)$_2$-(p)DN significantly inhibits human cancer cell proliferation. Thus, the above observations emphasize to design efficient nuclease catalyst for the treatment of G4-related diseases.

RNA Cleavage
Like DNA, RNA is another important therapeutic target (Pearson and Prescott, 1997). RNA, a genetic component of retroviruses, is involved in a virus life cycle that causes viral infections, including HIV and HCV. The virus RNA is a complex structure possessing a set of sub-domains that are attractive for potential therapeutic targets. The cleavage of RNA in HIV or HCV by M-ATCUN has been studied through oxidative pathway like DNA (Joyner and Cowan, 2011; Bradford and Cowan, 2012; Jin and Cowan, 2006; Ross et al., 2015). RNA has some degree of resistance to oxidative cleavage with respect to DNA due to the presence of additional 2'-hydroxyl (2'-OH) in sugar-phosphate backbones (Thorpe 2000). This 2'-OH attributes variety of secondary and tertiary structures, which originate a complicated reactivity pattern. However, the oxidative reactivity pattern of RNA has a firm relation with DNA, such as hydrogen abstraction. The oxidation of RNA is initiated by H-atom abstraction in presence of ROS, which is produced by catalytic metal center (Zhao et al., 2018; Crich and Mo, 1997). The selective RNA cleavage is also employed by an analogous strategy of DNA, where RNA cleavage domain (M-ATCUN) is coupled with RNA targeting domain.

HIV RNA Cleavage. HIV-1 gene expression is regulated by two virally encoded proteins, Rev and Tat. HIV-RNA binding protein (arginine rich peptide) is derived from the HIV-1 regulatory proteins, Rev that interacts with the Rev response element (RRE) of HIV-1 to generate the Rev-RRE complex (Battiste et al., 1996). The formation of this complex is an essential step in the viral replication cycle (Frankel and Young, 1998). To achieve this, Cowan et al. have designed antiviral metallodrugs, M-ATCUN-TQARRNRRRR-WRERQR (M = Cu/Ni/Co and ATCUN = GGH and KGHK) that selectively binds and oxidatively cleaves the RRE stem-loop of HIV RNA model (Joyner and Cowan, 2011; Jin and Cowan, 2006). The specific binding pocket (A$_{29}$ and U$_{30}$ nucleotides) and selective cleavage sites (C$_9$, G$_6$, and U$_5$ nucleotides) in RNA are also shown in Figure 11. The adjacent G-G base pair in backbone of RRE RNA creates a specific binding pocket that allows the incoming Rev peptide for specific recognition (Battiste et al., 1996). The oxidative RRE RNA cleavage by Cu-GGH-Rev or Cu-KGHK-Rev derivative is similar to DNA cleavage where C4'-H abstraction is the major product (Joyner et al., 2013a, 2013b). Rationally designed Cu-ATCUN-Rev derivative is a promising candidate for selective cleavage of the RRE site of HIV to inhibit the virus replication cycle. The selective RNA cleavage by designed M-ATCUN derivatives has been well documented but at the same time, the practical requirements of cell delivery and activity in vivo of metallopeptide are also needed. A series of metallopeptides have been designed and evaluated the intracellular delivery and cleavage activity toward
the target HIV-1 RRE RNA in both vitro and vivo (in Escherichia coli and mammalian cell) by cellular fluo-rescence assay.

Figure 12 shows that the plasmid encoded RRE RNA sequence is fused to C-terminal of GFP. N-terminus of Rev peptide (TRQARRNRRRWRERQR) is coupled with ATCUN (GGH) through a linker, Gx (glycine (G)x, x = 0, 1, 2, 4, 6 corresponding peptides are Rev1, Rev2, Rev3, Rev4, Rev5, and Rev6, respectively) (Jin and Cowan, 2007; Hocharoen and Cowan, 2009). Among them, Rev1, Rev2 peptides with shorter linker, and their copper derivatives significantly reduce the cellular expression of GFP (Plasmid encoded GFP-RRE), suggesting optimal RRE RNA cleavage activity. Interestingly, both metal bound or metal-free state of Rev1 and Rev2 derivatives show similar cellular activity, suggesting both metal free derivatives are able to recruit the metal ions from a cell by ATCUN motif.

Another important site-specific RNA cleavage target is the hairpin-loop (stem loop), trans-activation-responsive region (TAR) RNA from HIV-1 that serves as specific binding site for arginine-rich tat protein (RKKRRQRRRPPQ) or Tat derived peptide (Aboul-ela and Varani, 1998). Long et al. reported that all Ni II- Xaa-Gly-His (Xaa = Gly, Lys, or Arg) derivatives selectively cleaved the stem loop of TAR RNA of HIV-1 (Britten et al., 1998).

HCV RNA Cleavage. The designed catalytic metallodrugs, M-ATCUN derivatives also cleave the RNA of HCV selectively. The internal ribosomal entry site (IRES) of HCV RNA is an important component for the life cycle of the HCV. Therefore, HCV IRES RNA containing stem loop IIb and IV (SLIIb and SLIV) are promising therapeutic targets (Figure 13) (Kieft et al., 2001; Lukavsky et al., 2003). Based on previous approach, M-GGH-YrFK derivative (Lower case ‘r’ represents D-arginine) is designed, which selectively binds and cleaves the stem loop-IIb (SLIIb) of the HCV IRES (Bradford and Cowan, 2012). The tera-peptide, YrFK, a dermorphin analog known as DALDA, highly specific µ-opioid receptor agonist (Samii et al., 1994), binds to targeting domain, SL-IIb of HCV IRES (Bradford and Cowan, 2012). The designed metallodrug, Cu-GGH-YrFK-amide derivative is examined on model oligonucleotide (5'-fluorescein-GGCAGAAAGCGU-
CUAGCCAUGGCGUUAGUA UGCC-3 of SL-IIb in vitro and in HCV cell. The cellular replicon assay of CuGGH-YrFK-amide derivative shows enzyme-like turnover ($K_M$ of 0.85 mM and $k_{cat}$ of 0.53 min$^{-1}$). The incorporation of D-Arg (r) into YrFK sequence enhances the binding affinity and reactivity of Cu-GGhYrFK toward SLIIb of IRES RNA. In addition, a designed Cu-GGh-yrfk derivative (lower case indicates the D-form of amino acid) is also evaluated for binding and reactivity patterns against RNA of HCV. Both compounds show the similar HCV cellular replicon assay and high stability in vivo that meet to the US-FDA approved HCV replicon assay (Bradford et al., 2014). Interestingly, Cu-GGh-yrfk, exhibits major cleavage sites, U$_{14}$ and A$_{15}$, whereas Cu-GGH-YrFK exhibits a wider range of cleavage sites A$_6$-G$_9$ of HCV SLIIb RNA (Figure 13). This concept is expanded to design a series of L-analog of Cu-GGH-YrFK derivative where recognition domain, YrFK is shuffled with existing amino acid or substituted other amino acids like A, D, N, and K (YRFK, ARFK, YAKF, YRAK, YRFA, KFRY, DRFK, NRFK, KRFK, and FRFK) in order to understand the impact of each amino acid residue toward SLIIb RNA binding and cleavage efficiency (Ross et al., 2017). Binding affinity and reactivity patterns of each metallo-peptide are modulated by stereochemistry, charge, size and position of amino acids in the targeting domain. For instance, Cu-GGH-YRFK and Cu-GGH-FRFK show the low catalytic efficiency compared to Cu-GGH-YrFK and Cu-GGh-yrfk (Ross et al., 2017).

Another promising therapeutic intervention of metallopeptide is stem loop IV (SL-IV) of HCV IRES RNA. SL-IV possesses ribosome assembly GCAC domain and AUG start codon (Mondal et al., 2008) (Figure 14). The LaR2C peptide, analog of human La protein, binds with SL-IV HCV IRES resulting inhibition of HCV replication (Pudi et al., 2003). The HCV replicon assay of designed metallodrugs, Cu-GGH-KYKETDLLIFKDYFACKNEERK and
Cu-GGH-KYKETDL containing LaR2C peptide (KYKETDLLILFKDDYFAKKNEERK and truncated sequence, KYKETDL of La protein are shown in Figure 14), are examined on model oligonucleotide (5’-fluorescein-GGACCUGCACUAUGAGCAGAAUCC-3’) of HCV IRES SLIV RNA in vitro (Ross et al., 2015). In this replicon assay, both derivatives show significant efficacy. Cu-GGH-KYKETDLLILFKDDYFAKKNEERK exhibits most significant cleavage sites, G15 and A16, whereas Cu-GGH-KYKETDL exhibits the main cleavage site, G8, representing a part of GCAC domain. Still need to develop these drugs for the treatment of human HCV infection.

Inhibition of Protein Activity

Inhibition of enzyme activity by drugs is another novel approach to cure the human diseases. Basically, drug blocks the active site of enzyme to inhibit the enzyme activity, and thereby arrests the enzyme related diseases. The diseases associated proteins such as human angiotensin converting enzyme (ACE) (Gokhale and Cowan, 2005; Joyner et al., 2012; Hocharoen et al., 2013), carbonic anhydrase-I (Gokhale et al., 2008), and sortase-A (Fidai et al., 2014) are potential therapeutic targets. Herein, M-ATCUN is conjugated with substrate inhibitor to generate a cocktail complex that basically plays the dual role: (a) selective binding and (b) oxidative modification of amino acid residues in proteins/enzymes.

Human ACE. The ACE, a Zn-containing metalloenzyme, converts the natural deca-peptide (DRVYIHPFHL) angiotensin I to angiotensin II as a potent vasoconstricting octapeptide (DRVYIHPF) and degradation of vasodilator nanopeptide bradykinin (RPPGSPFR), leading to increase in blood pressure (Crackower et al., 2002). ACE inhibitor such as lisinopril is a potential drug candidate for the treatment of hypertension and heart failure. The binding mode of lisinopril (N2-[(S)-1-carboxy-3-phenylpropyl]-L-lysyl-L-proline) with Zn-site in ACE is shown in Figure 15A (Natesh et al., 2003). Interestingly, the backbone of lisinopril is very close to ACE substrate, Hip-His-Leu that contains benzene group at S1 site, His group at S1’ site, and dimethyl group at S2’ site of lisinopril (Figure 15B).

In general, lisinopril (inhibitor) makes inactive the somatic-ACE-1 (isoform of ACE family) activity by replacing the substrate from metalloenzyme-substrate complex to yield metalloenzyme-inhibitor complex, but it is a reversible process. The alternative of reversible inhibitor (lisinopril) is an M-ATCUN-inhibitor derivative that plays for irreversible inactivation of sACE-1 enzyme by modification of biomolecules through oxidative pathway (Figure 16).
Cowan et al. demonstrated the catalytic inactivation of ACE enzyme by using a designed Cu-KGHK derivative. The attachment of Lys in KGHK is mimicked to the lysine side chain of lisinopril. The inactivation assay of human somatic ACE-1 (Soubrier et al., 1998) is monitored by the cleavage of the fluorogenic peptide, Mca-RPPGFSAFK(Dnp)-OH (Mca; methyl coumarin, Dnp; dinitro-phenyl) (Johnson and Ahn, 2000) as a model of bradykinin. This assay shows the catalytic inactivation (rate constant, $k/C_2 = 2.9 \times 10^{-2}$/min$^{-1}$) of ACE enzyme with pre-incubation of Cu-KGHK derivative under oxidative conditions (Gokhale and Cowan, 2005). Based on this result, a new metallodrug, Cu-GGH-lisinopril is developed that selectively binds somatic ACE-1 with high affinity (Joyner et al., 2012).

The somatic-ACE harbors two homologous N- and C-domains (Hocharoen et al., 2013). The interest is to develop metallo-peptide that selectively binds the particular domain in somatic-ACE. Molecular modeling studies on Cu-GGH-lisinopril within sACE-1 reveals that the Cu in Cu-GGH-lisinopril coordinates nearby D140 in N-terminal domain, whereas E162 or D377 in the C-terminal domain of sACE-1 (Joyner et al., 2012). The catalytic inactivation of each N/C-domain of somatic ACE-1 is examined by designed M-ATCUN-lisinopril derivative (Hocharoen et al., 2013). The M-ATCUN-lisinopril complex shows significant higher catalytic inactivation rate in N-domain over C-domain of somatic ACE, suggesting optimal orientation of M-chelate-lisinopril complex within N-domain compared to C-domain (Hocharoen et al., 2013). These results provide a valuable drug development for other therapeutic targets.

Carbonic Anhydrase. Another Zn-containing metalloenzyme is carbonic anhydrase (CA), which is responsible for the rapid conversion of CO$_2$ to HCO$_3^-$ and protons, involving in various physiological and pathological processes (Supuran et al., 2003). Overexpression or elevated CA is related to various diseases, which include human diabetes, heart failure, cancer, and glaucoma (Supuran et al., 2003; Supuran 2008). So, CA is an important therapeutic target. The activity of CA is suppressed by a substrate inhibitor, sulfonamide derivative through CA(Zn)-sulfonamide complex formation (Figure 17) (Khadikar et al., 2005).
This advantage of sulfonamide derivative, p-aminobenzene-sulfonamide (SLN) is utilized as a recognition domain and it is conjugated with a catalytic domain, Cu-ATCUN (Cu-GGH) to yield a metallo-inhibitor, Cu-ATCUN-SLN. This derivative selectively binds at the active site of human CA-1 and oxidatively modifies the possible close proximity (5–20 Å) amino acid residues, including H 64, H 67, H 200, H 243, W 97, W 123. Interestingly, no modification of Zn coordinated His residues (H 94, H 96 and H 119), as well as no cleavage of protein are observed (Gokhale et al., 2008).

**Sortase A.** Sortase A (SrtA), a membrane-bound bacterial surface adherence protein found in gram positive bacteria (staphylococcus aureus), is a common cause of many serious hospital- and community-acquired infections and cleaves the amide backbone in-between Thr (T) and Gly (G) residues in LPXTG motif of the C-terminal cell-wall sorting signal (Cossart and Jonquieres, 2000). The bacterial infection can be halted by inactivation of SrtA protein (Aboul-ela and Varani, 1998). The designed metallopeptides, Cu-GGHLPETG, Cu-GGHLPET, Cu-GGHLPETG, and Cu-GGHLPET containing SrtA-targeting motif, LPET or LPETG (Fidai et al., 2014; Zong et al., 2004), selectively bind and suppress the activity markedly, resulting inactivation of SrtA protein. The mechanism of enzyme inactivation describes the oxidative modification of the enzyme active site by Cu-GGH catalytic domain. The head (G) of SrtA substrate, LPETG, is directed toward the active site containing conserved amino acid sequence, R 197 - C 184 - H 120. The tail (L) of LPETG is coordinated with Cu-ATCUN catalytic domain that can modify the close proximity amino acid residues, R 197 - C 184 - H 120, but there is no evidence for modification of Gly 167, Val 168, and Leu 169 residues (Figure 18) (Fidai et al., 2014). Interestingly, shortest (GGHLPET) and longest (GGHLPETG) sequences of metallopeptide show a significant higher rate of enzyme inactivation than the moderate-size (GGHLPET) of metallopeptide. The shortest peptide provides the better alignment of the catalytic domain, Cu-ATCUN toward the enzyme active site (R 197 - C 184 - H 120), resulting higher rate of enzyme inactivation, whereas longest peptide provides the catalytic center far away from the enzyme active site, but it gives more solvent exposed catalytic center that produces more ROS. Overall, the shortest metallopeptide shows higher rate of Cys oxidation in active site of enzyme (Bradford et al., 2014; Fidai et al., 2014).

**Peptide-Based Inhibitor of Viral Protease.** The viral protease, NS2B/NS3 (non-structural protein) is an essential for viral replication and maturation, which is a promising pharmaceutical target for flavivirus infections, including west Nile (WNV), japanese encephalitis, yellow fever (YFV), zika (ZKV), and dengue (DENV) viruses (Chappell et al., 2008). NS2B-NS3 protease belongs to serine protease, trypsin-like, with a classic catalytic triad, His-Asp-Ser (Kang et al., 2017; Bazan and Fletterick, 1989). The peptide-based inhibitor therapy is one of the most potential therapeutic strategies for treatment of various viral infections but currently...
no drugs are available in market. To achieve this, the peptide-based inhibitor is designed by introduction of ATCUN (GGH) motif into the N- or C-terminus of WNVP targeting peptides such as naphthoylated, benzoylated, or acetylated (Pinkham et al., 2018a, 2018b). All metallo-peptide derivatives show noticeable decrease in WNVP activity by oxidative modification of amino acids. Particularly, His_{51}-Asp_{75}-Ser_{135} triad in WNVP is essential for substrate binding and enzyme activity (Figure 19) (Pinkham et al., 2018a, 2018b). Overall, the naphthoylated metallo-peptide modifies the active site amino acid residue, Ser_{135} and additionally Thr_{132} and Thr_{134} residues whereas benzoylated metallopeptide modifies the active site amino acid residue, Asp_{75} and additionally Ser_{71}, Lys_{73} and Glu_{74} residues, which ultimately lead to the enzyme inactivation.

Overall, designed metallo-peptide inhibitors (Cu-ATCUN-substrate) give deeper insights on catalytically inactivation of enzymes activity through protein-substrate interactions, as well as oxidative damage the amino acid residues, but it remains in cellular replicon study yet. These results help to develop the drugs in future for antiviral activity in clinical treatments.

**Antimicrobial Activity**

Worldwide human health is threatened by multi-drug resistant pathogens. Currently, naturally occurring AMPs have great potential in drug development due to their wide range of activity toward multi-domain pathogens. Interestingly, the ATCUN motif is also found in naturally occurring AMPs, including histatin, hepcidin, and piscidin, which serve as natural antibiotics (Alexander et al., 2018; Hancock and Sahl, 2006; Zasloff 2002). M-ATCUN derivatives are to be a novel therapeutics agent against multi-domain pathogens. These peptides attack the cell membrane of many pathogens (Gram-negative and Gram-positive bacteria and fungi) through a common mode of action, such as cell membrane disruption (Brogden 2005; Dawson and Liu, 2006). Human saliva protein, histatin is one of the best natural examples of AMP that kills many pathogens, suggesting a promising candidate for drug development (McCaslin et al., 2019; Melino et al., 2006). Melino et al. designed a model of histatin-5 (DSHAKHHGHYKRKFHEHHSHRGY), known as ATCUN-C16 peptide (C16 representing 16 amino acid residues of the C-terminal of histatin-5; DSHAGYKRKFHEHHSHRGY), which shows similar activity of histamine (Melino et al., 2006). Several studies indicate that ATCUN combined AMPs show more active in antimicrobial activity compared to ATCUN-free AMP. For instance, clavanin, a naturally occurring AMP is isolated from the hemocytes of
the marine tunicate styela clava and has different forms, including clavanin-A, -B, -C, and -D. Among them, only clavanin-C contains ATCUN site that can speed up five-fold antimicrobial activity over ATCUN-free clavanin (Lee et al., 1997). Obviously, incorporation of ATCUN motif into AMPs gives extra mileage in mode of action of AMPs that facilitates to oxidatively damage the bacteria cell, unlike to simple electrostatic interaction between positive charge of AMPs and anionic bacterial membrane surface. The modes of action of AMP with and without ATCUN motif are different as shown in Figure 20. Thus, it motivates to develop and design ATCUN-coupled AMPs derivatives. For instance, the designed ATCUN-AMP derivatives (Cu-GGHGWRWYCRNH₂ and Cu-GGHWRWYCRGGK-NH₂) are more efficient compared to the parent AMPs (Joyner et al., 2013a, 2013b).

The antibacterial activity is also influenced by choice of amino acid in ATCUN motif. Three metal binding ATCUN motifs, GGH, VIH, and DAH are selected for study of antibacterial activity, in which Cu-DAH exhibits poor formation of ROS compared to others. The conjugation of Cu-GGH, Cu-VIH, Cu-DAH with AMP, the Cu-DAH-AMP is likely to be less active in antimicrobial activity than others two (Libardo et al., 2014). This approach is expanded to design various M-ATCUN-AMP derivatives (ATCUN: GGH, VIH, and DAH), where AMPs have different binding modes of action such as membrane-disrupting peptides, (anoplin; GLLKRIKTL-NH₂ sequence) (Ifrah et al., 2005; Konno et al., 2001) pro-apoptotic peptide (PAP; KLAKKLAKLAKLAK-NH₂) (Javadpour et al., 1996; McGrath et al., 2013; Kim et al., 2011), and non-membrane-active peptide (sh-buforin; RAGLQFPVGRVHRLRK-NH₂) (Park et al., 1996, 2000). All derivatives are tested on antimicrobial activity against four different bacterial strains. Among them, Cu-VIH-anoplin derivative shows the highest antibacterial activity over others.

This approach is expanded to design an RTH-sh-buforin derivative possessing more positively charged on ATCUN motif that allows high binding affinity toward DNA over neutral ATCUN motif, VIH-sh-buforin. The switching from L- to D-amino acid, the antibacterial activity is enhanced, like nuclease activity (Libardo et al., 2015). To expand the utility of metal binding domain in AMPs, the Cu-ATCUN-OV-3 (OV-3; ovispirin-3 from the cathelicidin family) derivative shows higher rate of antimicrobial activity toward wide range of bacteria relative to OV-3 alone. This derivative also shows the high level of membrane leakage and lipid peroxidation with respect to Cu-GGH or OV-3 alone. These results suggest that the Cu-GGH-OV-3 derivative oxidatively disrupts the bacterial cell membranes, resulting cell death, whereas OV-3 alone is able to permeabilization of cell membrane without lipid oxidation (Alexander et al., 2017). The other AMP, Sub5 is a linear synthetic peptide of bactenecin, which shows a broad range of antimicrobial activity against microbes (Mania et al., 2010). Similarly, Sub5 (RRWKIVVIRWRR-NH₂) lacks the metal binding domain,

Figure 16. Possible Reversible (Simple Inhibitor) vs. Irreversible (M-ATCUN-Inhibitor) Inactivation Mechanism of Metalloproteins-Substrate with Inhibitor
ATCUN motif. Therefore, the addition of Cu-ATCUN motif (GGH and GGHG) into Sub5 to yield Cu-GGHRRWKIVVIRWRR-NH₂ and Cu-GGHGRRWKIVVIRWRR-NH₂ derivatives that significantly enhance (~16 fold) the antimicrobial activity relative to parental Sub5 (Alexander et al., 2019).

Overall, the introduction of the M-ATCUN motif into AMP enhances the antimicrobial activity for a wide range of microbes relative to AMP alone. Therefore, M-ATCUN-AMP can be considered as a potential next-generation antibiotic for clinical treatments.

Modification of Amino Acid Residues

The redox-active amino acid residues, such as Met, Cys, His, Tyr, and Trp, are involved in different biological redox processes (Liu et al., 2014) that include (i) deleterious phenomena, as the one occurring under oxidative stress conditions, where the amino acids oxidation could modify (damage) the protein biological function (Stadtman 1993), and secondly, useful functions, as exemplified by stable or transient tyrosine radical or modified tyrosine radicals that are important for ribonucleotide reductase (Barlow et al., 1983), photosystem (Haumann et al., 1999), cytochrome c oxidase (Yu et al., 2012), and protein-protein cross-linkage (Zhang et al., 2016) and (ii) nitration of tyrosine residues in proteins represents a specific footprint of the formation of reactive nitrogen species (RNS) in vivo (Radi 2013). However, above discussion described that Ni-ATCUN derivatives are actively cleavage the DNA/RNA through NiII/NiIII redox couple. This redox chemistry of Ni-ATCUN derivative has been applied to other performances including protein-protein cross-linking (intermolecular tyrosine-tyrosine cross-linking) (Horowitz et al., 2012) and tyrosine nitration (Maiti et al., 2019).

Protein-Protein Cross-Linking. The protein-protein cross-linking is a useful method to study for probing the multi-protein architectures, including RNase (Gill et al., 1997), bacteriophage Qb (Meunier et al., 2004), adeno-associated viral (AAV) capsids (Horowitz et al., 2012; Campbell et al., 1998; Fancy and Kodadek, 1999). This type of cross-linking process has been also employed in many fields such as biochemical and biomedical research, food processing, tissue engineering (Meunier et al., 2004; Brown et al., 1995). One useful method of cross-linking is oxidation of tyrosine residues to yield di-tyrosine product. Oxidative tyrosine cross-linking is catalyzed by various metalloenzymes (Michon et al., 1997; Sánchez-Ferrer et al., 1995).
or metal-binding peptides (Gill et al., 1997; Brown et al., 1995; Fancy and Kodadek, 1998). For example, Ni-ATCUN with oxidant is a useful protein cross-linking reagent for the study of macromolecular protein assemblies. Brown et al. reported a useful cross-linking reagent, Ni-GGH and MMPP (monoperoxyphthalic acid) that efficiently produced protein-protein cross-linking product, Tyr-Tyr unit through tyrosyl radical pathway (Brown et al., 1995). To expand the usage of this reagent, Ni-GGH is fused with ecotin protein, yielding Ni-GGH-ecotin that oxidatively produces cross-linking product, homodimeric-ecotin as serine protease inhibitor, and it is found in the periplasm of *E. coli* (Chung et al., 1983; McGrath et al., 1991). To confirm the formation of bityrosyl cross-linking product, Person et al. used a model protein, mutage GGH-ecotin (D137Y) that enhanced the 4-fold cross-linking product formation in comparison with wild-type (Person et al., 2001). The Ni-GGH catalytic site in ecotin (ecotin-A) oxidizes the close proximity Tyr127 to yield tyrosine radical that finally reacts with close proximity engineered Tyr137 residue of another ecotin (ecotin-B) molecule to yield cross-linking product, ecotin-A-Tyr127-Tyr137-ecotin-B (Figure 21).

This tyrosine-tyrosine stitching reagent is also effectively used to investigate the structure of virus system where inter-subunit of virus is covalently stitched by the oxidation of adjacent tyrosine residues exclusively (Meunier et al., 2004). The utility of tyrosine cross-linking is expanded toward viral capsid for structural analysis of AAV during infection. Tyr-cross-linked modified AAV showed lower transduction efficiency compared to unmodified capsids (Horowitz et al., 2012). In addition, tyrosine-cross linking product is also found in insulin (Correia et al., 2012). This versatility of di-tyrosine cross-linking reagent is a useful candidate in the development of new biomaterial production for potential implication in medical research in coming years (Kodadek 2002; Petrik 2001; MacBeath and Schreiber, 2000).

**Tyrosine Nitration.** The formation of 3-nitrotyrosine in vivo represents a specific footprint of RNS that is often related to several pathophysiological disorders, such as Alzheimer’s and Parkinson’s diseases (Reyes et al., 2011). The 3-nitrotyrosine is formed in multiple pathways involving variable RNS such as peroxynitrite (ONOO−) or nitrogen dioxide (•NO2) and promoted by transition metal ions (Fe, Cu, and Mn) or metalloproteins (Ferrer-Sueta et al., 2018). Very recently, Maiti et al. have presented an alternative sulfur...
metabolism pathway formation of 3-nitrotyrosine, where Ni II-ATCUN motif is conjugated with ORP to yield NiII-ATCUN-ORP that catalyzes the nitration of tyrosine residue in ORP in the presence of nitrite and sulfite in vitro under biological conditions (Figure 22) (Maiti et al., 2019). To our knowledge, the 3-nitrotyrosine formation by Ni II-ATCUN in the presence of the NO2-/SO32-/CO/O2 system over peroxynitrite or H2O2/NO2 system is the first report. This study represents an alternative pathway formation of 3-nitrotyrosine, which may influence the progression of disease states in vivo involving sulfur metabolism.

Small Molecule Activation

Currently, more attention is given on small molecules activation for current sustainability issues (Meyer and Tolman, 2015; Gutsulyak et al., 2013; Hong et al., 2017). Nature is the master of small molecule activation and designs variety of machines such as nitrogen fixation (Hoffman et al., 2014), and photosynthesis (Yano and Yachandra, 2014), involving one-, two-, or multi-electrons and -protons process. Every transformation has great significance for sustainable energy. Inspired by nature, Chemists have designed peptide-based-metal catalysts that are an emerging catalysts for small molecule activations (Yu et al., 2014; Jones et al., 2007). The redox active M-ATCUN derivatives are primarily studied on biomolecules cleavage/modification. These catalytic systems also serve as small molecule activations, which include water oxidation, hydrogen evolution, and nitrite to ammonium (see below).

Water Oxidation. Water oxidation is a thermodynamically unfavorable chemical process, where oxygen evolution from water involves the loss of four protons and four electrons (2H2O = O2 + 4H+ + 4e−, E° = 1.23 V vs. NHE). This challenging chemical process is performed by the Nature, known as the terminal step of photosystem II (PSII) (Hurst 2010; Liu and Wang, 2012; McEvoy and Brudvig, 2006). The interest is basically on the formation of protons and electrons (production of H2 energy) instead of oxygen evolution, which has been considered as the water splitting process. In this field, the designed various metal-complexes such as peptide based metal complexes have progressed rapidly. For instance, Meyer et al. have reported a Cu-peptide complex (Cu-GGGG; non-ATCUN motif), a square-based with coordinated four N donor ligand like Cu-ATCUN system, which shows good water oxidation by electrochemically with turnover frequency, 33 s−1 but operates at high pH ~ 11 and high oxidation potential at 1.32 V vs. NHE (Zhang et al., 2013).
Switching to Cu-ATCUN, water oxidation is also observed. Malinka et al. reported two mononuclear Cu\textsuperscript{II} complexes with Dap-based peptides, G(Dap)GG-NH\textsubscript{2} (3G) and G(Dap)GH-NH\textsubscript{2} (2GH) (Dap = L-2,3-diaminopropionic acid) that catalyzed water oxidation. In these derivatives, the 2GH facilitates more proton-coupled electrons transfer (PCETs) over 3G derivative in the course of water oxidation, and also improves the turnover frequency (TOF), 53 s\textsuperscript{-1} (23 s\textsuperscript{-1} for 3G) at pH 11 with redox potential at 1.32 V vs. NHE. This PCETs reaction is facilitated by the presence of His residue (Pap et al., 2015; Szyrwiel et al., 2017). Deng et al. reported a series of ATCUN motifs, including neutral as well as positively or negatively charged tri-peptides (GGH, KGH, RTH, DAH), tetra-peptides (GGHG, KGHG, RTHD, DAHF), and C-terminally aminated tripeptides (GGH-NH\textsubscript{2}, KGH-NH\textsubscript{2}) for water oxidation. Interestingly, Cu\textsuperscript{II}-tripeptide derivatives exhibit higher TOF for water oxidation than Cu-tetrapeptide, as well as aminated tripeptides derivatives. Unlike other Cu\textsuperscript{II}-peptide derivatives, the Cu-ATCUN derivatives operate water oxidation at neutral pH and at lower oxidation potential (Szyrwiel et al., 2017).

**Hydrogen Evolution.** Nature utilizes an efficient enzyme, Hydrogenases, that cleave heterolytically the hydrogen molecule in a reversible mode (Maiti et al., 2017a; Lubitz et al., 2014). H\textsubscript{2} production (an alternative source of fossil fuel) from the reduction of aqueous proton is one of the potential sustainable energy routes. Numerous metal-complexes as well as peptide-based metal complexes are reported for H\textsubscript{2} production. Among them, the most promising artificial catalysts of hydrogen evolution are mainly cobalt complexes, which exhibit high turnover numbers (TONs), \( \sim 5 \times 10^4 \) (Sun et al., 2011; Kleingardner et al., 2014; Eckenhoff et al., 2013).
Surprisingly, water-soluble Co-ATCUN represents a new class of hydrogen evolution catalyst and has a similar coordination environment as reported Co-catalysts. Bren et al. reported a small Co-tripeptide model (Co-GGH) complex that produced H$_2$ from proton with 275 TON at nearly neutral pH and at $\sim$600 mV overpotential (Kandemir et al., 2016). This overpotential value is relatively lower than other water-soluble cobalt porphyrin-peptide (850 mV) at pH 7 (Kleingardner et al., 2014) but higher than the popular H$_2$ evolution Co-catalyst, “cobaloxime”, which has lower over-potential (250–300 mV) at nearly neutral pH (Jacques et al., 2009; Bacchi et al., 2014). This interesting feature of Co-GGH derivative is observed due to the presence of adjacent terminal amino group to Co-center, which plays as a proton shuttle during the catalysis process. The depornonation directly connects with pKa value that may influence the overpotential. So, this pKa value can be modulated by modification of ATCUN peptide (Neupane et al., 2013, 2014). At pH 6.5, the hydrogen evolution by Co-GGH derivative is higher, but the stability of the complex is lower, which influences the activity. Recently, Bren et al. have also reported a light-driven reaction, where a CoGGH derivative produces hydrogen from aqueous protons in water at near neutral pH by using a photosensitizer ([Ru(bpy)$_3$]$^{2+}$) and an electron donor candidate (ascorbate) (Chakraborty et al., 2019). The turnover number of hydrogen production is up to 2200 than previous system and this catalyst also shows exceptional longevity (more than 48 hr).

In this regard, the modification of GGH peptide in Co-GGH can be an attractive and promising hydrogen evolution candidate that may hold the current energy issues.

**Reduction of Nitrite to Ammonium.** The reduction of nitrite to ammonium is a multi-electron and multi-proton chemical task (Rosca et al., 2009). The six electrons reduction of nitrite to ammonium is one of the key steps in the biological nitrogen cycle that is catalyzed by cytochrome c nitrite reductase through dissimilatory pathway (Einsle et al., 1999) or the siroheme-containing nitrite reductase through assimilatory pathway (Crane et al., 1995). Important progress of bio-inspired small synthetic model systems for nitrite reductase has been developed but most of these catalysts perform partial reduction of nitrite to nitric oxide (1 electron game) (Timmons and Symes, 2015; Moore and Szymczak, 2015; Hematian et al., 2012) and/or nitrous oxide (2 electrons game) (Uyeda and Peters, 2013). In 2018, Guo et al. reported a Co-GGH model that performed selective reduction of nitrite to ammonium (six-electron and eight-proton game) by electrochemically in aqueous buffer at pH $\sim$7 (Guo et al., 2018). This result indicates that a simple Co-GGH model serves as multi-electrons and multi-protons functional activity of cytochrome c nitrite reductase and siroheme-containing nitrite reductase (Rosca et al., 2009; Einsle et al., 1999; Crane et al., 1995). NiGGH
and CuGGH derivatives are also tested under the same conditions but none of them generates ammonia from nitrite.

The field of small molecule activation by M-ATCUN is still in an early development stage. However, the results obtained, so far, are promising and motivating for the search of new chemical performances.

CONCLUSION AND FUTURE CHALLENGES

This review aimed to shed light on wide and selective applications of designed ATCUN-derivatives involved in multifunctional aspects such as: spectroscopic probes, sensors, target-based catalytic metallo-drugs, biomolecules stitching, inhibition of Aβ-peptide aggregation, and small molecules activation.

The redox chemistry of M-ATCUNs still rises questions but not yet fully clarified, particularly the redox couples, Cu II/CuI and Cu III/CuII involved in Cu-ATCUN derivatives during production of hydroxyl radical under ascorbate/H2O2 system. The Cu II/CuI and Cu III/CuII redox couples with high redox potential, as well as inherent coordination preferences by Cu I and Cu II make still question on redox mechanism as well as catalytic efficiency. The one step progress could be the incorporation of bis-His motif nearby ATCUN (bind site of Cu I), but the problem still remains due to Cu migration from ATCUN to bis-His motif upon ascorbate reduction that may decrease the catalytic efficiency. So, a design strategy needs to be developed, where Cu II and Cu I play redox chemistry in the same framework for enhancing redox activity. The redox chemistry of Cu-ATCUN-substrate adduct (substrate: ascorbate/H2O2/O2) can modulate the redox potential, an aspect to be accounted.

However, the design of hybrid M-ATCUN-recognition derivatives is a useful concept for therapeutic intervention. The interest is to drive the ATCUN derivatives toward cellular targets mainly cytosol and nucleus, and these should be reached to the target position in an active form. In cellular assays, like cis-Pt (Lasorsa et al., 2019) and TTM drug (Maiti and Moura, 2020), ATCUN drugs may also interact with intercellular
proteins, mainly, copper trafficking proteins such as metallothioneins (MTs; as a biological strong Cu(I)-chelators) and glutathione (GSH; as a biological copper reducer) (Boal and Rosenzweig, 2009). Recently, Santoro et al. have reported that Cu(I) and Cu(II) oxidation states of Cu-catalysts during O₂ activation under ascorbate are unstable against GSH/MT system (Santoro et al., 2020). Therefore, intercellular Cu-trafficking proteins are a challenge for the design of Cu-ATCUN complexes as oxidative cleavage catalysts in cells. The other analog, Ni-ATCUN is also a promising drug but problems arise in cellular compartments due to its thiophilic nature.

Recent studies have shown that the novel severe acute respiratory syndrome coronavirus 2, (SARS-CoV-2) directly binds to human ACE2 prior to entry into host cell, causing viral infection (Wrapp et al., 2020; Yan et al., 2020; Maiti 2020). ATCUN motif can be conjugated with a recognition domain (ACE2) of SARS-CoV-2 to yield ATCUN-recognition-domain, which can selectively bind and oxidatively modify the amino acid residues (such as tyrosine) of spike protein and hence it would be a promising drug/therapy against SARS-CoV-2 in future. However, more screenings are required for the development of more efficient drugs in clinical practice.

The M-ATCUN derivatives are also employed for novel applications, which include water oxidation, hydrogen evolution and ammonium synthesis. In these studies, the small M-ATCUN motifs (GGH) were used wherein the second-sphere of protein scaffold was fully absent, which could enhance the catalytic activity. This ATCUN-motif has the ability for second-sphere interactions with the metal active site via carboxylate moiety or N-terminus. In the past decade, the light-driven catalytic reactions have been significantly developing. The conjugation of photosensitizer into ATCUN motif can produce a bioinspired system that can operate the chemical reaction by light as a renewable energy source. Recently, Bren et al. reported a light-driven hydrogen production by using a photosensitizer ([Ru(bpy)₃]²⁺) (Chakraborty et al., 2019). In addition, incorporation of redox active metal ions (such as Ru, Au, and Mn) into the designed ATCUN motif can perform a wide range of vital and challenging chemical transformations. Therefore, there are many challenges in small molecule activation that can be overcome by incorporation of photosensitizer and/or redox active metal ions into the designed ATCUN motif.

In biology, HSA possessing ATCUN site plays a key role for copper transport in human blood but the mechanism of copper transport and extracellular copper delivery by ATCUN tag in HSA are still unclear in vivo. Recently, Bal et al. have demonstrated that a transient two N-coordinated Cu-ATCUN complex is formed within very very short life time (τ₁/₂~100 ms) during the initial stage of the interaction between Cu(II) and ATCUN site, which may be able to maintain the Cu(I)/Cu(II) redox couple and extracellular copper transport/delivery (Kotuniak et al., 2020). As ATCUN motif has high affinity for Cu(II), it can recruitment Cu(II) ions from the labile copper pool, which has clinically significance as potential markers of copper-related pathologies. Therefore, measurement of this labile copper pool in biological samples is a significant interest. Several ATCUN-conjugate fluorescent sensors are reported, which are mainly based on turn-off Cu(II) sensors but the rational design of turn-on Cu(II) sensors is still very challenging in biological medium.

Because of small size, the Cu(II)-ATCUN is likely to introduce into protein as paramagnetic NMR tag for understanding the structure/function relationship of many biomolecules. The paramagnetic relaxation enhancement can be improved by incorporation of lanthanoid ions such as Gd(III) or conjugation of Gd(III) chelates into ATCUN motif.

The redox silent (or high stable) M-ATCUN derivative is another important issue in vivo applications, such as spectroscopic probe, bio-imaging, copper chelation therapy. Thus, high stability and high copper binding affinity of a designed motif can be useful for copper chelation therapy by forming a stable M-ATCUN derivative that significantly prevents the hydroxyl radical formation as well as Aβ-peptide aggregation.

Compared with other metal-motif, the ATCUN motif is a naturally occurring small peptide, less toxic, soluble in biological buffer, and high affinity for Cu(I). It can be rationally designed by choice of amino acids that can tune-up the catalytic activity and redox inertness. Furthermore, the ATCUN motif can be conjugated with a recognition domain that provides target specificity and drug delivery, with reduced toxicity and minimum side effects. Bio-conjugation is not limited to the ATCUN motif. For instance, peptide is conjugated with organometallic groups such as metalloccenes (Chantson et al., 2006), hence representing another popular strategy to modulate activity and cellular uptake. However, the major challenge is intracellular delivery.
to cellular targets. So, ATCUN conjugation would be a promising strategy in future. The other metal-complexes such as [Fe(EDTA)]$^{3-}$ (EDTA = ethylenediaminetetraacetic acid) (Pogożelski et al., 1995), [Cu(phen)$_2$]$^{2+}$ (phen = 1,10-phenanthroline) (Sigman 1986), metalloporphyrins (Mestre et al., 1996), and metal-salen (Czlapinski and Sheppard, 2004) are well-known metalonucleases, but in practical application, these are less selective. Therefore, ATCUN motifs can be introduced over other metal complexes. However, rational design is an ultimate step to overcome the critical challenges encountered in various biological and chemical issues.

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