Studies on Copper and Aβ1-16-Induced Conformational Changes in CAG/CTG Trinucleotide Repeats Sequence

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Abstract. DNA conformation and stability are critical for the normal cell functions, which control many cellular processes in life, such as replication, transcription, DNA repair, etc. The accumulation of amyloid-β peptide (Aβ) and Copper (Cu) are the etiological factors for neurodegenerative diseases and hypothesized that they can cause DNA instability. In the current investigation, we studied copper and Aβ1-16 induced conformation and stability changes in CAG/CTG sequences and found alterations from B-DNA to altered B-conformation. Further, the interaction of the copper and Aβ1-16 with CAG/CTG sequences was studied by molecular docking modeling and results indicated that the interaction of copper and Aβ1-16 was through the hydrogen bond formation between adenine, guanine, and cytosine. This study illustrates the role of the copper and Aβ1-16 in modulating the DNA conformation and stability.

Keywords: Circular dichroism, conformation, neurodegeneration, trinucleotides

INTRODUCTION

DNA contains all the genetic information for cellular function and plays an important role in the life process [1]. DNA is polymorphic in nature that adopts different conformations like B, Z, C DNA, and also alterations in the geometry of the double helix, including the depth and width of the major and minor grooves, is variable in different conformations [1]. The repetitive DNA sequences like triplets are found to be dispersed throughout eukaryotic genomes and trinucleotide repeat regions in the genome have been shown to be unstable and polymorphic [1]. CAG trinucleotide repeats, a unique class of repeats that forms hairpin secondary conformations, and this alters DNA replication and repair [2]. Trinucleotide repeats namely CAG, CTG, CCG sequences are linked to various neurological disorders including Huntington’s disease and myotonic dystrophy [3–5]. It is hypothesized that genome containing long stretches of CAG repeats in the N-terminal region of the huntingtin protein, associated with neuronal cell death [6–9]. Further, studies evidenced that polypeptides such as amyloid-β (Aβ), tau, and prion have
been shown directly interacting with several metal ions in vitro and undergo conformational change [10, 11]. Earlier studies from our lab showed that, Aβ and metals bind to DNA and induce conformational change from B-DNA to an altered B-DNA and Ψ-DNA, which are one of the pathological features of Alzheimer’s disease (AD) [12]. The conformational change from B-DNA to altered B-DNA/Z-DNA/Ψ-DNA plays a critical role in the DNA instability of the genome and associated cellular dysfunctions in brain disorders [13–15]. There are limited reports on Aβ and copper (Cu) induced conformational changes in DNA. The current studies focused on Cu and Aβ induced conformational changes in triplet CAG/CTG sequences to get an insight for their role in neurodegenerative disorders.

MATERIALS AND METHODS

Repeats

(CAG)₅/(CTG)₅ sequence were purchased from MWG Biotech GmbH and dissolved the DNA with the MilliQ water. Further dilutions of DNA were prepared with 5 mM Tris-HCl (pH of 7.4) buffer for the circular dichroism (CD) and UV studies.

Copper chloride (CuCl₂·2H₂O)

Copper chloride dihydrate (Cu) was purchased from Merck Schuchard, and was used without purification. Stock solution (50 mM) was prepared in milliQ water and used for the interaction studies with DNA and also with peptides. The required dilutions were prepared as and when necessary for the experiments.

Aβ₁-₁₆ peptide

Aβ₁-₁₆ fragment with the sequence DAEFRHDS-GYEVHHQK having molecular weight 1956 Da was purchased from M/s USV Peptides and used without further purification.

Circular dichroism studies

CD is used to explore the conformations of nucleic acids. The Cu and Aβ₁-₁₆ induced conformational changes of (CAG)₅ and (CTG)₅ were recorded using a Jasco J-715 spectropolarimeter at 25°C. Quartz cuvets with path length of 1 mm were used for spectral recording in the range between 200 nm to 320 nm with sampling interval at every 1 nm. The spectra recorded were average of four repetitive scans in the absence and presence of different concentrations of the Cu and Aβ₁-₁₆.

UV/Vis absorption studies

The electronic absorption studies of (CAG)₅/(CTG)₅ and its binding ability to Cu, Aβ₁-₁₆, were investigated with a Jasco V-530. Spectrophotometer (Jasco, Japan) equipped with a peltier temperature controller. (CAG)₅/(CTG)₅ samples were dissolved in Tris–HCl buffer (5 mM, pH 7.4) in the absence and presence of Cu and Aβ₁-₁₆ were recorded between 220 nm to 320 nm with a matched set of 1 cm path length quartz cuvettes.

Thermal denaturation studies of CD and UV

Thermodynamic studies of (CAG)₅ and (CTG)₅ in the absence and presence of Cu were measured on a Jasco J-715 spectropolarimeter connected with Model PTC-348WI, Peltier type temperature control system. Samples were recorded in the wavelength between 200 nm and 320 nm by varying the temperature of 20°C to 100°C.

UV–Thermal melting studies of (CAG)₅ repeats in 10 mM HEPES buffer (pH 7.4) in the absence and presence of Cu were measured with Jasco V-530 spectrophotometer equipped with Jasco-ETC-505T temperature controller and cell holder that permits temperature control. Absorbance and Tm was measured at 253 nm from 25°C to 95°C, with a heating rate at 1.0°C/min. For each optically monitored transition Tm of (CAG)₅ was determined as the transition midpoint.

Molecular docking studies

Considering the results obtained in vitro study and to have a better understanding, it was assumed of more significance to perform molecular docking studies to identify the mode of interactions of the Cu and Aβ₁-₁₆ with DNA- CAG and CTG repeats, which further correlate both in silico and in vitro results. The Docking approaches were carried out using discovery studio 3.5 modelling program (DS 3.5); GOLD (Genetic Optimization for Ligand Docking) and Cdocker, A charmm-based molecular dynamics (md) simulated-annealing-based algorithm and a conventional molecular mechanics force field. Initial work involved the crystal structure retrieval and
preparation. The crystal structures, PDBID: 4YN6 of RNA model r (CAG) motif repeat sequence (UUGGGCCAGCAGCAGGUCC)2 and CTG repeat (ATGCTGCAT) PDBID:1MNV retrieved from the Research collaboration for structural bioinformatics Protein Data Bank (RCSB PDB) were used in this study. Crystallization water molecules were removed from the DNA molecules and are optimized for docking by the preparation and function edit Nucleic Acid tool of by DS 3.5. Similarly, the Cu (II) ion (PubChem CID:27099) retrieved form Pubchem database and the 3D coordinates of Aβ1-16 fragment was taken from the NMR structures elucidated [16]. The DNA, ligand molecules; metal ion and Aβ1-16 were minimized prior to docking by using CHARMm force field with potential energy (kcal/mol) of –978.23719. DNA is kept rigid while the ligands are treated as fully flexible and a final minimization step is used to refine the docked possess. The docking tool “GOLD” employs genetic algorithm to search the rotational flexibility of receptor hydrogen’s and ligand conformational flexibility.

The active site with a 6Å radius sphere was defined by selecting define and edit binding site function of DS 3.5. A set of top 10 solutions was explored for favorable and stable interactions. Gold Score function was used to evaluate the docking poses. The individual binding poses of each ligand were identified and their interactions with the DNA were studied. The best and most energetically favorable conformation of each ligand was selected and the interactions tabulated. To further illustrate, the influence of Cu and Aβ1-16 binding effect on conformation in B-DNA structure, Structural overlay approaches was carried between native DNA form and a docked Aβ-Cu bound DNA complexes. The outcomes and orientation of the nucleotide bases/DNA axis and the influences of the DNA major and minor grooves were further evaluated.

RESULTS

Interaction of DNA or nucleotide sequences with Cu and Aβ1-16

CD Studies on the effect of Cu and Aβ1-16 on the conformation of (CAG)5/(CTG)5 sequence

CD spectra’s (CAG)5 and (CTG)5 repeats were monitored in the absence and presence of Cu and Aβ1-16 peptides. The spectra depicted in Figs. 1 and 2 showed typical B-DNA conformation with a characteristic positive band at 275 nm for CAG and 285 for CTG sequence attribute to the base stacking and a negative band at 252 nm (CAG) and 258 nm (CTG) due to the right-handed helicity [17, 18]. Substantial changes in the CD spectra were observed upon interaction with Cu (250 μM) to CAG, the spectra shifted to higher wavelengths exhibited the largest change in the positive band to 288 nm (14 nm) and negative band to 261 nm (9 nm) as shown in Fig. 1. Similar results were noticed in (CTG)5 with 250 μM Cu (b), Cu alone (c).
the absence of Aβ peptide. In the presence of Aβ, the 250 nm and 272 nm bands diminished. The spectral variations in (CAG)$_5$ and (CTG)$_5$ sequence DNA is due to the interaction of Cu and Aβ$_{1-16}$ peptide to a specific binding site and may push the strands of DNA apart to accommodate the incoming Cu and Aβ$_{1-16}$ peptide. The decrease in the magnitude of both positive and negative band could be related to an increase in the winding angle of the DNA helix may be due to the super helical tension caused by Cu and Aβ$_{1-16}$ [19, 20]. The binding of Cu to DNA causes the perturbation of B-DNA and alter the base stacking pattern in the DNA structures of each strand resulting in unwinding of the helices. This process leads to altered B-DNA conformations. These results suggested that there was a DNA conformational transition during incubation with Aβ as well as Cu, and also indicate that the secondary conformation of DNA was disturbed. Earlier reports indicated DNA binding property of Aβ and synuclein leading to the modulation of DNA topology [21, 22]. There were reports on the nuclear localization of Aβ in AD brain and a possible mechanism for Aβ playing a role in DNA conformational change [23]. The present study provides a clue on the mechanism of Aβ$_{1-16}$ and Cu binding to triplet DNA and altering conformation.

**UV Absorption studies on the effect of Cu and Aβ$_{1-16}$ on the conformation of (CAG)$_5$/(CTG)$_5$**

Interaction of Cu with (CAG)$_5$ and (CTG)$_5$ sequences were monitored by UV absorption spectrum. Figure 5 showed the maximum absorbance of (CAG)$_5$ sequence at 253 nm, there was a negligible change in absorbance upon addition of Cu at lower concentration. At higher concentration of Cu, there was an increase in the absorbance (hyperchromicity) with a peak shift to lower wavelength to at 244 nm (blue shift) as a function of Cu concentration. Similar changes were also noticed in (CTG)$_n$ with Cu (Fig. 6). The spectral changes may be attributed to a strong interaction between $\pi-\pi^*$ electron cloud of the interaction of Cu with DNA base pairs.

The Absorption spectra of (CAG)$_5$ and (CTG)$_5$ sequence were monitored by the addition of varying concentrations of Aβ and found that there were significant changes in the absorbance at 260 nm, supporting Aβ and DNA interaction (Figs. 7 and 8). The changes observed in the spectra of DNA reveals that the Aβ interacts with DNA and the changes in conformation may be due to the interaction between the electronic state of the intercalating Aβ peptide with that of the DNA bases [23, 24]. The hypochromism
along with red shift observed in the spectra may be due to the interaction of Aβ to the DNA through intercalation resulting electronic perturbations and twist the helix(winding) in DNA to accommodate the incoming Aβ peptide.

Effect of temperature on the conformation changes of (CAG)/CTG sequence

To determine the effect of temperature on the DNA conformation and stability, we measured CD spectra of CAG and CTG sequences in the absence and presence of Cu from 20°C to 100°C. CD spectra of (CAG) show characteristics of B-DNA (Figs. 9–12). The decrease in the CD bands with complete loss of the negative band with the redshift provided evidence for the disruption of a base-paired structure as the temperature increased. It was saturated to at 80°C. In the presence of 50 μM Cu, similar decrease in the positive and negative bands with the redshift noticed and was saturated to at 70°C (Fig. 11). The effect of temperature on (CTG) spectrum was shown in Fig. 12. As temperature increased, there was a decrease in intensity of positive band and complete loss of the negative band with the blue shift was noticed. Similar pattern of changes were also observed with 50 μM Cu and an isodichroic point was observed at 265 nm. The result indicates that both the bands are sensitive to temperature and the conformational change showed
cooperative and biphasic transition. The melting profile provides insight into the physical attributes of the (CAG)$_5$ and (CTG)$_5$ sequence. At high temperature, the hydrogen bonds between the two complementary DNA strands are broken, and disturb the secondary conformation. The melting of strands ($T_m$) occurs over a narrow temperature range depends strongly on the percentage of the stronger hydrogen bonding present between G-C base pairs, therefore, DNA with a higher G-C content will have a higher $T_m$ [25, 26].

The results of UV absorption spectroscopy showed the $T_m$ of (CAG) sequence as 70°C under experimental conditions. Addition of 500 nM Cu increased the $T_m$ to 73°C, and in the presence of 50 μM, it increased to 80.6°C. However, at 100 μM Cu, $T_m$ decreased to 78.4°C). The increase in $T_m$ at lower concentration of Cu has been attributed to the stabilization of the DNA conformation by binding to phosphate backbone of the DNA, and decrease in $T_m$ at higher concentrations of Cu may be due to the binding of Cu to the bases of (CAG)$_5$ sequence and breaking the hydrogen bonds that held between the G-C bases of the complimentary strands of DNA. DNA-Cu (II) complex is sensitive to higher temperatures and involves complex binding sites between strands and is strong enough to compete effectively and the secondary conformations of each strand are destroyed due to unwinding of the helices. Cu ion perturbs the B-form residues in the base composition of DNA, resulting in destabilization of the DNA double helix [27]. The $T_m$ data of both CD and UV studies suggest that Cu ions bind to both phosphate and nucleotide bases of DNA, unwind the two strands at high temperature, and lower the stability.

**Docking studies**

The systematic molecular docking techniques used in the study may possibly provide a clue toward understanding the specific binding site of the DNA for Cu and Aβ$_{1-16}$. To determine the most possible binding site, interaction mode and binding affinity, docking studies have been performed on Cu and Aβ$_{1-16}$ with DNA- CAG repeat and CTG- DNA.

### Table 1

| Sl NO | Nucleotides in DNA A chain | Aβ$_{1-16}$ amino acids interacting with DNA repeats | Hydrogen Bonds distance in Å
|------|---------------------------|-----------------------------------------------------|------------------|
| 1    | CYT7                      | ARG5                                                | 18.5             |
| 2    | ADE8                      | GLN15                                               | 2.473            |
| 3    | GUA9                      | GLN15                                               | 2.528            |
|      |                           | GLN11                                               | 3.431            |
| 4    | CYT10                     | VAL12                                               | 2.827            |
| 5    | ADE11                     | HIS14                                               | 4.889            |
| 6    | GUA12                     | HIS13                                               | 4.500            |
| 7    | CYT13                     | ASP1                                                | 5.861            |

### Table 2

| Sl NO | Nucleotides in DNA B chain | Aβ$_{1-16}$ amino acids interacting with DNA repeats | Hydrogen Bonds distance in Å
|------|---------------------------|-----------------------------------------------------|------------------|
| 1    | CYT7                      | ASP1                                                | 4.080            |
| 2    | ADE8                      | GLU3                                                | 3.197            |
| 3    | GUA9                      | HIS13                                               | 3.722            |
|      |                           | GLN6                                                | 2.066            |
| 4    | CYT10                     | ASP7                                                | 3.163            |
| 5    | ADE11                     | HIS14                                               | 4.889            |
| 6    | GUA12                     | VAL12                                               | 3.960             |
| 7    | CYT13                     | GLU11                                               | 2.327             |
| 8    | ADE14                     | GLU11                                               | 3.999             |
Table 3
CTG repeat and Aβ₁₆ amino acid interaction data across A chain

| Sl NO | Nucleotides in DNA A chain | Aβ₁₆ amino acids interacting with DNA repeats | Hydrogen Bonds distance in Å
|-------|---------------------------|---------------------------------------------|-----------------
| 1     | CYT4                      | GLN15                                      | 3.071           |
| 2     | THY5                      | TYR10                                      | 5.450           |
| 3     | GUA6                      | GLU3                                       | 2.715           |
| 4     | CYT7                      | GLU3                                       | 3.083           |

Table 4
CTG repeat and Aβ₁₆ amino acid interaction data across B chain

| Sl NO | Nucleotides in DNA B chain | Aβ₁₆ amino acids interacting with DNA repeats | Hydrogen Bonds distance in Å
|-------|---------------------------|---------------------------------------------|-----------------
| 1     | CYT13                     | ASP1                                       | 2.743           |
| 2     | THY14                     | TYR10                                      | 4.956           |
| 3     | GUA15                     | HIS14                                      | 4.185           |
| 4     | CYT7                      | GLU3                                       | 3.083           |

As shown in Figs. 12 and 14 (Tables 1–4), the complexes interact with DNA through partial intercalation at the major groove. The Docking results have been established that, Aβ₁₆ and Cu have favorable DNA-CAG repeat binding affinities across major groove, with a resulting relative binding energy of $\sim$31.9763 kJ mol$^{-1}$ through, CYT7, ADE 8, GUA9, CYT10, ADE11, GUA12, CYT13, ADE14 of A chain, indicating that there are certain hydrogen bonding interaction between the complexes and DNA by forming total of 8 Hydrogen bonds with amino acids. Similarly, 11 hydrogen bonds are formed with CYT7, ADE8, GUA9, CYT10, ADE11, GUA12, CYT13, ADE14 of B chain. The more negative binding energy shows more potent DNA-binding affinity of the Aβ₁₆. Likewise, interaction of Cu specifically with guanine contributes to the changes in the stability of the DNA molecule. Hence, formation of certain hydrogen-bonding interaction and more number of hydrogen bond interactions with nitrogenous bases of B-DNA- (CAG) confirm the interaction of Aβ₁₆ in presence of Cu. Similarly, docking studies of CTG- DNA repeat (ATGCTGCAT) [PDBID: 1MNV] with Aβ₁₆ in the presence of Cu, as evident from the results (Fig. 14 and Tables 3 and 4), that the CTG-DNA repeats also exhibit the promising interaction with the Aβ₁₆ and Cu. To further illustrate the influence of Cu and Aβ₁₆ and their binding effect on conformation in B-DNA structure, the structural overlay was carried out for both the CAG and CTG DNA repeats between native DNA and docked Cu Aβ bound DNA complexes. As the overlay shown in Figs. 13II and 14II depicts that, the base pairs show significant rotations/deviations upon binding of Aβ₁₆ and Cu, while compared to the native form of DNA. The circled nucleotides (Figs. 13 and 14) exhibit the shift in position of the nucleotides. On the other hand, from Fig. 13II, Guanine residues reveal the interaction of Cu ions by forming covalent coordination bonds preferentially to N7 position of guanine and there was no indication of binding of Cu to phosphate oxygen has been observed.

Thus, our Insilico results furthermore confirm and support the perturbation of B-DNA and alter the base stacking pattern in the DNA structures of each strand resulting in altered B-DNA conformations. The binding of Cu to DNA probably causes disturbed complementary interactions of DNA. Thus, our Insilico results are in consistent with the DNA-binding properties as evidenced by CD and UV-Vis Spectroscopic studies.

**DISCUSSION**

The B-DNA conformation is the normal conformation of DNA in cells and is attributed with a majority of biological functions and change in the conformation will alter the major and minor groove properties [28, 29] and thus alters minor transcription binders leading to altered gene expression [30, 31].

Cu is an essential metal for normal CNS development and function. However, excess Cu possesses toxicity through interaction with DNA and proteins [32]. Cells in the specific regions of brain, the basal ganglia and parts of the cortex are very sensitive to the effects of the DNA change [33]. Several observations implicated that accumulation of Cu in the brain can modulate DNA stability/other physiological events, may possibly lead to brain pathology [10, 34–39]. In this study, we provide evidence that, conformational change of (CAG) and (CTG) induced by Cu and Aβ₁₆, may have relevance to the genomic biology of neurodegeneration.

Cu ions have a strong base-affinity and binds to G-C bases, specifically bind to N7 of guanine and N3 of cytosine of (CAG)/(CTG) sequence which was confirmed by docking studied. Coordination of the Cu(II) with N7 of guanine, which approaches the outer sphere of the DNA more closely in the Cu(II)-DNA complex and was related to the positions of the atom that is most susceptible to oxidative damage due to its lowest oxidation potential (dG 1.29 V,
Fig. 13. I) best docked conformations of Aβ_{1-16} and Cu with RNA model r (CAG) motif repeat sequence of (UUUGGCCAGCAGCAGGGUCC), [PDBID: 4YN6]. II) Structural overlay between native DNA form and a docked Aβ-Cu bound DNA complexes. The encircled nucleotide bases display the orientation of nucleotide base pairs/significant rotations across the base pairs/ base stacking pattern of Bound (dark orange) and Unbound forms of DNA helices (Green) indicating the shift in the position upon binding of the Aβ-Cu representing the changes in the altered B-DNA conformations.

Fig. 14. Docking of CTG repeat (ATGCTGCAT) with Aβ_{1-16} and Cu [PDBID: 1MNV]. II) Structural overlay between Native DNA form and docked Aβ-Cu bound DNA complexes. The encircled nucleotide bases display the orientation of nucleotide base pairs/significant rotations across the base pairs. Base stacking pattern of bound (lemon Yellow) and unbound forms of DNA helices (Green) are showing the shift in the position upon binding of the Aβ-Cu representing the changes in the altered B-DNA conformations.
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Conflict of Interest

All the authors hereby declare that they do not have any financial or any other conflict of interest related to the work in this publication.

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