Exploring the binding of resveratrol to an oncogene promoter DNA sequence d(CCAATTGG)$_2$ through multispectroscopic, nuclear magnetic resonance and molecular dynamics studies

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Abstract

We report the interaction of resveratrol with an octamer DNA sequence \( \text{d(CCAATTGG)}_2 \), present in the promoter region of many oncogenes, using a combination of absorption, fluorescence, calorimetric and nuclear magnetic resonance techniques to probe the binding. Resveratrol binds to the duplex sequence with a binding constant \( \text{2.20} \times 10^6 \text{ M}^{-1} \) in absorption studies. A ligand-duplex stoichiometry of \( 2.2:1 \) was obtained with binding constant varying from \( 10^9 \) to \( 10^6 \text{M}^{-1} \) with the concentration of DNA varied in fluorescence titration measurements. Spectral changes indicated external binding of resveratrol to duplex DNA. Circular dichroism data displayed minimal variation suggesting external binding. Melting temperatures of DNA and its 1:1 complex showed a difference of approximately \( 2.25^\circ \text{C} \), which supports the external binding. Nuclear magnetic resonance data showed resveratrol binds to the minor groove region near the AT basepair from the nuclear Overhauser effect spectroscopic cross peaks. Distance restrained molecular dynamics was employed in explicit solvent condition to obtain the lowest energy structure. The complex was stable and retained the B-DNA conformation. Findings in this study identify resveratrol as a minor groove binder to the AT region of DNA and pave the way for exploring resveratrol and its analogues as promising anticancer/antibacterial drug.

Keywords: Resveratrol; Deoxyribonucleic acid; Fluorescence; Nuclear Magnetic Resonance Spectroscopy; Molecular Dynamics
1. Introduction

Deoxyribonucleic acid is one of the most significant molecular targets of therapeutic drugs and plays a significant role in various cellular processes such as replication, transcription and translation. During reproduction, DNA replication takes place and genetic information transferred to the next generation. This process is highly controlled at the molecular level. The new traits formed are governed by the gene sequences of DNA, which varies from healthy to fatal diseases [1,2]. Life-threatening conditions like cancer are a massive threat due to uncontrolled cell growth and cell division. Targeting DNA is a crucial step to stop the replication and transcription which control or inhibit the gene regulatory pathways in various diseases [3,4]. The DNA-drug interaction is of great interest in the various research fields of pharmacological and molecular medicine. Investigating the structural and functional characteristics of biomolecules can provide more insight into the interaction between the two [2]. It can help in understanding the mechanism of action and mode of association between the two. Often, this strategy is used to develop new DNA targeted therapeutic molecules [5]. Many natural and synthetic molecules with pharmacological properties show active binding to DNA [6,7].

Resveratrol (trans-3,4,5-trihydroxystilbene) is a natural stilbene derivative found in plants such as mulberries, peanuts, grapes and also found in plant parts like seeds, petioles, woody parts. It is a polyphenolic phytoalexin, present in many food products like red wine, grape juice etc. [8,9,10]. The basic structure of resveratrol comprises of two phenol rings connected through a double styrene bond. The double bond character is responsible for the isomeric state that exists in cis- and trans-form of resveratrol. Trans-form is the most stable conformation of resveratrol and is widely used for various therapeutics and also occurs in medical preparations [10,11]. It can regulate several pathways which are involved in cell proliferation, apoptosis and inflammation [12]. Various cell culture studies reveal that resveratrol can reduce cell cycle progression and can
activate apoptotic cell death [9,13-15]. Many researchers demonstrated that resveratrol induces the apoptotic activity of p53 [16,17].

Interaction of resveratrol with different DNA sequences for sequence-specific binding mode has been reported. Spectroscopic studies showed resveratrol binds to calf thymus DNA through intercalation. While docking studies showed intercalation as well as groove binding. It is also said that in intercalative mode, it is surrounded by ATTATT segment while in groove binding, the surrounding nucleotides are AATAAT [8]. Resonance light scattering techniques showed that resveratrol could be used for the detection of traces of DNA, based on its DNA binding property [9]. It has been demonstrated that resveratrol inhibited the growth of preneoplastic lesion in carcinogen-treated mouse mammary glands. Also, it hindered tumorigenesis in mouse skin cancer model [18]. Usha et al. showed that resveratrol could modulate DNA intercalation of dyes and protect the DNA from the damage induced due to the planar dyes, thereby reducing the DNA-directed toxicity [19]. Binding constants were calculated using absorption, and Fourier transform infrared (FTIR) spectroscopy for the drug-DNA and drug-RNA complexes. Partial stacking and H-bond formation with DNA/RNA is responsible for the antineoplastic activity of resveratrol [20].

Short sequences such as TATA, CCAAT, GC etc. are present in specific regions of promoters and enhancers which activates polymerase II enzyme transcribed mRNA genes. Few such stretches like TATA, GC and CCAAT boxes occur at extremely high frequency [21]. The CCAAT box was the first element identified [22]. Later studies established that such pentanucleotide sequences are present in a wide variety of vertebrate, yeast and plant promoters and are essential for transcription. All five nucleotides are almost invariably conserved. CCAAT sequences can be found both in the direct and in the inverted orientation, and it is present in both TATA containing and in TATA-less promoters. Statistical analysis of 500 unrelated proteins
revealed that CCAAT pentanucleotide is present in 30% of them. Many sequence-specific regulators recognize these short nucleotides. NF-Y is a nuclear protein which recognizes the pentanucleotide CCAAT. The binding site of CCAAT-enhancer binding protein (c/EBP) contains the CCAAT nucleotides in the intervening sequences. CCAAT transcription factor (CTF/NF-1) binds to viral and cellular promoters, recognize sequences containing CCAA nucleotides [15,23]. Borghini et al. [24] pointed out the lack of conventional TATA boxes and the presence of two identical CCAAT boxes as the crucial elements involved in the transcription regulation of the human TLX3 gene which is overexpressed in T-cell Acute Lymphocytic leukaemia.

The palindromic CCAATTGG sequence is present in promoters of human, fly and rat genomes which regulate the activity of several growth factors, oncogenes and tumor suppressor genes. Fibulin-5 gene (FBLN5) promoter contains palindromic sequence CCAATTGG at upstream of the first translation initiation codon, which functions as a potential CCAAT/enhancer binding element [25]. The CCAATTGG motif occurs in BEN (BANP, E5R, and NAC1) domain promoter of drosophila insensitive (Insv) gene which is responsible for its neural development and is widely distributed in metazoans and viruses [26]. The mammalian BEN5/6, RBB and BTB domain protein also recognize a CCAATTGG palindromic motif with homologous Insvfactor[26-29]. The dyad-symmetric CCAAT core sequence is found in the CCAATTGG motif of rat argininosuccinatelyase (AL) promoter at the position of -80 base pair which involves in arginine biosynthesis and expressed in a high level in the liver [30]. Nuclear Factor-Y (NF-Y) similar to transcription factor contains two subunits that show good affinity towards CCAAT central motif in several promoters. Point mutations or spaces in the half-site of this motif inhibit the binding of NF-Y to CCAATTGG palindromic sites. This finding establishes that NF-Y recognizes the correct reading frame dyad symmetric CCAAT core motif [31]. Literature reveals
that the CCAAT sequence is vital in many cellular processes that involve cell proliferation and cell integrity [32-35].

Different groups have studied the interaction between DNA sequences comprising CCAAT segment with small molecules. In a sequence-specific interaction study, distamycin was found to bind at the central portion in the minor groove of duplex DNA d(GGCCAATTGG)_2 [36]. Crystallographic analysis of distamycin-d(CGCAAATTTGCG)_2 complex showed the preference of distamycin to the AT rich region, in the minor groove of DNA [37]. Another molecule netropsin displayed a strong affinity to the central CAATTG segment of the sequence d(GGCCAATTGG)_2 in the crystal structure [38]. NMR spectroscopic study on the binding of Hoechst 33258 revealed that two drug molecules bind independently at the 5'-TTTT and 5'-AAAA region of d(CTTTTGCAAAAG)_2 duplex DNA sequence with interactions limited to four A.T base pairs [39]. Crystallographic studies on the interaction of Hoechst 33258 with sequences d(CGCGAATTCGCG)_2 and d(CGCGATATCGCG)_2 demonstrate that Hoechst molecule binds with central AT rich regions of the duplex DNA through hydrogen bonding, electrostatic interactions and van der Waal interactions [40-42].

Neidle's group studied the sequence-specific binding of Hoechst with dodecamer duplex DNA (CGCAAATTTGCG)_2 sequence in which Hoechst molecule recognizes the central ATTTG base pair patch via minor groove binding in specific orientations, with high affinity [43]. Crystal structure of berenil with dodecamer DNA sequence d(CGCAAATTTGCG)_2 showed that the drug molecule binds to the 5'-AATT segment with hydrogen bonds formed between amidinium group of berenil and thymine bases of DNA [44]. Berenil also forms a complex with another dodecamer DNA sequence d(CGCGAATTCGCG)_2. It binds to the region 5'-AAT forming hydrogen bonds to adenine atoms [45]. These studies highlight the sequence-specific binding of small molecules to DNA.
In this paper, we report the interaction of DNA sequence $d$-(CCAATTGG)$_2$ with resveratrol to throw light on the binding mode and conformational changes to DNA upon binding. The DNA sequence $d$-(CCAATTGG)$_2$ was selected because of two reasons: firstly, due to its biological significance as a promoter site in various oncogenes [23,46] and secondly due to its AT rich regions, which are strongly over-represented in origins of replication in many organisms [47]. We tried to investigate the binding site and structural changes that took place in the DNA sequence $d$-(CCAATTGG)$_2$ upon interaction with resveratrol through absorption, fluorescence, circular dichroism (CD), differential scanning calorimetry (DSC) and nuclear magnetic resonance (NMR) studies. NMR derived distance parameters are employed to perform restrained molecular dynamics (MD) along with simulated annealing to obtain the final structural parameters of the DNA-resveratrol complex. Figure 1 represents the structure of resveratrol (A) and DNA sequence (B).

Fig. 1. Schematic representation of (A) resveratrol and (B) $d$-(CCAATTGG)$_2$DNA sequence.

2. Materials and Methods

The single-stranded oligonucleotide sequence CCAATTGG and resveratrol were purchased from Sigma Aldrich Co., USA. DNA stock solution was prepared in 20 mM sodium phosphate buffer containing 10 mM sodium chloride and 0.2 mM EDTA at pH 7.2. The DNA solution was heated and kept at 95°C for 5 minutes and annealed overnight to facilitate the formation of double-stranded DNA structure. Resveratrol was dissolved in 1:3 methanol-water. Concentrations of
DNA and resveratrol were determined spectrophotometrically using molar extinction coefficients 76300 M$^{-1}$ cm$^{-1}$ at 260 nm and 32,000 M$^{-1}$ cm$^{-1}$ at 318 nm respectively. All chemicals used were of analytical grade. Millipore water was used throughout the experiments.

2.1. UV–Visible absorption measurements

The absorption spectra of free DNA and complexes with resveratrol were carried out using CARY 60 spectrophotometer (Agilent Technologies). Absorption was recorded from 200 to 700 nm using a 1 cm pathlength quartz cuvette. Spectra were collected with different concentrations of DNA (N) added to a fixed amount of resveratrol (D). Appropriate blank solutions were used for baseline subtraction. Binding coefficient $K_b$ was calculated by using a double reciprocal plot as given below [48, 49].

$$\frac{[\text{DNA}]}{\varepsilon_a - \varepsilon_f} = \frac{[\text{DNA}]}{\varepsilon_b - \varepsilon_f} + \frac{1}{K_b(\varepsilon_b - \varepsilon_f)}$$

where $\varepsilon_f$, $\varepsilon_b$ and $\varepsilon_a$ are molecular extinction coefficients of the free, bound and apparent (observed) state respectively and [DNA] is the concentration of DNA.

2.2. Fluorescence measurements

Fluorescence experiments were performed using FluoroMax Plusspectrofluorometer (Horiba Scientific, USA). Fluorescence titrations were carried out in quartz cuvette with a fixed concentration of resveratrol at 10 µm. DNA solutions were gradually added in various D/N ratios at room temperature. The emission spectra were recorded from 330 nm to 600 nm, with excitation wavelength at 318 nm. The excitation and emission slits were kept at 5 nm. Binding constant ($K_b$) and the number of binding sites (n) were calculated from the double logarithmic plot using the equation [50].

$$\log \frac{F_0 - F}{F} = \log K_b + n \log [\text{DNA}]$$
Where \( F_0 \) and \( F \) are the fluorescence intensities of the fluorophore in the absence and presence of different concentrations of DNA, respectively.

### 2.3. Circular Dichroism measurements

Circular dichroism experiments were carried out by using Chirascan CD spectrometer (Applied Photophysics, Leatherhead, UK) attached with a temperature controlled cell holder. Quartz cuvette of 0.1 cm path length was used for the measurements. CD spectra were recorded in the wavelength a range of 190 to 700 nm. The concentration of the DNA sequence was kept constant at 100\( \mu \)M during the titrations and resveratrol solutions were added serially in various D/N ratios at 25°C in phosphate buffer at pH 7.2. Spectra were averaged over three scans, and the baseline was subtracted. All CD spectra were processed using Chirascan software.

### 2.4. Differential Scanning Calorimetry

The Differential Scanning Calorimetry (DSC) experiments were carried out using a MicroCal VP-DSC instrument (MicroCal, Northampton, MA, USA). In this method, excess heat capacity as a function of temperature was measured to observe the thermal transitions from folded to the unfolded state. The DNA sample was prepared in 20 mM of sodium phosphate buffer. All samples were degassed before the experiment. Each sample was scanned from 25°C to 120°C with a scan rate of 60°C/h at constant pressure 33 psi. The reference and sample cells were loaded with the buffer solution and repeatedly scanned until proper baseline was obtained. 100 \( \mu \)M free DNA solution was loaded in the sample cell and scanned to obtain the thermal profile of free DNA. The complexes of DNA (N) with resveratrol (D) at D/N = 1 and 2 were scanned to obtain the thermal profile of bound forms. All scans were subtracted from the baseline, and thermograms were analyzed using inbuilt Origin software. Thenon-two-state model curve fitting was applied to fit the raw data of each thermogram.

All spectroscopic data were analyzed and plotted using Origin software.
2.5. Nuclear Magnetic Resonance experiments

NMR samples of DNA were prepared by dissolving dry oligonucleotide to give 3.4 mM solution in 90% H$_2$O and 10% D$_2$O containing 10 mM sodium phosphate buffer, pH 7, 0.25 mM EDTA and 10 mM sodium chloride. Samples of resveratrol (28 mM) were prepared by dissolving the compound in d-ethanol. DNA-drug complexes at different Drug (D) to Nucleic acid (N) duplex ratio (D/N) were prepared by titrating different concentrations of DNA and drug. Trimethylsilyl propionic acid (TSP) was used as an internal reference.

All NMR data were collected on a Bruker Avance AMX-500 MHz FT-NMR spectrometer located at Nuclear Magnetic Resonance (NMR) Facility, Indian Institute of Technology Roorkee. The data obtained were processed with TOPSPIN version 1.3 software (Bruker). One dimensional (1D) NMR were collected at 278 K, 283 K and 298 K with 64 K data points, 64-128 number of scans, spectral width of 12/20 ppm. The removal of HOD signal was accomplished using watergate suppression. The two dimensional (2D) phasesensitive nuclear Overhauser Effect spectroscopy (NOESY) experiments were carried out at 283 K and 298 K with mixing times $\tau_m$ = 200 and 300 ms. Typical parameters for 2D experiments were: 2048 data points along $t_2$ dimension; 512 free induction decays in $t_1$ dimension; pulse width $\approx$ 9.5–12 $\mu$s; spectral width $\approx$ 5000 Hz; no. of scans = 64–128; digital resolution 2.30–4.60 Hz/point and relaxation delay $\approx$ 2.0 s. Hetero Nuclear Multiple Bonded Correlation spectroscopy (HMBC), TOtal Correlation SpectroscopY (TOCSY) were carried out at 283 K with 400/512 by 2048 data points. One dimensional $^{31}$P and Two dimensional phosphorous exchange ($^{31}$P-$^{31}$P) spectra were collected using mixing times of 150 ms and 200 ms for different D/N ratios at 283 K and 298 K. Distance restraints were obtained from the volume of experimental NOE cross peaks, calibrated with cytosine H5-H6, using a distance of 2.45 Å in Sparky software[51]. A range of $\pm$0.5 Å was provided to avoid any errors in integration.
Distance restrained Molecular Dynamics (rMD) was performed using AMBER 18 simulation package [52] and the protocols used were described in the relevant section.

All helical parameters, backbone torsional angles, and sugar conformations of the resulting rMD structures were analyzed using Curves+ software[53].

3. Results and Discussion

3.1. Absorption studies

The absorption spectrum of resveratrol showed a broad absorption spectrum with two kinks around 304 nm and 318 nm. The titration was carried out with a fixed concentration of resveratrol. The gradual addition of oligonucleotide d(CCAATTGG)2 to resveratrol (N/D) in a set of 25 experiments resulted in a sequential hyperchromic effect without any noticeable shift in the maximum absorbance wavelength as shown in Fig. 2A. Data were analyzed with absorption at 318 nm. A plot of absorbance at 318 nm against DNA concentration shows a linearly increasing curve until saturation around N/D = 2 (Fig. 2B). The binding constant was calculated using the double reciprocal plot (Fig. 3) obtained as $K_b = 2.20 \times 10^6$ M$^{-1}$. Earlier our studies showed that resveratrol binds to d(CGAAATCG)2 and d(CGTTAACG)2 sequences with binding constants of the order of $10^5$ [54]. The results of the present study, therefore indicate that resveratrol binds more strongly to the sequence CCAATTGG as compared to the above two sequences, pointing to the sequence specificity of the compound. Absence of any isosbestic point suggests that either no equilibrium state or multiple equilibrium states exist in the solution. Hyperchromism with no shift in absorption maxima usually results from an external binding of the molecule to the DNA sequence. Often hyperchromism with a bathochromic shift indicates an intercalative mode of binding. The observed hyperchromism with no change in absorbance maxima, therefore, indicate the external binding of resveratrol.
3.2. Fluorescence studies

Resveratrol gives intense emission around 388 nm upon excited at 318 nm. The broadband is peaked at 381 nm with two shoulders at 353 nm and 410 nm. These peaks are attributed to the emission from a noncharged form of the resveratrol as described elsewhere[55]. Fig. 4 illustrates the fluorescence emission spectra of resveratrol alone and in the presence of DNA at different nucleic acid to ligand ratios (N/D). Incremental addition of DNA to a fixed concentration of resveratrol resulted in a blue shift, from 388 nm to 381 nm. At the same time, the 353 nm band red shifted to 364 nm along with an increase in emission intensity. The spectral changes indicate the interaction between DNA and resveratrol. Resveratrol was added to DNA till saturation is
reached, around N/D = 3.0. The differences suggest that the hydroxyl groups of the ligand are in different environments [55]. In general, when the ligand is in a hydrophobic environment, its fluorescence intensity increases [56]. The fluorescence intensity almost doubled at N/D = 1 and increased by four times when DNA concentration is 20 µM (N/D = 2) (Fig. 5). Such kind of increase in fluorescence when a ligand binds to AT rich DNA has been reported. This can be due to the lowering of the conformational flexibility of the compound within the complex [57,58]. This property of the molecule can be exploited to develop DNA sequence selective probe, and our group is working to this goal.

The Binding constant (K_b) and the number of binding sites (n) were calculated using double logarithmic plot (inset of Fig. 4A) and obtained as 7.94×10^9 M^-1 and 1.8 respectively for N/D = 0 to 1.4. Whereas for N/D= 1.4 to 2.3, K_b was obtained as 8.0×10^4 M^-1) and n = 0.76 (Inset of Fig. 4B) respectively. N=2.3 indicates that there are at least two strong binding sites in the DNA sequence. The binding stoichiometry of the complex was measured using the Job plot.
(Continuous variation method). The total concentration of ligand and DNA was kept constant at 20 μM. The emission intensity of resveratrol at 388 nm was plotted against the mole fraction of resveratrol as shown in Fig. 6. The cross over point was obtained at mole fraction equal to 0.69 which indicates a stoichiometry (χ / 1 - χ, where χ is the mole fraction of ligand) of 2.2: 1 indicating that two molecules of resveratrol bind with one molecule of DNA.

3.3. Circular dichroism spectroscopy (CD) studies

Circular dichroism (CD) spectroscopy is an important technique which displays structural changes in DNA sequences. The CD spectrum of canonical right handed B-DNA exhibits a positive band around 269 nm, owing to base stacking and a negative band around 247 nm, owing to helicity [59]. Often changes in the CD signals of DNA on interaction with ligand is attributed to the corresponding changes in DNA structure. Resveratrol did not show any CD band. The changes in CD spectrum of DNA were monitored upon ligand binding, which is due to corresponding changes in DNA conformations.

Fig. 5. Relative fluorescence intensities of resveratrol as a function of DNA concentration.

Fig. 6. Job plot for the binding of resveratrol to DNA. The difference in fluorescence intensity at 388 nm as a function of mole fraction of resveratrol was plotted.
Fig. 7 describes the variation in CD bands when different concentrations of resveratrol were added to DNA (D/N ratio). The intensities of the positive band around 269 nm and negative band around 247 nm decreased till D/N = 5.0 without any noticeable shift. The 269 nm band decreased by 40%, while the 247 nm band decreased by 22%, indicating that the interaction between resveratrol with DNA caused a disturbance in the stacking of the bases as well as helicity. Absence of any significant spectral shift and induced CD band rules out the possibility of intercalation. The spectral changes observed suggest that resveratrol is binding externally and induce slight variations in the conformation of DNA.

3.4. Differential Scanning Calorimetry

Melting temperature (Tm) due to transition from complexed to uncomplexed (ordered to disordered) state can be estimated by evaluating the excess heat capacity as a function of temperature using differential scanning calorimeter (DSC). DSC thermograms were obtained for alone DNA and its complex with resveratrol at N/D =1 and 2 as presented in Fig. 8. In the case of alone DNA, the thermogram gave a good fit with two peaks, one major peak centred at $T_{m1} = 39.25^\circ C$ and another peak around $T_{m2} = 46.27^\circ C$. More than one $T_{m1}$ shows that the melting
pathway consists of intermediate species[60]. At N/D = 1.0, T_{m1} is shifted to 42.20˚C and T_{m2} to 51.23˚C. ΔT_{m1} = 2.95˚C observed from changing N/D = 0 to 1 indicates a minimal change, pointing to the external binding of the compound to DNA. This result correlates with the observations of absorption and fluorescence findings. At N/D = 2.0, the DSC thermogram fits with two peaks with T_{m1} = 42.07˚C and T_{m2} = 52.92˚C as indicated in Table 1. Positive enthalpy obtained from the DSC analysis is a sign of hydrophobic binding reaction[61]. The ratio \( n \) of van't Hoff enthalpy to calorimetric enthalpy gives an estimate of the effective number of base pairs in a cooperative melting system\[62,63\]. Apparent values of \( n \) show that at least 6 base pairs are there in the cooperative melting system.

3.5 Proton Resonance Assignments

3.5.1. d-(CCAATTGG)\(_2\)

One dimensional proton spectra were recorded for the sample containing DNA at 278K, 283K and 298K in 90% H2O and 10% D2O. The NMR resonances of the two strands of the octamer coincide due to the 2-fold symmetry of the duplex, and only one set of resonances was observed. The assignment of nucleotide protons (Supplementary material Table 1A) was carried out following the strategies available in the literature for sequential assignment in right handed B-DNA [64-66]. NOESY spectrum of DNA recorded at 298K with a mixing time 200ms was used.
Table 1. Thermodynamic parameters obtained from DSC

<table>
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<th>Sample</th>
<th>Tm1 (°C)</th>
<th>ΔH1a (kcal/mol)</th>
<th>ΔHV1b (Kcal/mol)</th>
<th>n</th>
<th>Tm2 (°C)</th>
<th>ΔH2a (kcal/mol)</th>
<th>ΔHV2b (kcal/mol)</th>
<th>ΔTm1 (°C)</th>
<th>ΔTm2 (°C)</th>
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<td>12.67</td>
<td>48.39</td>
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<td>-</td>
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<tr>
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<td>10.97</td>
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<td>6.09</td>
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<tr>
<td>DNA RES</td>
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<td>6.8</td>
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a - ΔH = Enthalpy of the transition, b - ΔHV = Van't Hoff enthalpy of transition

for resonance assignment. Fig. 9 shows the fingerprint region of base H6/H8–H1’ protons and connectivities of base H6/H8 and sugar H1’ protons are obtained by the method of sequential walk as indicated.

3.5.2. Resveratrol

The protons of resveratrol have been assigned with the help of 1D proton NMR, Rotating Frame Overhauser Effect Spectroscopy (ROESY) and Total Correlation Spectroscopy (TOCSY) experiments (not shown). The chemical shifts of resveratrol were obtained, as shown in supplementary material Table 2. The one dimensional proton spectra of resveratrol in deuterated ethanol and deuterium oxide (D2O) at 298K are shown in Fig. 10. The results were consistent with the literature on the NMR studies of resveratrol[67,68]

3.5.3. DNA-resveratrol complex

Upon addition of resveratrol to d-(CCAATTGG)2 in small steps up to D/N ratio 1.0, a set of new signals corresponding to H2 and H6 protons of resveratrol appear (Fig.11a).
Fig. 9. NOESY spectrum of d(CCAATTGG)$_2$ showing base H6/H8 - sugar H1' connectivities.

Fig. 10. One dimensional proton spectrum of resveratrol in deuterated ethanol (above) and in D$_2$O (below).
The change in chemical shift ($\Delta\delta$) of d-(CCAATTGG)$_2$ protons with increasing D/N ratio was gradual and small in magnitude (as shown in Supplementary material table 1A and figure 11). T6H1 and T6H5" shifted upfield by ~0.04 and 0.06 ppm while A3H2 and A4H2 showed a downfield shift of ~0.03 ppm on binding to resveratrol. A4H6 was downfield shifted by 0.06 ppm.

All the spectral lines of DNA were somewhat uniformly broadened upon binding as the internal motions are affected, and protons get immobilized. Fig. 12 shows the broadening of spectral lines in the H1′ region. The very small chemical shift changes indicate that the interaction between resveratrol and d-(CCAATTGG)$_2$ induced only a slight distortion in the DNA structure.

$^1$H NMR spectra of the resveratrol-d-(CCAATTGG)$_2$ complex in the temperature range of 278-298 K (Fig.11b) show that drug protons are more prominent at 298K.

The 2D NOESY spectra of resveratrol d-(CCAATTGG)$_2$ complex were obtained at different mixing times, D/N ratios and temperatures. Fig. 13 and Supplementary material Fig. 1A shows NOE connectivities observed in the resveratrol-d-(CCAATTGG)$_2$ complex at D/N=1, 298 K and the NOE peaks (all not shown) have been used in deriving the structure of the complex. The NOESY spectra were analyzed extensively at different mixing times. NOEs between pairs of proton G7NH$^b$ - C2NH$^b$, C2NH$_2^{nb}$; T6NH - A3NH$^b$, A3NH$_2^{nb}$, A3H2; and T5NH - A4NH$^b$, A4NH$_2^{nb}$, A4H2 were observed establishing Watson-Crick base pairing between C2…. G7, T6....A3 and T5....A4 base pairs in the complex. This is further supported by the presence of intramolecular sequential connectivity T5NH-A4NH$_2^{nb}$ and intermolecular sequential connectivities T6NH - C2NH$^b$, C2NH$_2^{nb}$ (Supplementary material Fig.1B). The sequential connectivities G7NH - T6NH, T6NH - T5NH were also observed. All the sequential connectivities between
Fig. 11. A) 1D 1H NMR spectrum of resveratrol-DNA complex at different D/N ratios, B) 1D 1H NMR spectrum of resveratrol-DNA complex at temperatures 278K, 283K and 298K. # denotes proton signals from resveratrol.
Base H8/H6/CH$_3$/H5 and sugar protons H’, H2’, H2” were observed at all basepair steps. The NOE data therefore proves the existence of B-DNA duplex with no possibility of opening of base pairs. Hence, the intercalation mode of interaction between resveratrol and DNA in the ligand-DNA complex is completely ruled out.

Interaction between two molecules often causes changes in the chemical environment of the atoms at the interface, leading to chemical shift changes in proton NMR spectra. As a result, analysis of chemical shift can help one to pinpoint the sites of local interactions between the molecules. If the resonances that shift upon binding correspond to the residue directly involved in binding, chemical shift change can provide valuable insights into the structural basis for

Fig. 12. Sugar H1’ of DNA when complexed with resveratrol at different D/N ratios showing broadening of the peaks.
recognition. Hence can lead the way for designing or screening efforts focused on the discovery of novel ligands. In the present study, the slight shifts in G7H6, G7H2’, G7H1’, T6H6, T6H5", A4H6, A4H2 and A3H2 protons suggests that binding affects the conformation of these bases of DNA. The shift in most of DNA protons is however insignificant, as has also been reported in earlier studies [69,70]. The distortions induced in DNA conformation appear to be relatively small in magnitude. The changes in resveratrol protons are not significant, ruling out the stacking mode of interaction. The NOE cross peaks were observed between the resveratrol H6/H2 protons and H6 of T6; NH2nb of A3; H2 of A3 and A4 (overlapped) as indicated in figure 13. The small chemical shift changes are indicative of external binding of the resveratrol molecule near the AT base pair. The detailed structural information was, however, derived by analyzing NOEs. No evidence like intermolecular cross peaks (between resveratrol) was obtained to establish the binding of two molecules of resveratrol. Also, because of the self-complementary nature of the sequence, it can be judged that resveratrol forms cross peaks with NH2nb of A3of first strand and H6 of T14 (i.e., T6 of the second strand). These intermolecular 1H-1H NOESY cross peaks (Supplementary material Table 1B) between resveratrol and DNA bases confirm that it binds near the AT base pair. Supplementary material Table 1C shows few intramolecular cross peaks observed for resveratrol in the complex spectrum.

3.6. Phosphorus-31 NMR Spectra

Phosphate resonances of uncomplexed DNA were assigned from the 1H-31P HMBC spectrum. C2pA3, G7pG8, C1pC2, T6pG7 and A3pA4 appear at -0.42, 0.52, -0.79, -0.85 and -1.01 ppm, respectively at 298 K while resonance of both A4pT5 and T5pT6 appear at -1.14 ppm. Upon
Fig. 13. Expansion of the specific regions of the NOESY spectra of resveratrol complexed with d-(CCAATTGG)$_2$ at D/N=1 at 298 K. ⭐ indicates intermolecular cross peaks between resveratrol and DNA protons, ▲ indicates intramolecular cross peaks of resveratrol.

Fig. 14. Proton decoupled $^{31}$P NMR spectra of resveratrol-(CCAATTGG)$_2$ complex as a function of added drug (D) to Nucleic acid duplex (N) stoichiometric ratios (D/N) of 0.0, 0.2, 0.4, 0.6, 0.8, 1.0 at 298 K in phosphate buffer.
binding of resveratrol to d-(CCAATTGG)_2, the resonances broadened without any significant shift, as shown in Fig. 14. The largest shift (upfield) of 0.053 ppm was observed for T6pG7 followed by 0.017 ppm for A3pA4 while the A4pT5/T5pT6 resonance upfield shifted by 0.013 ppm (Supplementary material Table 3). No additional peaks were observed.

It is known that ^31P chemical shifts vary in response to local, sequence specific and induced environmental distortions in the duplex geometry[71], occur with ester O-P-O bond angle distortions or modifications in backbone torsional angles. The intercalating molecules induce deformation of phosphodiester chain, to allow the opening of base pairs, by converting from energetically more favorable B_I conformation to a more flexible B_{II} conformation. This usually results in a downfield shift more than 1.5 ppm[72]. It is known that downfield shifts of ~1.5 ppm or more occur on the binding of DNA to classical intercalated[73,74], while ligands which bind externally to DNA such as pyridopurine[75], mitoxantrone[76], camptothecin[77], show much smaller shifts. Absence of any such large downfield shift in the present study excludes the possibility of opening of base pairs to allow intercalation. This result is in accord with the observation that the change in the absorption maximum on the binding of resveratrol to DNA sequence is insignificant as compared to that of 25-44 nm observed on the binding of typical intercalator to DNA[78,79]. Purely electrostatic associations between drug and nucleic acid, on the other hand, produce only small and generally upfield ^31P chemical shifts [80] which are not observed in the present study. Thus the association of resveratrol with octamer does not involve direct intercalation or electrostatic interaction. The minute chemical shifts found thus corroborate for the external binding of the compound. The small shifts in the positions of T6pG7, A3pA4 and A4pT5/T5pT6 indicate that resveratrol is binding near the AATT region.

3.7. Observations from NMR and other spectral data
Analysis of the shifts in proton NMR structure indicates no substantial changes in the DNA conformation upon binding with resveratrol. Chemical shift changes in the base protons of T5, T6, G7, A3 and A2, though small, bespoke the binding of the compound near these bases. The NOE cross peaks obtained with the resveratrol H2/H6 protons substantiate these findings and confirms that resveratrol binding to the AATT minor groove region. The features of observed 31P resonances also vouch for the external binding of the compound. The absorption, fluorescence and CD spectral data described an external binding of the compound to the DNA sequence. These findings support our earlier findings of the external binding of resveratrol to two DNA sequences with central AATT segment [54].

3.8. Structure-Conformation and Dynamics

The sequenced-(CCAATTGG)$_2$ was prepared from PDB data 1JTL. The terminal residues and ligand from the structure were deleted and used for further analysis. To obtain a complex structure, d-(CCAATTGG)$_2$ was first docked with resveratrol. The docking conformations showed three clusters. The lowest energy structure from the most populated clusters was taken as the starting structure of the complex. The complex structure from docking was curated for DNA and resveratrol structure using ambpdb and antechamber protocol for amber compatible structure. Modified starting structure of the complex was prepared. The structure was energy minimized and MD simulations were performed.

3.8.2. Molecular Dynamics (MD) simulations

MD simulations were carried out using the sander module in AMBER18 simulation package. Hydrogen atoms were added to both ligand and DNA. The leaprc force fields, general amber force field (GAFF) and bsc1DNA were used for ligand and DNA, respectively in the tleap module[81]. The complex structure was solvated in a truncated octahedron periodic cell centered
on DNA and explicit water molecules were added using the TIP3P water model [82]. tleap has added around 5710 water molecules within 15 Å of solvation between the solute (DNA) and the edges of the box. The solute was neutralized with 14 sodium ions making the box size 72.86 × 72.86 × 72.86 Å. Minimization and equilibration of the DNA-resveratrol complex were carried out followed by MD production run. Firstly, the DNA and the ligand were minimized with both the molecules restrained for 1000 steps, with a cut off of 10 Å. The minimized structure was subjected to another round of minimization without any restraint either on DNA or on resveratrol. This is followed by heating the system using the NVT ensemble and the SHAKE algorithm to constrain hydrogen bonds [83]. The temperature was increased from 100K to 300K during the first 10 ps (5000 steps) and kept constant at 300K for the next 90 ps. The system was then equilibrated after reducing the constraint to 0.5 kcal/mol in the NPT ensemble at constant temperature (300K) and pressure (1 atm) for 50ps. The production phase of the simulation was performed in the NVT ensemble at 300 K and Langevin thermostat with collision frequency 1 ps⁻¹. Distance restraints obtained from NOE intensities were introduced using make DIST_RST program in Amber and simulation was carried out for 1ns with a time step of 2 fs. A total of 223 distances were used. Root mean square deviation and backbone torsional angles were estimated using the cpptraj module [84].

The 1 ns trajectory was stable, as indicated by the steady root mean square deviation (rmsd) values, as shown in Fig. 15. The rmsd was between 2.5 Å and 3.5 Å considering all 8 residues and 2 Å to 2.5 Å for central residues (red line). The potential and kinetic energies of the system were stabilized after the initial 20 ps equilibration run. The minimum energy structures were generated using cpptraj, and the root mean square deviation (rmsd) of nine lower energy structures with the lowest energy structure was obtained as 1.709 Å and the superimposed minimum energy structures are shown in Fig. 15C. Pseudo puckering of atom C1′ through O4′ for
residues 3 to 6 is shown in Fig. 15B. Sugar puckering of residues 3, 5 and 6 fall in the range of 150 degrees while residue 4 shows a slightly higher value as displayed in the Fig. 15B. This could be due to the binding of resveratrol near this residue. These values are consistent with B DNA parameter. The lowest energy structure is stabilized with a total potential energy = -57746.5 kcal/mol. In the lowest energy structure, resveratrol 3OH forms four hydrogen bonds with DNA namely, O with G15H6 and A3NH2; while H with O4’ of G15sugar and O5’of T14sugar. All distances are within 2.7 Å. This reveals that resveratrol is accommodated to the AT region with one terminal fixed by making hydrogen bonds, thereby stabilizing the structure. The chemical shift changes observed for T6 and A3 base and sugar protons as well as phosphorus also support this. The binding of resveratrol near these bases induce the changes, though the shifts are very small. Increase in the fluorescence intensity of resveratrol upon binding, as described earlier, is because of the intrusion of the OH group into the minor groove region.

Structural parameters of DNA were analyzed using Curves+ software. The helical analysis shows an average twist of 20.2˚ and helical rise of 3.23 Å. All sugars except the ones for A4 on the first strand and T13 on the second strand are in C2’endo puckering, retaining the features of B-DNA. Minor groove width of DNA was increased as shown in Fig. 16. Minor groove depth increased slightly. Major groove width of AT base pair slightly increased while the depth decreased. This might have caused because of the binding of resveratrol in the minor groove. Similar features were indicated in the literature, in the case of minor groove binding agents [85-88].

4. Conclusion

The paper discusses the interaction of resveratrol with an octamer sequence which is present in the promoter region of various oncogenes and many nuclear factors. Absorption and fluorescence studies revealed that the compound is interacting externally. Fluorescence technique which is
more sensitive than absorption shows that resveratrol binds with moderate to high affinity to the octamer sequence. Also, the high fluorescence exhibited by resveratrol in the presence of DNA can be used as a practical DNA light probe. Circular dichroism data display moderate changes in the DNA conformation upon binding. The small changes in the melting temperatures of DNA and its 1:1 and 1:2 complexes with resveratrol upholds the spectroscopic observations that resveratrol binds externally to DNA. NOE cross peak between H2/H6 protons of resveratrol with A3, A4 and T6 base protons establishes the formation of resveratrol-DNA complex at the
minor groove. Subtle changes in the chemical shifts corroborate for the binding. Lowest energy structure obtained from the molecular dynamics analysis confirms that the DNA conformation changed subtly to accommodate the compound, supports with the findings of circular dichroism data. The complex structure was stabilized by hydrogen bonds formed between one terminal end of resveratrol with DNA. These results endorse resveratrol as a minor groove binding agent and pave the way to develop resveratrol and its analogues as suitable candidates as a new class of DNA sensing probes. Many minor groove binders are known for their antibacterial activity. The results obtained urge one to probe the effect resveratrol binding to specific nucleotide sequences which play an essential role in gene regulation and potential candidate as a novel antibacterial/anticancer drug.

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Authorship contribution

PK performed NMR experiments and analyzed data, SK performed absorption, fluorescence, CD, and DSC experiments. MN conceptualized, designed and analyzed the data and wrote the manuscript.

Conflict of Interest

Authors declare no conflict of interest.

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Supplementary material

Exploring the binding of resveratrol to an oncogene promoter DNA sequence d(CCAATTGG)$_2$ through multispectroscopic, nuclear magnetic resonance and molecular dynamics studies

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Department of Biotechnology, Indian Institute of Technology Roorkee, Roorkee, Uttarakhand, India.
Supplementary material Table 1A. Chemical shift values (ppm) of d-(CCAATTGG)$_2$ in 90% H$_2$O +10% D$_2$O at 298K.

<table>
<thead>
<tr>
<th>PROTONS</th>
<th>C1</th>
<th>C2</th>
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<th>A4</th>
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<tr>
<td></td>
<td>δb</td>
<td>δf</td>
<td>Δδ</td>
<td>δb</td>
</tr>
<tr>
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<td>2.05</td>
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<tr>
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<th>G8</th>
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<tr>
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<td>δb</td>
<td>δf</td>
<td>Δδ</td>
<td>δb</td>
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<td>-0.01</td>
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<tr>
<td>NH2nb</td>
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<td>13.75</td>
<td>13.76</td>
<td>-0.01</td>
<td>13.99</td>
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δb = Chemical shift values of bound form, δf =Chemical shift values of free form, Δδ = δb - δf. -ve Δδ indicates upfield shift, +ve Δδ indicates downfield shift.
**Supplementary material Table 1B. Intermolecular cross peaks between resveratrol and d-(CCAATTTGG)$_2$.**

<table>
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<tr>
<th>Resveratrol- DNA cross peaks</th>
<th>Calculated NOE distances (Å)</th>
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<tr>
<td>RES 2,6 H- A3NH$_2^{ab}$</td>
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<tr>
<td>RES 2,6 H -T6H6</td>
<td>3.32</td>
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<tr>
<td>RES 2,6 H –A3/A4 H2</td>
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Supplementary material Table 1C. Intramolecular cross peaks within resveratrol, observed in complex spectrum.

<table>
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<th>Intramolecular Resveratrol cross peaks</th>
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<tbody>
<tr>
<td>H2′/H6′</td>
</tr>
<tr>
<td>Hα′</td>
</tr>
<tr>
<td>H2′/H6′</td>
</tr>
<tr>
<td>H3′/ H5′</td>
</tr>
<tr>
<td>H2′/H6′</td>
</tr>
<tr>
<td>Hα</td>
</tr>
</tbody>
</table>
Supplementary material Table 2. Chemical Shifts of resveratrol protons in deuterated ethanol and D$_2$O.

<table>
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<tr>
<th>Resveratrol Protons</th>
<th>Chemical shift (ppm) in ethanol</th>
<th>Chemical shift (ppm) in D$_2$O.</th>
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<tbody>
<tr>
<td>2',6'</td>
<td>7.31</td>
<td>7.51</td>
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<tr>
<td>3',5'</td>
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<td>α'</td>
<td>6.79</td>
<td>7.17</td>
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<td>α</td>
<td>6.76</td>
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<td>2,6</td>
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<td>6.67</td>
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<tr>
<td>4</td>
<td>6.21</td>
<td>6.32</td>
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**Supplementary material Table 3.** Chemical shift of $^{31}$P resonances (ppm) of DNA octamer in d-(CCAATTGG)$_2$ complexed with resveratrol at 298K

<table>
<thead>
<tr>
<th></th>
<th>$\delta_f$</th>
<th>$\delta_b$</th>
<th>$\Delta \delta = \delta_b - \delta_f$</th>
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<tr>
<td>C1pC2</td>
<td>-0.785</td>
<td>-0.782</td>
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<td>C2pA3</td>
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<td>-0.426</td>
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<td>-1.033</td>
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<td>A4pT5</td>
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<td>T6pG7</td>
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<td>G7pG8</td>
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$\delta_b$ = Chemical shift values of bound form, $\delta_f$ = Chemical shift values of free form, $\Delta \delta = \delta_b - \delta_f$, $\Delta \delta$ indicates upfield shift, +ve $\Delta \delta$ indicates downfield shift.
Supplementary material Fig.1A. NOE connectivities of resveratrol-DNA complex showing base-H2'-H2'' region.
Supplementary material Fig.1B. NOE connectivities of resveratrol-DNA complex showing base-NH region.