Conformational Study on Pheromonotropin neuropeptide using NMR and Molecular Dynamics

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Abstract

Pheromonotropic neuropeptide, *Pseudaletia* pheromonotropin, is an 18 amino acid peptide with the sequence Lys-Leu-Ser-Tyr-Asp-Asp-Lys-Val-Phe-Glu-Asn-Val-Glu-Phe-Thr-Pro-Arg-Leu which is widely distributed in female moths of *Bombyx mori*. This peptide is structurally related to leucopyrokinin, an insect myotropic neuropeptide, responsible for melanization and reddish coloration hormone (MRCH) activity. A combined NMR and Molecular Dynamics (MD) methods were used to fathom the conformational behaviour in water which was used as a solvent. The structure was investigated using 1D-NMR and 2D-NMR (COSY, TOCSY and ROESY) experiments. The conformation was built by constrained MD simulations using distance and dihedral restraints from NMR data in the GROMACS simulation package. The peptide predominantly adopts a β-sheet structure in water.

Keywords: Pheromonotropin; Nuclear Magnetic resonance; Conformation; β-sheet; Molecular Dynamic simulations

Introduction

In lepidopteran insects, a vast number of neuropeptides are involved in regulation of various physiological events e.g. pheromone production, diuresis, myotropic activity, blood sugar level, mating, and pupal development [1]. These modifications happen in reaction to environmental conditions or to specific developmental stages [2]. In the past decades, vast studies were conducted on isolation and characterization of many insect neuropeptides. The first investigation of the function of neuropeptides in insects was carried out by Kopeč in 1917 [3]. Even though a great number of conformations of insect neuropeptides have been found, there has been no report of the NMR structure of pheromonotropin. The pyrokinin/pheromonotropin biosynthesis activating neuropeptide (PK/PBAN) multifunctional family play a pivotal part in the development in the insect life cycle [4]. Myotropins, pheromone biosynthesis activating neuropeptide (PBAN), melanization and reddish coloration hormone (MRCH), diapause hormone and pheromonotropin are many of the neuropeptides in PK/PBAN family. The functions of these neuropeptides involves gut hormone contraction, sex hormone production, melanin biosynthesis (defense mechanism), pupariation [5-12] and have been affirmed by both (in vivo and in vitro) experiments carried on various species signifying that they do not exhibit species specificity [13]. The novel pheromonotropin neuropeptide which was extracted from the head of *P. separata* larvae of *Bombyx mori* [14] bears a structural resemblance to PBAN and MRCH having a congruent sequence of Phe-Thr-Pro-Arg-Leu at its C-terminus. MRCH, a neurohormone which regulates sex hormone production, diuresis, myotropic activity, blood sugar level, mating, and pupal development were examined by NMR. 5 mg of the peptide dissolved in 0.6 ml of H2O/D2O (95:5) at pH 7.0 with 3-(Trimethylsilyl)propanoic acid (TSP) was used as the internal standard, which was obtained from Stohler Isotope Chemicals, USA. At this concentration the peptide showed no aggregation.

Nuclear magnetic resonance experiments

The NMR experiments were carried out on Bruker Avance (500,700 and 800 MHz) FT NMR spectrometers as per requirements. The 1D proton spectra were studied with a spectral width of 14000 Hz, 512 scans, and digitized with 32 K data points. Solvent suppression for the sample in H2O/D2O (95:5) was accomplished using the 3-9-19 WATERGATE [25] pulse sequence. The NMR data was processed and analyzed using TopSpin (Bruker Inc.) running on a Rocks Cluster Suite 6.1. The temperature coefficients for amide protons (NH) shifts were measured from 1D NMR spectra at temperatures ranging from 300K to 320 K.

Sample preparation

The 18 amino acid peptide, Lys-Leu-Ser-Tyr-Asp-Asp-Lys-Val-Phe-Glu-Asn-Val-Glu-Phe-Thr-Pro-Arg-Leu was purchased from Sigma Chemical Co., USA and was found to be in pure form when examined by NMR. 5 mg of the peptide dissolved in 0.6 ml of H2O/D2O (95:5) at pH 7.0 with 3-(Trimethylsilyl)propanoic acid (TSP) was used as the internal standard, which was obtained from Stohler Isotope Chemicals, USA. At this concentration the peptide showed no aggregation.

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The 2D NMR experiments included 'H'-H ROESY (Rotating frame Overhauser Effect Spectroscopy) [26] and total correlation spectroscopy (TOCSY) [27] which were used for spin system assignments. At the beginning of experiment, 32 dummy scans were collected to allow the system to reach thermal equilibrium. The residual water signal in D$_2$O samples was suppressed by low-power pre-saturation during the relaxation delay. The spectra were acquired with a sweep width of 14000 Hz. Typically, 512 FIDs of 2048 complex points were collected. The TOCSY experiment [27] was carried out using spin lock pulse of 150 ms with 512 experiments at 300 K and a spectral width of 9000 Hz for 4096 complex data points. The data were multiplied with $\pi/4$ and $\pi/8$ sine bell window functions along $t_1$ and $t_2$ axes prior to Fourier transformation. Coupling constants ($J'/J_{max}$) were extracted from 1D NMR spectra data. Identification of spin systems was done using TOCSY and ROESY sequential assignments which were made according to the methods described by Wüthrich et al. [28]. The reason ROESY spectra was recorded instead of a NOESY because there was an absence of sequential nOe’s in longer mixing time (500 ms) of 2D NOESY spectrum indicating that the rotational correlation time ($\tau_c$) of the peptide was closer to the cross-over point in the relationship between $\tau_c$ and nOe. It was thus necessary to obtain nOe’s from 2D ROESY spectra as nOe’s, as they are always positive in this 2D NMR spectrum.

ROE intensities and distance restraints

Approximate distance restraints were calculated from the ROESY cross peak intensities using the relationship $\eta_i/\eta_o = (r_i/r_o)^6$ where $\eta_i$ and $\eta_o$ are the rOe intensities for the atom pairs $i,j$ and $k,l$ separated by distances $r_i$ and $r_o$ correspondingly. The intensity of the ROE between side chain $\beta$ protons of Lys-7, $\beta$ protons of Phe-9 and Phe-14 with an interunial distance of 1.73 Å which was used as the reference. The distances calculated for methyl and methylene groups, were according to the rules formulated by Wüthrich et al. [28]. The strong, moderate, and weak rotating frame overhauser effect (rOe) cross-peaks were translated into distances with the lower boundary limits set at 1.9 Å, and the upper boundary limits set at 2.6 Å for strong, 3.1 Å for medium, and 4.2 Å for weak peaks [28]. The distances translated as restraints, were used with force constants that ranged from 1 to 30 kcal mol$^{-1}$ rad$^{-2}$ and a ± 0.5Å allowance put on the calculated distances, to represent the upper and lower bounds [28].

$J_{N\alpha\alpha}$-dihedral angle restraints

The $J_{N\alpha\alpha}$ coupling constants values were extracted using the modified Karplus equation [29].

$\quad J_{N\alpha\alpha} = A \cos^2(\phi) + B \cos(\phi) + C$

The values A= 6.52, B= -1.76 and C= 1.60. The coupling constants were converted to $\phi$ values, and introduced as dihedral restraints, allowing a range of ± 10° on the calculated restraint values. Force constants in the range of 1 to 30 kcal mol$^{-1}$ rad$^{-2}$, were used.

Chemical shift index (CSI)

The chemical shift index was developed by Wishart et al. [30]. It is calculated as a difference between the observed chemical shift and the chemical shift for that proton in the peptide when it is in a random coil structure. Residues belonging to the $\alpha$-helical region of the peptide are observed as a negative deviation from the chemical shift of random coil while residues in the $\beta$-sheet region show a positive deviation for the same.

Molecular dynamics modeling

Computational modeling technique is widely applicable for the analysis of biomolecular motion and interactions. This modelling methodology has been applied to determine the chemical, physical and biological properties of bio systems [31]. Molecular Dynamics (MD) simulation is suitable for small and large systems where individual atoms and or small clusters of atoms are involved and the phenomena influenced by the motion of individual atoms. These models explore a time scale of picoseconds ($10^{-12}$ s) to microsecond simulations. The determining equations in molecular dynamics follow classical Newtonian physics models. This method is derived from Newton’s equation of motion based on the selected force fields that defines the associated forces in the biosystems. This method is popularly used as a means to model biological structures to discern events like protein folding, interaction of drug candidates with proteins and catalytic transformation.

Modeling analysis methodology

The MD simulations were performed using the GROMACS 4.6.5 package of programs [32-34], using the OPLS-AA/L all-atom force field [35,36] running on Intel Xeon based 13-node high performance computing cluster operating on the Rocks Cluster Suite 6.1 to simulate the dynamics of the peptide in water under constant temperature (300 K) and pressure (1 atm) with NMR restraints. The peptide was built using the Maestro module 2014 in Schrodinger suite of program and converted to protein data bank (pdb) file for use in GROMACS simulations. The peptide was centered box of dimension of a 35 Å x 35 Å x 35 Å PBC box filled with 28438 Simple Point Charge (SPC216) water molecules and neutralized with 3 Na$^+$ ions [37]. The model was subjected to energy minimization with a combination of steepest descents and conjugate gradients ended with the Newton-Raphson method, to a gradient of 0.01 kcal mole$^{-1}$ Å of total of 50000 steps of 100 ps. This configuration was then constrained using LINCS [38] algorithm and equilibrated using NVT and NPT ensembles. The NVT equilibration was done with all bonds constrained and temperature coupled by a velocity rescale thermostat which is a modified Berendsen thermostat specific to GROMACS. Pressure constraint in NPT ensemble via the Parrinello-Rahmen barostat allowed the simulation cell box to change its shape accordingly. The particle mesh Ewald summation method was used for the treatment of long-range electrostatic interactions [38]. Both NVT and NPT equilibrations were carried out for 100 ps each with the temperature and pressure dynamic variations verified to ensure that the system was indeed equilibrated. The time step for the simulations was 2 fs. Lennard-Jones forces were treated using a cutoff of 1.2 nm. Long-range electrostatic forces (r>1.2 nm) were treated using the particle mesh Ewald method. The fully equilibrated system was used as the starting configuration for the MD production run where the peptide was constrained using distance and dihedral restraints for a total of 2500 ps. Post processing analysis tools were applied to visualize the trajectories and to quantify the configurations of the system using the Visual Molecular Dynamics (VMD) software and GROMACS package tools.

Results and Discussion

NMR structure of pheromonotropin

The 1D NMR spectrum of the peptide was well resolved and all the NH peaks except the terminal Lys were observed between 6.3 to 8.5 ppm. As a rule of thumb, amino acids in the peptide were identified by the pattern of the TOCSY cross peaks according to the spin system
of each amino acid in relation to the actual chemical shift values [28]. The unique spin systems of Thr (A_3MX) and Glu [AM (PT) X] were distinctly identified from the TOCSY spectra, four AMX spin systems were assigned to one Asn, two Asp residues and one Ser residue or long side-chain spin systems in which further scalar coupling to more peripheral protons was observed (i.e. Glu and Arg). The Asp residue was then identified based on the fact that its δ protons resonate downfield compared to those of Asn residues. The Pro residue was assigned from the typical chemical shift values of its δ protons (C_αH and C_δH signals). The unique spin systems of Val (A_3B_3MX) and Thr (A_3MX) help to identify them unambiguously. The two Lys residues were ascertained from the typical spin system [A_2(F_2T_2)MPX] seen in the TOCSY spectrum. The aromatic AMX spin system, Phe, was identified from the observation of nOe’s from the H_β protons observed to the nearest aromatic ring protons in the 2D ROESY spectrum. Once all the amino acids were identified a sequential self-walk assignments was done based on 2D TOCSY and 2D ROESY data shown as in Figure 1. The rOe cross-peaks corresponding to sequential d_αN distance led to the sequence-specific ¹H-¹H resonance assignments Figure 2 of all the amino acids from the N to the C terminal of the peptide except between Thr-15 and Pro-16, since Pro-16 since it has no HN proton. The rOe observed between H_δ protons of Pro-16 and Hα of Thr-15 helped in assigning the Thr-15 and-Pro-16 in the sequential walk. The sequential walk of the peptide sequence shown in Figure 1 with overlapping TOCSY and ROESY spectra’s.

Temperature coefficients of NH chemical shifts have been calculated from 1D spectra recorded in the temperature range of 300 to 320 K. Values below (–Δδ/ΔT) 3.00 ppb/K are indicative of a strong intramolecular H-bonds and values between 3.0-5.0 ppb/K are characteristic of dynamic equilibrium between intramolecularly hydrogen bonded form which are exposed to the solvent [39]. The values obtained fall in the range of 6-8.9 ppb/K indicating that the NH protons are freely exposed to the solvent. The results from the temperature coefficients of amide protons of the peptide KLSYDDKVFENVEFTPNL indicate that no amide proton (HN) is involved in hydrogen bonding Figure 3.

Positive CSI values for the C_αH protons of the peptide and lack of significant rOe’s (long range d_αN) and relatively high amide temperature coefficients for almost all residues, indicate a β-sheet or random coil like structure Figure 4. The J^NHα coupling constant is a function of the dihedral angle ф calculated using the Karplus equation which is helpful in identifying different peptide conformations [28]. The perceived J^NHα coupling constants for the peptide are higher than 7.0 Hz and were characteristic of a predominantly extended backbone conformation. Overall, 58 meaningful distance restraints could be obtained along with 15 dihedral angle restraints were incorporated in the restrained molecular dynamic simulation carried out for 2500 ps. The backbone torsion angle (ф,Ψ) averaged over the entire trajectory for the peptide is indicative of a very good homogeneity of the simulated structures showing a trend for a β-sheet. The structures were classified as converged consistent with the NMR data and maintained correct stereochemistry. All converged structures had few NOE constraints violations more than 0.3 Å. The superposition of the best lowest-energy NMR structures for the peptide is depicted in Figures 5 and 6. The average pairwise RMSD calculated for is 0.35 Å. Figures 7 and 8. Structural statistics tabulated in Table 1 and 2.

Overall, the peptide KLSYDDKVFENVEFTPNL had a predominantly extended backbone conformation (β-sheet like) in H_2O at 300 K.
Figure 5: 6-Best-fit backbone superposition for the best NMR structures resulting from NMR-restrained Molecular dynamics calculations of Pheromonotropin with GROMACS.

Figure 6: The NMR derived structure of Pheromonotropin in H$_2$O.

Figure 7: Using dssp program in GROMACS to depict the secondary structure of Pheromonotropin resulting from NMR-restrained Molecular dynamics.
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Figure 8: Ramachandran plot for the ensemble of best NMR structures from restrained molecular dynamics.

<table>
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<th>Distance restraints</th>
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<td>Inter-residue</td>
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<tr>
<td>Sequential</td>
<td>17</td>
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<td>NOE violations &gt;0.3Å</td>
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RMSD of backbone atoms of the ensemble against global minimum structure:

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<tr>
<td>Minimum</td>
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<tr>
<td>Average</td>
<td>0.35</td>
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Table 1: Structural statistics of Pheromonotropin in 95:5 H2O:D2O mixture

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<th>Hα</th>
<th>Hβ</th>
<th>Others</th>
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<td></td>
<td>-</td>
<td></td>
<td></td>
<td>γCH2-1.48</td>
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<td>δCH2-1.65</td>
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<td></td>
<td>εCH2-3.102</td>
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<td>K1</td>
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<td>4.38</td>
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<td>γH-1.469, δCH2-0.90.92</td>
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<td>4.407</td>
<td>1.589</td>
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<td>δCH2-1.65</td>
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<td>S3</td>
<td>8.420</td>
<td>4.24</td>
<td>3.811</td>
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<td>4.245</td>
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### Table 2: Observed values for Pheromonotropin in water.

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<th>4.414</th>
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<th>γCH₂-1.56</th>
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<th>δCH₁-1.69</th>
<th>εCH₃-3.202</th>
<th>-7.2</th>
<th>9.2</th>
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<td>3.996</td>
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<td>8.206</td>
<td>4.612</td>
<td>3.131,2.996</td>
<td>2.6H 7.30</td>
<td>4H 7.27</td>
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<tr>
<td>E10</td>
<td>8.229</td>
<td>4.670</td>
<td>2.749,2.613</td>
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<td>-7.2</td>
<td>8.8</td>
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<tr>
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<td>8.328</td>
<td>4.294</td>
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<td>V12</td>
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<td>2.063</td>
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<tr>
<td>E13</td>
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<td>4.649</td>
<td>2.23,2.53</td>
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<td>8.225</td>
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<td>3.104,3.052</td>
<td>2.6H 7.25</td>
<td>4H 7.20</td>
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<tr>
<td>T15</td>
<td>7.975</td>
<td>4.540</td>
<td>4.054</td>
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<td>4.36</td>
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<td>δH 3.56, 3.67</td>
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<td>L18</td>
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<td>δCH₃-0.925,0.870</td>
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<td>8.1</td>
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</table>

#### Conclusion

Computational modeling and simulations based on molecular dynamics provide an effective means to understand the structure and dynamics of biomolecules. Using the GROMACS molecular dynamics package, we investigated the conformation of the insect neuropeptide pheromonotropin based on NMR data. The conformation using NMR data in conjunction and molecular dynamics simulations in water yielded a β-sheet. The dynamical simulation results were visually and quantitatively analyzed for the conformation. The analysis and visualization of the generated peptide structure can give us meaningful insights into its pharmacological/physiological role aiding rational drug design for peptidomimetics and also unravelling other unknown neuropeptides in the (PK/PBAN) multifunctional family.

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