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The Structural Studies of Peptide P143 Derived from Apo B-100 by NMR

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Abstract Apolipoprotein B-100 (apo B-100), the main protein component that makes up LDL (Low density lipoprotein), consists of 4,536 amino acids and serves to combine with the LDL receptor. The oxidized LDL peptides by malondialdehyde (MDA) or acetylation in vivo act as immunoglobulin (Ig) antigens and peptide groups were classified into 7 peptide groups with subsequent 20 amino acids (P1-P302). The biomimetic peptide P143 (IALDD AKINF NEKLS QLQTY) out of C-group peptides carrying the highest value of IgG antigens were selected for structural studies that may provide antigen specificity. Experimental results show that P143 has β -sheet in Ile[1]-Asn[9] and α -helice in Gln[16]-Tyr[20] structure. Homonuclear 2D-NMR (COSY, TOCSY, NOESY) experiments were carried out for NMR signal assignments and structure determination for P143. On the basis of these completely assigned NMR spectra and proton distance information, distance geometry (DG) and molecular dynamic (MD) were carried out to determine the structures of P143. The proposed structure was selected by comparisons between experimental NOE spectra and back-calculated 2D NOE results from determined structure showing acceptable agreement. The total Root-Mean-Square-Deviation (RMSD) value of P143 obtained upon superposition of all atoms were in the set range.

The solution state P143 has a mixed structure of pseudo α -helix and β -turn(Phe[10] to Glu[12]). These results are well consistent with calculated structure

from experimental data of NOE spectra. Structural studies based on NMR may contribute to the prevent oxidation studies of atherosclerosis and observed conformational characteristics of apo B-100 in LDL using monoclonal antibodies.

Keywords Apolipoprotein B-100, Immunoglobulin, Molecular dynamic computation, NMR

Introduction

Apolipoproteins playing an important role in lipid transport and metabolic process in blood stream are components of LDL and HDL. Apolipoproteins are classified into A-I, A-II, B, C-I, C-II, D, E, respectively, based on the ABC nomenclature. There are three main lipoprotein classes according to density: very low density lipoproteins (VLDL), low density lipoproteins (LDL, d:1.019-1.063 g/mL), and high density lipoproteins (HDL, d:1.063-1.21 g/mL).¹ Studies relating to the structure and metabolism of LDL are important because of the direct correlation between atherosclerosis and high LDL levels in human plasma. LDL is the end product of VLDL catabolism and the major cholesterol transporting lipoprotein in human plasma.

Apolipoprotein B (apo B) is the largest and one of the most important proteins that cover the lipid surfaces of lipoproteins. Apo B exists in two forms, The majority of LDL particles contain a single apo called

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apo B-100.^{1,2} and apo B-48.³ which is the major protein component of plasma LDL. It plays functional roles in lipoprotein bio-synthesis in liver and intestine, and is the ligand recognized by the LDL receptor during receptor mediated endocytosis.⁴ After the elucidation of the role of apolipoproteins in

the regulation of lipoprotein metabolism, it became apparent that improvements in the characterization of apo B-100 were needed to facilitate the development of the linkage between LDL and atherosclerosis.¹

Apo B-100 which consists of 4,536 amino acids has a molecular mass of 513 kDa and its levels of both LDL-cholesterol and plasma apo B are correlated with coronary heart.⁵ The structure of human apo B has been analyzed in term of its functions in lipid binding, lipoprotein assembly and as the ligand responsible for LDL clearance by the LDL receptor pathway.⁶⁻⁸ In apo B-100 few of the predicted α -helices are truly amphipathic in terms of charge distribution on the polar surface except for one extended region which contains good examples of amphipathic α -helices, and may contribute to lipid binding. The secondary structure of apo B-100 has been suggested to consist of 43% α-helical, 21% β-sheet, 16% β-turn and 20% random structure.⁹⁻¹⁰ The β -structure of apo B-100 is thought to be responsible for its interaction with lipids, due to its high hydrophobicity, but is not confined to a particular region and various sections of the protein are buried in the lipid moiety.¹¹

Recently, computer modeling studies based upon biochemical analyses have shown that large segment of the apo B backbone have a high amphipathic structure predicted to bind lipid. If these amphipathic β -sheets and α -helices are not folded and associated with lipid in the proper temporal sequence, the structural model predicts that the hydrophobic surfaces would become unstable in the aqueous environment of the endoplasmic reticulum (ER) lumen, leading to improper folding of nascent apo B and eventual degradation.^{12,13}

Apo B-100 has 357 lysine residues per molecules, because this protein is intimately associated with lipid in the LDL particles.¹⁴ These lysine residues has highly conjugate with MDA (malondialdehyde)

during lipid peroxidation. During the peroxidative modification of LDL products to apo B occurs primarily covalent linkage to epsilon amino groups of lysine residues. If oxidized LDL contains lysine residues conjugated to a number of such fatty acid fragments, the occurrence of oxidatively modified LDL in vivo could be demonstrated by its recognition by mono specific antibodies directed against the various lysine adducts.^{6,15}

Conformational studies for biomimetic peptide P143 (IALDD AKINF NEKLS QLQTY) recognized by the monoclonal antibodies will be discussed from computer simulation analysis in this study.^{16,17}

2D-NMR signal assignments were accomplished by using 2D-NMR experiments. On the basis of these distance data from NOESY experiments, Distance Geometry (DG) and Molecular Dynamics(MD) were carried out to obtain the tertiary structure of biomimetic peptide P143.¹⁶⁻²⁰

Experimental Methods

Sample preparation- Apo B-100 biomimetic peptide P143 (IALDD AKINF NEKLS QLQTY) was obtained from Peptron Inc. Peptide was synthesized using solid-phase method. The water soluble P143 was dissolved at 3.22 mM concentration in 400 μ L DMSO-d₆ for NMR experiments.

NMR experiments- All NMR experiments were performed by using the 500 MHz (499.94 MHz) Varian spectrometer at 298 K. 2D NMR experiments included COSY, TOCSY and NOESY were performed with a 512×2048 data matrix size with 32 scans per t1 increment and spectra were zero filled of 2048×2048 data points. TOCSY spectrum was collected with a mixing time of 80 ms, MLEV-17 spin lock pulse sequence. Although NOESY experiments were recorded at two different 300 ms and 400 ms mixing time, the NOE spectrum at 300 ms mixing time was used for signal assignment and 2D NOE back-calculation. Data were processed and analyzed on a SGI Octane workstation using Felix and NMR ViewJ software. NMRViewJ was also used for sequential assignment of each amino acid.^{16,19}

Solution state structure determination- Biomimetic peptide structure determinations were carried out using HYGEOMTM and HYNMRTM. Sequential assignments of amino acid spin systems were made using TOCSY and NOESY spectra. Most important, direct way for secondary structure were determined based on qualitative analysis of NOE. The structures were calculated from the NMR data according to the standard HYGEOMTM simulated annealing and refinement protocols with minor modifications. Most of distance geometry (DG) algorithm accepts the input of proton-proton distance constraints from NOE measurements. The structure was calculated using the DG algorithm HYGEOMTM, and separated structures were generated using all of the constraints and random input. No further refinement by energy minimization was carried out on the output of the DG calculations. Root mean square distances (RMSD) deviations between the NMR structures were 0.33 Å for the backbone. Back calculation was assigned to HYNOE calculation in order to generate the theoretical NOEs. A consecutive serial files, obtained from HYNOE calculation, were incorporated into HYNMRTM to generate NOE back calculation spectra which can be directly compared with experimental NOESY spectra.16,19

Results and Discussion

NMR signal assignment- Connectivities derived from through bond J-coupling and through space coupling are important in NMR signal assignment and solution state structure determination. The proton-proton connectivities that identify the different amino acid type are established via scalar spin-spin coupling, using COSY and TOCSY.

Complete NMR signal assignments listed in Table 1 were accomplished by using homonuclear COSY and TOCSY experiments. Assignments using sequential NOEs can be obtained for proteins with natural isotope distribution. Relations between protons in sequentially neighboring amino acid residues i and i+1 are established by NOEs manifesting close approach among $d_{\alpha N}$, d_{NN} , $d_{\beta N}$.

Residue	NH	αH	βH	γH	Others
Ile[1]	-	3.612	1.756	0.876 0.833	δH ₃ : 1.107 γH ₃ : 1.476
Ala[2]	8.503	4.426	1.223	-	-
Leu[3]	8.110	4.299	1.442	1.591	$\begin{array}{l} \delta H_3 \colon 0.852 \\ \delta H_3 \colon 0.812 \end{array}$
Asp[4]	8.144	4.484	2.648 2.486	-	-
Asp[5]	8.080	4.491	2.627 2.451	-	-
Ala[6]	7.851	4.208	1.208	-	-
Lys[7]	7.967	4.212	1.648	1.297	δH_2 : 1.513 ϵH_2 : 2.753
Ile[8]	7.581	4.139	1.641	1.342 0.991	δH ₃ : 0.746 γH ₃ : 0.696
Asn[9]	8.054	4.510	2.504 2.369	-	δH ₂ : 7.433 δH ₂ : 6.954
Phe[10]	8.003	4.414	2.998 2.781	-	δH : 7.141 εH : 7.200 ζH : 7.690
Asn[11]	8.284	4.512	2.575 2.420	-	δH ₂ : 7.406 δH ₂ : 6.952
Glu[12]	7.907	4.215	1.912 1.757	2.235	-
Lys[13]	8.012	4.202	1.664	1.309	δH ₂ : 1.546
Leu[14]	7.865	4.289	1.455	1.580	δH ₃ : 0.863 δH ₃ : 0.813
Ser[15]	7.844	4.256	3.601 3.544	-	-
Gln[16]	8.026	4.206	1.910 1.741	2.100	$\delta NH_2 : 7.233$ $\delta NH_2 : 6.786$
Leu[17]	7.884	4.288	1.452	1.580	$\delta CH_3 : 0.860$ $\delta CH_3 : 0.815$
Gln[18]	8.028	4.259	1.900 1.753	2.100	$\delta NH_2: 7.271 \\ \delta NH_2: 6.788$
Thr[19]	7.613	4.190	3.916	0.991	-
Tyr[20]	7.867	4.354	-	-	δH : 6.976 εH : 6.631

 Table 1. ¹H-NMR chemical shifts of P143



Figure 1. Intraresidue cross peaks labeled in the NH- α H region (upper), NH-NH region (lower) of the NOE spectra of P143 ($\tau_m = 300$ ms).

The correlating signals of adjacent residues on the basis of dipolar connectivities obtained from 2D NOE spectra are listed in Table 2. Dipolar connectivities from amide protons to α - and amide protons were also used for sequential signal assignments, and the fingerprint region of the NOE spectra recorded at 300 ms are shown in Figure 1. Structure calculation was used to many inter and intra residue NOE connectivities.

Residue NOE connectivities δ(ppm) 4.650 $A^2_{\beta H}(m), L^3_{\delta H}(v.w)$ Ala[2]_{aH} Ala[2]_{NH} 8.728 $I_{\alpha H}^{1}(m), A_{\alpha H}^{2}(w), A_{\beta H}^{2}(w), L_{NH}^{3}(v.w)$ Leu[3]_{NH} 8.334 $A^{2}_{\alpha H}(m), A^{2}_{\beta H}(w), L^{3}_{\beta H}(s), L^{3}_{\gamma H}(w)$ $L^{3}_{\alpha H}(m), L^{3}_{\delta H}(v.w), D^{4}_{\alpha H}(s), D^{4}_{\beta H}(m)$ Asp[4]_{NH} 8.369 $D^{5}_{\alpha H}(s), D^{5}_{\beta H}(w), A^{6}_{NH}(w)$ Asp[5]_{NH} 8.305 $A^{6}_{\beta H}(s), K^{7}_{\gamma H}(m), K^{7}_{\delta H}(m)$ Ala[6]_{aH} 4.433 4.437 Lys[7] $_{\alpha H}$ $I^{8}_{\gamma H}(w)$ 8.191 $K^{7}_{\beta H}(w), K^{7}_{\gamma H}(w), K^{7}_{\delta H}(w), I^{8}_{NH}(m)$ Lys[7]_{NH} Ile[8]_{aH} 4.364 $I^{8}_{\ \beta}H(m), I^{8}_{\ \gamma H}(m), I^{8}_{\ \delta H}(w), N^{9}_{\ \alpha H}(w)$ Ile[8]_{NH} 7.806 $K^{7}_{\delta H}(w), I^{8}_{\alpha H}(m), I^{8}_{\gamma H}(w), N^{9}_{NH}(m)$ 8.279 $I^{8}_{\alpha H}(m), N^{9}_{\beta H}(m), N^{9}_{\delta H}(v.w)$ Asn[9]_{NH} $K^{7}_{\beta H}(v.w), F^{10}_{\alpha H}(w), F^{10}_{\beta H}(w)$ Asn[9]_{aH} 4.735 Phe[10]_{NH} $N_{\alpha H}^{9}(m), N_{\beta H}^{9}(w), F_{\alpha H}^{10}(s), N_{NH}^{11}(m)$ 8.228 Phe[10]_{aH} $N_{\alpha H}^{9}(w)$ 4.639 $N_{\alpha H}^{9}(w), F_{\alpha H}^{10}(s)$ Phe[10]_{βH} 3.222 Asn[11]_{NH} 8.509 $F^{10}_{\alpha H}(m), F^{10}_{\beta H}(w), N^{11}_{\alpha H}(s), E^{12}_{NH}(m)$ Asn[11] $_{\alpha H}$ 4.737 $N^{11}_{\beta H}(s), E^{12}_{\alpha H}(w)$ Glu[12]_{NH} 8.132 $F^{10}_{\ \alpha H}(w), N^{11}_{\ \alpha H}(m), N^{11}_{\ \beta H}(w), E^{12}_{\ \gamma H}(w)$ $Glu[12]_{\alpha H}$ 4.439 $N^{11}_{\beta H}(w), E^{12}_{\gamma H}(m)$ Lys[13]_{NH} 8.237 $E^{12}_{\gamma H}(w), K^{13}_{\alpha H}(s), K^{13}_{\gamma H}(w), K^{13}_{\delta H}(m)$ $F^{10}{}_{\beta H}(w),\,K^{13}{}_{\beta H}(s)$ Lys[13] αH 4.426 Lys[13]_{βH} 1.888 $K^{13}_{\nu H}(m)$ Leu[14]_{NH} 8.110 $K^{13}_{\beta H}(w), K^{13}_{\gamma H}(w), L^{14}_{\beta H}(w), L^{14}_{\gamma H}(s)$ $S^{15}_{\ \alpha H}(s), S^{15}_{\ \beta H}(m)$ Ser[15]_{NH} 8.069 $L^{17}_{\beta H}(v.w)$ Ser[15]_{BH} 3.826 Gln[16]_{NH} 8.251 $S^{15}_{\alpha H}(m), S^{15}_{\beta H}(w), Q^{16}_{\beta H}(s), Q^{16}_{\gamma H}(m)$ $Gln[16]_{\alpha H}$ 4.430 $Q^{16}_{\beta H}(s), Q^{16}_{\gamma H}(m), L^{17}_{\beta H}(m), L^{17}_{\delta H}(w)$ 2.135 $Q^{16}_{\ \gamma H}(s), L^{17}_{\ \delta H}(m)$ Gln[16]_{βH} Leu[17]_{NH} 8.105 $L^{17}_{\delta H}(w), Q^{18}_{NH}(m), Q^{18}_{\gamma H}(w)$ $L^{17}_{\beta H}(s), L^{17}_{\gamma H}(m), L^{17}_{\delta H}(m)$ Leu $[17]_{\alpha H}$ 4.493 $L^{17}_{\ \gamma H}(w), L^{17}_{\ \delta H}(s)$ Leu[17]_{βH} 1.678 $L^{17}_{\delta H}(w), Q^{18}_{\beta H}(s), Q^{18}_{\gamma H}(m), T^{19}_{NH}(m)$ Gln[18]_{NH} 8.253 Gln[18]_{aH} 4.483 $Q^{18}_{\ \gamma H}(m), T^{19}_{\ \gamma H}(v.w)$

Gln[18]_{βH}

Thr[19]_{NH}

Thr[19]_{aH}

Tyr[20]_{NH}

2.124

7.838

4.415

8.092

 $Q^{18}_{\gamma H}(s)$

 $L^{17}{}_{\beta H}(v.w), Q^{18}{}_{\alpha H}(m), Q^{18}{}_{\beta H}(w), Y^{20}{}_{NH}(m)$

 $T^{19}_{\ \beta H}(s), T^{19}_{\ \gamma H}(m)$

 $L^{17}{}_{\delta H}(m), T^{19}{}_{\gamma H}(w), Y^{20}{}_{\alpha H}(m), Y^{20}{}_{\beta H}(w)$

Table 2.	Important N	NOE	connectivities	used	for	the
structure	determination	on of	P143			



Figure 2. NOESY connectivities involving backbone protons for amino acids i and j. The height of the bars symbolizes the relative strength (strong, medium, weak) of the cross peaks in a qualitative way.

Table 2 indicates the important NOE connectivities used for the peptide structure determination. Specific conformation of peptide P143 are observed to be Ala[2]_{α H} - Asp[4]_{NH}, Ile[8]_{α H} - Glu[12]_{NH}, Ser[15]_{α H} -Gln[16]_{NH}, Gln[18]_{aH} - Thr[19]_{NH} Although relatively weak $d_{\alpha N}$ (i, i+2) dipolar connectivities from Phe[10]_{α H} - Glu[12]_{NH} were observed (Figure 2), it was apparent that the P143 has a characteristic conformation including a α -helice form and partial β -turn. In order to determine the DG structure, several variable velocities simulated annealing and conjugate gradient minimization steps were used in the refinement scheme. Addition of restraints to account for minor differences between experimental and back calculate spectra enabled the generation of new DG structures with substantially reduced penalties. To determine which of the DG structures most accurately reflect the experimental NOESY data, 2D NOESY back calculations were carried out. As illustrated in Figure 3, calculated spectrum of the P143 was generally consistent with the experimental NOESY data and best positioned DG structure are shown in Figure 4.

Discussion

NMR signal assignments of P143 peptide were made by using homonuclear proton correlation 2D NMR experiments including COSY, TOCSY and NOESY. Dipolar connectivities from amide protons to α -and amide protons were used for sequential signal assignments. NMR based solution structures were determined with important NOEs and 2D-NOE back calculation method. Out of P143 residues (IALDD AKINF NEKLS QLQTY) Phe[10] to Glu[12] clearly



Figure 3. Overlapped comparison of experimental and back-calculation NOESY spectrum of P143 collected at 300 ms mixing time. Each marked site in the figure indicate the similarity between the experiment and the back-calculated of P143 peptide structure



Figure 4. Stereo-views of P143 showing best-fit super positions of all atoms (excluding protons).

demonstrate a β -turn and β -sheet in Ile[1]-Asn[9] and α -helice in Gln[16]-Tyr[20] conformation as shown in Figure 4. As a result of NMR based solution state structure determination of Apo B-100 partial peptides P143. The final result of solution state peptide structure determination is presented as a best position of a group of conformers for pairwise minimum root mean square deviation (RMSD) relative to a predetermined conformer. It was found that peptides have a characteristic β -turn structure possibly derived from the phenylalanine in the middle of sequences as

compared. It is believed that glutamine, lysine and asparagine residue may play important role in the recognition of the monoclonal antibodies in via peroxidation with MDA. Further molecular docking studies with current results may provide the antigen specificity and regulation of surface conformational changes in the mechanism of apo lipoproteins playing an important role in lipid transport and metabolic process in blood stream.

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