

[15]. The protein band of Thy-1 (20–25 kDa) was excised, cut into pieces, and destained. The gel pieces were dehydrated with 50% acetonitrile. The dried gels were then equilibrated with 50 mM sodium phosphate buffer (pH 7.2) and incubated at 37 °C with 3 units of PNGase F. The released *N*-glycans were extracted three times from gel pieces by intermittent sonication for 30 min in water. All extracts were combined and lyophilized. The released *N*-linked oligosaccharides were reduced with NaBH₄, as previously reported [33], and subjected to GCC-LC/IT-MS-FT-ICR-MS.

2.3. *N*-linked oligosaccharide analysis by GCC-LC/IT-MS-FT-ICR-MS

GCC-LC/MS was carried out using a MAGIC 2002 system (Michrom BioResource, Auburn, CA, USA) connected to IT-MS instrument coupled with FT-ICR-MS instrument

(Finnigan LTQ FT, Thermo Electron Corp., San Jose, CA, USA). The eluents consisted of 5 mM ammonium acetate, pH 9.6, containing 2% CH₃CN (pump A), and 5 mM ammonium acetate, pH 9.6, containing 80% CH₃CN (pump B). The borohydride-reduced *N*-linked oligosaccharides were separated on Hypercarb (150 mm × 0.2 mm, 5 μm, Thermo Electron Corp.) as GCC with a linear gradient of 5–30% for pump B over a period of 60 min at a flow rate of 2 μl/min.

The MS^{*n*} experiment includes sequential scans, as follows: a full MS¹ scan (*m/z* 700–2000) by FT-ICR-MS in positive ion mode, data-dependent MS^{*n*} scans by IT-MS for most abundant ions regardless of their charge state, a full MS¹ scan (*m/z* 700–2000) by FT-ICR-MS in negative ion mode, and data-dependent MS^{*n*} scans by IT-MS for the most intense ions regardless of their charge state. For the data-dependent MS^{*n*}, the following settings were used: the isolation window for precursor masses, ±2.5 Da; collision energy, 35%; dynamic exclusion

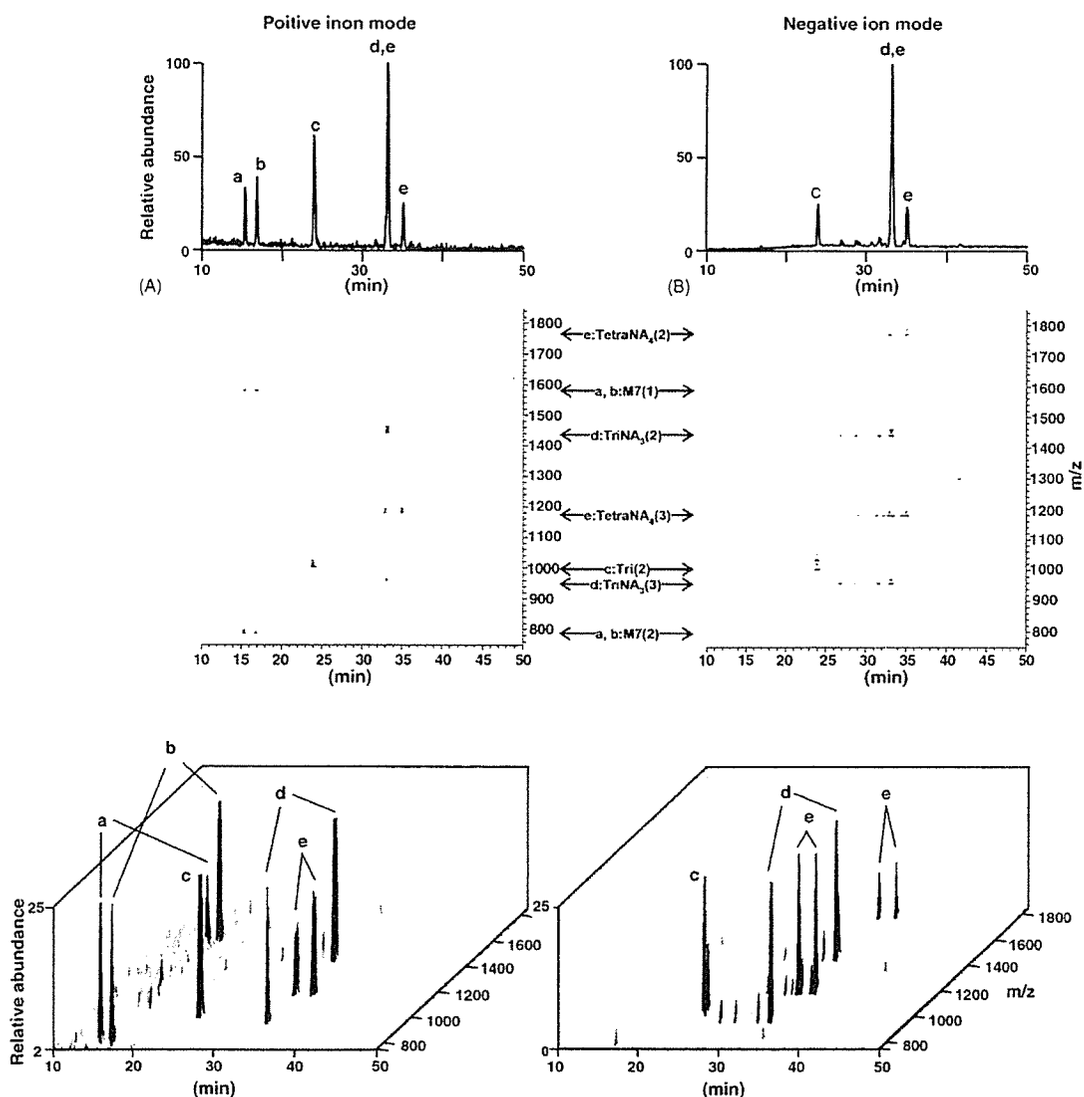


Fig. 2. Typical oligosaccharide profiles obtained by full MS¹ scans with FT-ICR-MS. (A) total ion chromatogram (TIC) (upper), two-dimensional (2D) display (retention time vs. *m/z*) (middle), and three-dimensional (3D) display (lower) in positive ion mode. (B) TIC (upper), 2D (middle) and 3D display (lower) in negative ion mode. Numbers in parentheses after the abbreviation of the model oligosaccharides refer to the charge state.

time, 15 s. The operating conditions employed for LC/MSⁿ were as follows: tube lens offset, 120 V; capillary voltage, 2.0 kV; and capillary temperature, 200 °C.

3. Results

3.1. GCC-LC/IT-MS-FT-ICR-MS of model oligosaccharides

By using the IT-MS-FT-ICR-MS instrument, the oligosaccharide profiling was shown to be more rapid, accurate, and informative. Man7/D1, Man7/D3, Tri, TriNA₃, and TetraNA₄, which were chosen as model neutral and acidic oligosaccharides (Fig. 1), were analyzed by alternative scans in positive and negative ion modes, which are consisting of a full MS¹ scan by FT-ICR-MS followed by data-dependent MSⁿ scans by IT-MS. Fig. 2A and B show the oligosaccharide profiles obtained by a full MS¹ scan with FT-ICR-MS (*m/z* 700–2000) in positive and negative ion modes, respectively. The monosaccharide compositions of individual oligosaccharides could be easily determined by accurate *m/z* values, and the major peaks of a, b, c, d, and e were assigned to Man7/D1 or D3, Man7/D3 or D1, Tri, TriNA₃ and TetraNA₄, respectively. Oligosaccharides detected at the same *m/z* values are positional isomers. Man7/D1 and D3 were detected in positive ion mode, but were only slightly detectable in negative ion mode. The major isomers of TriNA₃ and TetraNA₄ were detected in both ion modes, whereas their minor isomers were detected only in negative ion mode. These results demonstrate the advantage of alternative scans in both ion modes.

We confirmed the possibility of data-dependent MSⁿ scans for sequencing neutral and sialylated oligosaccharides. Man 7/D1 and D3 could be distinguished from each other by data-dependent MSⁿ (Fig. 3). Oligosaccharide eluted at 15 min could be assigned to Man7/D1 by the relatively intense ions at *m/z*

913 (Y_{3α}⁺) and 1237 (Y_{3β}⁺), which would be predominantly produced from Man7/D1 by the cleavage of the α1–6-linked or α1–3-linked branch arm of the core mannose (Fig. 3A) (nomenclature proposed by Domon and Costello [37]). Likewise, the oligosaccharide at 17 min could be Man7/D3 based on the intensity of Y_{3α}⁺ at *m/z* 1075 generated from Man7/D3 by the cleavages of both the α1–6-linked and α1–3-linked branch arms (Fig. 3B).

Fig. 4A and B show the product ion spectra of TetraNA₄ in positive and negative ion modes, respectively. In positive ion mode, the characteristic B ions such as *m/z* 454 (B_{2x}⁺), 657 (B_{3x}⁺), 1475 (B_{4x}⁺), and 1658 (B₆²⁺), and a ladder of several Y ions with intervals corresponding to Hex, HexNAc, and NeuAc were detected. B/Y ions were also detected at *m/z* 366, 527, 819 (B₅/Y_{3x}²⁺), and 1330 (B₆/Y_{4x}²⁺). In negative ion mode, only sialic acids were predominantly eliminated by MS² and MS³. The structural information was provided by MS⁴, whereby both B and Y ions were originated from TetraNA₂, together with the internal fragmentation ions and cross ring cleaved ions (Fig. 4B). In addition to the B and Y ions, which were predominantly produced in positive ion mode, fragment ions at *m/z* 470 (C_{2x}⁻), 1322 (Z_{6x}²⁻, [Y_{6x}⁻-H₂O]²⁻), 1241 (Z_{5x}²⁻, [Y_{5x}⁻-H₂O]²⁻), and 1057 (Y_{5x}⁻/Z_{4x}²⁻, Y_{4x}⁻/Z_{5x}²⁻, Y_{4x}⁻/Z_{5x}²⁻, Y_{5x}⁻/Z_{4x}²⁻, [Y_{4x}⁻/5x⁻-H₂O]²⁻, [Y_{4x}⁻/5x⁻-H₂O]²⁻) were detected in negative ion mode. These ions were also useful for the structural characterization of oligosaccharides.

3.2. Glycosylation analysis of Thy-1 by GCC-LC/IT-MS-FT-ICR-MS

The improved oligosaccharide profiling using IT-MS-FT-ICR-MS was applied to the glycosylation analysis of Thy-1. PIPLC-treated Thy-1 in rat brain was isolated by SDS-PAGE [31]. N-linked oligosaccharides were extracted from the gel after in-gel PNGase F digestion and were reduced

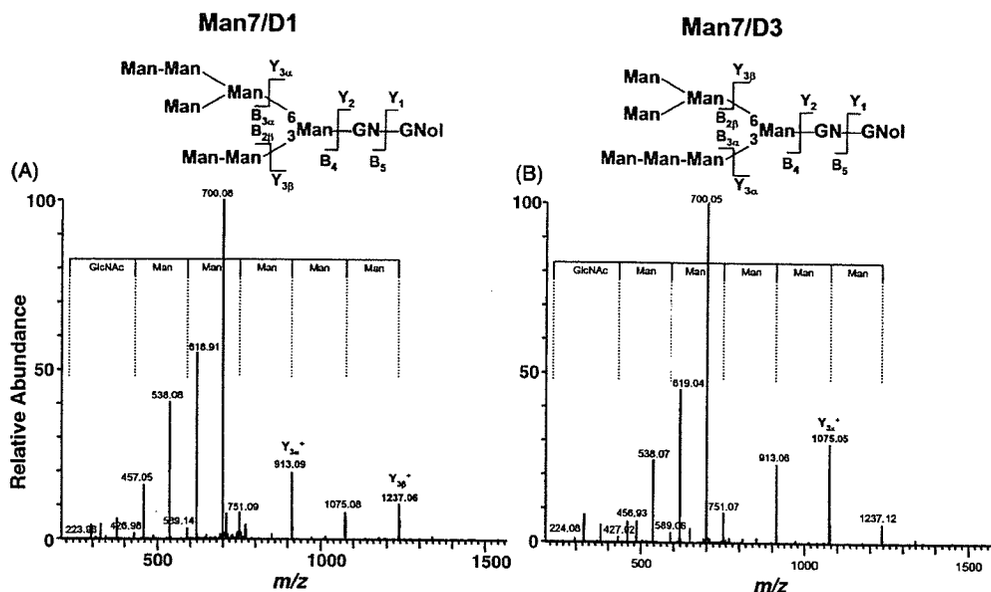


Fig. 3. Product ion spectra of oligosaccharide Man 7/D1 (A) and Man 7/D3 (B).

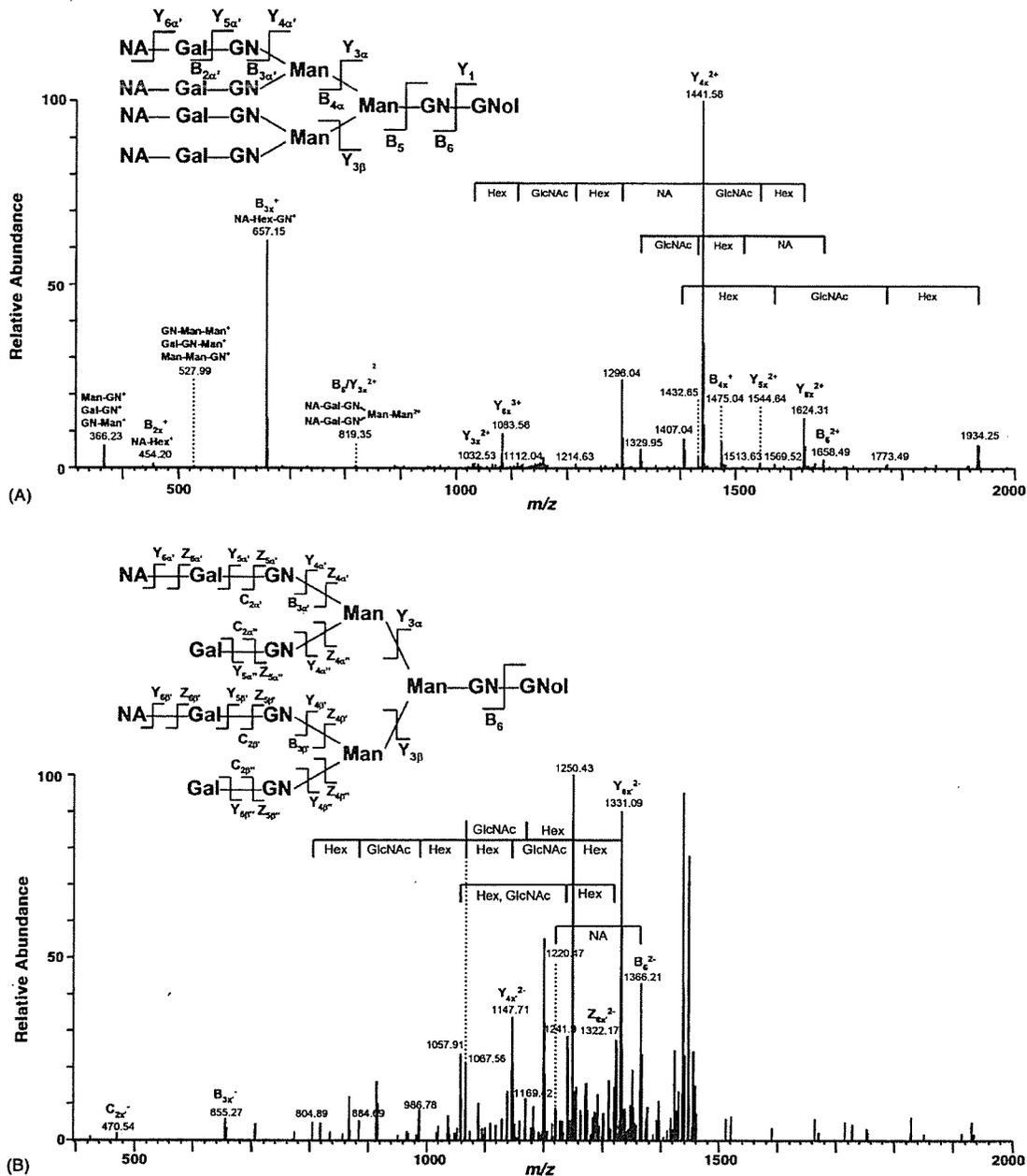


Fig. 4. Product ion spectra of model oligosaccharide, TetraNA₄. (A) MS² spectrum derived from [TetraNA₄]³⁺ at m/z 1180 in positive ion mode. (B) MS⁴ spectrum derived from [TetraNA₄]³⁻ at m/z 1178 → [TetraNA₃]³⁻ at m/z 1081 → [TetraNA₂]²⁻ at m/z 1476 in negative ion mode.

with NaBH₄. Fig. 5 shows total ion chromatogram (TIC) (A) obtained by GCC-LC/IT-MS-FT-ICR-MS of borohydrate-reduced oligosaccharides, and two-dimensional display of full MS¹ scans in positive ion mode (red) and negative ion mode (blue) (B), in which oligosaccharides appear as protonated and ammonium adducted forms along with fragment ions. Alternative scanning in positive and negative ion mode enables us to detect many oligosaccharides without missing less ionized oligosaccharides in either ion mode. For example, oligosaccharides at m/z 762 (2+) and 822 (2+) were detected only in positive ion mode, whereas those at m/z 1387 (2-), 1440 (2-), and 1542 (2-) were detected only in negative ion mode. Furthermore,

accurate m/z values acquired by FT-ICR-MS provide their monosaccharide composition, and subsequent data-dependent MSⁿ allows us to elucidate their monosaccharide sequence as follows.

3.2.1. Monosaccharide composition of oligosaccharides

Oligosaccharides in Thy-1 were assigned to NeuAc₀₋₃dHex₀₋₃Hex₃₋₉HexNAc₁₋₅HexNAc₀₁ based on their accurate m/z values (Table 1). Oligosaccharides bearing two Fuc residues, in which the m/z values of multiple charged ions are nearly identical to those of oligosaccharides bearing one NeuAc residue instead, could be determined

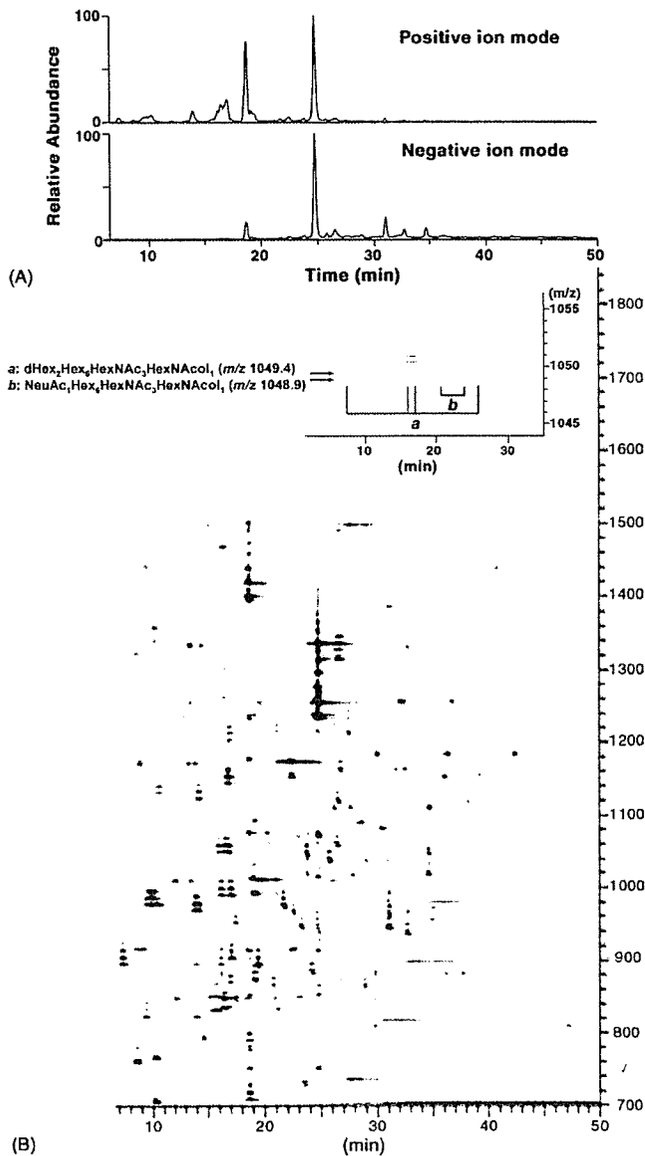


Fig. 5. N-Linked oligosaccharide profile of rat brain Thy-1 obtained by full MS¹ scans with FT-ICR-MS. (A) TIC, and (B) overlapped 2D display in positive (red) and negative (blue) ion modes.

by FT-ICR-MS. For instance, difucosylated oligosaccharides (dHex₂Hex₆HexNAc₃HexNAcol₁, theoretical molecular weight: 2096.78 Da) detected at 7.6, 16, 17, and 26 min (Fig. 5, inset, a) were clearly distinguished from monosialylated oligosaccharides (NeuAc₁Hex₆HexNAc₃HexNAcol₁, theoretical molecular weight: 2095.76 Da) detected at 21 and 24 min (Fig. 5, inset, b). The improved oligosaccharide profiling indicated that Thy-1 possesses a significant variety of N-linked oligosaccharides containing high-mannose-type (Man₅, Man₆, Man₇, Man₈, and Man₉) and many different complex- and hybrid-type oligosaccharides bearing NeuAc_{0–3} and Fuc_{0–3}. These results are coincident with those of our previous study, in which we carried out mass spectrometric analysis of Thy-1 glycopeptides.

3.2.2. Monosaccharide sequence of oligosaccharides

Monosaccharide sequences of oligosaccharides were elucidated based on the MS/MS spectra. One of the remarkable features of Thy-1 oligosaccharides is the attachment of multiple Fuc and NeuAc residues. We describe below some examples of assignment fucosylated and sialylated oligosaccharides.

3.2.2.1. Gal-(Fuc-)GlcNAc-(Lewis a/x type). Fig. 6A shows the product ion spectrum of a difucosylated oligosaccharide, dHex₂Hex₄HexNAc₃HexNAcol₁, at *m/z* 887 (24.3 min) in positive ion mode. There are two possible sites of fucosylation: GlcNAc at the non-reducing end and at the reducing end in the trimannosyl core. The ions detected at *m/z* 350 (Fuc-GlcNAc⁺, B_{2α}/Y_{5α}⁺), 370 (Fuc-GlcNAcol⁺, Y_{1α}⁺), and 512 (Gal-(Fuc-)GlcNAc⁺, B_{2α}⁺) indicate that Fuc residues link to both the non-reducing end like Lewis a/x, and the inner trimannosyl core GlcNAc. Other ions detected at *m/z* 1553 (Z_{3γ}⁺, [Y_{3γ}-H₂O]⁺) and 1570 (Y_{3γ}⁺) suggest a linkage of non-substituted HexNAc at the terminal end. From these characteristic ions together with a Y ion at *m/z* 938.03 (Y_{3α/3β}⁺), it can be deduced that this HexNAc is a bisecting GlcNAc that attaches to the core mannose residue via β1–4 linkage. On the basis of these product ions, the oligosaccharide is assigned to the structure indicated in Fig. 6A, inset.

3.2.2.2. Fuc-Gal-(Fuc-)GlcNAc-(Lewis b/y type). Fig. 6B is the product ion spectrum of a difucosylated oligosaccharide, dHex₂Hex₅HexNAc₄HexNAcol₁, at *m/z* 1070 (9.2 min). The characteristic ions at *m/z* 512 (Gal-(Fuc-)GlcNAc⁺, Fuc-Gal-GlcNAc⁺, B_{3α}/Y_{6α}⁺, B_{3α}/Y_{5α}⁺) and 1915 (B₆⁺) suggest the absence of Fuc at the reducing end GlcNAc; a B ion at *m/z* 658 (B_{3α}⁺), a B/Y ion at *m/z* 350, and a Y ion at *m/z* 1408 (Y_{3β/4α}⁺) suggest the attachment of two Fuc to Gal-GlcNAc at the non-reducing end, in a similar manner to the Lewis b/y antigen, Fuc-Gal-(Fuc-)GlcNAc-. A Y ion at *m/z* 1936 (Y_{4α}⁺) indicates a linkage of non-substituted HexNAc at the terminal end. A B/Y ion at *m/z* 877 (B_{4α}/Y_{5α}⁺, B_{4α}/Y_{6α}⁺) and a Y ion at *m/z* 1610 (Y_{3β}⁺) suggest that this non-substituted HexNAc residue is linked to the mannose residue attached to the Fuc-Gal-(Fuc-)GlcNAc- structure. These ions lead to assignment of this oligosaccharide as the structure indicated in Fig. 6B, inset.

3.2.2.3. NeuAc-Gal-(NeuAc-)GlcNAc-. Fig. 7A shows the product ion spectrum of a disialylated oligosaccharide, NeuAc₂dHex₁Hex₅HexNAc₂HexNAcol₁, at *m/z* 1085 (30.4 min). Characteristic fragment ions at *m/z* 495 (B_{3α}/Y_{5α}⁺), 948 (B_{3α}⁺), and 1110 (B_{4α}⁺), together with B ions at *m/z* 453 (B_{2α}⁺) and 657 (B_{3α}/Y_{5α}⁺, B_{3α}/Y_{6α}⁺) suggest the presence of a partial structure of NeuAc-Gal-(NeuAc-)GlcNAc-. Furthermore, detection of Y ions at *m/z* 370 (Y_{1α}⁺) and 1059 (Y_{3α}⁺, Y_{4α/4β}⁺) as well as a B ion at *m/z* 1799 (B₆⁺) indicate the linkage of a Fuc residue at the inner trimannosyl core GlcNAc. Based on these product ions, the oligosaccharide detected at *m/z* 1085 was assigned to the structure in Fig. 7A, inset.

Table 1
Summary of N-linked oligosaccharides of rat brain Thy-1

Monosaccharide composition ^a				Theoretical mass ^b	Positive ion mode		Negative ion mode		Deduced structure ^d
dHex	Hex	HX	NA		Observed m/z^c	Retention time (min)	Observed m/z^c	Retention time (min)	
1	3	2	0	1058.40	1059.46(1)	29.17			CoreF
0	5	2	0	1236.45	1237.47(1)	24.74	1235.45(1)	24.76	M5
0	3	4	0	1318.50	1319.57(1)	8.63			
0	6	2	0	1398.50	1399.53(1)	18.73	1397.50(1)	18.68	M6
0	5	3	0	1439.53	1440.59(1)	9.17			
1	3	4	0	1464.56	733.31(2), 1465.65(1)	23.44			
0	3	5	0	1521.58	761.80(2)	8.63			BisectGN
0	7	2	0	1560.55	781.29(2), 1561.60(1)	18.66			M7
1	5	3	0	1585.59	793.82(2)	14.59			Hybrid
1	5	3	0	1585.59	793.81(2)	19.13			
1	5	3	0	1585.59	793.83(2)	20.96			
0	5	4	0	1642.61	822.33(2)	9.48			Hybrid
0	5	4	0	1642.61	822.33(2)	14.02			
1	3	5	0	1667.64	834.83(2), 1668.69(1)	16.48	832.81(2)	16.44	CoreF
0	4	5	0	1683.63	842.85(2)	17.48			Hybrid
0	8	2	0	1722.61	870.83(2) ^e	17.07			M8
0	5	3	1	1730.62	866.34(2)	20.31			
0	5	3	1	1730.62	866.35(2)	28.91	864.31(2), 1729.64(1)	28.93	Hybrid
2	5	3	0	1731.64	866.86(2)	20.83	864.81(2)	20.85	Hybrid, CoreF, Lax
1	6	3	0	1747.64	874.84(2)	19.19			
2	4	4	0	1772.67	887.37(2)	23.84	885.33(2)	23.86	
2	4	4	0	1772.67	887.36(2)	24.25	885.33(2)	24.27	Hybrid, CoreF, BisectGN
1	5	4	0	1788.67	895.36(2)	7.37			Hybrid
1	5	4	0	1788.67	895.36(2)	13.90			
1	5	4	0	1788.67	895.35(2)	14.16			Hybrid, CoreF
1	5	4	0	1788.67	895.35(2)	19.44	893.33(2)	19.46	Hybrid, CoreF
0	6	4	0	1804.66	903.35(2)	17.07	901.33(2)	17.09	Hybrid, BisectGN
1	4	5	0	1829.69	915.88(2)	8.63			Hybrid
1	4	5	0	1829.69	915.88(2)	9.17			Hybrid
1	4	5	0	1829.69	915.89(2)	18.00			Hybrid
1	4	5	0	1829.69	915.89(2)	22.61	913.84(2)	22.63	
1	5	3	1	1876.68	939.37(2)	21.17	937.34(2)	21.12	
1	5	3	1	1876.68	939.36(2)	24.90	937.34(2)	24.92	
1	5	3	1	1876.68	939.39(2)	32.76	937.33(2)	32.78	Hybrid, CoreF
0	9	2	0	1884.66	951.88(2) ^f	17.53			M9
0	6	3	1	1892.68	947.39(2)	23.31	945.33(2)	23.33	Hybrid
0	6	3	1	1892.68	947.39(2)	31.09	945.33(2)	31.05	Hybrid
2	6	3	0	1893.70	947.87(2)	24.61	945.84(2)	24.70	
1	4	4	1	1917.71			957.85(2)	27.73	
1	4	4	1	1917.71			957.85(2)	28.86	
1	4	4	1	1917.71			957.85(2)	34.91	CoreF
1	4	4	1	1917.71			957.85(2)	35.55	
0	5	4	1	1933.70	967.89(2)	22.61	965.85(2)	22.63	Hybrid
0	5	4	1	1933.70	967.86(2)	24.61	965.82(2)	24.70	
2	5	4	0	1934.72	968.39(2)	13.97			Hybrid
1	6	4	0	1950.72	976.39(2)	9.93			Hybrid, Lax
1	6	4	0	1950.72	976.41(2)	21.76	974.35(2)	21.79	Hybrid, CoreF
2	4	5	0	1975.75	988.90(2)	16.21	986.85(2)	16.16	Complex
2	4	5	0	1975.75	988.90(2)	17.07	986.87(2)	17.09	Complex
0	5	3	2	2021.72			1009.86(2)	26.35	
0	5	3	2	2021.72			1009.85(2)	26.83	
1	6	3	1	2038.73	1020.40(2)	23.84	1018.36(2)	23.80	
1	6	3	1	2038.73	1020.44(2)	27.77	1018.37(2)	27.80	CoreF
1	6	3	1	2038.73	1020.42(2)	34.66	1018.36(2)	34.69	Hybrid, CoreF
1	5	4	1	2079.76	1040.92(2)	25.73	1038.87(2)	25.81	CoreF
1	5	4	1	2079.76	1040.92(2)	29.04	1038.88(2)	28.99	
3	5	4	0	2080.78	1041.42(2)	23.84	1039.39(2)	23.86	
0	6	4	1	2095.76	1048.94(2)	20.57			Hybrid
0	6	4	1	2095.76	1048.91(2)	23.84	1046.87(2)	23.80	Hybrid
2	6	4	0	2096.78	1049.42(2)	7.58			

Table 1 (Continued)

Monosaccharide composition ^a				Theoretical mass ^b	Positive ion mode		Negative ion mode		Deduced structure ^d
dHex	Hex	HX	NA		Observed <i>m/z</i> ^c	Retention time (min)	Observed <i>m/z</i> ^c	Retention time (min)	
2	6	4	0	2096.78	1049.42(2)	15.97			
2	6	4	0	2096.78	1049.42(2)	16.61			Hybrid, BisectGN
2	6	4	0	2096.78	1049.43(2)	25.73			
1	7	4	0	2112.77			1055.38(2)	34.62	
1	4	5	1	2120.79	1061.45(2)	20.43			Complex
1	4	5	1	2120.79	1061.45(2)	24.74	1059.39(2)	24.70	
1	4	5	1	2120.79	1061.45(2)	26.47	1059.39(2)	26.42	CoreF
2	5	5	0	2137.80	1069.94(2)	9.17			Lby
2	5	5	0	2137.80	1069.94(2)	21.30			
2	5	5	0	2137.80	1069.95(2)	23.09	1067.9(2)	23.04	
1	5	3	2	2167.78	1084.94(2)	30.41	1082.89(2)	30.37	Hybrid, CoreF
2	4	6	0	2178.83	1090.45(2)	26.08			
0	6	3	2	2183.77	1092.95(2)	28.63	1090.88(2)	28.60	Hybrid, diSia
0	5	4	2	2224.80	1113.45(2)	26.10			
2	5	4	1	2225.82	1113.95(2)	27.56			
2	5	4	1	2225.82	1113.98(2)	34.80			
1	6	4	1	2241.81	1121.95(2)	26.60	1119.90(2)	26.63	
1	6	4	1	2241.81			1119.90(2)	30.58	
1	6	4	1	2241.81			1119.91(2)	38.14	
3	6	4	0	2242.83	1122.46(2)	14.23			
2	7	4	0	2258.83	1130.46(2)	10.47			
3	5	5	0	2283.86	1142.96(2)	16.87			
1	4	6	1	2323.87	1162.98(2)	26.60			
1	6	3	2	2329.83			1163.91(2)	31.72	
1	6	3	2	2329.83			1163.91(2)	32.54	Hybrid, diSia
1	5	4	2	2370.86	1186.55(2)	29.89	1184.42(2)	30.00	Complex, CoreF
1	5	4	2	2370.86			1184.43(2)	36.00	
1	5	4	2	2370.86	1186.52(2)	36.31	1184.42(2)	36.40	Complex, CoreF
1	5	4	2	2370.86	1186.50(2)	42.47	1184.43(2)	42.43	Complex, CoreF
3	5	4	1	2370.86			1184.93(2)	30.99	
2	5	5	1	2428.90	1215.50(2)	21.17	1213.44(2)	21.25	
2	5	5	1	2428.90	1215.50(2)	23.84			
2	5	5	1	2428.90	1215.52(2)	26.32	1213.45(2)	26.28	
2	5	5	1	2428.90	1215.50(2)	27.50	1213.45(2)	27.53	
2	5	4	2	2516.91	1259.60(2)	32.23	1257.45(2)	32.19	Complex, Lax, CoreF, diSia
2	5	4	2	2516.91			1257.45(2)	36.72	
2	5	6	1	2631.98			876.32(3), 1314.99(2)	26.76	
3	5	4	2	2662.97			1330.49(2)	32.78	
1	5	6	2	2777.02			1387.50(2)	30.99	
0	6	5	3	2881.03			1439.50(2)	34.83	
0	6	5	3	2881.03			1439.49(2)	40.77	
2	6	5	2	2882.05			1440.05(2)	37.96	
2	6	6	2	3085.13			1541.55(2)	31.38	

^a dHex, deoxyhexose; Hex, hexose; HX, *N*-acetylhexamine; NeuAc, *N*-acetylneuramic acid.

^b Monoisotopic value.

^c Values in parentheses are charge state.

^d Structures are deduced by MSⁿ. Complex, complex-type-oligosaccharide; Hybrid, hybrid-type-oligosaccharide; M5-9, high-mannose-type oligosaccharide containing 5–9 mannose residues; BisectGN, bisecting GlcNAc; Lax, Lewis a/x structure; Lby, Lewis b/y structure; diSia, disialic acid.

^e Ammonium adducted form.

3.2.2.4. NeuAc-Gal-GlcNAc— Fig. 7B shows the product ion spectrum of a disialylated oligosaccharide, NeuAc₂dHex₁Hex₅HexNAc₃HexNAc₁, at *m/z* 1187 (42.5 min). Although B ions were detected at *m/z* 454 (B_{2x}⁺), 657 (B_{3x}⁺) and 819 (B_{4x}⁺), none of the fragment ions at *m/z* 495 (NeuAc-GlcNAc⁺), 948 (NeuAc-Gal-(NeuAc—)GlcNAc⁺), or 1110 (NeuAc-Gal-(NeuAc—)GlcNAc-Man⁺) were detected in the spectrum. This result suggests that the two NeuAc residues occupy both non-reducing ends of the biantennary

form. Fucosylation of the inner trimannosyl core GlcNAc was determined by the detection of Y ions at *m/z* 370 (Y_{1α}⁺) and 1059 (Y_{4α/4β}⁺). These product ions lead to assignment of this oligosaccharide the structure in Fig. 6B.

3.2.2.5. (v) NeuAc-NeuAc— Fig. 7C shows the product ion spectrum of a disialylated and difucosylated oligosaccharide, NeuAc₂dHex₂Hex₅HexNAc₃HexNAc₁, at *m/z* 1260 (32.2 min). The characteristic ions at *m/z* 583 (NeuAc-NeuAc⁺,

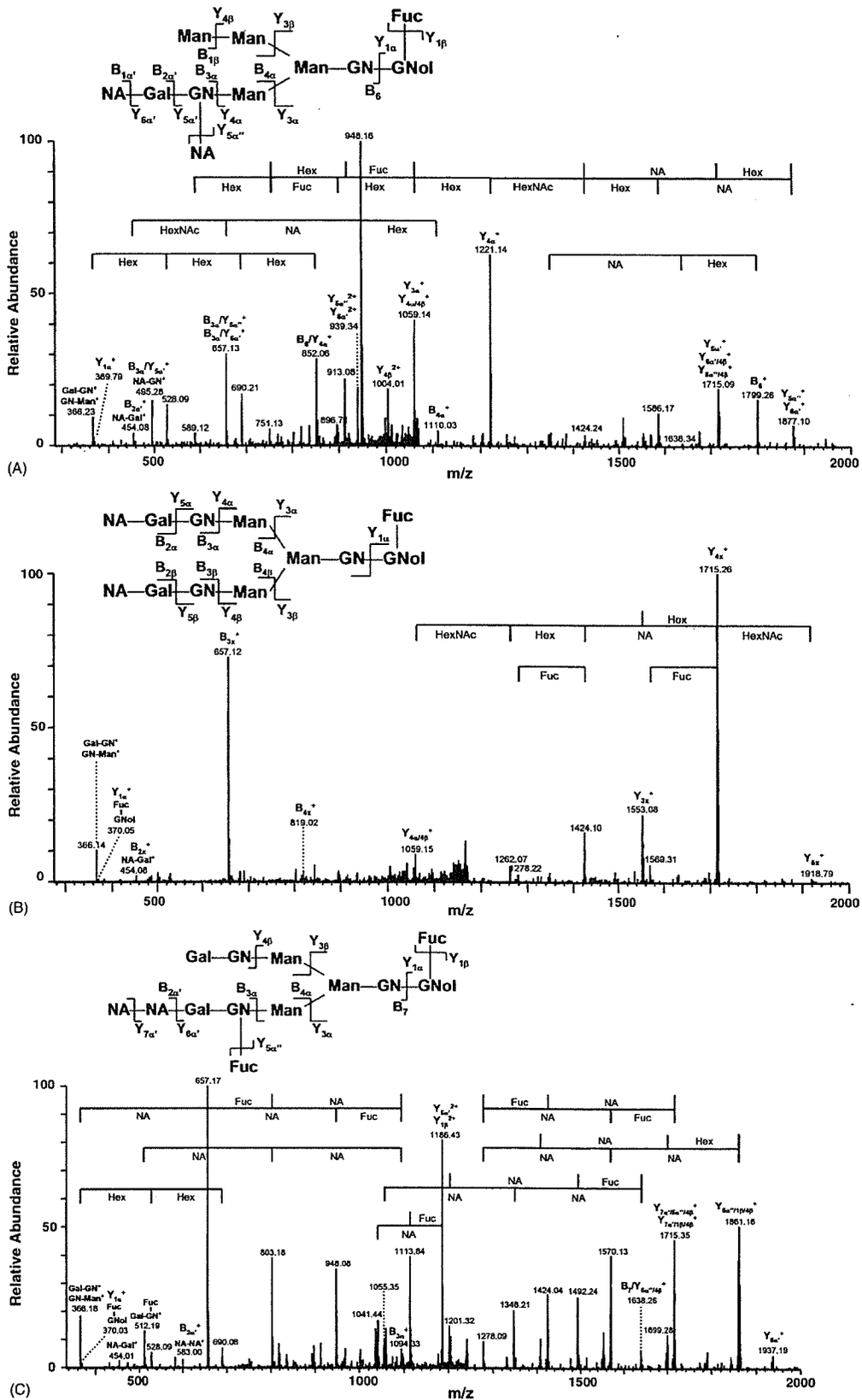


Fig. 7. Product ion spectra of N-linked oligosaccharides of rat brain Thy-1. (A) MS² spectrum of NeuAc₂dHex₁Hex₅HexNAc₂HexNAcol₁ at *m/z* 1085 (30.4 min). (B) MS² spectrum of NeuAc₂dHex₁Hex₅HexNAc₃HexNAcol₁ at *m/z* 1187 (42.5 min). (C) MS² spectrum of NeuAc₂dHex₂Hex₅HexNAc₃HexNAcol₁ at *m/z* 1260 (32.2 min).

In these two studies, we have demonstrated a strategy for glycosylation analysis of Thy-1, including identification of a glycoprotein, determination of glycosylation sites, site-specific glycosylation analysis, and structural analysis of oligosaccharide details. This strategy can be applied to glycosylation analysis of other glycoproteins. Specifically, the glycoprotein sample is divided into two. One is subjected to proteinase digestion followed by peptide/glycopeptide mapping, which provides information on glycosylation sites and site-specific heterogeneity. The other is subjected to PNGase F digestion followed by mass spectrometric oligosaccharide profiling, by which a detailed structure of *N*-glycans released from a glycoprotein could be provided. Recently, proteomic approaches, which are based on two-dimensional electrophoresis followed by mass spectrometry, have been used in various fields. Although glycosylation analysis of abundant glycoproteins in gel has been successful, that of a low-abundance glycoprotein in gel remains a great challenge. The proposed method consisting of peptide/glycopeptide mapping followed by oligosaccharide profiling with sequential scans by IT–MS–FT–ICR–MS will likely be a powerful tool for glycosylation analysis of low-abundance glycoproteins and for proteomics/glycomics.

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Site-specific N-glycosylation analysis of human plasma ceruloplasmin using liquid chromatography with electrospray ionization tandem mass spectrometry

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Abstract

Ceruloplasmin has ferroxidase activity and plays an essential role in iron metabolism. In this study, a site-specific glycosylation analysis of human ceruloplasmin (CP) was carried out using reversed-phase high-performance liquid chromatography with electrospray ionization tandem mass spectrometry (LC-ESI-MS/MS). A tryptic digest of carboxymethylated CP was subjected to LC-ESI-MS/MS. Product ion spectra acquired data-dependently were used for both distinction of the glycopeptides from the peptides using the carbohydrate B-ions, such as m/z 204 (HexNAc) and m/z 366 (HexHexNAc), and identification of the peptide moiety of the glycopeptide based on the presence of the b- and y-series ions derived from the peptide. Oligosaccharide composition was deduced from the molecular weight calculated from the observed mass of the glycopeptide and theoretical mass of the peptide. Of the seven potential N-glycosylation sites, four (Asn119, Asn339, Asn378, and Asn743) were occupied by a sialylated biantennary or triantennary oligosaccharide with fucose residues (0, 1, or 2). A small amount of sialylated tetraantennary oligosaccharide was detected. Exoglycosidase digestion suggested that fucose residues were linked to reducing end GlcNAc in biantennary oligosaccharides and to reducing end and/or α 1–3 to outer arms GlcNAc in triantennary oligosaccharides and that roughly one of the antennas in triantennary oligosaccharides was α 2–3 sialylated and occasionally α 1–3 fucosylated at GlcNAc.

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Keywords: Ceruloplasmin; Glycopeptide; Liquid chromatography-electrospray tandem mass spectrometry; Product ion spectrum; Exoglycosidase digestion

Ceruloplasmin (CP)¹ is a blue copper serum glycoprotein synthesized in the liver. CP has ferroxidase activity and plays an essential role in iron metabolism [1–4]. The primary structure of human CP has been determined by amino acid sequencing, and it is composed of a single poly-

peptide chain of 1046 amino acid residues [5]. The amino acid sequence was confirmed from complete cDNA sequence [6]. The major oligosaccharides in human CP were reported to be sialylated bi- and triantennary structures with or without a fucose residue [7,8]. Although four N-glycosylation sites (Asn119, Asn339, Asn378, and Asn743) were identified among seven potential sites [9], the heterogeneity of oligosaccharides was still unknown at each glycosylation site. CP is an acute phase reactant, and the serum concentration increases during inflammation, infection, and trauma [10]. It is known that the patterns of glycosylation are changed by inflammatory cytokines [11]. Several studies have reported that CP is a good diagnostic marker of solid malignant tumors [12,13] and that the CP glycoform might

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¹ Abbreviations used: CP, ceruloplasmin; LC-ESI-MS, liquid chromatography with electrospray ionization mass spectrometry; Hex, hexose; HexNAc, N-acetylhexosamine; LC-ESI-MS/MS, liquid chromatography with electrospray ionization tandem mass spectrometry; EDTA, ethylenediaminetetraacetic acid; TFA, trifluoroacetic acid; Q-TOF, quadrupole time-of-flight; TIC, total ion chromatogram; NeuAc, N-acetylneuraminic acid; GlcNAc, N-acetylglucosamine; Fuc, fucose.

be a valuable supplement [12]. Thus, it is important to conduct a site-specific glycosylation analysis of normal human CP.

One of the most effective techniques for determining the site-specific carbohydrate heterogeneity of glycoproteins is the mass spectrometric peptide mapping of proteolytic fragments of glycoproteins by liquid chromatography with electrospray ionization mass spectrometry (LC-ESI-MS) [14–19]. The specific detection of glycopeptides in a complex peptide mixture is generally achieved by monitoring specific carbohydrate fragment ions such as m/z 204 (HexNAc) and m/z 366 (HexHexNAc) produced by cone voltage fragmentation or by precursor ion scanning [15–19]. Because product ion spectra of glycopeptides show high abundant carbohydrate fragment ions and low abundant b- and y-series fragment ions derived from the peptide backbone [20,21], product ion spectra acquired data-dependently in liquid chromatography with electrospray ionization tandem mass spectrometry (LC-ESI-MS/MS) can be used for both the selection from the peptides and the identification of the glycopeptides [22]. MS in combination with specific exoglycosidase digestions allows us to obtain the site-specific information on anomericity and linkage of glycans [23]. In the current study, we conducted a site-specific glycosylation analysis of human CP and successfully determined glycosylation status and glycosylation profile at each N-glycosylation site.

Materials and methods

Materials

Acetonitrile, formic acid, and guanidine hydrochloride were purchased from Wako Pure Chemicals Industries (Osaka, Japan). Purified human CP was purchased from Calbiochem (San Diego, CA, USA). Modified trypsin was purchased from Promega (Madison, WI, USA). α 2–3 Neuraminidase (EC 3.2.1.18) of *Macrobodella decora*, a recombinant form, and α 1–3,4 fucosidase (EC 3.2.1.51) from *Xanthomonas* sp. were purchased from Calbiochem. α 2–3,6,8,9 Neuraminidase (EC 3.2.1.18) of *Arthrobacter ureafaciens*, a recombinant form, and β 1–4 galactosidase (EC 3.2.1.23) were purchased from Sigma Chemical (St. Louis, MO, USA). The water used was obtained from a Milli-Q water system (Millipore, Bedford, MA, USA). All other reagents were of the highest quality available.

Reduction and S-carboxymethylation of CP

CP (100 μ g) was dissolved in 270 μ l of 0.5 M Tris–HCl buffer (pH 8.5) that contained 8 M guanidine hydrochloride and 5 mM ethylenediaminetetraacetic acid (EDTA). After the addition of 2 μ l of 2-mercaptoethanol, the mixture was incubated for 2 h at 40 °C. Then 5.67 mg of monoiodoacetic acid was added, and the resulting mixture was incubated for 2 h at 40 °C in the dark. The reaction mixture was applied to a PD-10 column (Amersham Biosciences, Upp-

sala, Sweden) to remove the reagents, and the eluate was lyophilized.

Trypsin digestion of CP

Reduced and carboxymethylated CP was redissolved in 100 μ l of 0.1 M Tris–HCl buffer (pH 8.0). An aliquot of 1 μ l of trypsin prepared as 1 μ g/ μ l was added to 50 μ l of CP solution (1:50, w/w), and the mixture was incubated for 16 h at 37 °C. The enzyme digestion was stopped by storing at –20 °C before analysis.

HPLC of tryptic digest of CP

Tryptic digests (0.2 and 0.4 μ g) of human CP were analyzed by LC-ESI-MS/MS to identify the peptides and glycopeptides, respectively. HPLC was performed on a Paradigm MS 4 (Michrome BioResources, Auburn, CA, USA) equipped with a Magic C18 column (0.2 μ , 50 mm, Michrome BioResources). The eluents consisted of water containing 2% (v/v) acetonitrile and 0.1% (v/v) formic acid (pump A) and 90% acetonitrile and 0.1% formic acid (pump B). Trypsin-digested samples were loaded onto a microtrap (peptide captrap, Michrom BioResources). After a wash with 15 μ l H₂O/CH₃CN (98:2) with 0.1% trifluoroacetic acid (TFA), the trapping column was switched into line with the column. Samples were eluted with 5% of B for 10 min, followed by a linear gradient from 5 to 65% of B in 60 min at a flow rate of 2 μ l/min.

ESI-Q-TOF-MS/MS

Mass spectrometric analyses were performed using a quadrupole time-of-flight (Q-TOF) mass spectrometer (QSTAR Pulsar, MDS Sciex, Toronto, Canada) equipped with a nano-electrospray ion source. The mass spectrometer was operated in the positive ion mode. The nanospray voltage was set at 2500 V. Mass spectra were acquired at m/z 400–2000 or m/z 1000–2000 for MS analysis and at m/z 100–2000 for MS/MS analysis. After every regular MS acquisition, two MS/MS acquisitions against top two of the multiply charged molecular ions were performed (data-dependent acquisition). The precursor ions with the same m/z as acquired previously were excluded for 120 s. The collision energy was varied between 30 and 80 eV depending on the size and charge of the molecular ion. Accumulation times for the spectra were 1.0 and 2.0 s for MS and MS/MS, respectively. All peaks were resolved monoisotopically.

Tandem MS/MS data from LC-ESI-MS/MS runs were submitted to the search engine Mascot to identify the tryptic peptides of CP. One missed cleavage was allowed, and tolerances of 2.0 and 0.8 u mass were used for precursor and product ions, respectively. From the data for LC-ESI-MS/MS at m/z 1000–2000, glycopeptide precursor ions were selected manually based on the presence of oligosaccharide oxonium ions such as m/z 204 (HexNAc) and m/z 366 (HexHexNAc). The glycopeptide ions were assigned based on

the presence of b- and y-series fragment ions of peptides of putative glycopeptides or molecular weight difference of sugar unit. The molecular weight of the carbohydrate in the glycopeptide was calculated from the molecular weights of the glycopeptide and the suggested peptide. The oligosaccharide composition and type were deduced from the molecular weight of the carbohydrate.

Oligosaccharide sequencing by exoglycosidase digestions

Trypsin in the digest of human CP was inactivated by boiling for 5 min at 100 °C. Aliquots of the digest (4 µg) were digested in a volume of 20 µl for 12 h at 37 °C in 50 mM sodium phosphate buffer (pH 5.0) using the following exoglycosidases alone or in combination: α2–3 neuraminidase, 20 mU/ml; α2–3,6,8,9 neuraminidase, 100 mU/ml; α1–3,4 fucosidase, 20 mU/ml; and β1–4 galactosidase, 30 mU/ml. Aliquots (0.08 µg) before and after exoglycosidase digestions were subjected to LC-ESI-MS at *m/z* 700 to 2000 in which MS/MS acquisition was not performed.

Results

Peptide mapping of tryptic digest of human CP (LC-ESI-MS/MS in *m/z* range of 400–2000)

The amino acid sequence of human CP (National Center for Biotechnology Information protein database: P00450) is shown in Fig. 1. The tryptic peptides, including potential N-glycosylation sites, are shown in bold type. Trypsin can digest human CP into seven glycopeptides containing only one potential N-glycosylation site. To determine the glycosylation state at each glycosylation site, we performed mass spectrometric peptide mapping of the tryptic digest of CP. An aliquot of 0.2 µg of the tryptic digest was analyzed by

LC-ESI-MS/MS in the *m/z* range of 400–2000 (data not shown). When molecular ions with more than a single charge were detected, the product ion spectrum was acquired automatically. Peptide identification of each product ion spectrum was done using the Mascot search engine. More than 70% of the amino acid sequence was identified; identified amino acids of CP are underlined in Fig. 1. Three peptides containing the potential N-glycosylation site (Asn208, Asn569, and Asn907 [residues 197–218, 558–579, and 895–917, respectively]) were detected, whereas peptides containing the other N-glycosylation sites were not detected. Thus, Asn119, Asn339, Asn378, and Asn743 might be glycosylated.

Glycosylation analysis of human CP (LC-ESI-MS/MS in the *m/z* range of 1000–2000)

N-glycosylated peptides have relatively high molecular weights due to their oligosaccharide moiety. Because ions at lower *m/z* values can be detected in the *m/z* range of 400–2000, glycopeptide ions with higher *m/z* values might be missed to obtain product ion spectra. To detect glycopeptide ions preferentially, another LC-ESI-MS/MS analysis was carried out in the *m/z* range of 1000–2000 using an aliquot of 0.4 µg of the tryptic digest. Fig. 2A shows a total ion chromatogram (TIC) of a TOF-MS scan for the full scan *m/z* 1000–2000. Fig. 2B shows a TIC of the product ion scan. Because product ion spectra of glycopeptide precursor ions have abundant carbohydrate B-ions, *m/z* 204 (HexNAc), *m/z* 186 (HexNAc-H₂O), *m/z* 366 (HexHexNAc), and *m/z* 292 (NeuAc), the extracted ion chromatogram at *m/z* 204.05–204.15 (HexNAc, 204.08) of the product ion scan is illustrated in Fig. 2C. The extracted ion chromatogram at *m/z* 204.05–204.15 of product ion spectra provides a useful indication of the selection of glycopeptide precursor ions. The glycopeptide ions were assigned based on an examination of product ion spectra using the information on amino acid sequences of the peptides containing a putative N-glycosylation site.

Identification of Asn119 glycopeptide

The product ion spectrum of 1366.6 (+3) at 26 min, labeled by A in Fig. 2C, is shown in Fig. 3A. There were abundant oligosaccharide oxonium ions such as *m/z* 204 (HexNAc), *m/z* 366 (HexHexNAc), *m/z* 186 (HexNAc-H₂O), *m/z* 168 (HexNAc-2H₂O), *m/z* 274 (NeuAc-H₂O), and *m/z* 292 (NeuAc). Thus, this precursor ion was assigned as a glycopeptide. Several fragment ions consistent with b- and y-series fragment ions [24] derived from the peptide EHEGA IYPDNTTDFQR (residues 110–125) were detected together with several deamidated (–17) or dehydrogenated (–18) b- and y-series ions and y-series ions with the GlcNAc residue. Thus, the peptide moiety EHEGA IYPDNTTDFQR was suggested. The carbohydrate's molecular weight, 2223.0, was calculated by subtracting the theoretical molecular weight of the peptide (1891.8) from

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KEKHYIIGII ETTWDYASDH GEKKLISVDT EHSNIYLONG PDRIGRLYKK ALYLQYTD119
FRITIEKPVW LGFLGPIIKA ETGDKVYVHL KNLASRPYTF HSHGITYYKE HEGAIYPDNT
TDFQRADDKV YPGEQYTYML LATEEQSPGE GDGNCVTRII Y HSHIDAPKDI ASGLIGPLTI
CKKDSLDEK EKHIDREBFVY MFSVVDENFS WYLEDNIKTY CSEPEKVDK NEDFQESNRM
YVNGYTFGS LPGLSMCAED RVKWLFCMG NEVDVHAAFF HGOALTNKNY RIDTNLFPA
TLFDAYMVAQ NPGEWMLSCQ NLNHLKAGLQ AFFQVQECNK SSSKDNIRGK HVRHYIIAAE
EIIWNYAPSG IDIFTKENLT 208 APGSDSAVFF EQGTTRIGGS YKLVYREYT DASFTNRKER
GPEEHLGLIL GPVIWA569EVGD TIRVTFNHNG AYPLSIEPIG VRFNKNNEGT YVSPNYNPOS
RSVPPSASHV APTETFTYEW TVPKEVGETN ADPVCLAKMY YSAVDPTKDI FTGLIGPMKI
CKGSSLHANG RQKDV907DK743EFY LFPTVFDENE SLLLEDNIRM FTTAPDQVDK EDEDQESNK
MHSNMGFMYG NOPGLTMCKG DSVVWYLFSA GNEADVHGIY FSGNTYLWRG ERDRTANLFF
QTSLLHMMWP DTEGTFNV743EC LTTDHYTGGM KQKYTVNOCR RQSEDSTFYL GERTYIIAAV
EVEWDYSPQR EWEKELHLQ EQVSNAPFLD RGEFYIGSKY KKVYRQYTD STFRVPVERK
AEEHLGILG POLHADVGDK VKIIFKNMAT RPYSIHAHVQ QTESSTVTPT LPGETLTYVW
KIPERSGAGT EDSACI907PWAY YSTVDQVKDL YSGLIGLIV CRRPYLKVFN PRRKLEFALL
FLVFDENBSM YLDDN907IKTYS DHPKVNKDD EEFIESNKMH AINGRMFGNL QGLTMHVGD907E
VNWYLMGMGN EIDLHTVHFH GHSFOYKHRG VYSSDVFDIF PGTYQTLEMF PRTPGIWLLH
CHVDHIHAG METTYTVLQNG EDTKSG

```

Fig. 1. Primary amino acid sequence of human CP (P00450). The tryptic peptides, including potential N-glycosylation sites, are shown in bold type. Tryptic peptides identified in the LC-ESI-MS/MS analysis are underlined. Cysteine residues are carboxymethylated. Identified N-glycosylation sites are indicated by arrow.

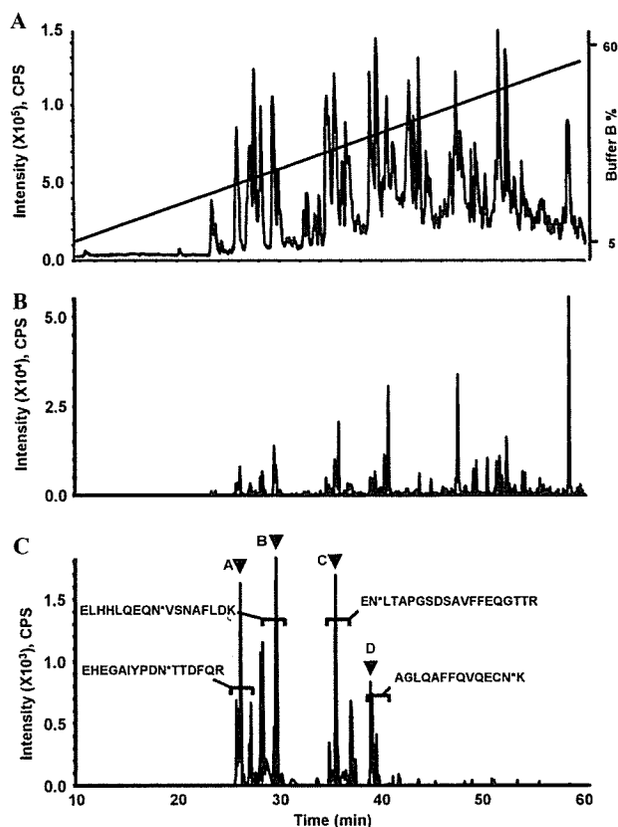


Fig. 2. LC-ESI-MS/MS in the m/z range of 1000–2000 of the tryptic digest of human CP. (A) TIC of the TOF-MS scan for the full-scan m/z 1000–2000 and the HPLC gradient. (B) TIC of the product ion scan acquired data-dependently. (C) Extracted ion chromatograph at m/z 204.05–204.15 of the product ion spectra. Brackets denote glycopeptide fraction and peptide sequences of the glycopeptides. Product ion spectra indicated by A–D are shown in Fig. 3.

the calculated molecular weight of the glycopeptide (4096.7) and adding the molecular weight of H₂O (18.0). The presence of product ions at m/z 274 (NeuAc-H₂O) and m/z 292 (NeuAc) suggested sialylation of the oligosaccharide. Thus, the carbohydrate's composition, [HexNAc]₄[Hex]₅[NeuAc]₂, was deduced.

Identification of Asn743 glycopeptide

The product ion spectrum of 1628.4 (+3) at 29 min, labeled by B in Fig. 2C, is shown in Fig. 3B. This precursor ion was assigned as a glycopeptide due to the presence of abundant oligosaccharide oxonium ions such as m/z 204 (HexNAc), m/z 366 (HexHexNAc), and m/z 292 (NeuAc) in the product ion spectrum. Several fragment ions were consistent with theoretical b- and y-series fragment ions derived from the peptide ELHHLQEQN⁷⁴³VSNAFLDK (residues 735–751). Doubly charged ions of peptide (m/z 1011.7), peptide + HexNAc (m/z 1113.1), peptide + 2HexNAc (m/z 1214.6), peptide + 2HexNAc + Hex (m/z 1295.5), peptide + 2HexNAc + 2Hex (m/z 1376.7), and peptide + 2HexNAc + 3Hex (m/z 1457.5) showed the sequential fragmentation of the pentasaccharide carbohydrate core. The

carbohydrate's molecular weight, 2879.1, was calculated from the theoretical molecular weight of the peptide (2021.0) and the calculated molecular weight of the glycopeptide (4882.1). The carbohydrate's composition, [HexNAc]₅[Hex]₆[NeuAc]₃, was deduced from the molecular weight.

Identification of Asn378 glycopeptide

The product ion spectrum of 1444.6 (+3) at 35 min, labeled by C in Fig. 2C, is shown in Fig. 3C. Abundant oligosaccharide oxonium ions were detected, as were several fragment ions consistent with b- and y-series fragment ions derived from the peptide EN³⁷⁸LTAPGSDSAVFFEQGTTR (residues 377–391). The carbohydrate's molecular weight, 2222.9, was calculated from the theoretical molecular weight of the peptide (2126.0) and the calculated molecular weight of the glycopeptide (4330.9). Thus, the peptide moiety ENLTAPGSDSAVFFEQGTTR and the carbohydrate's composition, [HexNAc]₄[Hex]₅[NeuAc]₂, were suggested.

Identification of Asn339 glycopeptide

The product ion spectrum of 1282.6 (+3) at 39 min, labeled by D in Fig. 2C, is shown in Fig. 3C. The spectrum contains abundant oligosaccharide oxonium ions, and several fragment ions consistent with b- and y-series fragment ions derived from the peptide AGLQAFFQVQE³³⁹CN³³⁹K (residues 327–340) were detected. The product ion spectrum contains the ions of the peptide (m/z 1640.8) and peptide + HexNAc (m/z 1843.9) and several y-series fragment ions of the peptide with a GlcNAc residue. The carbohydrate's molecular weight, 2223.0, was calculated from the theoretical molecular weight of the peptide (1639.7) and the calculated molecular weight of the glycopeptide (3844.7). Thus, the peptide moiety AGLQAFFQVQE³³⁹CN³³⁹K and the carbohydrate's composition, [HexNAc]₄[Hex]₅[NeuAc]₂, were suggested.

Heterogeneity of oligosaccharides at each glycosylation site

Glycopeptides with the potential N-glycosylation sites Asn119, Asn339, Asn378, and Asn743 were detected, whereas no glycopeptides containing the other sites (Asn208, Asn569, and Asn907) could be detected in this LC-ESI-MS/MS analysis. These findings suggest that Asn119, Asn339, Asn378, and Asn743 of human CP are glycosylated and that Asn208, Asn569, and Asn907 are not. Once a glycopeptide was identified, the other glycopeptides with the same peptide could be easily assigned because they were eluted at a similar retention time in the order of the number of NeuAc and had similar product ion spectra and molecular weight difference of sugar units. The oligosaccharide heterogeneity at each four N-glycosylation sites was determined by mass spectrum. For a representative example, the mass spectrum of the glycopeptides containing

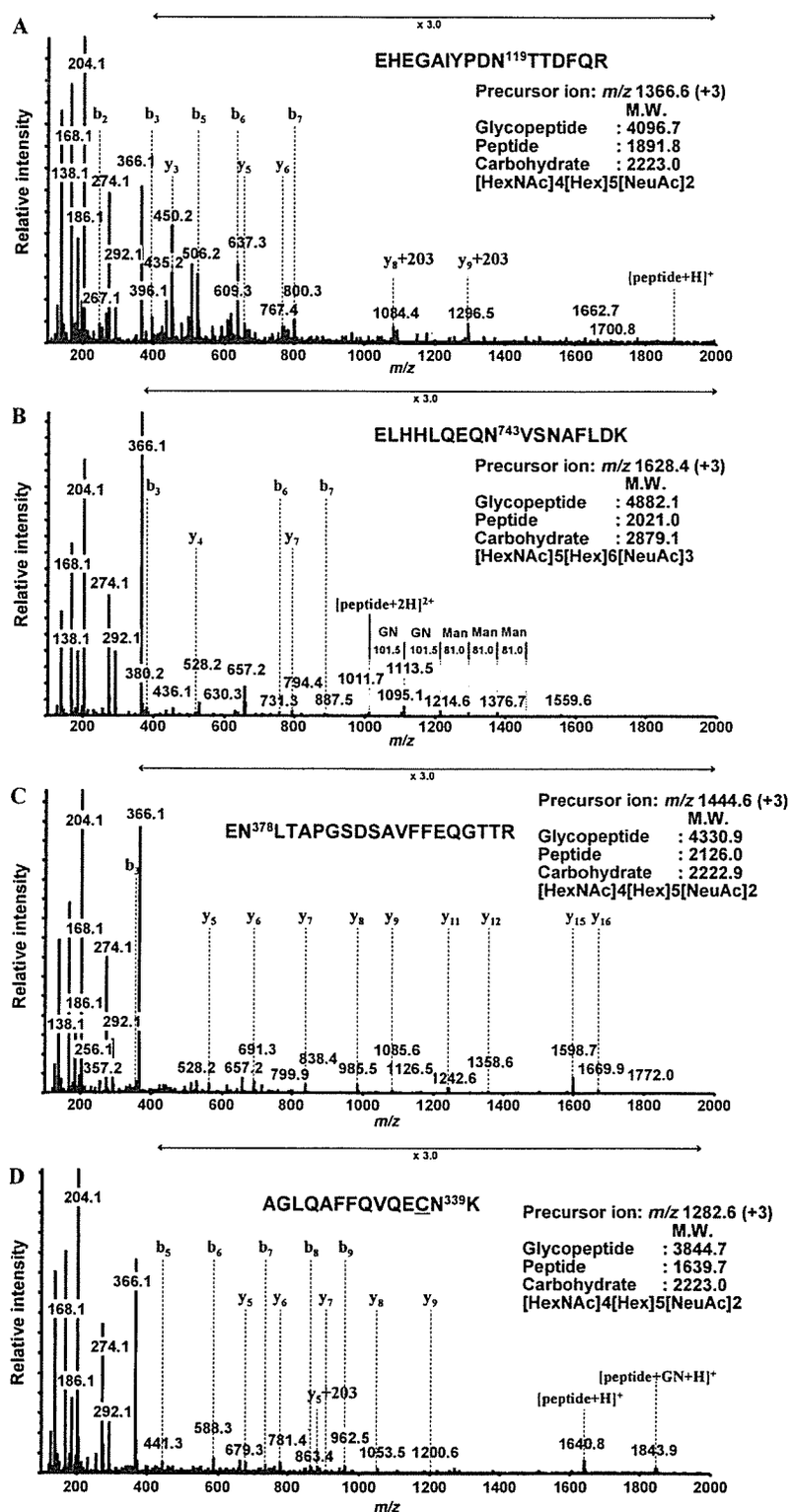


Fig. 3. Product ion spectra of *m/z* 1366.6 (+3) at 26 min (A), *m/z* 1628.4 (+3) at 29 min (B), *m/z* 1444.6 (+3) at 35 min (C), and *m/z* 1282.6 (+3) at 39 min (D) labeled by A, B, C, and D, respectively, in Fig. 2C. These spectra show abundant carbohydrate-derived ions at *m/z* 168 (HexNAc-2H₂O), *m/z* 186 (HexNAc-H₂O), *m/z* 204 (HexNAc), *m/z* 366 (HexHexNAc), *m/z* 274 (NeuAc-H₂O), and *m/z* 292 (NeuAc). The b- and y-series fragment ions [24] derived from the peptide moiety were observed. The molecular weights of the oligosaccharide were calculated from the molecular weights of the glycopeptide and peptide, and the deduced oligosaccharide composition is presented. Cystein residue was carboxymethylated.

Asn743 at 27.5 to 31.5 min is shown Fig. 4. The results of glycosylation analysis are summarized in Table 1. Deduced compositions of the oligosaccharides are estimated based on the calculated molecular weights of the oligosaccharides. Relative peak intensity was calculated by comparing triply charged glycopeptide ions. All glycosylation sites were occupied by at least three kinds of oligosaccharides, namely disialobiantennary structures ([HexNAc]₄[Hex]₅[NeuAc]₂), disialobiantennary structures with fucose ([HexNAc]₄[Hex]₅[NeuAc]₂[Fuc]₁), and trisialotriantennary structures ([HexNAc]₅[Hex]₆[NeuAc]₃). Trisialotriantennary structures with one fucose or two fucoses ([HexNAc]₅[Hex]₆[NeuAc]₃[Fuc]₁₋₂) were also detected at Asn119 and Asn743; furthermore, tetrasialotetraantennary structures with no fucose or one fucose ([HexNAc]₆[Hex]₇[NeuAc]₄[Fuc]₀₋₁) were detected at Asn743.

Linkage analysis of oligosaccharides by exoglycosidase digestion

To elucidate the oligosaccharide structure in terms of sequence and linkage, aliquots of the tryptic digest were further digested with exoglycosidases. As a representative example, Fig. 5 shows integrated mass spectra during the periods at which Asn119 glycopeptides were eluted in LC-ESI-MS analyses before and after digestion with exoglycosidase arrays. Treatment with α 2–3 neuraminidase removed one NeuAc residue from most of the triantennary structures ([HexNAc]₅[Hex]₆[NeuAc]₃[Fuc]₀₋₂) and a small amount of biantennary structures ([HexNAc]₄[Hex]₅[NeuAc]₂[Fuc]₀₋₁) (Fig. 5B). A minor amount of triantennary structures removed two NeuAc residues. Thus, it appears that most triantennary structures contain one α 2–3-linked NeuAc. Treat-

ment with α 2–3 neuraminidase + β 1–4 galactosidase removed all terminal galactose residues from the desialylated glycans without fucose residues but only partially digested terminal galactoses from the desialylated glycans with fucoses (Fig. 5C). The addition of α 1–3,4 fucosidase to α 2–3 neuraminidase + β 1–4 galactosidase treatment completely digested the remaining terminal galactose by releasing one fucose and one galactose (Fig. 5D). Thus, galactose residues are linked β 1–4 to GlcNAc, and undigestion of terminal galactose by β 1–4 galactosidase is due to attachment of fucose [25,26]. Because galactose was linked to GlcNAc in the β 1–4 position, the fucose removed with α 1–3,4 fucosidase may be linked α 1–3 to GlcNAc but not α 1–4 to GlcNAc. These data strongly suggested that sialyl Lewis X structure was present in human CP. Sialyl Lewis X structure was present predominantly in triantennary oligosaccharides, but a small amount seemed to be present in biantennary oligosaccharides as well. The remaining fucose residue may be linked α 1–6 to reducing end GlcNAc (core fucose).

Fig. 6 shows integrated mass spectra of Asn119, Asn743, Asn378, and Asn339 glycopeptides in LC-ESI-MS analysis following digestion with α 2–3,6,8,9 neuraminidase + β 1–4 galactosidase. Treatment with α 2–3,6,8,9 neuraminidase + β 1–4 galactosidase removed all NeuAc and then removed terminal galactoses in the outer arms without fucose. Thus, this treatment could differentiate glycoforms based on the location of fucose residues. Fucosylation occurred predominantly at reducing end GlcNAc in biantennary oligosaccharides and occurred at reducing end GlcNAc and/or outer arm GlcNAc in triantennary oligosaccharides. Mass spectra of Asn119 and Asn743 glycopeptides showed higher oligosaccharide heterogeneity, and a minor amount of tetraantennary glycans could be detected. The glycosylation profile

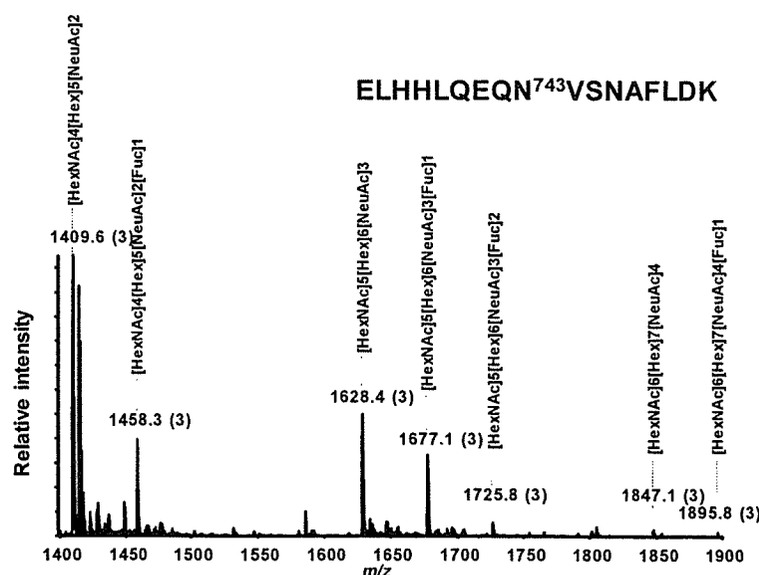


Fig. 4. Mass spectrum of the glycopeptides containing Asn743 eluting at 27.5–31.5 min from Fig. 2A. Deduced composition of the oligosaccharides is indicated based on the molecular weights of the oligosaccharides.

Table 1
Results of site-specific glycosylation analysis of human CP

Retention time (min)	Glycopeptides		Relative peak intensity ^a (%)	Peptide Sequence	Oligosaccharide		
	m/z	Charge			Calculated MW	Theoretical MW	Composition ^{b,c}
26	1415.3	+3	52	EHEGAIYPDN ¹¹⁹ TTDFQR	4242.8	1891.8	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂ [Fuc] ₁
26	1366.6	+3	100	EHEGAIYPDN ¹¹⁹ TTDFQR	4096.7	1891.8	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂
27	1682.7	+3	6	EHEGAIYPDN ¹¹⁹ TTDFQR	5045.1	1891.8	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃ [Fuc] ₂
27	1634.0	+3	21	EHEGAIYPDN ¹¹⁹ TTDFQR	4899.0	1891.8	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃ [Fuc] ₁
27	1225.8	+4			4899.0		
27	1585.3	+3	24	EHEGAIYPDN ¹¹⁹ TTDFQR	4753.0	1891.8	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃
27	1189.3	+4			4753.0		
28	1458.3	+3	35	ELHHLQEQN ⁷⁴³ VSN AFLDK	4372.0	2021.0	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂ [Fuc] ₁
28	1409.6	+3	100	ELHHLQEQN ⁷⁴³ VSN AFLDK	4225.9	2021.0	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂
28	1057.5	+4			4225.9		
29	1725.8	+3	5	ELHHLQEQN ⁷⁴³ VSN AFLDK	5174.5	2021.0	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃ [Fuc] ₂
29	1294.6	+4			5174.2		
29	1677.1	+3	29	ELHHLQEQN ⁷⁴³ VSN AFLDK	5028.2	2021.0	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃ [Fuc] ₁
29	1258.1	+4			5028.2		
29	1628.4	+3	43	ELHHLQEQN ⁷⁴³ VSN AFLDK	4882.1	2021.0	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃
29	1221.5	+4			4882.1		
31 ^d	1895.8	+3	2	ELHHLQEQN ⁷⁴³ VSN AFLDK	5684.4	2021.0	[HexNAc] ₆ [Hex] ₇ [NeuAc] ₄ [Fuc] ₁
31 ^d	1422.1	+4			5684.3		
31	1847.1	+3	3	ELHHLQEQN ⁷⁴³ VSN AFLDK	5538.4	2021.0	[HexNAc] ₆ [Hex] ₇ [NeuAc] ₄
31	1385.6	+4			5538.3		
35 ^d	1493.3	+3	6	EN ³⁷⁸ L TAPGSDSAVF FEQGTTR	4477.0	2126.0	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂ [Fuc] ₁
35	1444.6	+3	100	EN ³⁷⁸ L TAPGSDSAVF FEQGTTR	4330.9	2126.0	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂
37	1712.1	+3	8	EN ³⁷⁸ L TAPGSDSAVF FEQGTTR	5133.2	2126.0	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃ [Fuc] ₁
37	1284.3	+4			5133.2		
37	1663.4	+3	23	EN ³⁷⁸ L TAPGSDSAVF FEQGTTR	4987.1	2126.0	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃
37	1247.8	+4			4987.2		
39	1331.3	+3	14	AGLQAFFVQECN ³³⁹ K	3990.8	1639.7	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂ [Fuc] ₁
39	1923.4	+2			3844.7		
39	1282.6	+3	100	AGLQAFFVQECN ³³⁹ K	3844.7	1639.7	[HexNAc] ₄ [Hex] ₅ [NeuAc] ₂
41	1501.3	+3	6	AGLQAFFVQECN ³³⁹ K	4500.8	1639.7	[HexNAc] ₅ [Hex] ₆ [NeuAc] ₃

Note. All masses are monoisotopic. Cysteine residue was carboxymethylated.

^a Relative peak intensity was calculated by comparing same charge state glycopeptide ions. The intensity of the glycoform with maximum at each glycosylation site was taken as 100%.

^b The oligosaccharide composition was deduced from the molecular weight of the oligosaccharide.

^c The glycopeptide ions adducted by NH₄⁺ or Na⁺ were excluded.

^d Product ion spectra of these molecular ions were not acquired. However, these were considered glycopeptides because of a molecular weight difference of 146 (Fuc) and the same retention time as other glycopeptides.

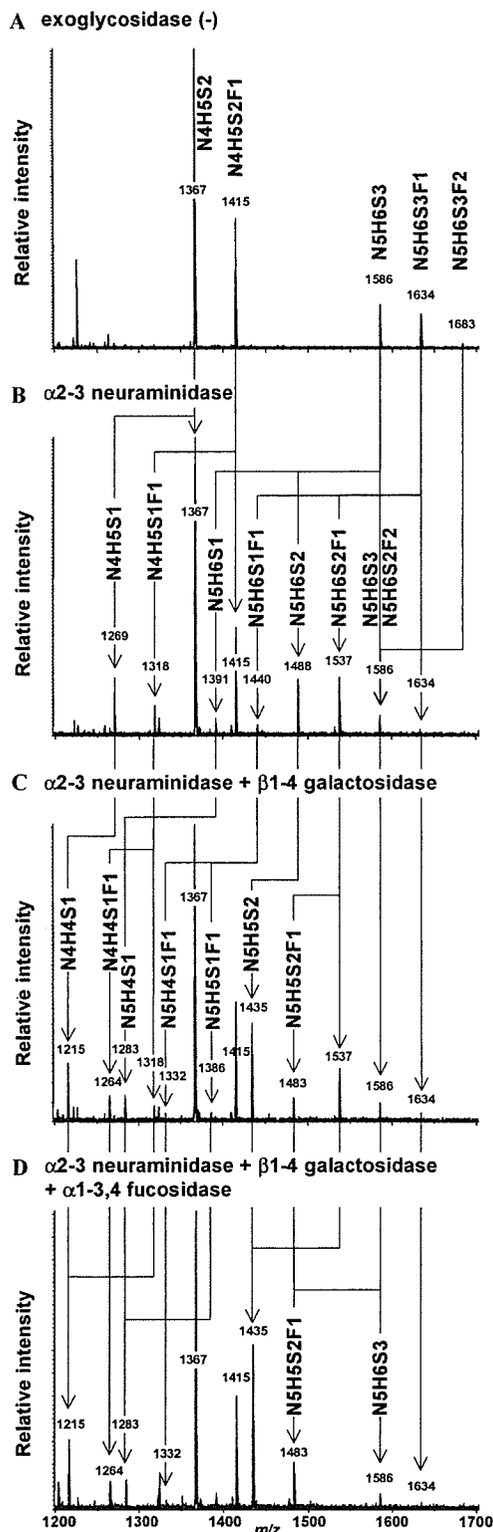


Fig. 5. LC-ESI mass spectra of the glycopeptides containing Asn119 digested with the following exoglycosidases: (A) exoglycosidase (-); (B) α 2-3 neuraminidase; (C) α 2-3 neuraminidase + β 1-4 galactosidase; (D) α 2-3 neuraminidase + β 1-4 galactosidase + α 1-3,4 fucosidase. Arrows between panels A and B, panels B and C, and panels C and D correspond to the digestion of NeuAc, Gal, and Gal+Fuc, respectively. H, hexose; N, N-acetylhexosamine; F, fucose; S, N-acetylneuraminic acid.

of Asn378 glycopeptides showed lower core fucosylation, and that of Asn339 glycopeptides showed lower branching. These glycosylation profiles provided the heterogeneity of fucose linkage and the number of arms at each glycosylation site in human CP.

Discussion

A site-specific glycosylation analysis of human CP was conducted using LC-ESI-MS/MS, where product ion spectra were acquired in a data-dependent manner. The collision energy for the product ion scan was adjusted from 30 to 80 eV depending on the size and charge of the precursor ion. Under these conditions, peptide precursor ions were degraded and produced b- and y-series fragment ions derived from the amino acid sequence. Glycopeptide precursor ions produced abundant carbohydrate ions (m/z 204, 186, 168, and 366) together with several low intensity b- and y-series fragment ions derived from the amino acid sequence [20,21]. Thus, product ion spectra of glycopeptides are readily distinguishable from those of peptides by such carbohydrate marker ions, and the peptide moiety in the glycopeptide could be deduced from the product ions that were consistent with the expected fragment ions derived from the peptide containing the N-glycosylation site. It is known that the glycopeptide ions are more labile than peptide ions and produce consecutive monosaccharide/polysaccharide losses at much lower collision energy, and this would provide information about branching and fucose location [18]. However, we used relatively high collision energy in this site-specific glycosylation analysis to identify the peptide ions in parallel with the detection and identification of the glycopeptide ions.

Protein coverage of more than 70% in human CP was obtained in the LC-ESI-MS/MS analysis with the m/z range of 400–2000 (for peptide mapping). The heterogeneity at four potential N-glycosylation sites was determined in the m/z range of 1000–2000 (glycosylation analysis). We could detect all of the potential glycosylation sites as either glycopeptides or nonglycosylated peptides. Peptides containing the potential N-glycosylation site Asn208, Asn569, or Asn907 were detected in nonglycosylated but not glycosylated forms. Peptides with the potential N-glycosylation site Asn119, Asn339, Asn378 or Asn743 were detected in glycosylated but not nonglycosylated forms. These findings indicate that Asn119, Asn339, Asn378, and Asn743 of human CP are glycosylated and that Asn208, Asn569, and Asn907 are not. Human CP was reported to have no O-linked glycosylation [8]. No information on O-glycosylation was obtained from this analysis. These results are consistent with a previous study determining the glycosylation sites of human CP [9].

Heterogeneity of oligosaccharides was determined at each of four glycosylation sites. Disialobiantennary structures with no fucose or one fucose ($[\text{HexNAc}]_4 [\text{Hex}]_5 [\text{NeuAc}]_2 [\text{Fuc}]_{0-1}$) and trisialotriantennary structures

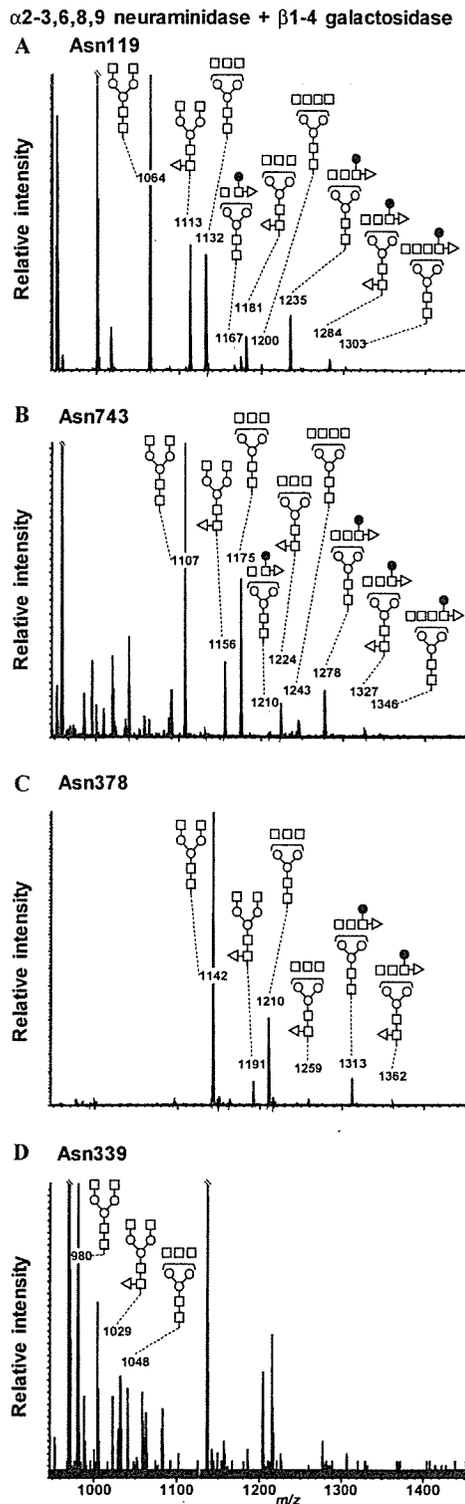


Fig. 6. LC-ESI mass spectra of the glycopeptides containing Asn119 (A), Asn743 (B), Asn378 (C), and Asn339 (D) after digestion with α 2-3,6,8,9 neuraminidase + β 1-4 galactosidase. Glycosylation profiles showed different degrees of branching and fucosylation at core GlcNAc and outer arm GlcNAc between glycosylation sites. Open circles, mannose; closed circles, galactose; open squares, *N*-acetyl glucosamine; open triangles, fucose.

([HexNAc]₅[Hex]₆[NeuAc]₃) were observed at all sites. These dominant oligosaccharides were consistent with structures published previously [7,8]. Furthermore, we detected trisialotriantennary structures with one fucose ([HexNAc]₅[Hex]₆[NeuAc]₃[Fuc]₁) at Asn119, Asn378, and Asn743, trisialotriantennary structures with two fucoses ([HexNAc]₅[Hex]₆[NeuAc]₃[Fuc]₂) at Asn119 and Asn743, and tetrasialotetraantennary structures with no fucose or one fucose ([HexNAc]₆[Hex]₇[NeuAc]₄[Fuc]₀₋₁) at Asn743.

To determine the linkage of fucose and NeuAc, exoglycosidase digestions were performed. Treatment with α 2-3 neuraminidase suggested that roughly one antenna of triantennary glycans was linked by NeuAc in the α 2-3 position. This is consistent with the previous findings that NeuAc is linked α 2-3 to the Gal β 1-4GlcNAc β 1-4Man α 1-3Man β 1-4GlcNAc β 1-4GlcNAc group in the triantennary glycan in human CP [7,8]. Results from α 2-3 neuraminidase + β 1-4 galactosidase treatments with or without α 1-3,4 fucosidase suggested that fucose residues were linked to reducing end GlcNAc and/or outer arm GlcNAc in the α 1-3 position in the antenna where NeuAc is linked to galactose in the α 2-3 position. These findings indicated that human CP contains a certain amount of sialyl Lewis X structure in triantennary glycans. Treatment with α 2-3,6,8,9 neuraminidase + β 1-4 galactosidase reveals the heterogeneity of the location of fucosylation as well as the number of arms. Although relative peak intensity does not express the relative amount of each glycan due to the different ionization efficiencies, the mass spectra showed the difference in fucosylation pattern and number of arms among sites.

No asialo oligosaccharides were detected in this analysis. It is known that desialylated CP is rapidly cleared from the circulation by the asialoglycoprotein receptor within the parenchymal cells of liver [27,28]. It is possible that desialylated CP might be cleared immediately by the liver.

Although the N-linked carbohydrate structures linked to human CP have been studied, only a few carbohydrate structures have been reported and site-specific characterization of these oligosaccharides has not been described. To determine the glycosylation state at each glycosylation site, the tryptic digest was examined by LC-ESI-MS/MS, where product ion spectra were acquired data-dependently. Glycopeptide ions were assigned based on the product ion spectra. Fucose and NeuAc linkages were determined by exoglycosidase digestions. Our data successfully provided comprehensive information on the site-specific N-linked oligosaccharides in human CP. This method is a powerful technique for elucidating the glycosylation of a biological sample.

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Site-specific glycosylation analysis of human apolipoprotein B100 using LC/ESI MS/MS

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Human apolipoprotein B100 (apoB100) has 19 potential *N*-glycosylation sites, and 16 asparagine residues were reported to be occupied by high-mannose type, hybrid type, and monoantennary and biantennary complex type oligosaccharides. In the present study, a site-specific glycosylation analysis of apoB100 was carried out using reversed-phase high-performance liquid chromatography coupled with electrospray ionization tandem mass spectrometry (LC/ESI MS/MS). ApoB100 was reduced, carboxymethylated, and then digested by trypsin or chymotrypsin. The complex mixture of peptides and glycopeptides was subjected to LC/ESI MS/MS, where product ion spectra of the molecular ions were acquired data-dependently. The glycopeptide ions were extracted and confirmed by the presence of carbohydrate-specific fragment ions, such as *m/z* 204 (HexNAc) and 366 (HexHexNAc), in the product ion spectra. The peptide moiety of glycopeptide was determined by the presence of the *b*- and *y*-series ions derived from its amino acid sequence in the product ion spectrum, and the oligosaccharide moiety was deduced from the calculated molecular mass of the oligosaccharide. The heterogeneity of carbohydrate structures at 17 glycosylation sites was determined using this methodology. Our data showed that Asn2212, not previously identified as a site of glycosylation, could be glycosylated. It was also revealed that Asn158, 1341, 1350, 3309, and 3331 were occupied by high-mannose type oligosaccharides, and Asn 956, 1496, 2212, 2752, 2955, 3074, 3197, 3438, 3868, 4210, and 4404 were predominantly occupied by mono- or disialylated oligosaccharides. Asn3384, the nearest *N*-glycosylation site to the LDL-receptor binding site (amino acids 3359–3369), was occupied by a variety of oligosaccharides, including high-mannose, hybrid, and complex types. These results are useful for understanding the structure of LDL particles and oligosaccharide function in LDL-receptor ligand binding.

Key words: apolipoprotein B100/glycopeptide/liquid chromatography electrospray mass spectrometry/product ion spectrum/*N*-linked oligosaccharide

Introduction

Low-density lipoprotein (LDL) is the main cholesterol carrier in human plasma, and a high serum level of LDL is involved in the development of atherosclerosis. LDL is originally secreted as very low-density lipoprotein (VLDL). VLDL is converted to LDL and then removed from the circulation. Apolipoprotein B100 (apoB100) is the only protein component of LDL and is the ligand recognized by the LDL receptor. The amino acid sequence of human apoB100 has been deduced by analysis of the apoB100 cDNA sequence (Chen *et al.*, 1986; Knott *et al.*, 1986; Law *et al.*, 1986; Yang *et al.*, 1986). Mature apoB100 consists of 4536 amino acids, and its molecular weight has been calculated to be 513 kDa. ApoB100 has 19 potential *N*-glycosylation sites (Asn-X-Ser/Thr), of which 16 asparagine residues are found to be glycosylated (Yang *et al.*, 1989). The carbohydrate moieties were linked to asparagine residues at the following 16 positions: 158, 956, 1341, 1350, 1496, 2752, 2955, 3074, 3197, 3309, 3331, 3384, 3438, 3868, 4210, and 4404. The carbohydrate structures of the *N*-linked sugar chains of human apoB100 were reported to be high-mannose, hybrid, and mono- and disialylated complex type oligosaccharides (Garner *et al.*, 2001; Taniguchi *et al.*, 1979).

The role of carbohydrate moieties of apoB100 has been investigated by several laboratories. The *N*-linked oligosaccharides at the amino terminus of human apoB100 are important for the assembly and secretion of VLDL (Vukmirica *et al.*, 2002). Seven of the *N*-glycans are predicted to occur close to the LDL-receptor binding region of apoB100 and seem to have an important role (Yang *et al.*, 1986, 1989). The carbohydrate composition of apoB100, particularly sialylation, has been considered to contribute to the atherogenic properties of LDL. However, Shireman and Fisher (1979) reported that they do not appear to play a significant role in the binding of apoB100 to the LDL receptor. Furthermore, the distribution and diversity of human apoB100 oligosaccharides isolated from normolipidemic, hypercholesterolemic, and hypertriglyceridemic diabetic subjects were highly conserved even when characterized in LDL subfractions (Garner *et al.*, 2001). The potential function of apoB100 carbohydrates posthepatic secretion is not well understood. Glycoproteins have a variety of sugar chains at each glycosylation site. Because of the individual functions at each site, a comparison of glycosylation among various sites is important. Therefore, to investigate the role of carbohydrate moieties of apoB100, we attempted to determine the carbohydrate heterogeneity site-specifically.

To determine the site-specific carbohydrate heterogeneity of glycoproteins, the glycoprotein must be digested into

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