

CERTIFICATE OF ANALYSIS

PRODUCT NAME:	AdvanceBio Human IgG N-Linked Glycan Library
PRODUCT CODE:	GKLB-005
LOT NUMBER:	DP21H5022A
PACK SIZE:	20 µg Glycan Library (qualitative chromatographic standard for N-glycan identification)
PACKED WITH:	0.5 mg Human IgG Glycoprotein (WS0162, lot W170145B), source for the Glycan Library
PURITY:	Suitable for mass spectrometry
FORM:	Glycan Library: Dry solid Human IgG Glycoprotein: Dry solid
STORAGE:	Store at -20°C in the dark before and after reconstitution
EXPIRATION:	December 2026, may be used for 1 year after reconstitution
STRUCTURE:	The Human IgG N-Linked Glycan Library consists of complex biantennary oligosaccharides consistent with N-glycans on normal human IgGs ^{1,2,3} (see Table 1). Human IgG Glycoprotein: Predominantly IgG1, IgG2, and IgG3 ⁴

Quality Control:

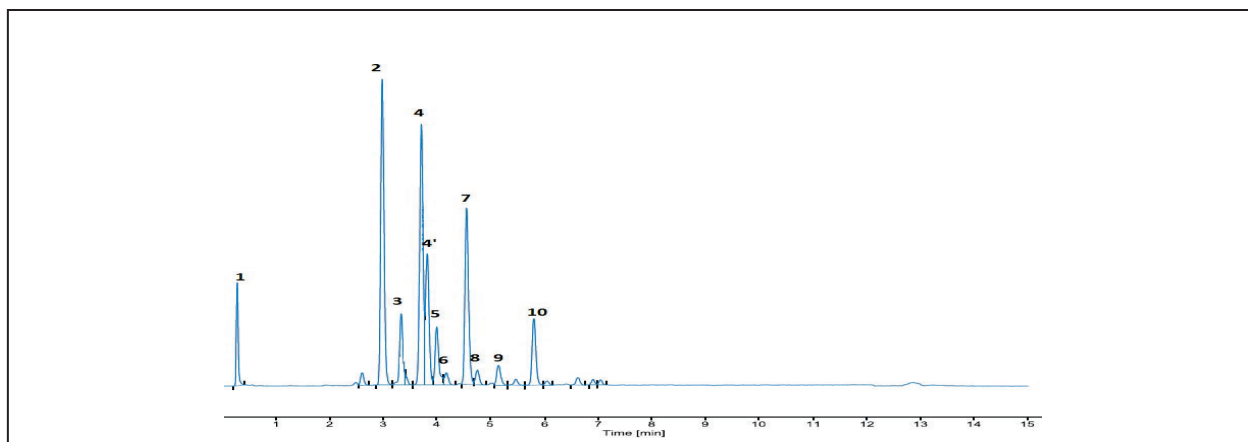


Figure 1 - UHPLC Results: 10 pmol (1 μ l, aqueous) of the 2-AB-labeled library was injected on a UHPLC System (Agilent 1290 Infinity II) utilizing a 15-minute method under the conditions below:

Time (min)	Flow	%ACN	%Buffer
00.0	1.0	75.0	25.0
12.0	1.0	52.5	47.5
12.1	0.5	40.0	60.0
12.5	0.5	40.0	60.0
12.6	0.5	75.0	25.0
12.7	1.0	75.0	25.0
15.0	1.0	75.0	25.0

Column: Waters ACQUITY UPLC BEH Glycan Column (1.7 μ m, 2.1 x 100 mm)

ACN: Acetonitrile

Buffer: 100 mM ammonium formate, pH 4.4

Flow rate: As stated in table, in ml/min

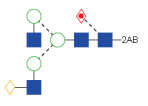
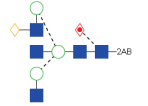
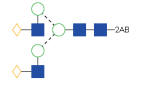
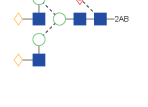
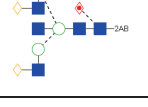
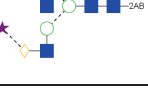
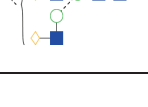
Temperature: 60° C

Max Pressure: 15,000 psi

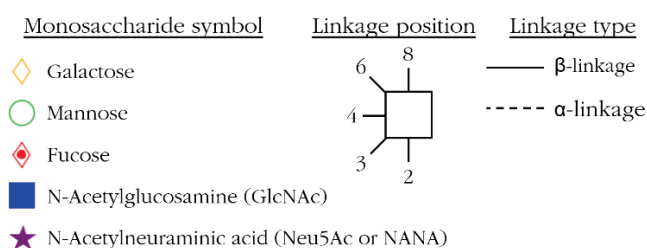
Fluorescence Detection: λ_{ex} = 330 nm, λ_{em} = 420 nm

Table 1 - Peak Identification of 2-AB Labeled Human IgG N-Linked Glycan Library.

Peak Number	Glycan Identification	ProZyme	Common	Oxford (New)	Structure ^{4,5}
1	Free Dye (2-AB)				
2	Asialo-, agalacto-biantennary with core fucose	NGA2F	G0F	F(6)A2	
3	Asialo-, agalacto-biantennary with core fucose and with bisecting N-Acetylglucosamine	NGA2FB	G0FB	F(6)A2B	
4	Asialo-, monogalactosylated biantennary with core fucose	NA2G1F	G1F[6]	F(6)A2[6]G(4)1	

Peak Number	Glycan Identification	ProZyme	Common	Oxford (New)	Structure ^{4,5}
4'	Asialo-, mono-galactosylated biantennary with core fucose	NA2G1F	G1F[3]	F(6)A2[3]G(4)1	
5	Asialo, mono-galactosylated biantennary, core substituted with fucose and bisecting N-Acetylglucosamine	NA2G1FB	G1FB	F(6)A2[6]BG(4)1	
6	Asialo, galactosylated biantennary	NA2	G2	A2G(4)2	
7	Asialo, galactosylated biantennary with core fucose	NA2F	G2F	F(6)A2G(4)2	
8	Asialo, galactosylated biantennary with core fucose and bisecting N-Acetylglucosamine	NA2FB	G2FB	F(6)A2BG(4)2	
9	Mono-α(2-6)-sialylated, mono-galactosylated biantennary with core fucose	NA2G1FS1	G1FS1	F(6)A2[3]G(4)1S(6) ₁	
10	Mono-α(2-6)-sialylated, galactosylated biantennary with core fucose	A1F	G2FS1	F(6)A2G(4)2S(6)1	

Structure Key^{5,6}:



Preparation: Human polyclonal IgG (containing >99% IgGs; IgG1 being at least 70% of the total IgG content, balance is IgG2 and IgG3)^{1,4} was digested with N-Glycanase[®] PLUS (ProZyme product code GKE-5010). Released N-linked oligosaccharides were

then labeled with 2-AB and purified from excess labeling reagents.

Structural Analysis: The purity and structural integrity of the glycan library was assessed by UHPLC^{7,8} (GU values) and MALDI-TOF mass spectrometry^{9,10} or LC-MS. Good

agreement was found between the results from mass spectrometry and UHPLC.

Application: Confirmation of identification of specific N-glycans by UHPLC analysis for samples prepared on the GlykoPrep® Sample Preparation Platform.

Handling & Reconstitution: The labeled oligo-saccharide is shipped as a dried solid. Use ultra-pure water or an aqueous buffer to dissolve the materials (see Directions for Use for suggested volumes). Allow the unopened vial to reach ambient temperature and tap on a solid surface to ensure that most of the material is at the bottom of the vial. Gently remove the cap, add the desired volume of ultra-pure water or aqueous buffer, re-cap and mix thoroughly to redissolve all the material.

For maximal recovery, ensure that the cap lining is also rinsed. Centrifuge the reconstituted vial briefly before use.

Make sure that any glassware, plasticware, solvents or reagents used are free of glycosidases and carbohydrate contaminants.

Minimize exposure to elevated temperatures or extremes of pH. Store the reconstituted glycan at -20° C. Allow the vial to equilibrate to ambient temperature before use.

Directions For Use: The amount of 2-AB-labeled library standard injected on a UHPLC column is typically 6-9 pmol of total glycan. For our Quality Control testing, one vial was dissolved in 30 µl of water and 1 µl injected on the ACQUITY column. For larger injection volumes or other LC systems we recommend further dilution as necessary for compatibility with your mobile phase. For suggested methods see Rapid UHPLC Methods for Screening Labeled N-Glycans at:

www.prozyme.com/protocols/

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