Join the sweet revolution in biopharmaceuticals

- Intact Glycoproteins
- Glycopeptides
- Free Glycans
- Monosaccharides
Biotherapeutic proteins, which include antibodies, are becoming the fastest growing category of drugs due to their efficacy for patient health. The complex nature of these large molecules brings major new challenges to the pharmaceutical scientist. The industry is undergoing a scientific revolution as it adapts to meet the challenges of characterizing biotherapeutics.

Glycosylation of biotherapeutic medicines plays a critical role in the efficacy of the drug. In-depth characterization of these glycans (complex sugars) bound to the drug entity is mandatory and challenging. To overcome this challenge, we have developed a series of four simple workflow solutions based on leading technology platforms.

**MEETING THE CHALLENGES of Glycan Characterization**

1. **WHAT IS MY GLYCOSYLATION PATTERN?**
   Intact Glycoprotein analysis

2. **WHERE DO MY GLYCANS RESIDE?**
   Glycopeptide profiling

3. **WHICH GLYCANS ARE IN MY DRUG AND WHAT IS THEIR CONCENTRATION?**
   Glycan structure determination

4. **WHAT IS THE COMPOSITION OF SUGARS IN MY DRUG ENTITY?**
   Monosaccharide analysis
**Intact Glycoprotein Analysis Workflow**

**WHAT IS MY Glycosylation Pattern?**

**Deconvolute Protein Spectra**

Screen, identify, and characterize intact proteins with higher productivity and confidence using Thermo Scientific™ Protein Deconvolution software. The software uses not one, but two deconvolution algorithms. One algorithm to take full advantage of the high-quality, high-resolution accurate-mass (HRAM) data produced by Thermo Scientific™ Orbitrap™ based mass spectrometers. The second algorithm is for lower resolution spectra where the mass spectral peaks are not isotopically resolved.

**Analyze Intact Proteins**

Many biopharmaceutical and biological research applications require analysis of proteins in their native (intact) state. The Thermo Scientific™ Exactive™ Plus EMR mass spectrometer combines unsurpassed high-resolution accurate-mass analysis with an extended mass range (EMR) to create an outstanding tool for investigating the structure, topology, and architecture of native-like tertiary and quaternary protein structures. A superb tool for high-performance screening of monoclonal antibodies, antibody-drug conjugates (ADC), PEGylated proteins, oligomerized protein-based drugs and glycoforms.

**Separate Antibody Variants & Fragments**

Thermo Scientific™ MAbPac™ columns are designed for the separation of monoclonal antibodies (mAbs) and related biologics by hydrophobic interaction (HIC), ion exchange (IEC), size exclusion (SEC) and reverse phase (RP) chromatography. This suite of columns have unique chemistries to provide high resolution, excellent bio-compatibility, and complementary selectivity suitable for a broad range of assays for mAbs and related substances.

**Analyze with Mass Spectrometry**

Intact glycoprotein profiling is used to ascertain the pattern and degree of glycosylation. Due to the heterogeneity of the attached glycan moieties, intact glycoprotein profiling is best performed using high resolution/accurate mass (HR/AM) mass spectrometry together with chromatographic separation to gain full insight into the various glycoforms present on a protein or antibody.

**Separate Glycoproteins**

Advanced ion exchange or hydrophobic interaction chromatography columns allow full separation of antibody fragments.

**Intact Glycoprotein**

Isolated antibodies and large proteins are often chemically reduced prior to analysis.

**Liquid Chromatography**

Reproducible, bio-compatible chromatography is required to simplify variants.

**Confirm Mass of Glycoprotein**

High resolution and mass accuracy are required together with deconvolution software to interpret spectra.

**Extended mass range (EMR)** of up to 20,000 and improved detection of high-mass ions.

**Screen, identify, and characterize intact proteins** with higher productivity and confidence using Thermo Scientific™ Protein Deconvolution software. The software uses not one, but two deconvolution algorithms. One algorithm to take full advantage of the high-quality, high-resolution accurate-mass (HRAM) data produced by Thermo Scientific™ Orbitrap™ based mass spectrometers. The second algorithm is for lower resolution spectra where the mass spectral peaks are not isotopically resolved.

**Analyze Intact Proteins**

Many biopharmaceutical and biological research applications require analysis of proteins in their native (intact) state. The Thermo Scientific™ Exactive™ Plus EMR mass spectrometer combines unsurpassed high-resolution accurate-mass analysis with an extended mass range (EMR) to create an outstanding tool for investigating the structure, topology, and architecture of native-like tertiary and quaternary protein structures. A superb tool for high-performance screening of monoclonal antibodies, antibody-drug conjugates (ADC), PEGylated proteins, oligomerized protein-based drugs and glycoforms.

**Deconvolute Protein Spectra**

Screen, identify, and characterize intact proteins with higher productivity and confidence using Thermo Scientific™ Protein Deconvolution software. The software uses not one, but two deconvolution algorithms. One algorithm to take full advantage of the high-quality, high-resolution accurate-mass (HRAM) data produced by Thermo Scientific™ Orbitrap™ based mass spectrometers. The second algorithm is for lower resolution spectra where the mass spectral peaks are not isotopically resolved.

**Analyze Intact Proteins**

Many biopharmaceutical and biological research applications require analysis of proteins in their native (intact) state. The Thermo Scientific™ Exactive™ Plus EMR mass spectrometer combines unsurpassed high-resolution accurate-mass analysis with an extended mass range (EMR) to create an outstanding tool for investigating the structure, topology, and architecture of native-like tertiary and quaternary protein structures. A superb tool for high-performance screening of monoclonal antibodies, antibody-drug conjugates (ADC), PEGylated proteins, oligomerized protein-based drugs and glycoforms.

**Deconvolute Protein Spectra**

Screen, identify, and characterize intact proteins with higher productivity and confidence using Thermo Scientific™ Protein Deconvolution software. The software uses not one, but two deconvolution algorithms. One algorithm to take full advantage of the high-quality, high-resolution accurate-mass (HRAM) data produced by Thermo Scientific™ Orbitrap™ based mass spectrometers. The second algorithm is for lower resolution spectra where the mass spectral peaks are not isotopically resolved.

**Analyze Intact Proteins**

Many biopharmaceutical and biological research applications require analysis of proteins in their native (intact) state. The Thermo Scientific™ Exactive™ Plus EMR mass spectrometer combines unsurpassed high-resolution accurate-mass analysis with an extended mass range (EMR) to create an outstanding tool for investigating the structure, topology, and architecture of native-like tertiary and quaternary protein structures. A superb tool for high-performance screening of monoclonal antibodies, antibody-drug conjugates (ADC), PEGylated proteins, oligomerized protein-based drugs and glycoforms.

**Deconvolute Protein Spectra**

Screen, identify, and characterize intact proteins with higher productivity and confidence using Thermo Scientific™ Protein Deconvolution software. The software uses not one, but two deconvolution algorithms. One algorithm to take full advantage of the high-quality, high-resolution accurate-mass (HRAM) data produced by Thermo Scientific™ Orbitrap™ based mass spectrometers. The second algorithm is for lower resolution spectra where the mass spectral peaks are not isotopically resolved.

**Analyze Intact Proteins**

Many biopharmaceutical and biological research applications require analysis of proteins in their native (intact) state. The Thermo Scientific™ Exactive™ Plus EMR mass spectrometer combines unsurpassed high-resolution accurate-mass analysis with an extended mass range (EMR) to create an outstanding tool for investigating the structure, topology, and architecture of native-like tertiary and quaternary protein structures. A superb tool for high-performance screening of monoclonal antibodies, antibody-drug conjugates (ADC), PEGylated proteins, oligomerized protein-based drugs and glycoforms.

**Deconvolute Protein Spectra**

Screen, identify, and characterize intact proteins with higher productivity and confidence using Thermo Scientific™ Protein Deconvolution software. The software uses not one, but two deconvolution algorithms. One algorithm to take full advantage of the high-quality, high-resolution accurate-mass (HRAM) data produced by Thermo Scientific™ Orbitrap™ based mass spectrometers. The second algorithm is for lower resolution spectra where the mass spectral peaks are not isotopically resolved.

**Analyze Intact Proteins**

Many biopharmaceutical and biological research applications require analysis of proteins in their native (intact) state. The Thermo Scientific™ Exactive™ Plus EMR mass spectrometer combines unsurpassed high-resolution accurate-mass analysis with an extended mass range (EMR) to create an outstanding tool for investigating the structure, topology, and architecture of native-like tertiary and quaternary protein structures. A superb tool for high-performance screening of monoclonal antibodies, antibody-drug conjugates (ADC), PEGylated proteins, oligomerized protein-based drugs and glycoforms.
**Glycopeptide Mapping Workflow**

1. **INTACT GLYCOPROTEIN** - Isolate antibody or glycoprotein.
2. **DIGEST TO GLYCOPROTEIDES** - Reproducible, fast digestion in less than one hour.
3. **SEPARATE GLYCOPROTEIDES** - Glycopeptides can be separated faster using advanced UHPLC columns.
4. **ANALYZE & MAP** - Routine peptide mapping provides protein sequence including glycopeptides. Advanced MS fragmentation provides exact glycopeptide location information.

**Glycan analysis can also be performed at the peptide level, with the goal of obtaining both glycan composition and peptide sequence at the site of glycosylation.**

**DIGEST**

Thermo Scientific™ SMART Digest™ kits are designed for biopharmaceutical applications that require highly reproducible, sensitive and fast analysis, often in high-throughput workflows. The kit uses unique optimized heat-stable immobilized trypsin technology.

SMART Digest allows complete antibody digestion in just 60 minutes; a saving of up to 23 hours versus in-solution digestion methods.

**SEPARATE PEPTIDES**

The glycopeptides can then be separated in a few minutes with the highest speed and resolution using the biocompatible Thermo Scientific™ Vanquish™ UHPLC system together with Thermo Scientific™ Accucore® C18 or Acclaim 120 RS UHPLC columns.

**Simple, Advanced Peptide Mapping Software**

Thermo Scientific™ PepFinder™ software provides accurate identification, in-depth characterization, and relative quantitation of biotherapeutic and other proteins from mass spectrometric data. It provides an automated workflow for glycopeptide identification, disulfide bond mapping, and quantification of PTMs. PepFinder software automates previously time-consuming manual processes, processing complex data and integrating the results into concise, informative reports.

**Determine the Exact Location of Glycosylation**

The Orbitrap Fusion Lumos Tribrid MS is the most sensitive Orbitrap instrument available and can perform a wide variety of analyses, from in-depth discovery experiments to characterization of complex PTMs to comprehensive qualitative workflows. The availability of multiple fragmentation techniques - CID, HCD, ETD, combined-ration spectra such as EThcD, and ultra product dependent acquisition such as HCDapETD, which generates a pair of HCD and ETD spectra. At any stage of MSn, with fragment ion detection in either the ion trap or Orbitrap mass analyzer, offers a new level of versatility and performance for the most challenging applications. Such as, the identification and location of glycosylated peptides.
ANALYZE LC
ANA
LYZE LC-M
S
INTACT GLYCOPROTEIN
Isolate antibody or glycoprotein.
RELEASE GLYCANS
Glycans are released enzymatically using glycosidase.
LABEL GLYCANS
Chemically react to add either a fluorophore or aminoxy tandem mass tag (TMT).
SEPARATE
Selective chemistries using combined HIC/WAX phases provide exceptional glycan separations.
ANALYZE & QUANTIFY
Separate and detect fluorescence labelled glycans.

GlycanPac AXH-1 & AXR-1 Columns
Thermo Scientific™ GlycanPac™ AXH-1 and GlycanPac AXR-1 columns are high-performance, silica-based HPLC columns designed for high-resolution glycan separations. They provide industry-leading resolution with unique selectivities for biologically important glycans, either labeled or native, using fluorescence and/or mass spectrometry (MS) detection. Both columns separate glycans by charge and size. Additionally, the GlycanPac AXH-1 columns are superior for separating glycan isomers. The GlycanPac AXH-1 columns are also used for fast, accurate, and simple quantification of glycans by charge.

Characterize Free Glycans
You can predict the structure of released labelled or unlabelled glycans using tandem mass spectrometry with SimGlycan® (PREMIER Biosoft). The software automatically matches experimental mass spectra against a comprehensive database and generates a scored list of candidate structures.

Revolutionize Your UHPLC Experience
The Vanquish UHPLC system takes biomolecule separations to a new level. The system offers better, faster, separations than previously possible. The system has exceptional control and stability giving you unrivalled reproducibility for meaningful answers.

- Maximized separation power, flow and gradient precision
- Controlled separations with reproducible Thermostating
- Analyze more samples with Vanquish Charger module
- Compliant Thermo Scientific™ Chromeleon™ CDS software

Monitoring of specific glycan species or determination of relative quantities of a particular set of glycans provides important information for the development of biotherapeutics. Due to the complexity of glycan structures, quantification and identification is performed upon release of glycans from the protein. N-linked glycans are released by enzymatic treatment, whereas O-linked glycans need to be released by chemical method as no enzyme exists for this purpose.

Glycans have no chromophore and have a poor response with conventional LC-UV detection. Glycans can be labelled with fluorescent tags prior to high sensitivity analysis by LC-Fluorescence detection. The most common label being 2-aminobenzamide (2-AB). Mass spectrometry has emerged as one of the most powerful tools for glycan structure elucidation. However, as most glycans do not ionize efficiently 2-AB labelling can also be performed to improve sensitivity.

Thermo Scientific™ aminoxyTMT™ (Tandem Mass Tag™) reagents can also be used for labeling proteins or released glycans. This enables multiplexing experiments to be performed for quantification of glycans by mass spectrometry.

Fluorescence Counts
Separation of labeled N-glycans from human IgG using a GlycanPac AXH-1 column.

Characterize Free Glycans
You can predict the structure of released labelled or unlabelled glycans using tandem mass spectrometry with SimGlycan® (PREMIER Biosoft). The software automatically matches experimental mass spectra against a comprehensive database and generates a scored list of candidate structures.

Revolutionize Your UHPLC Experience
The Vanquish UHPLC system takes biomolecule separations to a new level. The system offers better, faster, separations than previously possible. The system has exceptional control and stability giving you unrivalled reproducibility for meaningful answers.

- Maximized separation power, flow and gradient precision
- Controlled separations with reproducible Thermostating
- Analyze more samples with Vanquish Charger module
- Compliant Thermo Scientific™ Chromeleon™ CDS software

Monitoring of specific glycan species or determination of relative quantities of a particular set of glycans provides important information for the development of biotherapeutics. Due to the complexity of glycan structures, quantification and identification is performed upon release of glycans from the protein. N-linked glycans are released by enzymatic treatment, whereas O-linked glycans need to be released by chemical method as no enzyme exists for this purpose.

Glycans have no chromophore and have a poor response with conventional LC-UV detection. Glycans can be labelled with fluorescent tags prior to high sensitivity analysis by LC-Fluorescence detection. The most common label being 2-aminobenzamide (2-AB). Mass spectrometry has emerged as one of the most powerful tools for glycan structure elucidation. However, as most glycans do not ionize efficiently 2-AB labelling can also be performed to improve sensitivity.

Thermo Scientific™ aminoxyTMT™ (Tandem Mass Tag™) reagents can also be used for labeling proteins or released glycans. This enables multiplexing experiments to be performed for quantification of glycans by mass spectrometry.

Fluorescence Counts
Separation of labeled N-glycans from human IgG using a GlycanPac AXH-1 column.

Characterize Free Glycans
You can predict the structure of released labelled or unlabelled glycans using tandem mass spectrometry with SimGlycan® (PREMIER Biosoft). The software automatically matches experimental mass spectra against a comprehensive database and generates a scored list of candidate structures.

Revolutionize Your UHPLC Experience
The Vanquish UHPLC system takes biomolecule separations to a new level. The system offers better, faster, separations than previously possible. The system has exceptional control and stability giving you unrivalled reproducibility for meaningful answers.

- Maximized separation power, flow and gradient precision
- Controlled separations with reproducible Thermostating
- Analyze more samples with Vanquish Charger module
- Compliant Thermo Scientific™ Chromeleon™ CDS software

Monitoring of specific glycan species or determination of relative quantities of a particular set of glycans provides important information for the development of biotherapeutics. Due to the complexity of glycan structures, quantification and identification is performed upon release of glycans from the protein. N-linked glycans are released by enzymatic treatment, whereas O-linked glycans need to be released by chemical method as no enzyme exists for this purpose.

Glycans have no chromophore and have a poor response with conventional LC-UV detection. Glycans can be labelled with fluorescent tags prior to high sensitivity analysis by LC-Fluorescence detection. The most common label being 2-aminobenzamide (2-AB). Mass spectrometry has emerged as one of the most powerful tools for glycan structure elucidation. However, as most glycans do not ionize efficiently 2-AB labelling can also be performed to improve sensitivity.

Thermo Scientific™ aminoxyTMT™ (Tandem Mass Tag™) reagents can also be used for labeling proteins or released glycans. This enables multiplexing experiments to be performed for quantification of glycans by mass spectrometry.

Fluorescence Counts
Separation of labeled N-glycans from human IgG using a GlycanPac AXH-1 column.
High performance anion-exchange chromatography with pulsed amperometric detection (HPAE-PAD) is a well-established analysis method to characterize released Glycans. The separation is exceptionally selective for sugars and resolves glycans based on charge, size, composition, isomers and linkages. It is even possible to separate glycans based on sialic acid linkage, providing valuable information not only about the sugar sequence of a glycan, but also the subtle linkage differences that may indicate disease states.

**INTACT GLYCOPROTEIN**
Isolated antibody or glycoprotein.

**RELEASE GLYCANS**
Glycans are released enzymatically using glycosidase.

**ANALYZE USING ION CHROMATOGRAPHY**
Simple, sensitive glycan analysis without labelling.

**HPAE-PAD**
High performance anion-exchange chromatography with pulsed amperometric detection (HPAE-PAD) is a well-established analysis method to characterize released Glycans. The separation is exceptionally selective for sugars and resolves glycans based on charge, size, composition, isomers and linkages. It is even possible to separate glycans based on sialic acid linkage, providing valuable information not only about the sugar sequence of a glycan, but also the subtle linkage differences that may indicate disease states.

**Unique CarboPac Chemistry**
Resins of Thermo Scientific™ CarboPac™ columns consist of nonporous beads covered with a fine latex of functionized Thermo Scientific™ Dionex™ MicroBead™. This particular resin structure permits excellent mass transfer, resulting in high-resolution chromatography and rapid reequilibration.

**Label-free Glycan Workflow**
**INTACT GLYCOPROTEIN**
Isolated antibody or glycoprotein.

**RELEASE GLYCANS**
Glycans are released enzymatically using glycosidase.

**ANALYZE USING ION CHROMATOGRAPHY**
Simple, sensitive glycan analysis without labelling.

**IS THE COMPOSITION OF sugars in my drug entity?**

**WHAT TYPES OF COMPLEX GLYCANS ARE attached to my drug?**

**RELEASE MONOSACCHARIDES**
Glycans are released enzymatically using glycosidase. Glycans are then hydrolyzed to monosaccharides using acid.

**ANALYZE USING ION CHROMATOGRAPHY**
Ultra sensitive monosaccharide and sialic acid analysis.

**Monosaccharide composition, namely Fucose, Galactosamine, Glucosamine, Galactose, Glucose and Mannose, is routinely determined as their number and composition bound to the protein can impact efficacy of biotherapeutics. Monosaccharides are weak acids and can be separated by anion-exchange chromatography under basic conditions. Samples are acid hydrolyzed to release monosaccharides and analyzed by HPAE-PAD after chromatographic separation with CarboPac columns.**

**High Performance Ion Chromatography™ for HPAE-PAD**
The Thermo Scientific™ Dionex ICS-5000+ and ICS-4000 HPLC™ systems are ideal for HPAE PAD ion chromatography of mono-, di-, poly saccharides and sialic acids. With completely metal free flowpaths we eliminate the possibility of metal contamination and improve robustness for biomolecules. The analysis can be performed using Reagent-Free™ (RFIC™) systems with eluent generation that only require deionized water to electrolytically generate the eluent. HPAE-PAD only detects those compounds that contain functional groups that become oxidized at the detection voltage employed. Detection is sensitive and highly selective for sugars, because many potentially interfering species cannot be oxidized or reduced, and are not detected.