Case Study

A single data processing platform that enables novel MS-based methods for in-depth characterization of biopharmaceuticals

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Background

The Merck Serono structural characterization laboratory in Guidonia is part of the Characterization and Innovative Analytics Unit of the Analytical Development Biotech Department. Our role is to advise and support Merck worldwide in the development and production of novel biopharmaceuticals by providing a comprehensive structural overview of drug molecules and supporting process development and validation through advanced characterization approaches.

The ability to deliver information-rich data at a molecular level makes mass spectrometry (MS) the technology of choice for applications such as (glyco)peptide mapping, disulfide-bridge and intact-mass analysis, top-down sequencing, and released glycan analysis. Our MS facility uses instruments from several different manufacturers, which are used on a case-by-case basis to perform a wide range of MS experiments. This case study presents the development and optimization of MS-based methods for charge-variant analysis (CVA) and glycosylation profiling and subsequent data processing using Genedata Expressionist, and describes the advantages over assays that had previously been performed using other technologies and data-processing approaches (Table 1).

<table>
<thead>
<tr>
<th>Characterization method</th>
<th>Conventional method</th>
<th>Novel method with Genedata Expressionist data processing workflows</th>
<th>Time savings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charge-variant analysis (CVA)</td>
<td>Ion exchange (IEX) chromatography, imaged capillary isoelectric focusing (icIEF)</td>
<td>Strong-cation exchange (SCX)-ultra high-performance liquid chromatography-mass spectrometry (SCX-UHPLC-MS)</td>
<td>80%</td>
</tr>
<tr>
<td>Glycoform pairing analysis</td>
<td>Papain digestion, reduction, reverse-phase high-performance liquid chromatography-mass spectrometry (RP-HPLC-MS)</td>
<td>FabALACTICA® digestion, hydrophobic interaction liquid chromatography-mass spectrometry (HILIC-MS)</td>
<td>50%</td>
</tr>
<tr>
<td>Released glycan analysis</td>
<td>Reverse-phase high-performance liquid chromatography-mass spectrometry (RP-HPLC-MS)</td>
<td>Rapid™ PNGase deglycosylation RapidFluor™ labeling, hydrophobic interaction liquid chromatography-fluorescent detection-mass spectrometry (HILIC-FLR-MS)</td>
<td>80%</td>
</tr>
</tbody>
</table>

Table 1: Conventional and optimized MS-based biopharmaceutical characterization methods and resulting time savings.
Main Challenges

Developing novel and efficient MS-based characterization methods
Optimization of sample prep and separation procedures was the first challenge we addressed when developing novel MS-based analytical methods. However, to bring the overall efficiency of the end-to-end analytical process to the level required at Merck, we also required a platform that streamlined MS data processing, analysis, and management.

Harmonizing analysis and preventing bottlenecks
Acquiring data with instruments from different manufacturers means dealing with multiple software packages that create data silos and hinder comparison of results. In addition, the analysis of large volumes of complex data typically generated during MS experiments represents a serious bottleneck when using non-scalable desktop applications.

Increasing data quality and confidence in results
The large volumes of information-rich data generated during MS experiments require expert interpretation and dedicated application-specific algorithms to extract the maximum amount of information and deliver the most comprehensive characterization results possible. Where possible, error-prone manual evaluation (e.g., peak boundary determination) should be replaced by unbiased algorithms to improve result accuracy.

Breaking down data barriers and improving knowledge sharing
After data has been analyzed, the knowledge obtained should be freely accessible to all stakeholders to improve decision-making. Presenting complex information in a form that is quickly and readily understood is vital to the dissemination of knowledge within and across labs.

The Solution

Optimized sample preparation and flexible, automated data processing
We developed novel sample preparation methods that used alternative chromatographic matrices, MS-compatible buffers, and reagents that generate analytes that are amenable to efficient MS analysis. During method development, the flexibility of Genedata Expressionist enabled us to comprehensively analyze results and iteratively optimize procedures. After method optimization was complete, the workflow-based system enabled us to automate data analysis, further streamlining the overall analytical process.

Vendor-neutral, highly scalable data analytical system
The ability of Genedata Expressionist to load data from any MS instrument enables us to process data from all our instruments using a single platform. As a centralized server-based enterprise solution, Genedata Expressionist scales to the needs of large organizations and allows us to quickly and efficiently analyze the large amounts of MS data that we generate.

Streamlined MS analysis using custom libraries
In CVA, we use the flexibility of Genedata Expressionist to perform two approaches within a single data processing workflow. In a targeted approach, custom molecular libraries containing entries corresponding to commonly encountered species are used to provide an overview of highly abundant, expected variants across the entire elution profile (Figure 1). In an untargeted approach, more extensive libraries are used to fully characterize time-resolved elution ranges. Together with intuitive and insightful visualizations, the ability to search sample-specific custom libraries and perform advanced iterative review streamlines the analysis of released glycan and glycopeptide mapping results.

Enterprise software with advanced reporting
We can fully integrate Genedata Expressionist into our existing data systems to create a knowledge database that allows existing product knowledge (for example, the nature of variants found in specific clusters) to be leveraged in subsequent analyses. Structured, highly configurable reports and visualizations provide at-a-glance interpretation that speeds up analysis and aids decision-making. For example, an XIC offset view of Fab subunit glycoforms demonstrates that they elute in order of increasing complexity, which aids their identification (Figure 2).

Figure 1: Upper panel: Elution profile of a UHPLC SCX separation of Cetuximab charge variants; Lower panel: Identification of principal species found in the targeted approach.
Higher-quality results

Using Genedata Expressionist not only provided significant time-savings but also increased the quality of results in all our analyses. Targeted MS analysis using custom libraries and consolidated scores provided high-confidence results and significantly reduced the number of false-positive identifications in CVA and glycosylation profiling following HILIC-MS analysis. In released glycan analysis—after their structure was assigned and confirmed by MS/MS analysis—glycans were automatically quantified based on their fluorescence detector signal, increasing the overall specificity of the method.

Efficient dissemination and leverage of knowledge

Reports generated using Genedata Expressionist can be configured by adding or removing individual elements to provide as much or as little detail as required. An overview report can be generated for quick comparison of samples within a batch, or a complete report detailing all results and data processing parameters can be created for analyses in regulated environments.

Curated results can be quickly leveraged to optimize subsequent analyses. For example, the custom-built library that reduced false-positives in middle-down glycosylation profiling was derived from released glycans identified in previous experiments. In a similar fashion, a subset of modifications found in a charge variant cluster were saved as a custom library to reduce combinatorial complexity and data processing time when analyzing corresponding clusters in subsequent experiments.

Figure 2: Upper panel: HILIC-MS total ion chromatogram (TIC) profile of WS 2014/01 digested with FabALACTICA; Lower panel: Extracted ion chromatogram (XIC) of Fab subunit glycoforms of increasing complexity.

Figure 3: Annotated FLR trace; Inset: Typical report table listing identified released glycans with consolidated MS/MS score.
Summary

Genedata Expressionist enabled us to rapidly develop novel methods and automated data workflows that were specifically customized and optimized for our novel MS-based biopharmaceutical characterization approaches. Utilizing a scalable platform allows us to rapidly process the large volumes of complex data that our labs generate—thereby increasing our overall productivity—and automated reporting facilitates collaboration and knowledge sharing across our organization.

Outlook

We are in the process of establishing a Good Manufacturing Practice (GMP) environment at our site. The compliance capabilities of Genedata Expressionist—which include detailed client and server logs, result review and approval, stringently defined project-specific user/role profiles, and the ability to “lock down” optimized Genedata Expressionist workflows—will facilitate harmonized and reproducible MS data processing.

"Automated data processing and report generation provide remarkable time-savings in all our MS-based biopharmaceutical characterization applications." Mauro Sassi

"Genedata Expressionist provides high-quality qualitative and quantitative results and its flexible workflow-based approach makes it the ideal platform for processing and analyzing MS data generated using our innovative methods." Lucio Manzi

About Genedata Expressionist

As a comprehensive software solution for transforming raw data from any mass spectrometry instrument into insights, Genedata Expressionist offers built-in functionalities that serve all MS applications for biopharmaceutical characterization, proteomics, and metabolomics in a single enterprise platform. Custom-built workflows address specific data processing, analysis, and reporting needs and enable harmonization and standardization between and across organizations. Complete automation, unbiased data interrogation with best-in-class algorithms and cutting-edge statistics, and intuitive visualizations deliver high-quality results with significant time and cost savings. Integrated data and project management enables organizations to streamline methods and efficiently manage data, results, and reports.

Genedata Expressionist is part of the Genedata Biopharma Platform for capturing, organizing, integrating, processing, and analyzing data to increase enterprise-wide productivity and R&D process efficiency.