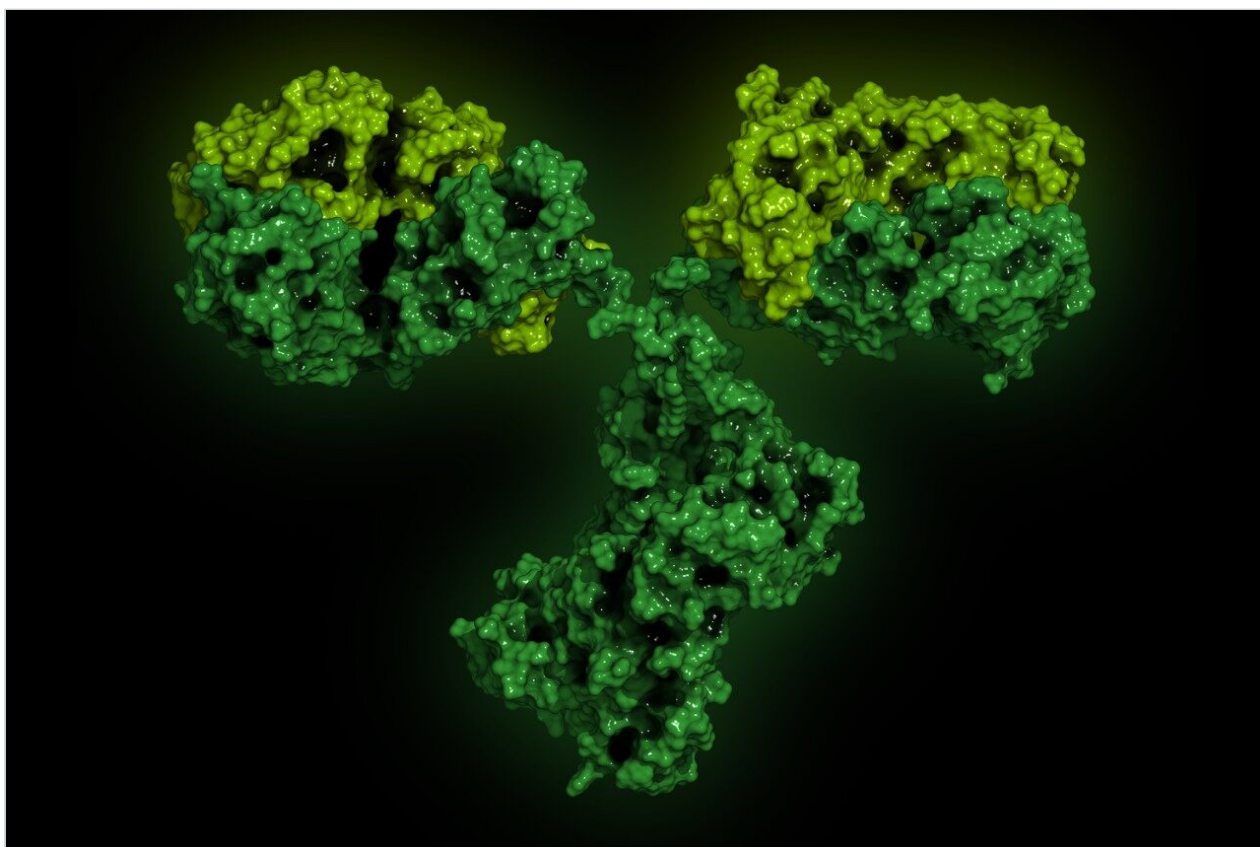


Application Note

Structural Comparability Assessment of Innovator and Biosimilar Rituximab Using the Biopharmaceutical System Solution with UNIFI

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Abstract

This application note demonstrates how an integrated biopharmaceutical LC-MS system utilizing the UNIFI Scientific Information System addresses these challenges by integrating and automating data acquisition, data processing, and result reporting into a seamless workflow for in-depth biotherapeutic structural characterization.

Benefits

For comparability studies performed with biosimilars, the integration of a fit-for-purpose UPLC/Tof-MS system with GxP-friendly data management, available with the UNIFI Scientific Information System, facilitates the development of a biotherapeutic product. This system solution enables complex biosimilar development to be carried out using routine analytical methodologies that are streamlined by efficient, workflow-based data management and reporting.

Introduction

Biopharmaceutical companies are challenged to design efficient analytical strategies for detailed assessment of structural comparability between biosimilar and innovator products. Extensive characterization increases confidence that a biosimilar product is safe and will meet regulatory compliance requirements for abbreviated approval pathways. Here, we demonstrate how an integrated biopharmaceutical LC-MS system utilizing the UNIFI Scientific Information System addresses these challenges by integrating and automating data acquisition, data processing, and result reporting into a seamless workflow for in-depth biotherapeutic structural characterization.

Comparability studies between an innovator, rituximab monoclonal antibody (mAb), and two biosimilar candidates were performed at the levels of intact protein, subunits (partially reduced antibody), and peptides using the Biopharmaceutical System Solution with UNIFI, shown in Figure 1. Differences in Critical Quality Attributes, such as primary structure (mutation), glycan fucosylation, and terminal amino acid heterogeneity were compared, quantified, and reported in a seamless workflow.

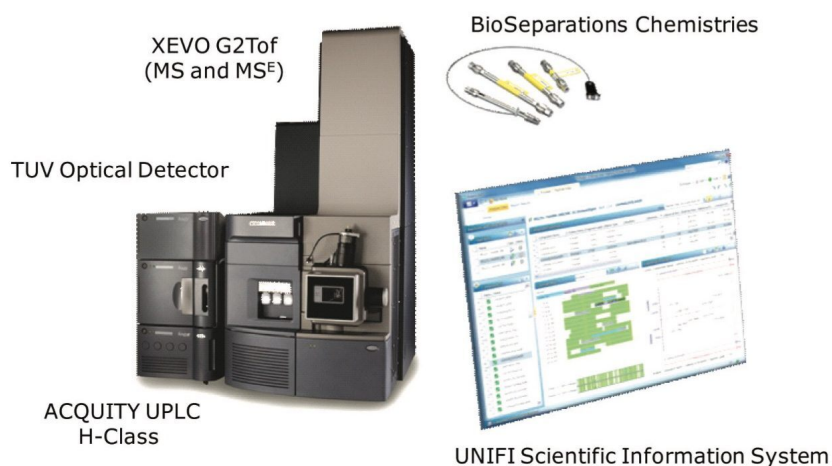


Figure 1. Biopharmaceutical System Solution with UNIFI.

Experimental

Sample description

Intact mass analysis: Innovator and both of the biosimilar mAb samples were diluted to 0.5 mg/mL using 25 mM ammonium bicarbonate, pH 7.9 for injection and analysis.

Reduced mAb analysis: The samples were diluted to 1 mg/mL in a reduction buffer (25 mM NaCl, 25 mM Tris, pH 7.5), and a concentrated DTT solution was added to the sample to obtain the final DTT concentration of 1.0 mM. The solution was then incubated at 37 °C for 20 min. The reduced samples were further diluted using a dilution buffer of 5% acetonitrile, 0.1% TFA to 0.2 mg/mL for LC-MS analysis.

Protein digestion: The samples were mixed with a denaturing buffer (8 M guanidine chloride, 1 M Tris, pH 7.5) to 1.0 mg/mL, reduced with 3 mM DTT, and alkylated with 7 mM iodoacetamide before buffer exchange over a NAP-5 column (GE Healthcare) to a digestion buffer of 100 mM Tris, pH 7.5. The samples were digested individually using either trypsin or chymotrypsin (S:E = 20:1) for 4 hrs. The digested samples were diluted with 3% acetonitrile, 0.1% TFA to 0.2 mg/mL for injection.

Method conditions

Biopharmaceutical System Solution with UNIFI:

ACQUITY UPLC H-Class with Peptide

Separation Technology (PST) and Protein
Separation Technology (PrST) UPLC
Chemistries Xevo G2 Tof, ACQUITY UPLC TUV
Optical Detector UNIFI Scientific Information
System

Intact protein LC-MS conditions

Column: ACQUITY UPLC BEH300 C₄, 2.1 x 50 mm

Column temp.: 80 °C

Mobile phase A: water

Mobile phase B: acetonitrile

Mobile phase C: 1% formic acid (aqueous)

Detection: UV 280 nm

Time (min)	Flow (mL/min)	%A	%B	%C	%D	Curve
Initial	0.3	85	5	10	0	Initial
2.00	0.3	85	5	10	0	6
2.10	0.2	85	5	10	0	6
5.00	0.2	10	80	10	0	6
6.00	0.3	10	80	10	0	6
6.50	0.3	85	5	10	0	6

Time (min)	Flow (mL/min)	%A	%B	%C	%D	Curve
10.00	0.3	85	5	10	0	6

MS conditions

Capillary:	2.5 kV
Sampling cone:	50 V
Extraction cone:	4 V
Source temp.:	150 °C
Desolvation temp.:	350 °C
Cone gas flow:	0 L/Hr
Desolvation gas flow:	800 L/Hr

Partially reduced protein LC-MS conditions

Column:	ACQUITY UPLC BEH300 C ₄ , 2.1 x 50 mm
Column temp.:	80 °C
Mobile phase A:	water
Mobile phase B:	acetonitrile
Mobile phase C:	1% formic acid (aqueous)
Detection:	UV 280 nm

Time (min)	Flow (mL/min)	%A	%B	%C	%D	Curve
Initial	0.3	85	5	10	0	Initial
2.00	0.3	85	5	10	0	6
2.10	0.2	85	5	10	0	6
3.00	0.2	65	25	10	0	6
13.00	0.2	60	30	10	0	6
13.10	0.3	10	80	10	0	6
15.00	0.3	10	80	10	0	6
15.50	0.3	85	5	10	0	6
25.00	0.3	85	5	10	0	6

MS conditions

Capillary:	3.0 kV
Sampling cone:	30 V
Extraction cone:	4 V
Source temp.:	120 °C
Desolvation temp.:	350 °C
Cone gas flow:	0 L/Hr

Desolvation gas flow: 700 L/Hr

Tryptic digest LC-MS conditions

Column: ACQUITY UPLC BEH300 C₁₈, 2.1 x 150 mm

Column temp.: 65 °C

Flow rate: 0.2 mL/min

Mobile phase A: water

Mobile phase B: acetonitrile

Mobile phase C: 1% formic acid (aqueous)

Detection: UV 214 nm

Time (min)	Flow (mL/min)	%A	%B	%C	%D	Curve
Initial	0.2	89	1	10	0	Initial
10.00	0.2	82	8	10	0	6
85.00	0.2	61	29	10	0	6
90.00	0.2	50	40	10	0	6
91.00	0.2	10	80	10	0	6
94.00	0.2	10	80	10	0	6
95.00	0.2	89	1	10	0	6

Time (min)	Flow (mL/min)	%A	%B	%C	%D	Curve
105.00	0.2	89	1	10	0	6

MS conditions

Capillary:	3 kV
Sampling cone:	30 V
Extraction cone:	4 V
Source temp.:	100 °C
Desolvation temp.:	250 °C
Cone gas flow:	0 L/Hr
Desolvation gas flow:	500 L/Hr

Chymotryptic digest LC-MS conditions

Column:	ACQUITY UPLC BEH300 C ₁₈ , 2.1 x 150 mm
Column temp.:	60 °C
Mobile phase A:	0.1% formic acid (aqueous)
Mobile phase B:	0.1% formic acid in acetonitrile
Detection:	UV 214 nm

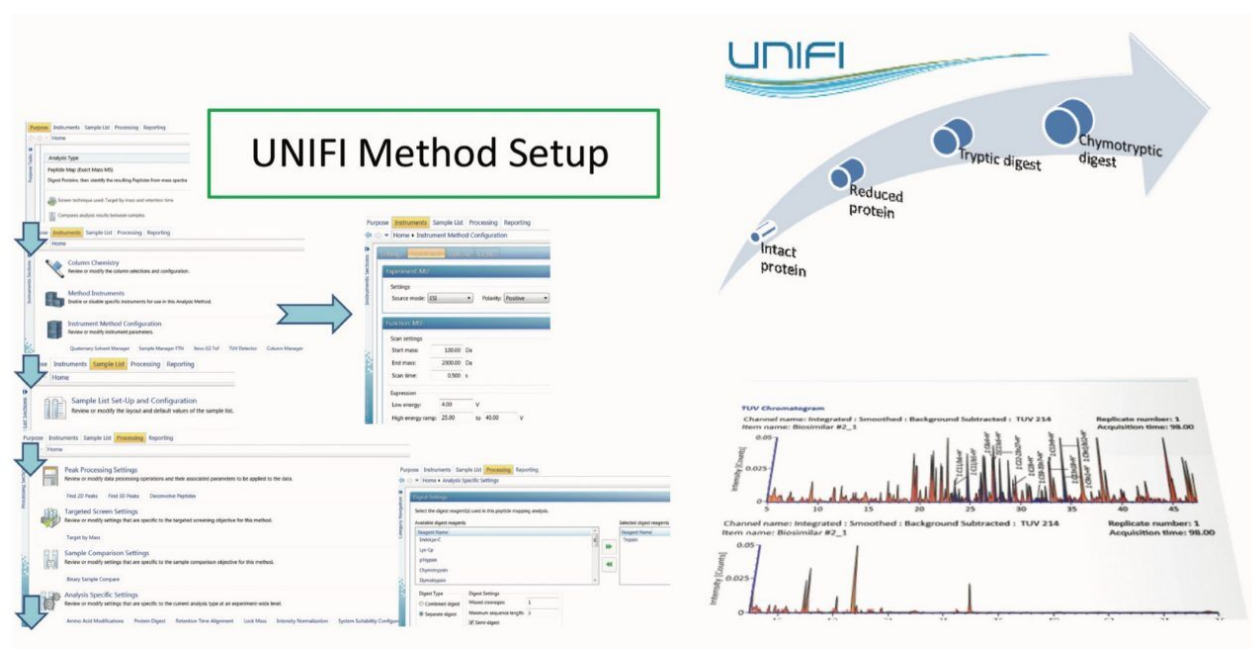
Time (min)	Flow (mL/min)	%A	%B	%C	%D	Curve
Initial	0.2	97	3	0	0	Initial
1.00	0.2	97	3	0	0	6
91.00	0.2	57	43	0	0	6
91.10	0.2	25	75	0	0	6
94.10	0.2	25	75	0	0	6
95.00	0.2	97	3	0	0	6
98.00	0.2	97	3	0	0	6

MS conditions

Capillary:	3 kV
Sampling cone:	30 V
Extraction cone:	4 V
Source temp.:	120 °C
Desolvation temp.:	350 °C
Cone gas flow:	0 L/Hr
Desolvation gas flow:	600 L/Hr

Results and Discussion

The therapeutic protein comparability workflow started with mAb sample analysis at the intact protein level, followed by the analysis of heavy and light chains after protein reduction, and finally addressed local post-translational modifications (PTMs) and mutations with LC-MS^E peptide map methodology. Such comprehensive workflow is managed by UNIFI for a regulatory environment by integrating data acquisition, data processing, and reporting in a highly automated fashion. The analysis method is completely defined prior to acquisition with the instrument settings, data processing parameters, and a reference to a reporting template included. Each analysis type focuses on a particular application need, such as intact protein analysis or peptide mapping experiment, facilitating the design of a method workflow, as shown in Figure 2. The report templates are composed of the objects that can be entirely configured by the user. The standard report templates include total ion chromatogram (TIC), mass spectra for all or selected ions in a form of either raw, deconvoluted, or centroid data format, and a tabulated summary of the interpreted LC-MS(MS^E) data.



Workflow 1: Intact protein MW determination and composition

For a quick assessment of the possible differences among the innovator, rituximab, and two biosimilar samples (Biosimilar 1 and Biosimilar 2), the intact protein mass analysis was performed. UNIFI has a built-in MaxEnt1 deconvolution capability for protein MW calculation and comparison. Figure 3 shows the distribution of glycoforms on the deconvoluted mAb spectra presented as the mirror plots. A systematic mass shift of 56 Da was observed in Biosimilar 2 glycoforms with respect to the innovator mAb; whereas, the Biosimilar 1 glycosylation profile displayed inconsistent mass difference (except G0F/G0F glycoform). The intact protein analysis data can also be viewed as raw, centroid spectra, or as a component summary, and can be used for the first-round evaluation of mAb sample heterogeneity.

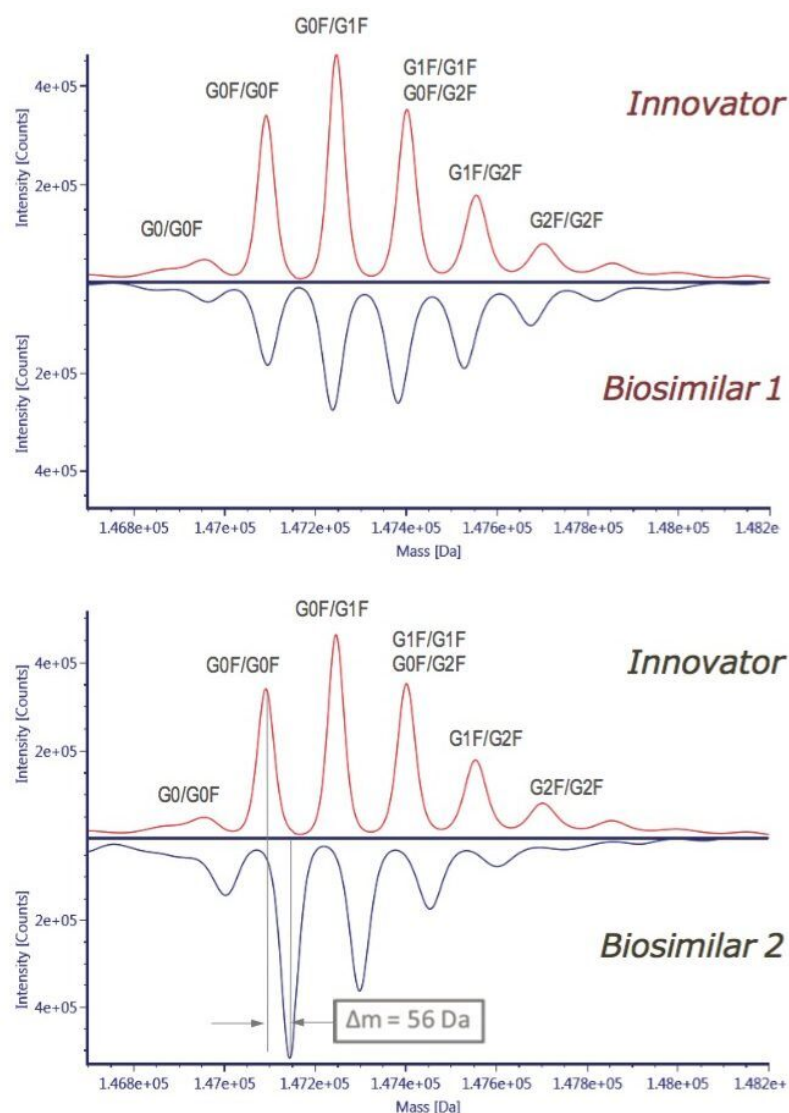


Figure 3. Intact protein mass analysis: MaxEnt1 deconvoluted mass spectra in compare mode. Biosimilar 2 glycoforms have a systematic mass shift of +56 Da with respect to the innovator mAb; whereas, Biosimilar 1 components do not display a systematic mass difference.

Workflow 2: Reduction of mAb

A closer look at the reduced form of rituximab allowed users to confine the structure heterogeneity to the individual heavy or light protein chains. Partially reduced mAb analysis measured and compared PTM and glycosylation profile among the innovator and both biosimilar mAbs, as seen in Figure 4. Consistent with 56 Da mass shift observed from intact protein data, our data suggest that 28 Da mass difference, possibly an amino acid sequence variation, belongs to the heavy chain of Biosimilar 2. N-terminal pyroglutamination Q

(PyrQ) levels were measured and reported for heavy and light chains. C-terminal Lys variants on the heavy chain as well as glycoform variants were automatically assigned in the UNIFI Review panel and plotted across all the samples of the innovator and biosimilars.

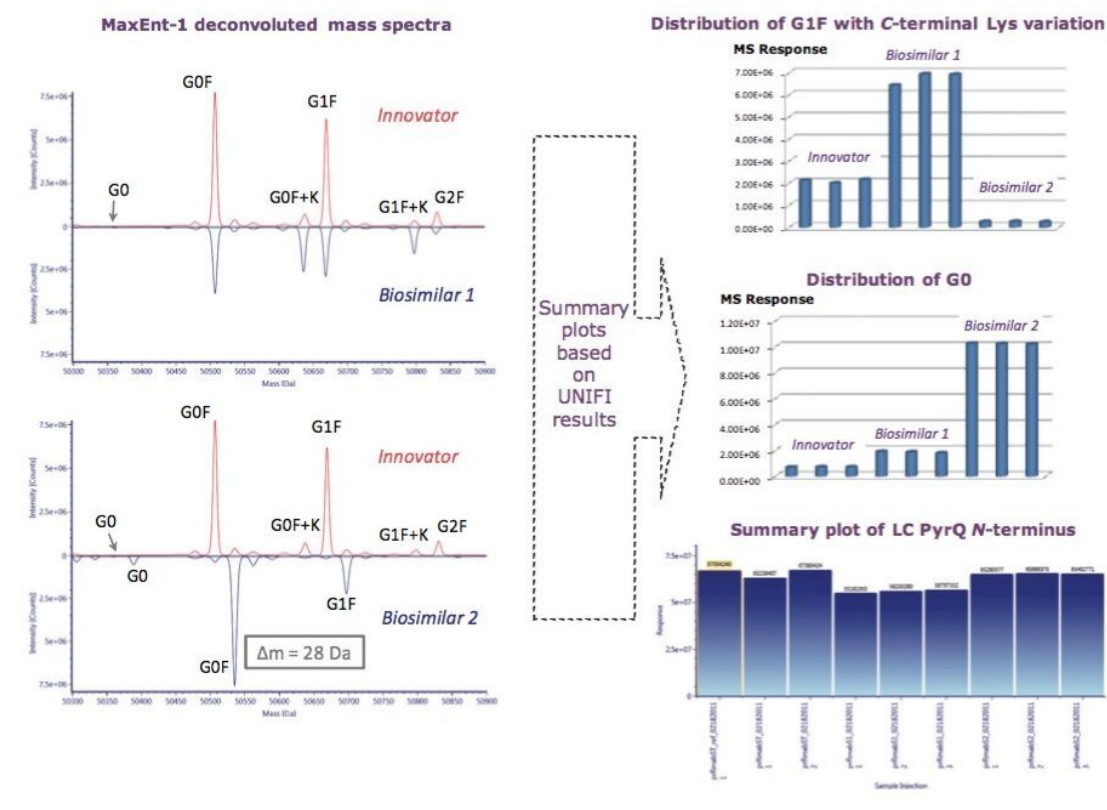


Figure 4. Partially reduced protein analysis. The following examples were measured and compared among three replicate injections of innovator and two biosimilar mAbs: C-terminal Lys variants on the HC, glycoform variant (G0) between biosimilars, amino acid sequence variant (+28 Da) on the heavy chain of Biosimilar 2, and N-terminal pyroglutamination Q (PyrQ) levels.

In comparison with the innovator, the obvious difference displayed in Biosimilar 1 in the deconvoluted HC spectrum is the higher degree of C-terminal Lys variation, which contributes to the inconsistent mass shift observed in the intact mass analysis. One of the remarkable differences between the two biosimilars was the relative abundance of G0 glycoform, which is known to correlate with antibody-dependent cellular cytotoxicity,¹ and is believed to affect drug safety and efficacy.

The summary plot tool allows users to select any observable data, such as response, mass error, retention time, etc., and trend it across all the injections, which is one of the UNIFI assets of the automatic and efficient data reviewing.

Workflow 3: Peptide mapping

To localize the difference among the three mAb samples, peptide mapping data were collected. A mirror plot of the tryptic digest demonstrated C-terminal Lys variant exists only in the Biosimilar 1 peptide map, shown in Figure 5, which was consistent with the glycosylation profiling data at the reduced protein level.

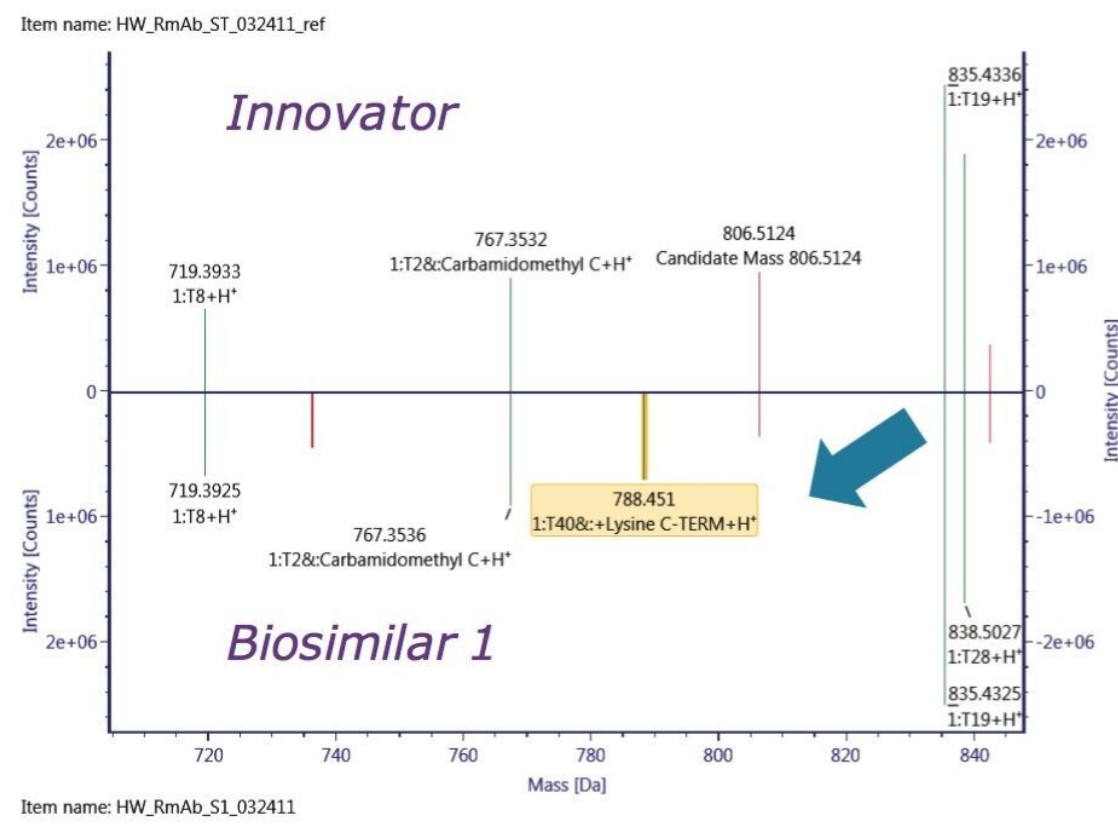


Figure 5. Tryptic digest comparison where components plot in compare mode. C-terminal Lys variant was observed only in the Biosimilar 1 peptide map.

The ultimate inquiry was localizing the amino acid mutation contributing to 28 Da mass shift of the Biosimilar 2. Based on published information,² an additional targeted sequence with Lys₂₁₈ → Arg₂₁₈ mutation was submitted to the method search. Compare mode view of the tryptic digest chromatogram, or component summary did not show a significant difference between the innovator and Biosimilar 2 mAb; therefore, no conclusion could be drawn about the primary sequence difference. The answer came with use of an alternative, non-specific enzyme, chymotrypsin. The chymotryptic map clearly showed the mass shift in a component view, as seen in Figure 6, and the peptide with a mutation site was automatically highlighted in the chromatogram, peptide map, and the component summary in the Review panel. Filtering the results in the Review by “showing unknown unique components” makes it easy to display the differences between the

innovator ("reference") and the biosimilar samples ("unknown").

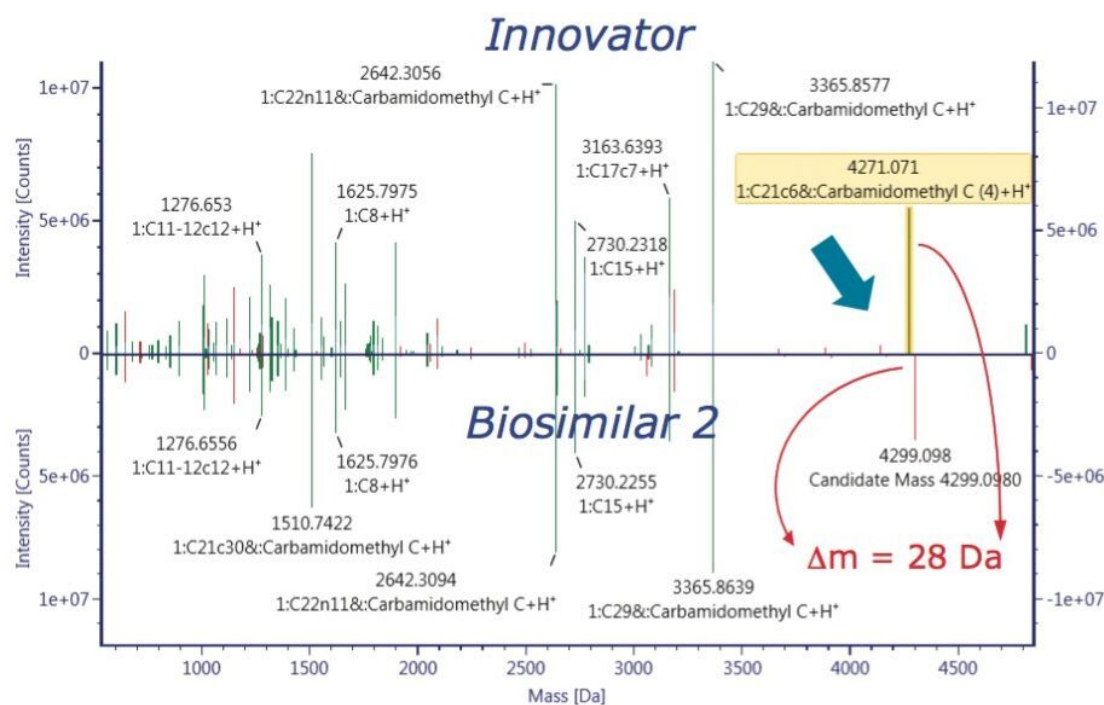


Figure 6. Chymotryptic peptide map analysis where components plot in compare mode, revealing 28 Da mass shift of the chymotryptic fragment in Biosimilar 2.

The reason that the tryptic map failed to pinpoint the amino acid substitution is that proteolytic cleavage occurs at Lys₂₁₇, Lys₂₁₈ or Arg₂₁₈. So, the very amino acid of question gets cleaved as a single amino acid entity. Chymotryptic digest, on the other hand, captures the mutation within a single peptide. Finally, Lys₂₁₈ → Arg₂₁₈ substitution was confirmed with MS^E data, as seen in Figure 7, which displayed a₁₆-ion fragment characteristic of Arg. UNIFI peptide map workflow proved the capability to confirm sequence mutation or other suspected PTMs.

Rituximab HC

Chymotryptic coverage of the Biosimilar 2 heavy chain

Identified: 98%

1: 1 to 60	QVQLQQPGAE	LVKPGASVKM	SCKASGYTFT	SYNMHWVKQT	PGRGLEWIGA	IYPNGDTSY
1: 61 to 120	NQKFKGKATL	TADKSSSTAY	MQLSSLTSED	SAVYYCA	YYGGDWYFNV	WGAGTTVTVS
1: 121 to 180	AASTKGPSVF	PLAPSSKSTS	GGTAALGCLV	KDYFPEPV	SWNSGALTSG	VHTFPAVLQS
1: 181 to 240	SGLYSLSSVV	TVPSSSLGTQ	TVICNVNHKP	SNTKVDKRAE	PKSCDKTHTC	PPCPAPELLG
1: 241 to 300	GPSVFLFPPK	PKDTLMISRT	PEVTCVVVDV	SHEDPEVKFN	WYVDGVEVHN	AKTKPREEQY
1: 301 to 360	NSTYRVVSVL	TVLHQDWLNG	KEYCKVSNK	ALPAPIEKTI	SKAKGQPREP	QVYTLPPSRD
1: 361 to 420	ELTKNQVSLTCLVKGFYPSD	IAVEWESNGQ	PENNYKTPPP	VLDSDGSFEFL	YSKLTVDKSR	
1: 421 to 451	WQQGNVFSCS	VMHEALHNHY	TQKSLSLSPG	K		

MS^E spectrum of the highlighted peptide with amino acid substitution

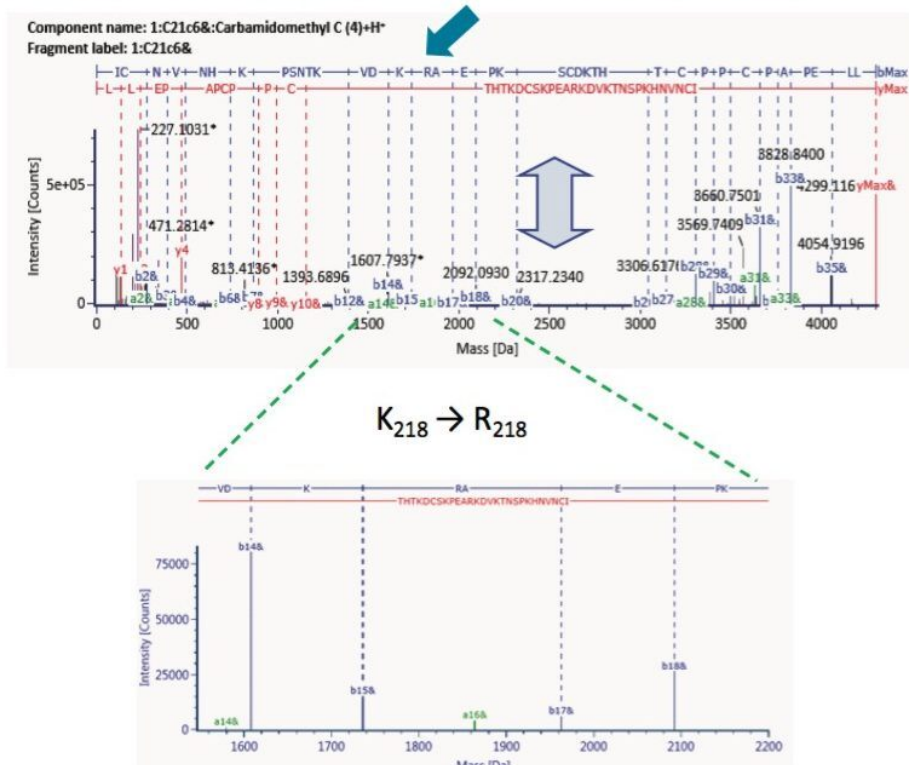


Figure 7. Chymotryptic digest analysis shows MS^E data confirm a single amino acid substitution (K → R) in Biosimilar 2.

Conclusion

UPLC/TOF-MS analysis at intact mAb, reduced mAb, and peptide map levels enabled the detection of primary structural differences, and quantitative assessments of these variations. An integrated biopharmaceutical LC-MS system utilizing the UNIFI Scientific Information System with automated data

acquisition, processing, and reporting for multiple analytical workflows enabled the efficient assessment of critical product attributes with minimal manual intervention.

The K → R mutation found in the Biosimilar 2 (of rituximab) study is not readily detectable under tryptic digest analysis. It demonstrates the need to routinely employ alternative digestion enzymes for product characterization. The integrated workflow of protein characterization at different levels, combined with intelligent methods and tools of UNIFI, will improve productivity and cut the cost of biosimilar drug development.

References

1. Schiestl M, Stangler T, Torella C, Cepeljnik T, Toll H, and Grau R. Acceptable changes in quality attributes of glycosylated biopharmaceuticals. *Nature Biotechnology*. 2011; 29 (4): 310-312.
2. Jefferis R, Lefranc MP. Human immunoglobulin allotypes. *mAbs*, 2009; 1 (4): 1-7.

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