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Multi-dimensional LC-MS: the next generation characterization of antibody-based therapeutics by unified online bottom-up, middle-up and intact approaches

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Accelerated development of new therapeutics in an increasingly competitive landscape requires the use of high throughput analytical platforms. In addition, the complexity of novel biotherapeutic formats (e.g. fusion proteins, protein-polymer conjugates, co-formulations, etc.) reinforces the need to improve the selectivity and resolution of conventional one-dimensional (1D) liquid chromatography (LC). Liquid chromatography-mass spectrometry (LC-MS)-based technologies such as native LC-MS for intact mass analysis or peptide mapping (also called bottom-up approach)-based multi-attribute methods (MAM) have already demonstrated their potential to complement the conventional analytical toolbox for monoclonal antibody (mAb) characterization. Two-dimensional liquid-chromatography (2D-LC-MS) methods have emerged in the last ten years as promising approaches to address the increasing analytical challenges faced with novel antibody formats. However, off-line sample preparation procedures are still required for conventional 1D and 2D-LC-MS methods for the in-depth variant characterization at the peptide level. Multi-dimensional LC-MS (mD-LC-MS) combine sample preparation and multi-level (i.e. intact, reduced, middle-up and peptide) analysis within the same chromatographic set-up. This review presents an overview of the benefits and limitations of mD-LC-MS approaches in comparison to conventional chromatographic methods (i.e. 1D-LC-UV methods at intact protein level and 1D-LC-MS methods at peptide level). The current analytical trends in antibody characterization by mD-LC-MS approaches, beyond the 2D-LC-MS workhorse, are also reviewed, and our vision on a more integrated multi-level mD-LC-MS characterization platform is shared

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1. Introduction

Therapeutic antibodies have been effectively used for treatment of a wide range of diseases, including oncology and auto-immune conditions.¹ The acceleration of biologics license applications (BLAs) approved in the last ten years is fueled by a robust late stage pipeline with a number of antibody candidates that has more than tripled between 2010 and 2019, with 79 antibody candidates evaluated just in 2019.²

Only 40% of antibody-based therapeutics approved in 2019 were monoclonal antibody (mAb) products, indicating the surge of alternative antibody formats. For example, a bispecific antibody was approved in 2017 for the treatment of hemophilia (Helimbra®/emicizumab) and one new nanobody modality was approved for the first time in 2019 (Cablivi®/caplacizumab)¹ for the treatment of thrombotic thrombocytopenic purpura (aTTP). The biopharmaceutical landscape is a highly competitive environment and the immunotherapy field in particular, where for example 2975 clinical trials involving only PD-1/PD-L1 mAbs were active in September 2019.³

The quest to drug the undruggable biological targets and bring innovative first-in class therapeutics faster to the patients is constantly challenging conventional analytical methods. These analytical methods are routinely used in quality control environments (QC) to support batch release⁴ and to assess potential critical quality attributes (pCQAs) at various stages of the drug lifecycle, during production and shelf-life. Antibody size and charge variants are usually quanti-

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fied by size exclusion chromatography (SEC) and ion exchange chromatography (IEC), respectively. Ideally, size variants are separated according to their difference in hydrodynamic radius in SEC,⁵ while the surface charge distribution of charge variants allows their separation by IEC.6,7 Capillary electrophoresis sodium dodecyl sulfate (CE-SDS)^{8,9} and imaged capillary isoelectric focusing (icIEF)^{10,11} techniques provide an orthogonal separation to SEC and IEC, respectively. Oxidations and glycoforms are commonly characterized using hydrophobic interaction chromatography (HIC)¹² and hydrophilic interaction liquid chromatography (HILIC), 13 respectively.

The rise of MS-based techniques for the characterization of antibody-based therapeutics has driven the development of MS-compatible mobile phases in order to couple native MS with non-denaturing SEC, IEC, and HIC modes.14 2D-LC-MS approaches have been developed to mimic the chromatographic separations achieved by QC methods with non-volatile salts in the first dimension and the use of a second dimension to desalt and remove non-volatile salts before MS analysis. The characterization of antibody variants at the bottom-up level is performed by peptide mapping, which involves multiple manual sample preparation steps, such as (i) chemical reduction of the antibody; (ii) alkylation of the reduced species, (iii) buffer exchange and (iv) an enzymatic digestion. The coupling with high resolution mass spectrometry (HRMS) allows the simultaneous sequence confirmation and determination of multiple post-translational modifications (PTMs) in a single analysis.15

However, conventional peptide mapping-based MAM can be time consuming and the multiple sample preparation steps can affect the reproducibility of the method. 16 In order to characterize the peaks separated at the intact level (by IEC or SEC for example) and identify corresponding PTMs, an additional off-line fractionation step is needed. As an example, the characterization of five charge variants separated by IEC using off-line procedures can require up to 52 hours. 17 In order to streamline this process, multi-dimensional LC-MS approaches have been developed and implemented to perform online fractionation followed by an in-line peptide mapping to

characterize antibody variants separated by IEC or SEC. 17-20 The automated characterization of antibody variants by mD-LC-MS can be performed with a much faster turnaround (typically 9 vs. 52 hours¹⁷) compared to conventional procedures (manual and/or off-line sample preparation and fraction collection), and has the potential to be used as a more integrated analytical platform for both upstream and downstream applications.

In the first part of this review, the identification of antibody variants at the intact level by native MS and 2D-LC-MS is described. Promising 2D-LC applications for the characterization of antibody higher order structures, antibodies present in cell culture supernatant or process analytical technology (PAT) are provided. In the second part, the characterization of PTMs at the peptide level using MAM peptide mapping, is discussed. In the last part, automated mD-LC-MS approaches allowing the online characterization of mAb variants at multiple levels, i.e. combination of intact and peptide levels, are reviewed.

Intact mass analysis

Native mass spectrometry (nMS)

Electrospray ionization (ESI)-MS analysis performed in native conditions (native MS) is of particular interest for the characterization of recombinant mAbs. 21,22 Native MS (nMS) technology minimizes sample preparation and preserves the protein higher-order structure and noncovalent interactions. 23,24 For instance, nMS is commonly used for the characterization of cysteine-linked ADCs, which consist of a broad population of noncovalent assemblies. 25-29 Some of the typical critical quality attributes (CQAs) such as (i) the average drug-to-antibody ratio (DAR); (ii) the distribution of drug-loaded species; and (iii) the amount of unconjugated mAbs can be determined by nMS.30

The use of volatile mobile phases allows the hyphenation of IEC, 31-34 SEC, 35-37 HIC, 38,39 RPLC, 40 and affinity chromatography (AC)⁴¹⁻⁴⁴ techniques to native MS. LC-nMS separates



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mAb variants prior to the entrance into the MS system, significantly improving sensitivity, dynamic range, and thus the detection of minor variants by limiting competition effects during the MS ionization process. For instance, the direct hyphenation of IEC to MS utilizing low ionic strength eluents on a strong cation exchange (SCX) stationary phase enabled MS data generation from minor mAb variants, and intact mass analysis of various PTMs, such as lysine truncation, glycosylation, and deamidation (mass shift of +1 Da relative to the main peak).33 To perform analysis under native conditions, the authors highlighted the importance of optimizing MS parameters since the ESI process is performed with an aqueous mobile phase, and recommended a low MS resolution to capture the protein signal with the highest sensitivity. However, adduct species related to the presence of salt required by the IEC chromatography are not resolved with a low resolution, which results in broader and asymmetrical spectral peaks and decreased mass accuracy following deconvolution, further limiting the applications to complex mixture samples.

Direct hyphenation of SEC to native MS has been facilitated by the development of SEC columns with improved chemical inertness¹⁴ and adopted for the characterization of mAb size variants by simply replacing non-volatile, high ionic strength buffers (e.g., phosphate salts) with volatile (e.g., ammonium acetate) mobile phase conditions.35-37 For instance, Yan et al. reported a mixed-mode SEC-nMS method using an ammonium acetate and ammonium bicarbonate mixture for highly sensitive detection and quantitation of homodimer impurities for four BsAbs with limits of detection (LOD) from 0.1 to 0.01% relative abundance.³⁵ However, the use of volatile salt mobile phases at relatively low concentrations resulting in poor peak resolution of the different variants. In a comparison study performed on 30 therapeutic proteins including 24 mAbs, 2 ADCs, and 4 Fc-fusion proteins, it was demonstrated that only acidic therapeutic proteins (pI < 7) could be successfully analyzed with a MS-compatible mobile phase containing 100 mM ammonium acetate for SEC.36 More recently, Ventouri et al., reported the potential risk that the separation conditions used for SEC-nMS can impact the protein structure.45 The authors have demonstrated that ammonium acetate (kosmotropic) at ionic strengths above 0.1 M effectively eliminated undesired secondary interactions with the stationary phase and did not induce denaturation, whereas ammonium formate and bicarbonate (both chaotropic) caused structural changes. These findings highlight the limited choice of volatile buffers that can be used for SEC-nMS analysis, as well as the challenge of having to operate at high salt concentrations (i.e. above 0.1 M), whereas MS manufacturers typically recommend the use of a 10-50 mM concentration range in order to reduce ion suppression and prevent the MS source contamination.

The introduction of new hydrophobic resins enabled HIC to be coupled to MS using ammonium acetate in combination with one or more organic modifiers in the mobile phase, for the characterization of both mAbs⁴⁶ and ADCs.⁴⁰ As an example, Wei et al. reported the development of a HIC-nMS method for the determination of minor mAb variants including oxidation, glycation, and free thiol variants by utilizing 150 mM ammonium acetate on a polypropyl UHPLC wide-pore HIC column, which both ensure retention and adequate MS sensitivity for the characterization of low-abundance peaks.³⁸ In another recent study, Chen et al. described the development of a native RPLC-nMS method using a 50 mM ammonium acetate and water/isopropanol mobile phase for the separation of intact ADCs. 40 The stationary phase used in this study provides weaker drug-surface interactions compared to the noncovalent interactions holding the ADC conjugates together. The hydrophobicity of the bonded phase is varied, and the least hydrophobic bonded phase in the series, poly(methyl



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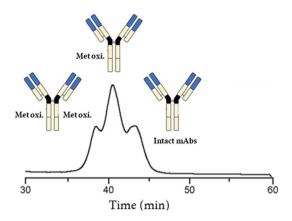


Fig. 1 Total ion chromatogram (TIC) recorded during analysis of oxidized mAb1 by native FcRn AC-nMS. Reproduced from R. Gahoual, A.-K. Heidenreich, G. W. Somsen, P. Bulau, D. Reusch, M. Wuhrer, et al., Detailed Characterization of Monoclonal Antibody Receptor Interaction Using Affinity Liquid Chromatography Hyphenated to Native Mass Spectrometry, Anal. Chem., 2017, 89, 105404-5412. With permission from American Chemical Society, copyright 2017.

methacrylate), resolved the intact species of two ADCs. However, the dissociation of noncovalent ADC complexes (some with DAR-6 and all with DAR-8) is substantial and extensive under these conditions.¹⁴

Affinity chromatography hyphenated to native MS (AC-nMS) has been also implemented for the characterization of therapeutic mAbs. 41,43,44 Gahoual et al., reported the hyphenation of a FcRn stationary phase with nMS to investigate methionine oxidation⁴¹ (Fig. 1). This FcRn affinity LC-nMS method proved to be suitable to identify and quantify PTMs and their impact on the binding to the FcRn receptor. In another study, the hyphenation of FcyRIIIa AC columns to nMS enabled the assessment of the major Fc glycoforms at the intact level,



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which is highly relevant to characterize antibody-dependent cellular cytotoxicity (ADCC).43,44

Overall, the hyphenation of conventional LC methods with native MS for intact mass analysis has been implemented over the last few years to complement the existing analytical toolbox for mAb characterization. However, some challenges still need to be overcome, such as avoiding the formation of adducts, enhancing sensitivity, and the optimization of the chromatographic separation for a broader implementation of this technology in the biopharmaceutical field.

2.2. Two-dimensional liquid chromatography/mass spectrometry (2D-LC-MS)

The value of 2D-LC in the biopharmaceutical industry became obvious in the last ten years and its implementation has been facilitated by the efforts of various research groups. 47-51 Schoenmakers, Stoll, and Pirok have listed 160 2D-LC methods between 2016 and 2018 time frame for example. 47 According to the FDA, there are 100 000 000 potential variants for one single mAb product.⁵² The greater complexity of large molecules in comparison to small ones has accelerated the development of 2D-LC methods to overcome the limits of (i) peak capacity with 1D-LC method; and (ii) nMS with regards to solvent compatibility. There are several beneficial aspects when analyzing large molecules by 2D-LC in comparison to small ones. It has been reported that resolution is independent from column length for RPLC gradient elution of proteins. 53,54 Therefore, the gradient slope has a greater contribution than column length on the chromatographic separation, allowing the use of short columns and analyte peak focusing on the ²D column inlet.⁵⁴ Second, the under sampling issue, which is well known with small molecules, 48 is less critical with large ones due to broader peaks caused by slower mass transport due to their larger size.55 Overall, 2D-LC is an elegant approach to improve the chromatographic peak capacity within a reasonable analysis time. Pirok et al. described a variety of orthogonal chromatographic modes that can be coupled, and weighted associated challenges in ensuring a solvent compatibility between the two dimensions.56

Multiple modulation strategies have been described in order to ensure solvent compatibility between chromatographic dimensions (typically RPLC-HILIC), including the popular active solvent modulation (ASM) introduced by Stoll,⁵⁷ or the stationary-phase-assisted modulation (SPAM).⁵⁸ While the compatibility of aqueous SEC to RPLC is ideal, the use of HILIC in the second dimension generally requires the use of active modulation strategies.⁵⁶ Several 2D-LC modes can be selected depending on the complexity of the sample, such as the heart-cutting (LC-LC), multiple-heart cutting (LC-LC), selective comprehensive (sLC × LC) and comprehensive (LC × LC) modes. 48 The heart cutting and multiple heart cutting modes are simple 2D-LC strategies allowing the fractionation of a limited number of peaks in the first dimension. The multiple-heart cutting core is largely used in the biopharmaceutical industry at the protein intact level as it allows the simultaneous characterization of multiple impurities.

The sLC \times LC and LC \times LC separation modes are generally preferred for complex mixtures because they provide a more complete picture of the sample. The sLC \times LC allows the collection of multiple fractions across one chromatographic peak, while the full 1D separation will be fractionated in the LC \times LC mode, which allows the fractionation and analysis of untargeted impurities. However, with the comprehensive 2D-LC mode, the 2D separation must be fast (less than 1 min) to allow collecting enough fractions in the first dimension, which may affect the 2D

separation performance. In addition, the data treatment can

become tedious and time consuming.

Non-denaturing LC techniques are often incompatible with MS due to the non-volatile salts (e.g. phosphate for SEC) used with these LC separations. Numerous 2D-LC applications for biopharmaceutical proteins have been described, where RPLC was used for desalting of samples coming from the first dimension separations that involve the use of non-volatile salts or high volatile salt concentrations. 59-62 In particular, SEC-RPLC and IEX-RPLC methods have been published for the characterization of size and charge variants of antibody at intact level using denaturing conditions in the second dimension. 63,64 In addition, online on-column reduction strategies have been developed to generate antibody fragments in order to improve the mass accuracy when compared to intact mass analysis. 64,65 Beside the use of 2D RPLC denaturing conditions, the use of ²D SEC columns allowed an online desalting using ammonium acetate under non-denaturing conditions, which are required for the analysis of samples with weak inter-molecular interactions (non-covalent aggregates)⁶⁰ or inter-molecular interactions (cysteine-linked ADCs). 59,65

Beyond the improvement in the peak capacity, 2D-LC has also gained a major interest for PAT, enabling the integration of this analytical technology in production and purification environments. Since recombinant mAbs are mainly produced in mammalian host cell lines,66 the presence of processrelated impurities such as host cell proteins (HCPs) in the harvested cell culture fluid (HCCF) may interfere with the analysis of mAb PTMs. 67,68 Therefore, the targeted mAb must be extracted from cell culture media prior to analysis by LC. Affinity chromatography is the most selective technique used in the biopharmaceutical field as it enables the separation of a protein based on a reversible interaction between the protein and a specific ligand. 67,69 Due to the high-affinity of Protein-A (ProA) for the Fc region of IgG-type antibodies, the affinity chromatography using Protein-A (ProA-AC) is conventionally used as an effective capture step in purification processes. 69,70 However, the isolation of antibody in cell culture by off-line procedures, prior to analytical testing, presents a considerable workload. The use of 2D-LC to fractionate and isolate the target antibody from process impurities such as HCPs before their in-line analysis represent a promising technology to streamline the purification and characterization process in a time efficient way. 71-73 For instance, Dunn et al., have recently reported the development of a 2D-LC ProA-SEC method to measure titer and aggregation of a target mAb from HCCF in under 5 min.⁷¹ Similarly, a fully automated 2D-HPLC method

was developed for characterizing protein aggregation of crude in-process bioreactor samples, by combining ProA purification and SEC separation, which has the potential to be employed at-line within a bioprocessing system. The authors utilizes an in-line fraction collection device that can collect up to twelve fractions from a single sample.⁷³ Alternatively, Sandra et al. described several 2D-LC approaches, combining ProA in the first dimension with either SEC-UV, IEC-UV or RPLC-MS in a heart-cutting mode in the second dimension.⁷² These strategies enable performing clone selection based on titer and structural aspects such as aggregation and fragmentation (SEC), charge variants (IEX). In addition, molecular weight, amino acid sequence, and glycosylation can be determined by RPLC-MS. Fig. 2 shows an example of the use of heart-cutting 2D-LC (Protein A-SEC) method for the analysis of size product variants for (a and b) Herceptin (trastuzumab) and (c and d) trastuzumab in the supernatant derived from different CHO clones. The 2D-LC method allowed the quick identification of high producing clones with a low tendency to produce aggregate.

The success of 2D-LC in the biopharmaceutical industry was first fueled by its inherent improvement of separation performance and more recently boosted by its possibility to be integrated into production and purification environment, allowing the automation of analytical characterization. However, 2D-LC approaches can only provide complementary information at the intact and sometimes reduced levels of mAbs, but more detailed information at the peptide level is often needed at the very early stages of development.

Therefore, the automation of the off-line sample preparation steps within the same chromatographic system is highly desirable. Despite the success of 2D-LC and bio-compatible 1D-LC systems in the biopharmaceutical industry, there is currently no commercial fully biocompatible 2D-LC system. When running SEC methods in the first dimension of 2D-LC setups with mobile phases containing a high amount of chloride anions, regular stainless steel systems will be corroded over time and release metals in the sample flow path. In the meantime, the release of iron has been shown to induce mAb adsorption on CEX columns.74 In addition, non-specific interactions between the stainless steel parts of LC systems and phosphate-containing buffers are well-known, but may occur with proteins too.⁷⁵ The lack of fully-biocompatible 2D-LC system might be explained by the limited mechanical stability of the truly biocompatible PEEK material, and only the partial removal of non-specific interactions with titanium or MP35N materials (iron free but not metal free). However, PEEK-lined titanium capillaries have been recently commercialized. These materials present the advantage of tolerating high pressure and being truly biocompatible. The improvement of chromatography data systems (CDS) software will further push 2D-LC to the next level with regards to its adoption beyond 2D-LC experts as discussed elsewhere. 51,76 For example, some CDS software now allows the control of both mD-LC setups and HRMS instrument using a single software, preventing possible mismatch between the LC and MS sequences.

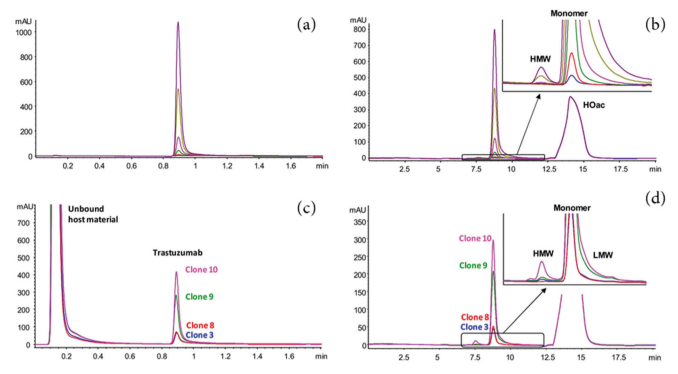


Fig. 2 2D-LC (Protein A-SEC) analysis of a dilution series of trastuzumab (a and b) and supernatant collected from CHO clones producing trastuzumab (c and d). (a) ¹D Protein A chromatograms at UV 280 nm of dilution series (0.02*, 0.05, 0.1, 0.2, 0.5 and 1 mg mL⁻¹). (b) ²D SEC chromatograms at UV 214 nm of dilution series of trastuzumab and zoomed view showing the detection of HMW variants. (c) ¹D Protein A chromatograms of the supernatant collected from four CHO clones producing trastuzumab. (d) ²D SEC chromatograms of fractionated trastuzumab and zoomed view showing the detection of HMW and LMW variants. Reproduced from K. Sandra, M. Steenbeke, I. Vandenheede, G. Vanhoenacker, P. Sandra, The versatility of heart-cutting and comprehensive two-dimensional liquid chromatography in monoclonal antibody clone selection, J. Chromatogr. A., 2017, 1523, 283-292. With permission from Elsevier, copyright 2017.

Bottom-up analysis

In 2009, the FDA recommended a quality-by-design (QbD) approach for manufacturing therapeutics.⁷⁷ The QbD guidelines require direct monitoring of pCQAs at the peptide level, thus ensuring the desired product quality is achieved at the end of the production process.77,78 This involves a better understanding early-on of both the product and process, as well as the deployment of analytical strategies to monitor and maintain high quality throughout the lifecycle of biotherapeutic products. As a result, despite the challenges posed by manual off-line steps and data analysis, LC-MS/MS peptide mapping techniques for quality testing into regulated (GxP) laboratories have generated significant interest in recent years. 69,79

3.1 Conventional multi-attribute methods (MAM)

The LC-MS/MS peptide mapping-based approach is a multiattribute method (MAM) as it can measure simultaneously multiple protein modifications in a single analysis. This approach was recently introduced for quality testing during mAb development. 15,80,81 Fig. 3 presents the standard peptide mapping-based MAM workflow for therapeutic mAb analysis. Briefly, mAb is digested using a protease, generally using

trypsin, followed by an in-depth characterization of obtained peptides using MS/MS experiments. Afterward, wild-type peptides as well as site-specific modifications including PTMs, sequence variants, clippings, etc. are identified from protein database search. Product quality attributes (PQAs) of interest are then selected, acceptance criteria are set for each attribute, and an MS1-based extracted ion chromatogram (EIC), together with the automated processing method is created. Relative abundance values for each PQAs are calculated based on the ratio of modified to total peptide abundance.⁶⁹

In addition to the targeted attribute quantification, new peak detection (NPD), also referred to as Non-Targeted MS Processing (NTMS), is a data processing commonly used in MAM approach, allowing to compare two LC-HRMS data files.⁷⁸ From this, it is possible to determine any differences between a test sample and that of a reference standard, which may indicate unexpected changes in product quality. NPD can also detect both the presence of new peaks and the loss of existing peaks.

Compared to historical methods (Table 1), which typically assess the intact molecule or its subunits, this LC-MS/MS peptide mapping-based approach provides information on site-specific CQAs (e.g., charged modifications) and a more detailed understanding of the product.

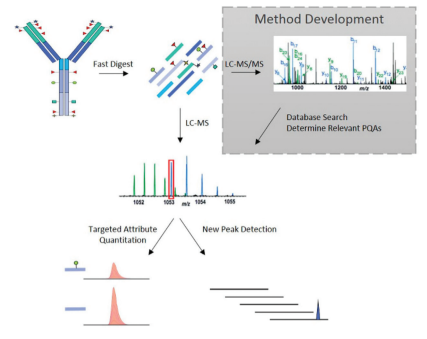


Fig. 3 Standard peptide mapping-based MAM workflow for therapeutic mAb analysis. After digestion of mAb, the obtained peptides are analyzed by LC-MS/MS, followed by a database searching, to characterize the mAb molecules and determine relevant PQAs for further testing. Reproduced from S. Rogstad, H. Yan, X. Wang, D. Powers, K. Brorson, B. Damdinsuren, et al., Multi-Attribute Method for Quality Control of Therapeutic Proteins, Anal. Chem., 2019, 91, 2214170-14177 with permission from American Chemical Society, copyright 2019.

Table 1 Comparison of historical techniques and peptide mapping-based MAM for the characterization of therapeutic proteins at the intact and peptide levels, respectively. Reproduced from S. Rogstad, H. Yan, X. Wang, D. Powers, K. Brorson, B. Damdinsuren, et al., Multi-Attribute Method for Quality Control of Therapeutic Proteins, Anal. Chem., 2019, 91, 2214170-14177 with permission from American Chemical Society, copyright 2019

Attribute	Historical technique	Peptide mapping-based MAM
Charge variant	• cIEF and IEX	• Specific post-translational modifications (asparagine deamidation, Lys-glycation, proline amidation, isomerization) and N- and C-terminal variants
Size variant	SEC (aggregates)CE-SDS (fragments)	Specific clipped species
 Glycan 	 HILIC (released N-glycans profiling) 	• Both <i>N</i> - and <i>O</i> -glycopeptides
Oxidation	• HIC	Specific methionine, tryptophan and cysteine oxidation

Because of these benefits, peptide mapping-based MAM approach has recently been suggested as a replacement of conventional batch release methods, which typically are designed to test one attribute at a time (e.g. IEX for charge variant analysis, reducing gel electrophoresis (rCE-SDS) for clipped variant analysis, or HILIC for glycan profiling). In this context, multiple comparisons of peptide mapping-based MAM workflows with conventional methods were performed. 15,80,82,83 For instance, Rogers et al. demonstrated the performance of a MAM based method using a Orbitrap MS for automated detection and relative quantification of PTMs including oxidation, deamidation, glycosylation, and clipped variant, by showing consistent results with the conventional QC methods. 15 In another study, Wang et al. reported the application of a MAM method for the assessment of multiple glycosylation sites of an IgG1 and compared with the traditional HILIC analysis of released and labeled glycans.83 The quantification of the glycopeptides using the MAM approach has shown a good correlation with the conventional method with similar accuracy and precision. Overall, these studies demonstrated the broad interest in MAM across the MS and biopharmaceutical fields, but still, some challenges need to be further addressed for implementing MAM for QC testing of therapeutic proteins.

One main challenge of MAM is to correlate peptide level with protein level information. For example, the quantification of low abundant PTMs such as glycated and oxidized amino acids at the protein intact level might not be confirmed at the peptide level due to the lower sensitivity of the MAM approach.84 In addition, the analysis at the peptide level usually requires off-line and time-consuming sample preparation procedures, which may generate artifacts such as oxidation, isomerization, and deamidation. 85-87 To limit possible artificial degradation, Wang et al. reported the development of an ultrafast peptide mapping-based approach (uLC-MS/MS)

with a 5 min tryptic digestion.⁸⁸ Although these results are promising, some improvements still need to be done such as the reduction of missed cleavages.

3.2 Correlating peptide with protein level information using mD-LC-MS methods

Automated peptide mapping approaches have been investigated via two main strategies with (i) the use of robotic liquid handling systems⁸⁹⁻⁹¹ and; (ii) the use of HPLC systems.⁹²⁻⁹⁵ The use of robotic liquid handling systems allows the simultaneous preparation of samples in the 96-well plate format using in-solution reduction and digestion, 90,91 while the use of HPLC systems allow generally the preparation of one sample at a time when the digestion is performed using an immobilized trypsin cartridge. 93-96 Robotic liquid handling systems are typically used for the sample preparation of a large number of samples, whereas HPLC-based methods present a valuable approach with a smaller number of samples as the digestion time is significantly reduced when it is performed on-cartridge (minutes) compared to in solution (hours). HPLC-based methods allow the automated online fractionation of antibody variants by conventional LC methods, followed by sample preparation and analysis. 92 Various automated approaches, using either on-cartridge⁹⁵ or liquid handling platforms,⁹¹ have been developed. These automated approaches aim at improving the throughput of conventional off-line peptide mapping procedure. However, the characterization of antibody variants separated at the intact level by SEC, CEX, or HIC or the isolation of a targeted antibody from complex matrices such as cell cultures still require a time-consuming manual off-line fractionation.

In 2018, Gstöttner et al. developed a 4D-LC-MS/MS workflow for the automated characterization of antibody charge variants by an online peptide mapping procedure involving inline fraction collection¹⁷ (see Table 2). The multiple-heart cutting 4D-LC-MS approach allowed to characterize up to 9 peaks with the use of a home-made 4D-LC-MS system with commercial LC modules and controlled via two software (one for the 4D-LC system and another one for the Q-TOF instrument) and an additional in-house macro. 17 The 4D-LC-MS setup allowed a significant time saving when characterizing 5 charge variants (9 hours in total) in comparison to 52 hours using the conventional off-line procedure. In addition, the authors reported almost similar sequence coverage achieved by the online and off-line approaches (94% LC, 86% HC vs. 94% LC, 94% HC). A schematic representation of 4D-LC-MS setup is reported in Fig. 4.

More recently, an optimized 4D-LC setup based on the work by Gstöttner *et al.*¹⁷ was performed using bio-compatible LC modules from another instrument supplier.¹⁸ With the biocompatible 4D-LC setup, a maximum of five fractions could be collected, which is sufficient for most CEX separations, and two software were also needed to control the various modules, including a QExactive Orbitrap instrument. A washing procedure was developed with the optimized 4D-LC system to limit sample carry-over issues,¹⁸ while the published 4D-LC

setup in 2018 is based on a commercial 2D-LC system that automatically washes the loops. Thorough optimization of the different steps of the online peptide mapping was performed with (i) a BioResolve RP mAb Polyphenyl column known to limit antibody adsorption issues in comparison to C4 and C18 RPLC columns⁹⁷ and increased length (50 mm vs. 12.5 mm) to limit ²D column overloading issues, (ii) a Poroszyme cartridge that increased sequence coverage in comparison to the StyrosZyme TPCK-Trypsin cartridge, and (iii) alternative RPLC column chemistry (CSH C18 vs. BEH C18) and dimensions to improve the chromatographic separation of peptides. In order to use the system for multiple days, a thorough optimization of the ²D on-column reduction conditions was performed with the use of TCEP, which is known to be more efficient and stable than the DTT reagent previously employed 17,98 (50% of 2 mM DTT prepared in 50 mM Tris at pH 8.0 is oxidized in 2.5 hours 98). Deamidation, oxidation, and glycation could be quantified using lower amounts of protein in comparison to the conventional off-line approach (60 µg online vs. 250 µg offline). Overall, the online approach showed similar sequence coverage of 90-95% in comparison to the previous study 17 and measured similar amount of oxidations in comparison to offline approaches. The precision of the 4D-LC method for the measurement of PTMs was similar to the one achieved by the conventional off-line approach. 18 Interestingly, a difference in the deamidation level of ~16% was observed for an asparagine residue between the off-line and online procedure. 18 The authors concluded on an artificial deamidation induced by the off-line procedure due to longer digestion times (4 hours vs. 0.4 min).18,99

In another study, a 4D-LC-MS/MS workflow for the characterization of antibody-drug conjugate (ADC) size variants was developed using the same 4D-LC setup previously published, demonstrating its versatility^{18,19} (Fig. 5). An optimization of the ²D RPLC gradient allowed the separation of various LC and HC drug-loaded species, which enabled the determination of the drug to antibody ratio (DAR) of the size variants (see Table 2). In addition to proving the presence of higher DAR species in the fraction containing high-molecular weight species (HMWS), the online peptide mapping also showed different amounts of an oxidized methionine for the HMWS in comparison to the monomeric species when the ADC was submitted to oxidative and light stress conditions. Deamidated asparagine residues have been shown to possibly compromise antibody colloidal stability, and enhance antibody aggregation. 100 The authors further studied the impact of methionine and tryptophan oxidation on antibody stability and aggregation under different conditions.101 To our knowledge, the SEC-Reduction-Digestion-RPLC 4D-LC-MS method was the first to characterize ADC by online peptide mapping and provides a specific example on how this analytical tool could be used to perform a high-throughput screening of clinical candidates and eventually use these data to assess whether a specific amino acidic residue is responsible for protein instability (e.g. aggregation) and reengineer the antibody based on this information.

Table 2 Published mD-LC-MS methods for the multi-level characterization of therapeutic proteins

mAb format	Injection (µg)	mD-LC-MS method	Analysis level	Analytical columns (dimensions)	Analysis time (min)	Targeted by mD-LC-MS	General comments	Ref.
mAb (IgG1)	175	4D – (IEX × Red. × Dig. × RPLC)/impact II ESI Q-TOF MS	Intact – peptide ^a	1 D ProPac WCX-10 (4.0 × 250 mm, 10.0 µm) 2 D AdvancedBio Desalting-RP (2.1 ×	¹ D IEX: 56 min ² D, ³ D, ⁴ D Pep. Map	Oxidation, deamidation, glycation	Online vs. offline Sequence coverages: 94% LC, 86% HC vs. 94% LC, 94% HC	17
				12.5 mm) ³ D StyrosZyme TPCK-Trypsin (2.1 × 100 mm) ⁴ D UPLC ACQUITY BEH C18 (2.1 × 50 mm, 1.7 µm)	96 min frac ⁻¹ . For 5 charge variants ~9 hours		9 hours vs. 52 hours	
3 mAbs (IgG1, IgG2, IgG4), BsAb	09	4D - (IEX × Red. × Dig. × RPLC)/ QExactive TM Orbitrap MS	Intact – peptide ^a	¹ D MAbPac SCX-10 (4.0 × 50 mm, 5.0 μm) ² D BioResolve TM RP mAb Polyphenyl (2.1 × 50 mm, 2.7 μm, 450 Å)	¹ D IEX: 30 min ² D, ³ D, ⁴ D	Oxidation, deamidation, isomerization	Reduction of artifact formation with the online approach (deamidation and isomerization)	18
				³ D Poroszyme® Immobilized Trypsin (2.1. × 30 mm) ⁴ D RPLC XSelect TM CSH C18 (2.1. × 100 mm, 3.5 μm, 130 Å)	Pep. Map.: 80 min per fraction For 5 charge variants ~7 hours			
ADC	09	4D – (SEC × Red. × Dig. × RPLC)/	Intact – peptide ^a	¹ D TSKgel UP-SW3000 UHP-SEC (4.6. × 300 mm,	¹ D SEC: 20 min	Size variants, aggregation, average	Complementarity information between the 2	19
		MS Contractive Of Dittap		2.0 pin, 230 A) 2D BioResolver™ RP mAb Polyphenyl (2.1 × 50 mm, 2.7 µm, 450 Å) 3D Poroszyme® Immobilized Trypsin (2.1 × 30 mm) 4D RPLC XSelect™ CSH C18 (2.1 × 100 mm, 3.5 µm, 130 Å)	² D, ³ D, ⁴ D Pep. Map.: 75 min per fraction	Dark, Oxudation, deamidation	approaches	
mAb (IgG1)	50	$2D - (HIC \times SEC)/UV$	Intact	1 D TSKgel Butyl-NPR (4.6. \times 35 mm, 2.5 μ m)	¹ D SEC: 20 min ¹ D			103
				² D TSKgel UP-SW3000 UHP-SEC (4.6. × 300 mm, 2.0 μm, 250 Å)	HIC: 40 IIIII			

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mAb format	Injection (µg)	mD-LC-MS method	Analysis Ievel	Analytical columns (dimensions)	Analysis time (min)	Targeted by mD-LC-MS	General comments	Ref.
	10	4D – (<i>IdeS</i> × Red. × Dig. × RPLC)/ QEsactive TM Orbitrap MS	Intact – peptide ^a	¹ D Ides-HPLC (2.1 × 50 mm) ² D BioResolve TM RP mAb Polyphenyl (2.1 × 50 mm, 2.7 µm, 450 Å) ³ D Poroszyme® Immobilized Trypsin (2.1 × 30 mm) ⁴ D RPLC XSelect TM CSH C18 (2.1 × 150 mm, 3.5 µm, 130 Å)	¹ D IdeS: 10 min ² D, ³ D, ⁴ D Pep. Map.: 105 min per fraction	Oxidation, deamidation, isomerization, glycosylation	Targeted bottom-up: improvement of both online digestion and peptide separation Sequence coverages: 98.5% LC, 97% HC Characterization of <i>N</i> -glycopeptides	
mAb (IgG1)	10	4D – (ProA × Red. × Dig. × RPLC) /QExactive TM Orbitrap MS	Peptide	¹ D Bio-Monolith Protein-A (4.95 × 5.2 mm) ² D BioResolve TM RP mAb Polyphenyl (2.1 × 50 mm, 2.7 µm, 450 Å) ³ D Poroszyme® Immobilized Trypsin (2.1 × 30 mm) ⁴ D RPLC XSelect TM CSH CLR (2.1 × 30 mm) ⁵ E CLL × 30 mm) ⁶ D RPLC XSelect TM CSH CLR (2.1 × 150 mm, 2.5 × 1.1 × 100 mm, 3.5 × 100 mm	Total analysis time: 100 min	Oxydation, succinimide formation, deamidation	Characterization of mAbs directly from HCCF materials 2 workflows (reduced and peptide mapping) within the same system	102
		3D – (Prod × Red. × HILIC)/QExactive TM Orbitrap MS	Reduced	July 130 A) 1D Bio-Monolith Protein- A (4.95 × 5.2 mm) 2D BioResolve ^{IIM} RP mAb Polyphenyl (2.1 × 50 mm, 2.7 µm, 450 Å) 3D ACQUITY UPLC glyco- protein BEH Amide (2.1 × 150 mm, 1.7 µm, 300 Å)	Total analysis time: 80 min	Glycosylation		
2 mAbs (IgG1)	10	3D – (IdeS × Red. × HILIC)/QExactive TM Orbitrap MS	Middle- up	¹ D Ides-HPLC (2.1 × 50 mm) ² D BioResolve TM RP mAb Polyphenyl (2.1 × 50 mm, 2.7 μm, 450 Å) ³ D ACQUITY UPLC Glycoprotein BEH Amide (2.1 × 150 mm, 1.7 μm, 300 Å)	Total analysis time: 85 min	Glycosylation, glycation	Characterization of the N-glycosylation patterns	20

Abbreviations: ADC, anti-drug conjugate; BsAb, bispecific monoclonal antibody; DAR, drug antibody ratio, Dig., digestion; "D; dimension; Frac., fraction; HLLC, hydrophilic interaction liquid chromatography; IEX, ion-exchange chromatography; IgG, immunoglobulin G; Red., reduction; RPLC, reversed-phase liquid chromatography; LC, liquid chromatography; Mab, monoclonal antibody; MS, mass spectrometry; Pep. Map, peptide mapping; ProA, protein A; SEC, size exclusion chromatography. All mD-LC-MS instruments are home-made and configured with multiple standard modules. Intact – peptide, correlation peptide level with protein level information.

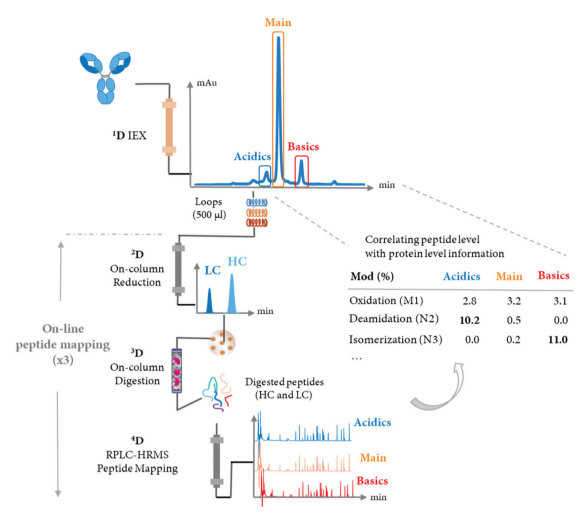


Fig. 4 Schematic representation of 4D-LC-MS setup for the automated characterization of antibody charge variants by an online peptide mapping procedure, enabling the separation and fractionation of charge variants by IEX (¹D) followed by on-column reduction by RPLC (²D), trypsin digestion in flow-through mode (³D), and peptide mapping analysis by RPLC (⁴D). IEX peaks assigned in one single analysis without off-line fraction collection and/or sample prep, while multiple online peptide mapping analyses are performed allowing the identification and quantification of PTMs under each IEX peaks (*i.e.* acidics, main, and basics).

Overall, the digestion performed using an immobilized trypsin cartridge in flow through mode significantly decreases the digestion time in comparison to off-line, in-solution protocols (minutes νs . hours). However, these shortened digestion-time for an intact mAb can cause incomplete peptide cleavages, thereby resulting in lower sequence coverage and poor repeatability. A study evidenced a higher number of misscleaved peptides with the online procedure digestion, approximately 30% (compared to 10% generally obtained with conventional off-line digestion), and this can complicate the data treatment for few PTMs. 102

Recently, a novel strategy aimed at minimizing the missed cleavages was developed (see Table 2). This innovative approach decreases the complexity of intact mAbs, by digesting and reducing the product into small fragments of around 25 kDa, followed by an online peptide mapping analysis of each fragment. This approach considerably improved digestion efficiency with a higher sequence coverage in comparison to

the previously published online workflow (>97% vs. 94%) (Fig. 6a). This improvement can also be explained by a better chromatographic separation of the peptides, since only a fragment of the total mAb is digested and analyzed at one time, and a better ionization (undesirable ionization competition may occur for co-eluting peptides). Notably, the sequence coverage obtained by the novel approach was also slightly higher in comparison to the conventional off-line peptide mapping procedure (>97% vs. 96%) (Fig. 6a). Based on these features, the strategy was successfully applied for the quantification of common PTMs such as oxidation, deamidation, and isomerization located in CDR regions, with similar amounts measured in comparison to the standard off-line approach (see Fig. 6b). The authors also evidenced that by digesting separately the various mAb fragments, the missed cleavages due the steric hindrance of the N-glycans are minimized and their relative abundance levels are similar to the ones obtained with the standard off-line peptide mapping approach (Fig. 6c).

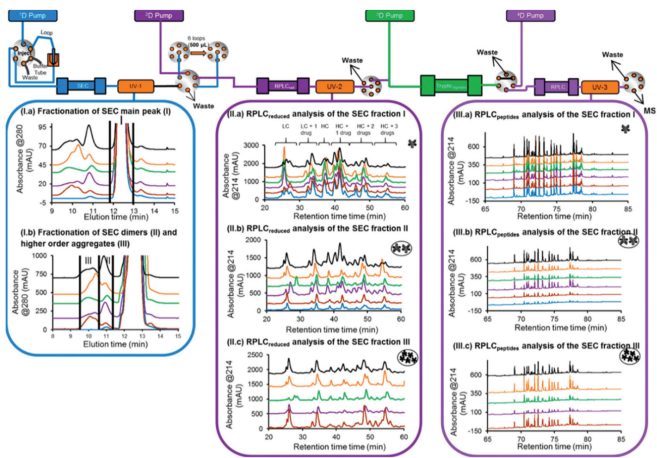


Fig. 5 Schematic representation of an automated 4D-LC-MS workflow for SEC coupled to online peptide mapping, enabling the separation and fractionation of size variants by SEC (¹D) followed by on-column reduction by RPLC (²D), trypsin digestion in flow-through mode (³D), and peptide mapping analysis by RPLC (⁴D). (I, a, b) SEC of ADC product reference, and samples stressed thermally, by light, at basic pH, at acidic pH, and under oxidative conditions (blue, brown, purple, green, orange, gray, and black traces, respectively). (II, a-c) ²D RPLC of the SEC heart-cut fractions that have been reduced online and then eluted. (III, a-c) ⁴D RPLC of the SEC heart-cut fractions that have been reduced and trypsin digested online. Reproduced from A. Goyon, M. Kim, L. Dai, C. Cornell, F. Jacobson, D. Guillarme, et al., Streamlined Characterization of an Antibody–Drug Conjugate by Two-Dimensional and Four-Dimensional Liquid Chromatography/Mass Spectrometry, Anal. Chem., 2019, 91, ²³14896–14903. With permission from American Chemical Society, copyright 2019.

In another study, a 4D-LC-MS (Protein-A \times reduction \times digestion \times RPLC-MS) was developed for the analysis of mAbs in cell culture media, further demonstrating the versatility of this analytical approach. In this proof of concept study, a Protein-A column was used in the first dimension for the isolation of the targeted mAb from HCCF materials, followed by an online peptide mapping analysis using an MS/MS experiment for the determination of multiple common PTMs including oxidation, succinimide formation, and deamidation (see Table 2). The data generated through this 4D-LC-MS setup showed the increase of most of PTM levels during production process. Fig. 7a shows the difference of PTM levels at two different time points (7 and 12 days).

In addition, a 3D-LC-MS (protein A – reduction – HILIC/MS) workflow was incorporated into the same chromatographic system to measure glycation and glycosylation at the reduced level. An accurate and comprehensive profiling of glycosylation patterns was obtained, showing the relative intensities

of mAb glycoforms throughout the production process (see Fig. 7b). Overall, the total acquisition time using both mD-LC setups (3D and 4D-LC-MS) for a multi-characterization level can be completed in just 3 hours. Based on these results, mD-LC-MS can be successfully implemented to perform fast and reliable monitoring of PTMs during the manufacturing.

4. Middle-up analysis

4.1 Conventional middle-down/up analysis

The middle-up level analysis is gaining interest due to its complementarity to the intact mass and bottom-up analysis. 85,104-106 The middle-up approach refers to the analysis of mAbs after their cleavage into large fragments or subunits following limited proteolysis. IgG-degrading enzyme of Streptococcus pyogenes (*IdeS*) protease, which specifically cleaves between the two consecutive glycine residues under the

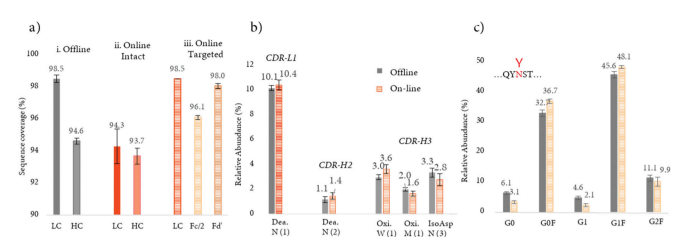


Fig. 6 Targeted bottom-up approach for the improvement of online digestion. (a) Sequence coverages for LC and HC subunits obtained for a mAb with (a) a standard off-line and (b) online approaches for intact mAb, and (c) LC, Fc/2, and Fd' obtained with the targeted approach. (b) Average levels (n = 3) of deamidation (Dea.), oxidation (Oxi.), and isomerization (IsoAsp) located in the CDRs; CDR-L1, CDR-H2, and CDR-H3 (from right to left), were depicted in full and hachured bars for the off-line and online approaches, respectively. Asparagine (N), tryptophan (W), and methionine (M) single codes are followed by numbers to distinguish the different residues. (c) Average levels (n = 3) of N-glycopeptides located in the Fc/2 fragment. Reproduced from J. Camperi, D. Guillarme, C. Stella, Targeted bottom-up characterization of recombinant monoclonal antibodies by multi-dimensional LC/MS, Anal. Chem., 2020, acs.analchem.0c02780. With permission from American Chemical Society, copyright 2020.

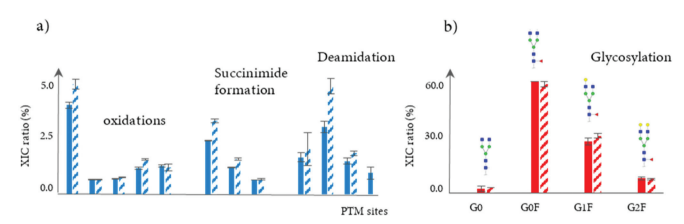


Fig. 7 Analysis of mAbs directly from HCCF materials using of a Protein-A affinity chromatography column (1 D), followed by online bottom-up analysis or HILIC/MS for subunit separation. (a) Determination and quantitation of oxidation, succinimide formation, and deamidation levels of a mAb for different modification sites, obtained by the online peptide analysis (n = 3). (b) Relative abundance of main N-glycans obtained by the reduced level analysis (n = 3), with the assignment of different N-glycans structures. The full and hatched bars correspond to two different days of the production process, days 7 and 12. Reproduced from J. Camperi, L. Dai, D. Guillarme, C. Stella, Fast and automated characterization of monoclonal antibody minor variants from cell cultures by combined Protein-A and multi-dimensional LC/MS methodologies, *Anal. Chem.*, 2020, acs.analchem.0c01250. With permission from American Chemical Society, copyright 2020.

hinge region yielding (Fab')2 and Fc/2 fragments, is typically used. ¹⁰⁴ The subsequent chemical reduction of *IdeS*-generated fragments results in smaller species of around 25 kDa: the LC and Fd. The off-line sample preparation comprising a digestion and reduction steps can be performed within one hour, and requires a low amount of material. ¹⁰⁴ The 25 kDa fragments can then be easily separated by LC¹⁰⁷⁻¹⁰⁹ or CE¹¹⁰⁻¹¹³ and simultaneously identified by HRMS instruments such as QTOFs and Orbitrap analyzers that can resolve the isotopic pattern of 25 kDa proteins with a mass accuracy around 5 ppm. The middle-down approach, which refers to MS/MS-based experiments, can also be performed for sequencing

these fragments.¹¹⁴ Middle-up/down approaches have been successfully applied for: (i) the characterization of mAb PTMs;^{84,113} (ii) the study of specific molecular properties such as mispairing of subunits for BsAb molecules;¹¹⁵ (iii) the mapping of multiple modifications disulfide linkages;¹¹⁶ and (iv) the determination of drug load profile and distribution of ADCs.¹¹⁷ The middle-up approach was particularly interesting for the qualitative profiling of mAb glycosylation patterns.^{108,118,119}

Fornelli *et al.* reported an effective middle-down analysis approach for the identification of oxidized mAb methionine residues using an *IdeS* digestion step, followed by a RPLC-ETD

analysis using an high-field Orbitrap Fourier transform mass spectrometry (FTMS). 109 The developed method allowed the quick detection and localization of IgG oxidation sites, making it ideal for mAbs' oxidation assessment. A similar strategy was applied to investigate both cysteine and lysine conjugated ADCs (i.e., BV; AdcetrisTM and T-DM1; KadcylaTM) at the subunit level, and evaluate multiple analytical attributes including average DAR, positional isomers, conjugation sites, site occupancy and microvariants. 117 D'Atri et al. described a middle-up approach to compare the glycosylation patterns of originators and biosimilars mAbs by using HILIC coupled to HRMS. 108 The HILIC mode was the most suitable for the elucidation of glycosylation patterns, due to the high resolving power and compatibility with MS. 87,120,121 Similar strategies were successfully applied for the characterization of an ADC, 122 Brentuximab Vedotin, and a recombinant Fc fusion protein, Etanercept. 107

In this context, the use of 2D-LC can be particularly beneficial for the analysis of complex mixtures such as antibodies digested at the middle-up or bottom-up levels. Several 2D-LC methods have been published for the comparison of originator and biosimilar antibodies, 123 high-resolution characterization of Fc-fusion proteins¹⁰⁷ or more fundamental applications.¹²⁴ For example, Sorensen et al. reported the use of a CEX \times RPLC 2D-LC method coupled to a TOF instrument in order to compare the originator and biosimilar products of 3 mAbs at the middle-up level. 123 The authors indicated an excellent retention time precision achieved with the 2D-LC-MS method, which could allow to track characteristic mAb fragment and quickly identify minor differences between a biosimilar and originator products.

Other IgG hinge-specific proteases such as IgdE(FabALACTICA®) and Kgp (GingisKHAN®) are also available for in-solution digestions, in order to generate fragments. 116,125-127 In contrast with IdeS protease, which cleaves below the hinge region, IgdE and Kgp cleaves above the hinge region, generating intact Fab fragments from human IgG1.104 The complementarity of both IdeS- and IgdE/Kgpdigested mAbs was demonstrated in many applications. 34,116,125 For instance, Faid et al. reported a middle-up analysis of mAbs after combined IgdE and IdeS hinge proteolysis for the analysis of mAbs disulfide bridges by LC-MS, where the three fragments as well as their corresponding free sulfhydryl forms were well separated. 116 Although the digestion sites of the three proteases IgdE, Kgp, and IdeS are in close proximity in the hinge region of mAbs, different retention profiles can be expected for the differently cleaved Fc and Fd' fragments. As an example, a middle-up approach using a 2D-LC-MS (RPLC-RPLC) method, where the first dimension was merely used for online sample reduction, followed by RPLC-MS analysis of protein fragment separation, was performed for the analysis of a mAb and an ADC (unpublished results). While only minor retention time shifts for Fc/2 and Fd' fragments were observed for the mAb (Fig. 8a), the retention profiles for ADC fragments were completely different due to the distribution of drugs on the Fc/2 and Fd' fragments (Fig. 8b).

4.2 Automated middle-up analysis and multi-level characterization

An immobilized IdeS-HPLC column was recently commercialized, enabling the full automation of mAb digestion into (Fab')2 and Fc/2 fragments within only 10 min. An innovative 2D-LC-MS workflow was developed to characterize a mAb and a BsAb using the immobilized IdeS-HPLC cartridge in the first dimension to generate fragments subsequently separated on a ²D RP column and identified by HRMS. ¹¹⁵ The reduction of spectral complexity of IdeS-generated ~25 kDa fragments allowed an accurate profiling of Fc/2 N-glycans, as well as the identification of mAb variants including C-terminal lysine cleavage, N-terminal pyroglutamination, and glycation. For the BsAb, the same 2D-LC-MS setup also enabled the detection of product impurities due to chain mispairing in one single analysis, with a total run time of only 85 min. This is an example of how a 2D-LC-MS based middle-up analysis can be used to successfully characterize both mAbs and BsAbs.

immobilized*IdeS*-HPLC cartridge implemented in a 3D-LC-MS setup to automatize the off-line sample preparation steps, allowing a fast (85 min per run) and effective characterization of N-glycosylation patterns of mAbs at the middle-up level²⁰ (Table 2). The schematic representation of the 3D-LC-MS workflow is represented in Fig. 9.

Following online ¹D *IdeS*-digestion combined with a ²D reduction step, the three ~25 kDa fragments including Fc/2, LC, and Fd' were successfully analyzed by ³D HILIC coupled to a QExactive MS instrument. As expected, the elution profile of mAb fragments by HILIC was reversed compared to ²D RPLC separation, and enabled the separation of N-glycoforms present on the Fc/2, thus facilitating the glycosylation patterns assessment by MS. Perchepied et al. demonstrated that increased level of confidence can be achieved for the identification of N-glycoforms when they are chromatographically separated since artificial fragmentation may occur during the ESI process. 128 In addition, other PTMs including C-terminal lysine truncation or glycation could also be identified. A mixing tee was required before the HILIC separation in order to improve the solvent compatibility by diluting the flow coming from the ²D RP column containing ~60% aqueous mobile phase with a high amount of acetonitrile (from the 3D pump) to inject mAbs under HILIC conditions (acetonitrile \approx 80%). This reported 3D-LC-MS setup could potentially characterize other mAb or complex antibody formats, especially those with a high number of glycosylation forms such as Cetuximab, where different N-glycosylation sites are present on both Fc and Fd' fragments, 129 or Fc-fusion proteins. 130

Several studies highlighted the advantages of combining the analysis of antibodies at multiple levels, i.e. intact, reduced, middle-up, and peptide level, to allow a fast and extensive characterization. 111,131-133 For instance, Giorgetti et al. reported the combination of intact, middle-up, and bottom-up analysis using a CE-MS instrument for the analysis of 7 FDA-approved mAbs¹¹¹ with: (i) the identification of major N-glycoforms at the intact level; (ii) additional PTMs as K-clip,

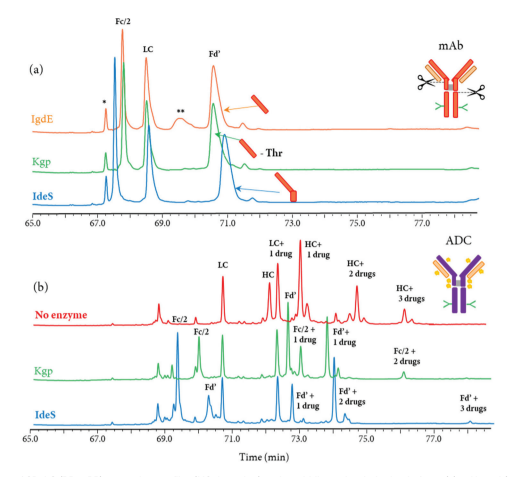


Fig. 8 Comparison of 2D-LC (RP × RP) separation profiles (MS detection) at the middle-up level obtained after a (a) mAb and (b) ADC digestion by different IgG hinge-specific proteases. For the in-solution digestion of mAb, IgdE, KgP, and IdeS-treatment were performed (from top to bottom). The indicated 'Thr' corresponds to the one amino acid difference on the Fd' fragment between IgdE- and KgP-generated fragments. For the ADC, an reduced analysis, as well as middle-up analysis using KgP- and IdeS-protease treatment were represented. mAb, ADC and relative fragments are represented. Incomplete digestion is represented by (**) and peak related to the gradient conditions by (*). Unpublished results.

oxidations or deamidations observed at the middle-up level; and (iii) a precise location and relative quantitation of each PTM at the bottom-up level. This study demonstrated the utility of profiling a mAb at multiple levels to achieve a comprehensive identification of the CQAs. Developed CE methods for the characterization of therapeutic antibodies at multiple levels typically involve similar experimental conditions (use of same separation buffer, of capillary coatings, injection buffer *etc.*) and can be used with a single instrument.

Multi-dimensional LC-MS technologies have the potential to combine the analysis of therapeutic proteins at multiple levels too. As an example, the 3D-HPLC-MS workflow previously described (Fig. 9) can be used for mAb characterization at both middle-up and reduced level within the same system, by bypassing the IdeS-HPLC column. Fig. 10 presents the LC separation profiles obtained for the analysis of an aglycosylated BsAb at reduced and middle-up levels, represented in blue and orange colors, respectively. Using the same mobile phases *i.e.* 0.1% formic acid and 0.05% trifluoroacetic acid in water (A) and acetonitrile (B), two RPLC and HILIC modes can

be simultaneously used in the third dimension to separate the fragments. At the reduced antibody level, RPLC selectivity was not sufficient to separate the two pairs of light (L1 and L2) and heavy (H1 and H2) chains (Fig. 10a), while that HILIC mode allowed to resolve the four BsAb species (Fig. 10b), enabling their assignment by HRMS. Additional middle-up characterization was performed for a deeper BsAb characterization. Fig. 10c shows the ³D RPLC separation of most of the 25 kDa fragments, including the two Fc/2 and Fd', but the two LCs were co-eluting. Conversely, the Fd and Fc/2 fragments could not be separated on the ³D HILIC column, but the two LCs were partially resolved (Fig. 10d). The results demonstrate the high complementary of the HILIC and RPLC separation modes and how the reduced and middle-up level analysis of this BsAb can be performed on such analytical workflow.

An alternative 3D-LC-MS setup was developed with the addition of an immobilized trypsin cartridge to perform PTM analysis at the peptide level. This novel 4D-LC-MS method can be used to localize and quantify PTMs such as oxidation, deamidation, or isomerization present in the CDRs. ¹⁰³

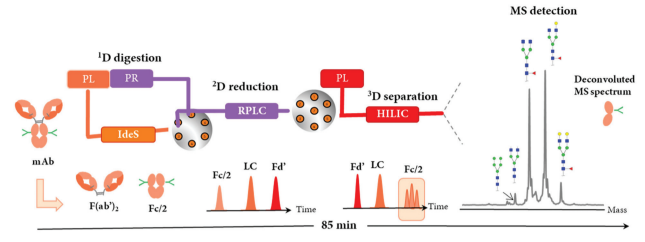


Fig. 9 Schematic representation of the online 3D-HPLC workflow for automated middle-up analysis of mAbs and assignment of major glycoforms of Fc/2 fragment. An *IdeS*-HPLC column was used as first dimension (¹D) followed by on-column reduction by RP-HPLC (²D) and HILIC (³D)/HRMS analysis of the proteolytic fragments Fd', LC, and Fc/2. Reproduced from J. Camperi, L. Dai, D. Guillarme, C. Stella, Development of a 3D-LC/MS workflow for fast, automated and effective characterization of glycosylation patterns of biotherapeutic products, *Anal. Chem.*, 2020, acs.analchem.9b05193 with permission from American Chemical Society, copyright 2019.

Overall, the versatility of the mD-LC-MS setup offers the possibility to perform multi-level of analysis, including reduced, middle-up, and peptide levels for a full comprehen-

sive characterization of therapeutic protein with rapid data acquisition times, within 90 min and without any tedious offline steps. In the future, the immobilization of additional

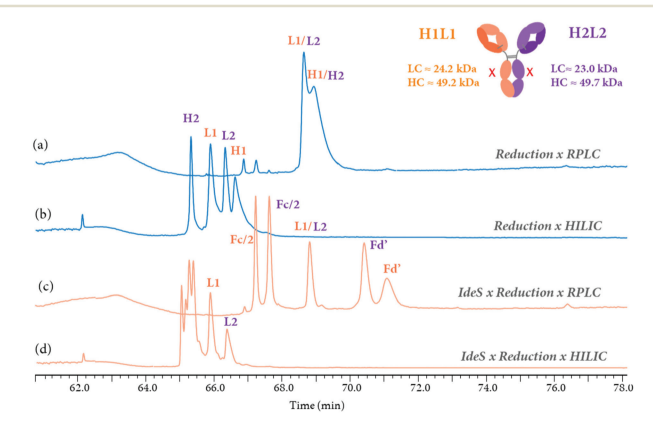


Fig. 10 Reduced and middle-up analysis of an aglycosylated BsAb performed by an online 3D-HPLC (IdeS \times reduction \times HILIC or RPLC-MS) workflow. UV chromatograms obtained at the reduced level (in blue) by (a) RPLC or (b) HILIC by bypassing the IdeS-HPLC column. Middle-up analysis (in orange) using a (3 D) (c) RPLC or (d) HILIC separation (from top to bottom). H1L1 and H2L2 were represented in orange and purple, respectively. The molecular weight of each subunits (LC and HC) were added. The corresponding fragments Fc/2, LC, and Fd' were represented on each chromatogram. Unpublished results.

specific enzymes such as the IgdE, Kgp ones presented in the section 4.1 would be beneficial for the automation of the middle-up/down approach using mD-LC-MS systems. A wider choice of immobilized enzymes would bring the middle-up approach to the next level, with a wider implementation and the possibility to solve analytical challenges inherent to the introduction of novel therapeutic antibody formats.

5. Conclusion

The present review highlighted applications where multi-dimensional LC-MS complement and automatize conventional methods for the characterization of therapeutic antibodies. State of the art LC-MS technologies used to characterize therapeutic proteins at the intact, middle-up and bottom-up levels have been discussed. Native MS and 2D-LC methods are now well established approaches and are implemented throughout the development of conventional therapeutic proteins. In the meantime, automation and high throughput analytical platforms have become a necessity in the fast-paced biopharmaceutical environment. The need to automatize conventional, time consuming off-line fraction collection and multi-steps peptide mapping has never been that urgent. To overcome the limits of conventional sample preparation and analytical methods, the recent development of mD-LC-MS methods, beyond the use of 2D-LC instruments, has brought the technique to the next level. With the data generated by the mD-LC-MS platforms, a direct correlation between peptide level and protein level information can be obtained. The capability of mD-LC-MS methods has been demonstrated with the online characterization of charge and size variants by in-line fraction collection and peptide mapping. Due to the minimal amount of required sample and significant time-saving, the mD-LC-MS technology can be also potentially integrated into production and purification environments for earlier characterization of antibody variants. As an example, the use of protein A chromatography in the first dimension of a multidimensional LC-MS setup allows the online characterization of PTMs, and could therefore enable a continuous monitoring of cell cultures. Therefore, mD-LC-MS has the potential to be an in-process control (IPC) method to better understand and improve biological processes. The versatility of mD-LC-MS platforms also offers the possibility to perform antibody characterization at multiple levels, i.e. reduced, middle-up, and bottom-up levels, within the same chromatographic system. The recent commercialization of a proteaseimmobilized HPLC cartridge has unleashed the potential of online middle-up analysis to provide complementary information to the conventional intact and bottom-up approaches. The development of additional immobilized enzyme cartridges would provide analytical scientists a complete toolbox to overcome future analytical challenges with more complex therapeutics.

Conflicts of interest

There are no conflicts to declare.

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