

mAbs Testing in Inflammation Using Mass Spectrometry – Combination of mAbXmise™ Kits and XEVO™ TQ Absolute XR Mass Spectrometer

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Disclaimer:

The data presented in this application note combines the use of two kits dedicated to the preparation of samples and the use of liquid chromatography and mass spectrometry instrumentation to perform quantitative analyses. The mAbXmise kits described have not been market-authorized by any regulatory entity for diagnostic purposes outside of Europe. Furthermore, the Waters Xevo TQ Absolute XR Mass Spectrometer has not been market-authorized by any regulatory entity. Per the IFUs for the ITDM2 and IADA1 kits, analytical performance depends on the instrument characteristics and its settings. The analytical method shall be validated according to your internal practices. The end user is responsible for completion of the method development and validation. Promise Proteomics mAbXmise kits are not available for sale in all countries. For information on availability, please contact your local sales representative.

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Abstract

Providing a mass spec workflow for monitoring inflammation monoclonal antibodies (mAbs) and anti-drug antibodies levels, combining sample preparation, analyses, and data processing that can be rapidly implemented in clinical labs willing to use mass spec for mAbs testing.

Introduction

For 25 years, mAbs have revolutionized the management of inflammatory bowel diseases. The original anti-TNF drugs, such as infliximab and adalimumab, and more recently additional biologics, including ustekinumab and vedolizumab, have demonstrated their efficacy for the induction and maintenance of remission in Crohn's disease and ulcerative colitis.^{1,2}

Therapeutic drug monitoring (TDM) has also been incorporated into daily clinical practice, especially for patients losing response to therapeutics (reactive monitoring). Hence, measuring serum drug levels and anti-drug antibodies is useful for clinicians to explain drug failure and adapting the medication regimen for patients.

For the 4 mAbs cited above, minimal therapeutic target trough levels in maintenance have been published by international groups of experts (ECCO, ACG, AGA) and are: infliximab >5 µg/mL, adalimumab >7,5 µg/mL, vedolizumab >15 µg/mL, and ustekinumab >1 µg/mL.³⁻⁵

The difficulties for a laboratory willing to offer TDM of mAbs reside in the limited offer of assays for the concerned mAbs and the variability and lack of robustness of these assays. The challenge is to offer a more robust approach with optimal turnaround time and cost-effectiveness.

Liquid chromatography - mass spectrometry (LC-MS/MS) is a method of choice for this application, now described and used by multiple clinical teams.⁶⁻⁸ LC-MS/MS enables direct, peptide-level detection and quantification and thus provides optimal specificity, with reduced susceptibility to matrix interference, enabling unambiguous identification of drug-derived peptides and immune responses even in patients receiving multiple biologics. By doing so, false-negative and false-positive rates are largely reduced, enabling users to drastically reduce errors and samples reanalysis.⁹⁻¹¹ Due to its dynamic range, LC-MS/MS offers an extended measuring range and enables multiplex quantification, which is useful to simultaneously monitor two biologics after a switch or in the case of combination therapies. Finally, a key benefit is the addition of internal standards, which correct the variability (sample variability, variability during technical operations or analyses) and improve the standardization across laboratories.

This application note presents a streamlined workflow dedicated to inflammation mAbs testing using mass

spectrometry. This workflow combines ready-to-use kits that largely facilitate the implementation of the assay in the clinical laboratory and use for routine clinical care, and the analytical sensitivity, precision, and accuracy of Waters Xevo TQ Absolute XR Tandem Quadrupole Mass Spectrometer. The analytical methods have been set up and optimized for 4 inflammation mAbs, commonly used to treat inflammatory bowel disease (IBD) patients – infliximab, adalimumab, ustekinumab, and vedolizumab – as well as two anti-drug antibodies – anti-IFX and anti-ADL. This global solution can be implemented rapidly in clinical labs that are willing to perform mAbs testing and looking for an alternative to immunoassays.

Experimental

Materials and Methods

Materials

- Sample Preparation Kits

1. ITDM2 Monoclonal Antibodies Quantification Kit – Multiplex infliximab, adalimumab, ustekinumab, vedolizumab, obinutuzumab, secukinumab, and tocilizumab (PROMISE Proteomics, France).
2. IADA1 Anti-Drug Antibodies Quantification Kit – multiplex ANTI-Infliximab and ANTI-Adalimumab (PROMISE Proteomics, France).

- Equipment

Equipment recommended: Multichannel pipettes multi-volumes, orbital shaker, vacuum manifold and centrifuge for 96-well plates or positive pressure manifold, nitrogen evaporator or speed-vacuum, pre-slit seal mat for 96 well plates.

- Mass Spec Analysis

1. ACQUITY™ Premier Peptide CSH™ C₁₈, 130 Å, 1.7 µm Column (p/n : 186009488 < <https://www.waters.com/nextgen/global/shop/columns/186009488-acquity-premier-peptide-csh-c18-column-130a-17--m-21-x-100-mm-1-.html>>)
2. Sample analyses were performed on a Xevo TQ Absolute XR Mass Spectrometer
3. Data processing was performed using MassLynx™ Software version 4.2 SCN1050

Methods

Sample Preparation

CALs, QCs, and patient serum samples were prepared for LC-MS analysis using the mAbXmise kits according to the kit instructions. For drug-mAbs quantification, 10 μ L of serum samples were used with an ITDM2 kit and 100 μ L of serum samples were used for antidrug mAbs quantification using an IADA1 kit. After sample preparation with the kits, samples were digested with CutXmise protease, provided in the kits and incubated at 37 °C overnight.

LC Conditions for ITDM2 and IADA1

Chromatographic separation was achieved using an ACQUITY Premier System equipped with a flow-through needle injector (FTN), a binary pump (BSM), and a column manager compartment (CM-A). The separation was performed within 8 min on an ACQUITY Premier Peptide CSH C₁₈ Column (100 x 2.1 mm, 130 Å, 1.7 μ m) with 0.1% formic acid in water and in ACN as mobile phases A and B, respectively. The gradient is shown in Table 1.

Time [min]	Flow [μ L/min]	%A	%B	Curve
0	400	98	2	1
1.0	400	98	2	6
4	400	60	40	6
4.1	500	5	95	6
5	500	5	95	6
5.1	400	5	95	6
5.3	400	95	5	6
5.5	400	5	95	6
5.7	400	95	5	6
5.9	400	5	95	6
6	400	5	95	6
6.1	400	98	2	6
8	400	98	2	6

Table 1. Analytical gradient for LC separation.

MS Conditions for ITDM2 and IADA1

The mass detection of the signature peptides was performed in MRM mode on a Xevo TQ Absolute XR Tandem Quadrupole Mass Spectrometer, equipped with an ESI source. The detection was achieved in positive polarity

with 1.0 kV capillary voltage, 650 °C desolvation temperature, and 1100 L/h desolvation gas flow. The MRM transitions for detecting the peptides and their corresponding internal standard are listed in Tables 2a and 2b.

mAb	Signature peptide	Precursor mass	Cone voltage [V]	Fragment mass	CE [V]
Infliximab	IFX SIN	469.6	30	546.8	13
				603.8	13
				833.4	13
	IFX SIN-SIL	472.2	30	550.8	13
				607.8	13
				841.4	13
	IFX ASQ	598.6	20	631.3	15
				680.8	15
				754.4	15
	IFX ASQ-SIL	602.0	20	636.3	15
				685.8	15
				759.4	15
	IFX SAV	648.8	20	763.4	22
				876.5	22
				1039.5	22
	IFX SAV-SIL	653.8	20	773.4	22
				886.5	22
				1049.5	22
IFX DIL	948.5	20	731.4	22	
			844.5	22	
IFX DIL-SIL	953.5	20	741.4	22	
			854.5	22	
Adalimumab	ADL APY	535.3	30	499.8	18
				738.4	18
				901.4	18
	ADL APY-SIL	539.3	30	503.8	18
				746.4	18
				909.5	18
	ADL GLE	888.2	40	903.9	25
				960.5	25
				1089.0	25
	ADL GLE-SIL	891.5	40	908.9	25
1044.5				25	
1094.0				25	
Ustekinumab	USTE GLD	887.4	20	801.4	23
				1019.5	23
				1189.6	23
	USTE GLD-SIL	892.5	20	811.4	23
				1029.5	23
				1199.6	23
Vedolizumab	VDZ LEW	1119.2	20	429.3	40
				542.3	40
				728.4	40
				1281.4	40
	VDZ LEW-SIL	1123.2	20	429.2	40
				542.3	40
				728.4	40
				1089.6	40

Table 2a. MRM transitions for ITDM2.

mAb	Signature peptide	Precursor mass	Cone voltage [V]	Fragment mass	CE [V]
Infliximab	IFX SIN	469.6	30	546.8	13
				603.8	13
				833.4	13
	IFX SIN-SIL	472.2	30	550.8	13
				607.8	13
				841.4	13
	IFX ASQ	598.6	20	631.3	15
				680.8	15
				754.4	15
	IFX ASQ-SIL	602.0	20	636.3	15
				685.8	15
				759.4	15
	IFX SAV	648.8	20	763.4	22
				876.5	22
				1039.5	22
IFX SAV-SIL	653.8	20	773.4	22	
			886.5	22	
			1049.5	22	
Adalimumab	ADL APY	535.3	30	499.8	18
				738.4	18
				901.4	18
	ADL APY-SIL	539.3	30	503.8	18
				746.4	18
				909.5	18

Table 2b. MRM transitions for IADA1.

Results and Discussion

ITDM2 Kit for Infliximab, Adalimumab, Ustekinumab, and Vedolizumab Drug Levels

Analytical Sensitivity and Calibration Linearity - ITDM2

For all peptides, calibration samples were prepared in triplicates and analyzed in a concentration range between 0.5 µg/mL and 100 µg/mL. The MRM chromatogram for QC high at 50 µg/mL is presented in Figure 1.

The calibration lines were found to be linear with $r^2 > 0.99$ for each of the peptides. The calibration curves are shown in Figures 2a and 2b. All individual calibration point deviations were within the acceptance criteria ($\leq \pm 15\%$). LOQ and r^2 values for each peptide are shown in Table 3.

Peptide	IFX SIN	IFX ASQ	IFX SAV	IFX DIL	ADL APY	ADL GLE	USTE GLD	VED LEW
LOQ [$\mu\text{g/mL}$]	0.5	0.5	0.5	2	0.5	2	0.5	0.5
r^2	0.9983	0.998	0.9978	0.998	0.997	0.9938	0.9986	0.9926

Table 3. LOQ level and r^2 values for ITMD2.

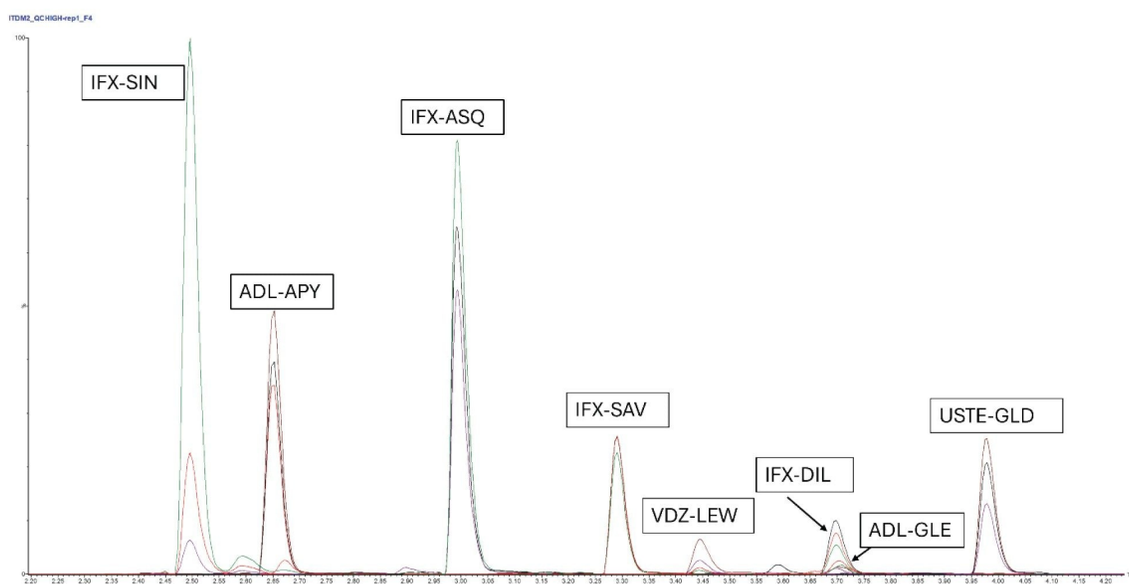
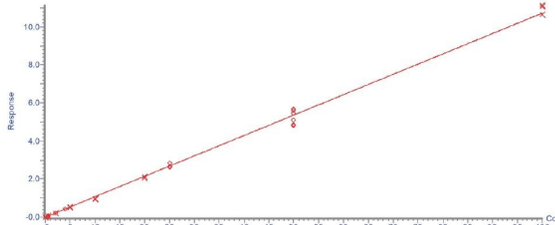
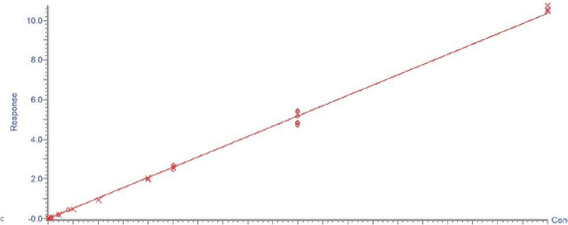


Figure 1. MRM chromatograms of IFX, ADL, USTE, and VDZ peptides at 50 $\mu\text{g/mL}$.

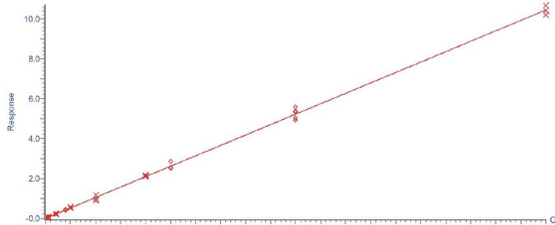
Compound name: Infliximab SAV light
 Correlation coefficient: $r = 0.999207$, $r^2 = 0.997815$
 Calibration curve: $0.107248 * x + -1.09981e-005$
 Response type: Internal Std (Ref 9), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None



Compound name: Infliximab ASQ light
 Correlation coefficient: $r = 0.999206$, $r^2 = 0.996014$
 Calibration curve: $0.103494 * x + 0.00604849$
 Response type: Internal Std (Ref 11), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None



Compound name: Infliximab DLL light
 Correlation coefficient: $r = 0.999224$, $r^2 = 0.998049$
 Calibration curve: $0.104462 * x + 0.00644212$
 Response type: Internal Std (Ref 9), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None



Compound name: Infliximab SINSA light
 Correlation coefficient: $r = 0.999150$, $r^2 = 0.998300$
 Calibration curve: $0.0225668 * x + -0.0058414$
 Response type: Internal Std (Ref 15), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None

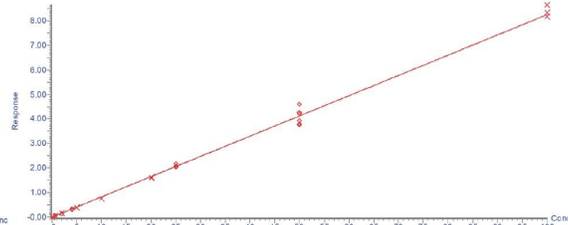
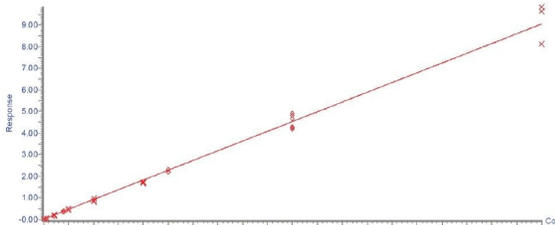
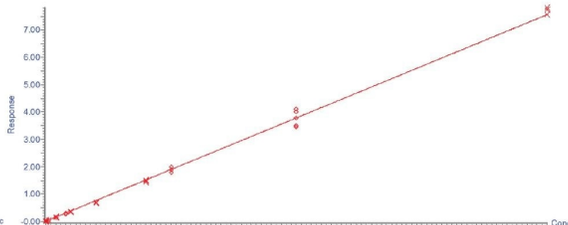


Figure 2a. Calibration curves for infliximab SAV, ASQ, DIL, and SIN peptides.

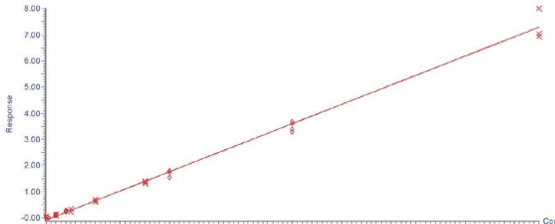
Compound name: Vedolizumab LEW light
 Correlation coefficient: $r = 0.998308$, $r^2 = 0.992630$
 Calibration curve: $0.0902328 * x + 0.0246326$
 Response type: Internal Std (Ref 1), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None



Compound name: Ustekinumab GLDW light
 Correlation coefficient: $r = 0.998661$, $r^2 = 0.997324$
 Calibration curve: $0.0754746 * x + 0.0133836$
 Response type: Internal Std (Ref 9), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None



Compound name: Adalimumab GLEW light
 Correlation coefficient: $r = 0.998326$, $r^2 = 0.993880$
 Calibration curve: $0.0738599 * x + -0.0728791$
 Response type: Internal Std (Ref 7), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None



Compound name: Adalimumab APY light
 Correlation coefficient: $r = 0.998514$, $r^2 = 0.997029$
 Calibration curve: $0.0853458 * x + 0.030624$
 Response type: Internal Std (Ref 15), Area * (IS Conc. / IS Area)
 Curve type: Linear, Origin: Exclude, Weighting: 1/x, Axis trans: None

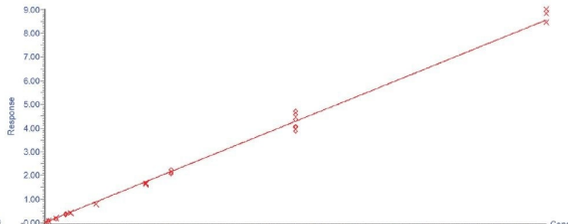


Figure 2b. Calibration curves for vedolizumab LEW, ustekinumab DLD, and adalimumab GLE and APY peptides (ITDM2 kit).

Precision and Accuracy - ITDM2

Method precision and quantitative accuracy were estimated based on various injections of QC samples at different concentrations. The relative standard deviation and accuracy of QC sup (0.5 µg/mL; n=3), QC low (4 µg/mL; n=3), QC med (25 µg/mL; n=3) and QC high (50 µg/mL; n=6) were calculated for each peptide and sample group individually. All peptides show an RSD <10% and are accurate within a range between 94.2% and 114.2%. Results are shown in Table 4.

	Infliximab				Adalimumab		Ustekinumab	Vedolizumab
	SAV	ASQ	SIN	DIL	GLE	APY	GLD	LEW
QC low (4 µg/mL) n=3								
Mean	4.1	4.3	3.9	4.0	4.6	3.9	3.8	3.8
SD	0.05	0.00	0.15	0.10	0.15	0.10	0.15	0.25
RSD %	1.2	0.0	3.8	2.4	3.3	2.5	4.0	6.5
Accuracy	101.7	106.7	98.3	103.3	114.2	98.3	94.2	95.8
QC med (25 µg/mL) n=3								
Mean	25.2	24.9	25.5	25.2	24.1	24.7	24.9	24.8
SD	1.10	0.55	0.90	1.60	1.90	0.75	0.70	0.65
RSD %	4.2	2.2	3.5	6.3	7.7	3.0	2.8	2.6
Accuracy	100.8	99.6	101.9	100.9	96.4	98.9	99.7	99.3
QC high (50 µg/mL) n=6								
Mean	49.0	48.8	49.6	49.9	48.2	49.8	49.3	49.8
SD	3.20	2.70	3.64	2.26	2.16	3.57	3.48	3.26
RSD %	6.5	5.5	7.3	4.5	4.5	7.2	7.1	6.5
Accuracy	98.0	97.6	99.2	99.8	96.5	99.5	98.5	99.6
QC sup (0.5 µg/mL) n=3								
Mean	0.5	0.5	0.6	n/a	n/a	0.5	0.5	0.6
SD	0.05	0.00	0.00	n/a	n/a	0.05	0.00	0.05
RSD %	9.4	0.0	0.0	n/a	n/a	9.4	0.0	8.8
Accuracy	106.7	100.0	113.3	n/a	n/a	106.7	93.3	113.3

Table 4. Analytical precision and quantitation accuracy at different concentrations levels for each individual peptide (ITDM2 kit).

IADA1 Kit for Anti-Infliximab and Anti-Adalimumab Levels

IADA1 Kit Results

Analytical Sensitivity and Calibration Linearity – IADA1

For all peptides, calibration samples were prepared in duplicates and analyzed in a concentration range

between 10 ng/mL and 400 ng/mL. The MRM chromatogram for QC at 75 ng/mL is presented in Figure 3.

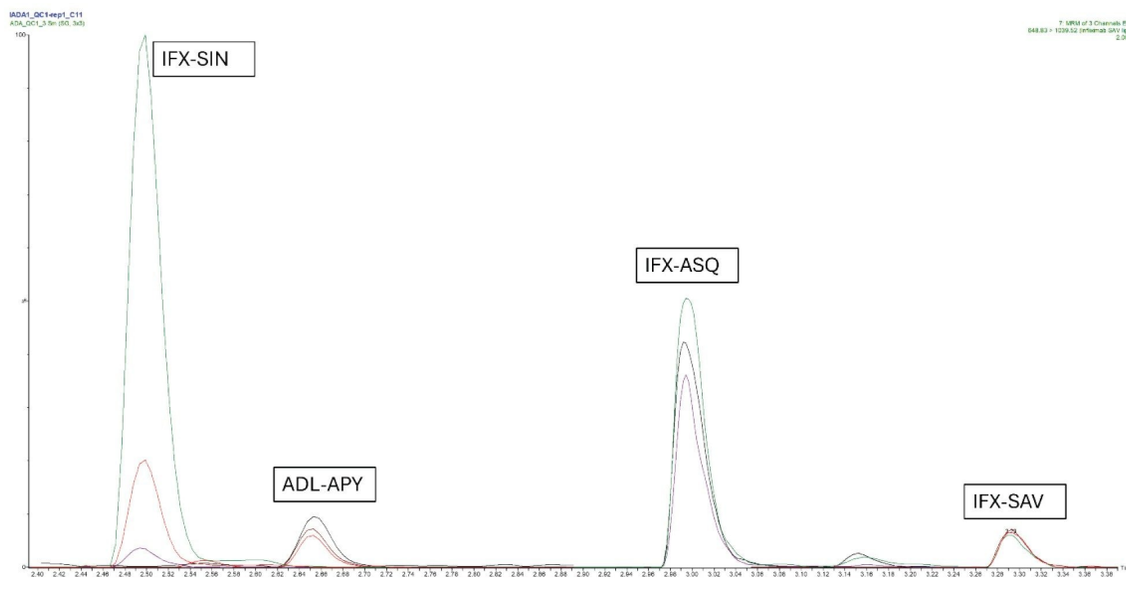


Figure 3. MRM chromatograms of IFX and ADL peptides at 75 µg/mL.

Data evaluation was performed using Arigo Biolaboratories software with 4PL regression.

The linearity plot is calculated based on the following parameters:

x = concentration [µg/mL]

y = (MS response labelled)/(MS response unlabelled) x100

The calibration curves were found quadratic as expected, where $y=d+(a-d)/(1+(x/c)^b)$, with a= Theoretical response at zero concentration, b= Slope factor, c=Inflection point (EC50/IC50), and d=Theoretical response at infinite concentration.

For each peptide, LOQ and r^2 values are presented in Table 5.

The calibration curves for adalimumab and infliximab peptides are shown in Figure 4.

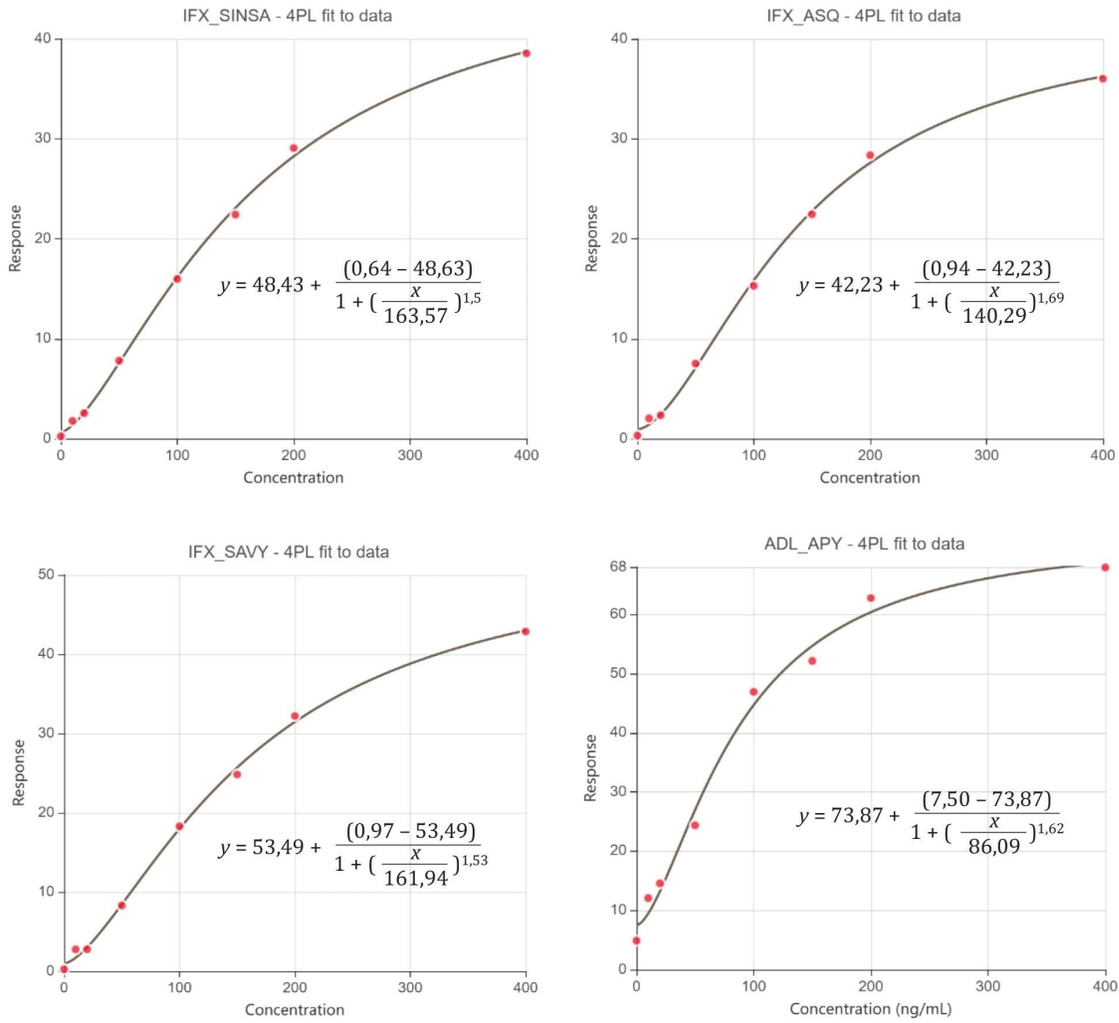


Figure 4. 4PL regression calibration curves for antibodies anti-infliximab and antibodies anti-adalimumab (monitored with IADA1).

LOQ and r^2 values for each peptide are shown in Table 5.

Peptide	IFX SIN	IFX ASQ	IFX SAV	ADL APY
LOQ [$\mu\text{g/mL}$]	10	10	50	10
r^2	0.999	0.998	0.998	0.991

Table 5. LOQ level and r^2 values (IADA1).

Precision and Accuracy IADA1

Method precision and quantitative accuracy were estimated based on 4 injections of individually prepared QC samples at 75 ng/mL. The relative standard deviation and accuracy of the QC samples (n=4) were calculated for each peptide individually. All peptides show an RSD <10% and are accurate within a range between 90.5% and 105.6%. Results are shown in Table 6.

mAb	QC (75 ng/mL) n=4			
	Infliximab			Adalimumab
Peptide	IFX-SAV	IFX-ASQ	IFX-SIN	ADL-APY
Mean	68.7	69.9	68.6	82.1
SD	4.4	2.65	2.79	16.92
RSD %	6.4	3.8	4.1	20.6
Accuracy %	90.5	92.6	90.6	105.6

Table 6. Analytical precision and quantitation accuracy for IADA1.

Quantitation of Patient Samples

Seven patient samples were analyzed using the Promise Proteomics ITDM2 kit and cross-compared to values assigned by immunoassay. Patients were under treatment of adalimumab or infliximab. No ustekinumab or vedolizumab were present in these samples. The results are in good accordance with the reported values achieved by immunoassays and shown in Table 7. As expected, in some cases the absolute values differ in direct comparison of both detection techniques. These findings have been already reported in literature.^{12, 13} However, good correlation between mAbXmise kits and immunoassay has already been reported in the

literature.⁸

Sample	Concentrations measured in µg/mL									
	mAbXmise + LC-MS/MS								Reference Immunoassay	
	IFX-DIL	IFX-SAV	IFX-ASQ	IFX-SIN	ADL-APY	ADL-GLE	Mean 4 peptides INFLIXIMAB	Mean 2 peptides ADALIMUMAB	INFLIXIMAB (IA)	ADALIMUMAB (IA)
Sample 1	25.3	26.7	26.3	19	<0,5 µg/mL	<2 µg/mL	24.3	<0,5 µg/mL	28	no data
Sample 2	4.1	6.9	6.7	6.5	<0,5 µg/mL	<2 µg/mL	6.1	<0,5 µg/mL	2.2	no data
Sample 3	<2 µg/mL	1.2	1.1	1.4	<0,5 µg/mL	<2 µg/mL	1.2	<0,5 µg/mL	<0,5 µg/mL	no data
Sample 4	<2 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<2 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<0,5 µg/mL	no data
Sample 5	<2 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<0,5 µg/mL	15.2	13.5	<0,5 µg/mL	14.4	no data	10.7
Sample 6	<2 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<0,5 µg/mL	5.2	6.8	<0,5 µg/mL	6.0	no data	3.6
Sample 7	<2 µg/mL	<0,5 µg/mL	<0,5 µg/mL	<0,5 µg/mL	interfered	<2 µg/mL	<0,5 µg/mL	<2 µg/mL	no data	<0,5 µg/mL

Table 7. Concentration of adalimumab and infliximab in µg/mL found in patient samples estimated by mass spectrometry (ITDM2 kit – Promise Proteomics) and immunoassay.

In parallel to ITDM2 analysis, the seven patient samples were analyzed with the IADA1 kit. The presence of antidrug antibodies were observed for two patients treated with infliximab and for one patient treated with adalimumab, in good accordance with the results of the ELISA antidrug antibodies. Both methods show the presence of anti-drug antibodies. Some values were above their specific method ULOQ.

Sample	Concentrations measured in ng/mL							
	mAbXmise + LC-MS/MS						Reference Immunoassay	
	Anti-INFLIXIMAB			Anti-ADALIMUMAB	Anti-INFLIXIMAB	Anti-ADALIMUMAB	Anti-INFLIXIMAB	Anti-ADALIMUMAB
	IFX-SAV	IFX-ASQ	IFX-SIN	ADL-APY				
Sample 1	0	0	0	0	0	0	<2,5	no data
Sample 2	0	0	0	0	0	0	<2,5	no data
Sample 3	>400	>400	>400	0	>400	0	>125	no data
Sample 4	205.6	147.6	157.0	0	170.1	0	58.9	no data
Sample 5	<LLOQ	0	0	0	0	0	no data	0
Sample 6	<LLOQ	0	0	0	0	0	no data	0
Sample 7	<LLOQ	0	0	>200	0	>200	no data	>125

Table 8. Concentration of anti-IFX and anti-ADL in µg/mL found in patient samples estimated by mass spectrometry (IADA1 kit – Promise Proteomics) and immunoassay.

Conclusion

This application note highlights a comprehensive suite dedicated to the quantification of monoclonal antibodies and anti-drug antibodies. The solution combines a range of kits for sample preparation (mAbXmise,

Promise Proteomics, France) and associated analytical methods, optimized for the Waters Xevo TQ Absolute XR Mass Spectrometer.

The results demonstrate excellent analytical performance (accuracy, precision). The key advantage of the described solution lies in its rapid implementation in clinical laboratories already familiar with mass spectrometry. The transition from small-molecule analysis to antibodies is made easier and faster, thanks both to the kits that simplify sample preparation for this application and to the analytical methods that are ready to use on the instrument.

Acknowledgements

The authors would like to thank Markus Mallek (MVZ Dortmund, Germany) who kindly provided anonymized samples for comparison.

Specific mentions

The analytical data presented in this material are intended to demonstrate the robustness of a clinical research method. These data in no way substitute for independent method validation required by any applicable legal or laboratory standards.

In Europe, mAbXmise kits are in vitro diagnostic medical devices for professional laboratory use. mAbXmise kits determine the plasma concentration of monoclonal antibodies. Analytical performance depends on the instrument's characteristics and its settings. A validation of the analytical method shall be conducted according to internal practices. Consult the specific instructions for more information. Promise Proteomics products are distributed globally, so uses, applications, and availability of product in each country depend on local regulatory registration status. Products are not registered outside of Europe. Manufacturer PROMISE PROTEOMICS SAS.

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