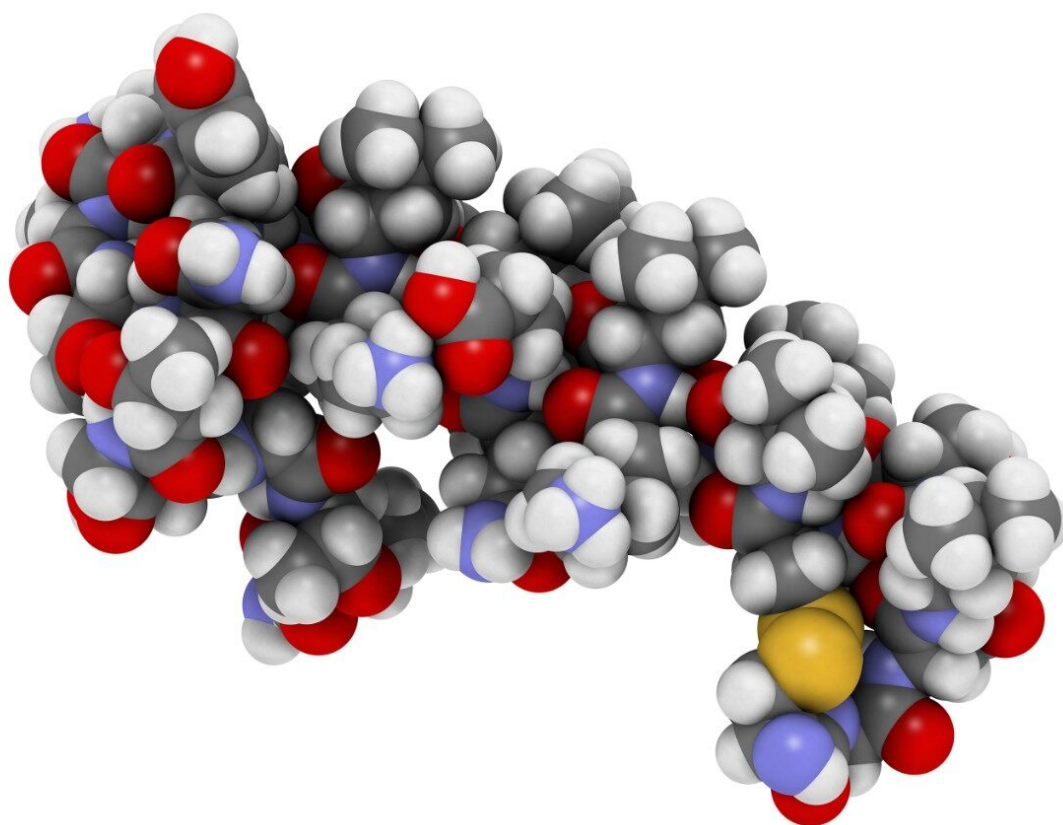


Synthetic Peptide Characterization and Impurity Profiling Using a Compliance-Ready LC-HRMS Workflow

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Abstract

This application note demonstrates how LC-HRMS coupled with UNIFI Scientific Information System can be used for qualitative and quantitative analysis of synthetic peptide impurities, from confirmation of impurities to relative % abundance calculation in a compliance-ready environment.

Benefits

- A compliance-ready solution for peptide characterization and impurity profiling
- UNIFI Scientific Information System with automated workflow methods for synthetic peptide API and impurity characterization
- A custom impurity library in UNIFI for impurity profiling using the accurate mass screening workflow
- Quantitative measurement of impurities combined with custom calculations for impurity limit checks

Introduction

Peptide-based therapeutics are alpha amino acid polymers containing less than one hundred amino acids,¹ represent a new class of highly evolving drugs in the market generating billions of dollars of revenue each year.² These peptide drugs exhibit relatively low toxicity, high biological activity, and potential applications for many medical challenges compared to the most conventional drug products.³⁻⁴ A majority of these therapeutic peptides are produced by chemical synthesis, however, they can be also rDNA-derived or natural source-based.⁵ Each production method is equally challenged by impurities; for synthetic peptides, these can be process-related or product-related that are likely to be biologically active.⁶ With the emergence of Reference Listed Drugs (RLD) versus generic peptide drugs, strict regulatory measures became a necessity for synthetic peptide drugs to ensure their efficacy and safety. The International Council for Harmonization (ICH) has quality guidelines applicable to impurity identification and specification and these have been adopted by regulatory agencies in the US, Europe, and Japan.⁶⁻⁹ Enforcing these quality guidelines in design, development and production processes of synthetic peptides has presented an opportunity for more sensitive analysis methods to be a part of the analytical toolbox.

The conventional synthetic peptide impurity profiling methods are mostly LC-optical-based assays that rely on chromatographic separation of impurities followed by molecular weight confirmation of each peak.¹⁰ Recent studies have shown that incorporating high-resolution mass spectrometry (HRMS) into the analytical workflow can obtain accurate mass-based identification of peptide API and impurities which also benefit from availability of fragment ions for peptide sequence verification.¹¹ The most challenging task in an HRMS workflow is data interpretation to confirm and assign the impurities. In this application note, we introduce the UNIFI Scientific Information System with compliance-ready workflows for this purpose.

Eledoisin, a vasodilator, is a synthetic therapeutic peptide with 11 amino acid residues used as a case study to demonstrate LC-UV-HRMS workflow with UNIFI Scientific Information System. The compliance-ready UNIFI workflows enable data acquisition, data processing to confirm impurities, and calculate the % relative amount, and reporting. In addition, UNIFI Software has custom library features to archive API and impurities. Once the library is built, it can be used for high throughput targeted impurity profiling for the same drug product. The overall strategy used for eledoisin impurity profiling can be applied for other peptide therapeutics during the production process and is ideal for regulated laboratory environments.

Experimental

The synthetic peptide Eledoisin (pEPSKDAFIGLM-NH₂) was purchased from New England Peptides Inc (Gardner, MS) and calcitonin (salmon) (CSNLTCVLGKLSQELHKLQTYPRNTGSGTP-NH₂) was purchased from Bachem (Torrance, CA). A 2 µg/µL stock solution in water was further diluted to a final concentration of 0.2 µg/µL for analysis.

LC conditions

LC system:	ACQUITY UPLC H-Class Bio
Detector:	ACQUITY UPLC Tunable Ultraviolet (TUV)
Wave length:	214 nm
Column:	ACQUITY UPLC Peptide CSH C ₁₈ , 130 Å, 1.7 µm, 2.1 mm × 100 mm
Column temp.:	65 °C
Mobile phase A:	0.1% Formic acid in H ₂ O
Mobile phase B:	0.1% Formic acid in acetonitrile
Gradient:	Eledoisin: 16–24% B over 30 min,

Calcitonin:	14–34% B over 20 min
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MS conditions

MS instrumentation:	Vion IMS QToF Mass Spectrometer
Capillary voltage:	2.8 kV
Cone voltage:	50 °C
Source offset:	50 °C
Source temp.:	80 °C
Desolvation temp.:	300 °C
Cone gas:	20 L/hr
Desolvation gas:	500 L/hr

Informatics

UNIFI 1.9.2 Scientific Information System

- Peptide impurities identification uses the Peptide Mapping Workflow
- Impurity profiling uses the Accurate Mass Screening Workflow

Results and Discussion

UNIFI Processing Method for Peptide API And Impurities Identification

Here, we demonstrate how UNIFI Scientific Information System, a compliance-ready software, comprised of built-in workflow methods with data acquisition, processing, and reporting can be utilized in synthetic peptide characterization and impurity profiling. This work has been done in parallel to our previous study using MassLynx and ProMass Software.¹² For UNIFI-based data processing, the primary amino acid sequence of the synthetic peptide can be imported from the scientific library or created by the user. The

challenge is to identify amino acid modifications causing the impurities. The UNIFI modification database is comprised of custom chemical modifications that can be easily incorporated into the workflow method for locating the impurities based on MS and MS/MS profiles. Figure 1 (*) shows the modifiers of synthetic impurities used in Eleodoisin impurity analysis such as: pyroglutamic acid modification (Figure 1, line 1), insertion and deletion of amino acids (line 4–21), addition of Fmoc, and t-butyl groups due to incomplete deprotection (line 24, 25), and peptide degradation-related products (line 3,23).

The screenshot displays the UNIFI software interface. On the left, the 'Manage Components' window shows a table with columns: Component name, Expected mass (Da), Expected wavelength (nm), Formula, and Description. The first entry is 'Eleodoisin' with an expected mass of 1207.3958 Da and an expected wavelength of 214.0 nm. Below this, the 'Modify Component' window is open, showing the peptide sequence 'EPSKDAFIGL M' with a mass of 1207.3958 Da. On the right, a panel titled 'Selected amino acid modifiers' lists 25 modifiers with columns: Order, Modifier name, Type, and Maximum modification. The modifiers include Pyroglutamic Acid E N-TERM, Amidation C-TERM, Oxidation M, and various amino acid modifications like -Ala, +Ser, -Ser, -Glu, -Gly, -Met, +Gly, +Pro, +Phe, +Ala, +Glu, +Lys, +Asp, +Leu, -Pro, -Phe, -Lys, -Leu, Acetyl N-TERM, Oxidation x2 M, +FMOC, and +t-butyl.

Figure 1. UNIFI processing method set up for product and process related peptide impurities. The component list contains the peptide Eleodoisin sequence without any modifications. The software generates isotopic mass (M and $M+H$) information for the specified peptide sequence. The panel on the right (*) shows the potential modifications of the amino acids sequence (in the method editor) that produces multiple impurities of the given peptide. These are user defined and can be product API-related or impurity-related modifications.

LC-HRMS-Based Impurity Identification

The goal of the analysis is to confirm the API sequence (e.g. Eleodoisin) and identify major impurities. In brief, the peaks were separated using a 16–24% acetonitrile in 0.1% formic acid gradient over 30 minutes using an ACQUITY UPLC CSH C_{18} Column. Even at optimal chromatographic performance, obtaining baseline resolved peaks for optical detection of low abundance impurities is a challenge. Adding HRMS to the analysis resolves this by improving the detection and identification of low level impurities based on the accurate mass and MS/MS information. The TUV (214 nm) in Figure 1A indicates that only a few low abundance peaks are present in the sample. The comparison of sample chromatogram to the reference

using UNIFI mirror plot feature in Figure 1B further confirms this observation. However, the summed mass spectrum for the API peak contains multiple monoisotopic m/z 's. The most abundant co-eluting ions in Figure 1C are +2 charged and based on the Eledoisin sequence the MS spectrum confirms that only m/z 594.8083 belongs to the API. The extracted ion chromatograms (XIC) of selected impurity peaks co-eluting with Eledoisin API are given in Figure 1D.

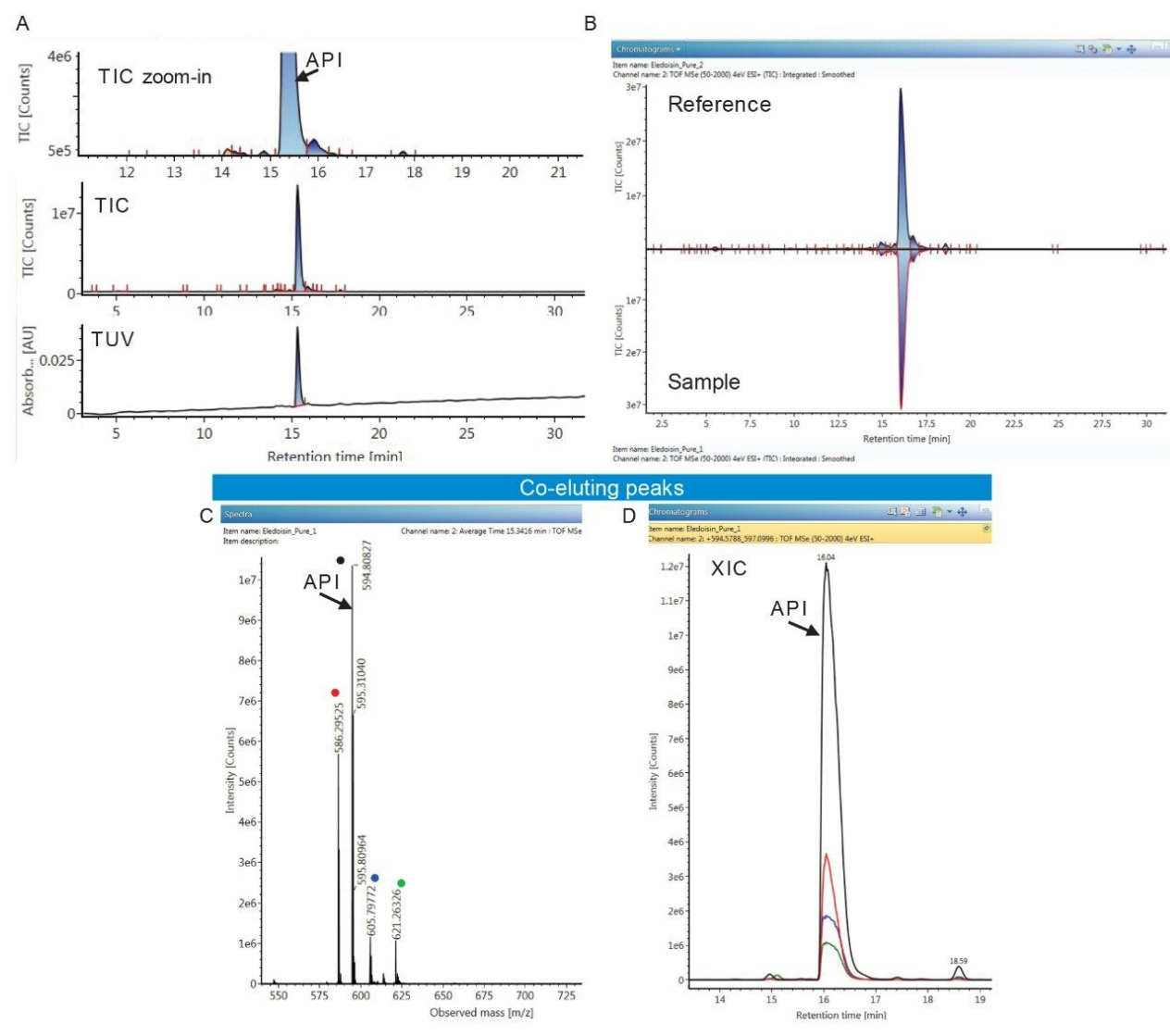


Figure 2. A) TUV, TIC, and zoomed-in TIC spectra are shown for the Eledoisin API peak. B) A mirror plot generated for the chromatograms acquired for Eledoisin pure sample and the reference. This feature can be used to visually compare and contrast two chromatograms, MS, or MS/MS spectra. C) The HRMS spectrum shows the peaks co-eluting with Eledoisin API peak. D) The extracted ion chromatograms (XIC) of the API and the impurities are given in Figure 1D. Each XIC peak is color coded to match the m/z values given.

The deconvoluted mass spectra were used to identify these co-eluting impurities. The UNIFI peptide

mapping workflow identified these peaks to be impurities containing insertions and deletions of amino acids, oxidation, and amino acid protection groups. The MS/MS fragment ions generated using data independent acquisition (MS^E) was used to further confirm the sequence of the Eledoisin API and impurities. Below is the XIC (Figure 3A) and MS/MS spectrum for Eledoisin API peak confirming the amino acid sequence and respective modifications of its most abundant form: N-terminal pyroglutamic modification and C-terminal amidation (Figure 3B). The key features in the UNIFI workflow include automated data processing, chromatographic peak and fragment ion annotation, and customizable scientific library. The components of interest identified in the peptide mapping workflow can be imported into a dedicated custom impurity library to be used in high throughput analysis later in the production process using the UNIFI screening workflow.

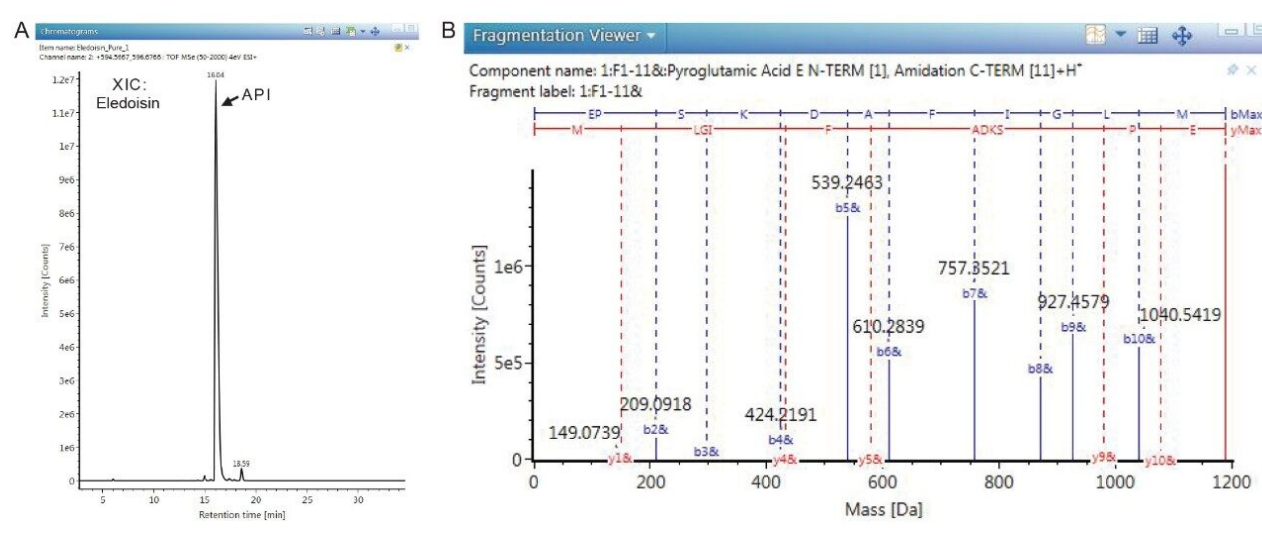


Figure 3. A) The extracted ion chromatogram (XIC) of Eledoisin +2 charge peak (m/z 594.56). B) The MS/MS spectrum for Eledoisin API confirms the peptide sequence containing two modifications; N-terminal pyroglutamic acid and C-terminal amidation.

Scientific Library Feature for Impurity Profiling

Not all impurities are biologically active or harmful; however, identifying undesirable synthetic byproducts is critical in maintaining the quality of the final therapeutic product. Profiling impurities at drug development and production using a streamlined workflow can significantly reduce impurity levels and analysis time spent on assessing the product at the end. The key feature is being able to use the custom impurity library to facilitate impurity screening. Impurities identified in the characterization stage by peptide mapping workflow can be added to the library or created as a new entry based on prior knowledge. Those imported from the peptide mapping data processing workflow contains information about the m/z , charge state, retention time, neutral mass, formula, and fragment ion spectra associated with each component peak. When batch selected, these components can be retrieved and imported back into the

screening workflow with their formula, neutral mass, and retention time information for impurity profiling. The schematic in Figure 4 displays this process.



Figure 4. The schematic shows the path for utilizing the scientific library feature in UNIFI for impurity profiling.

Qualitative and Quantitative Impurity Profiling with Limit Checks and Data Reporting

Using Eledoisin as an example, we demonstrate the use of screening workflow to profile impurities related to oxidation, deletion of Gly/Ala/Pro/Met, and insertion of Lys/Ser/Fmoc. It is noteworthy that these modifications were imported from the impurity library built for Eledoisin.

In addition to detecting these impurities in a large batch of synthetic Eledoisin samples, the screening workflow illustrated here was used to obtain the %purity of each peptide based on UV and MS responses and the %relative abundance to Eledoisin API MS response. The purity levels determined from MS and UV response are different due to co-eluting peptides only detected in the MS-based purity assessment. The reported optical purity for Eledoisin API was 94.7% while MS was 74.8% (the FDA drafted guidelines for synthetic peptides state that any impurity with 0.1% abundance¹³ or higher relative to the API should be identified; the sum of impurity levels of an eligible drug product must not exceed 0.5%¹³). By modifying the limit check parameters the users are able to deploy thresholds for potentially harmful impurities in the sample. Similarly, an example is given in Figure 5A, where %(impurity/API) levels are highlighted in red or yellow based on whether they exceed error limit (maximum threshold) or warning levels respectively. The user-defined error limit used in here is 0.5%. The pass/fail limit checks for % (impurity/API) and retention time are given by using a simple Boolean type custom formula in the method. In addition, the workflow provides sample statistics validating the quality of the analytical method and instrumentation. The examples given in Figure 5B and C show the XIC and %RSD for Eledoisin API and a selected low abundance impurity, API-Proline. Even at 0.08% level API-Pro has a %RSD as low as 2.4% (Figure 5C i, ii).

A report has been generated as shown in Figure 5D following the automated processing using a template to

summarize the final results. The built in automated reporting capability in UNIFI allows the users to create a report based on an existing, modified or a user-defined template. Moreover, the data can be filtered and copied from the analysis method directly on to the template.

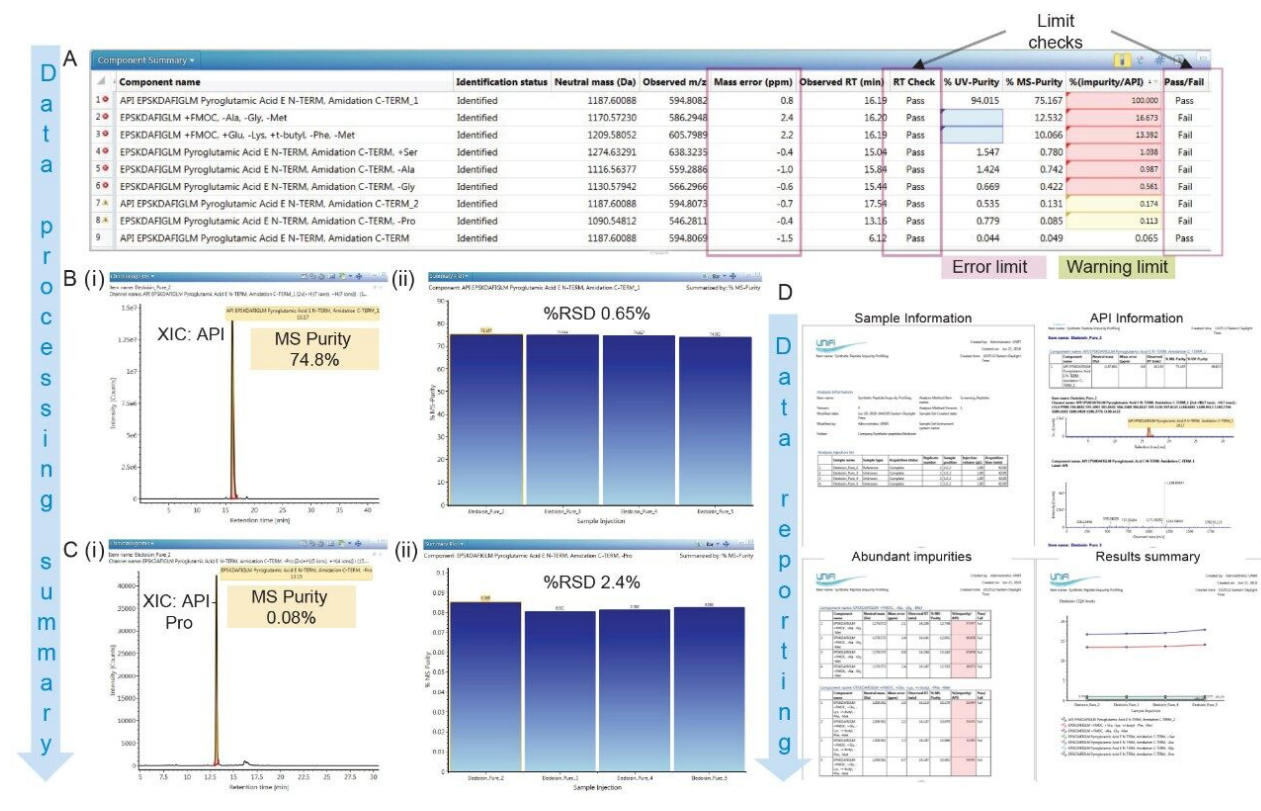


Figure 5. The figure shows the impurity profiling method in the screening workflow followed by the final report generated for the analysis. A) The panel shows the screening workflows' component summary displaying the purity level of API and each pre-selected impurity using both MS and UV detection. Using the features in UNIFI workflow method a pass/fail criteria display has been created to improve data readability. B) This figure displays the XIC (i) and the summary plot (ii) for Eleodoisin API. C) The plot presents the XIC (i) and the summary plot (ii) for the impurity, API-Proline. The %RSD for both peptides are <3%. D) The reporting feature in UNIFI generates a consolidated report for the analysis. The reporting template can be user defined.

Peptides with Unnatural Amino Acids and Cyclic Peptides in Impurity Profiling

Many synthetic peptides contain unnatural amino acid(s) or have a cyclic/semi-cyclic structure. Data acquired for these types of samples are challenging to annotate since most informatics platforms are designed for analyzing natural or recombinant proteins and peptides. To demonstrate UNIFI-based data analyses in the above scenarios, we present the following examples: generating unnatural amino acids in UNIFI for data analysis (Figure 6A), identifying the fragments ions of a linear peptide containing an amino acid "X" with a formula $C_9H_{11}O_2N$ (Figure 6B) and fragmentation spectrum of semi-cyclic peptide

A Analyzing peptides containing unnatural amino acid(s)

Unnatural Amino Acid X (Default Custom Library)

Property	Value
Item type	Amino Acid
Item description	Unnatural Amino Acid X
IUPAC name	
Formula	CBH13N2O2
SMILES formula	CBH13N2O2
Average molar mass	163.1891
Monoisotopic mass	163.0780
Item tag	
SMILES	

Create Library Item

Item name: Unnatural Amino Acid X

Library: Default Custom Library

OK Cancel

Properties

Synonyms	Identifiers										
<table border="1"> <thead> <tr> <th>Synonym</th> <th>Synonym type</th> </tr> </thead> <tbody> <tr> <td>X</td> <td>One letter code</td> </tr> <tr> <td>BB</td> <td>Three letter code</td> </tr> </tbody> </table>	Synonym	Synonym type	X	One letter code	BB	Three letter code	<table border="1"> <thead> <tr> <th>Identifier</th> <th>Value</th> </tr> </thead> <tbody> <tr> <td>#</td> <td></td> </tr> </tbody> </table>	Identifier	Value	#	
Synonym	Synonym type										
X	One letter code										
BB	Three letter code										
Identifier	Value										
#											

B Fragmentation Viewer

E-P-S-K-D-A-X-I-G-L-M

Intensity [Counts]

Mass [Da]

188.069

149.0742, 209.0921, 424.2193, 539.2467, 560.2844, 757.3525, 870.4366, 927.4579, 1040.5427

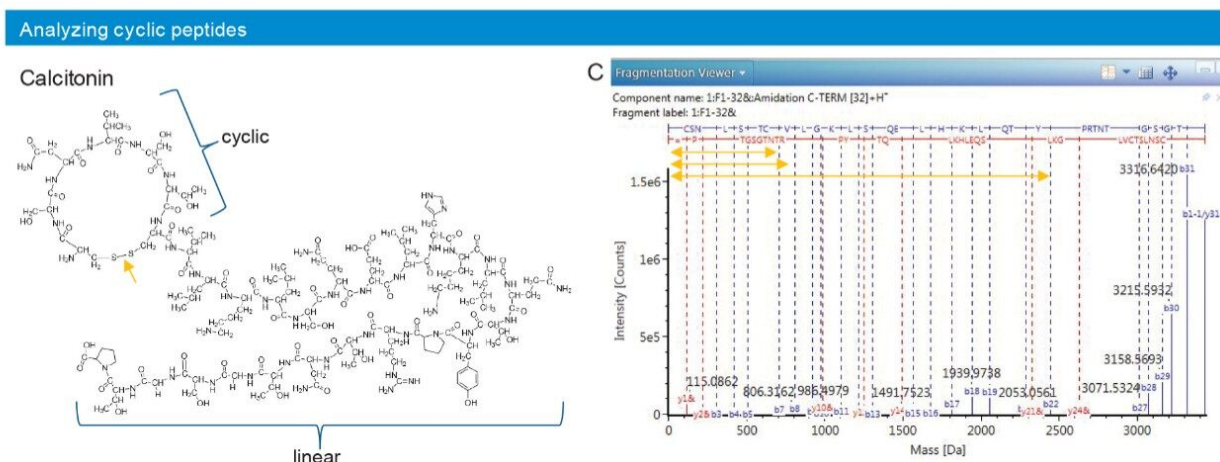


Figure 6. A) UNIFI Scientific Information System has a simple interface that enables the user to generate modifications such as an unnatural amino acid for data analysis. Each of these user defined amino acid will be included in peptide sequence and fragment ion identification. B) The fragmentation spectrum generated for the peptide sequence given contains unnatural amino acid 'X'. The respective fragment ions containing X are labeled (red arrow). C) Calcitonin, a semi-cyclic synthetic peptide cyclized through disulfide bond formation is shown in the structure (yellow arrow). The b series ions, b7 and above contains disulfide bonded fragment ions and a few are shown in yellow.

Conclusion

This application note demonstrates how LC-HRMS coupled with UNIFI Scientific Information System can be used for qualitative and quantitative analysis of synthetic peptide impurities, from confirmation of impurities to relative % abundance calculation in a compliance-ready environment. With automated data processing, impurity archiving and retrieving, custom calculations, limit check capability, and result statistics, this platform provides an ideal compliance-ready solution for profiling and reporting impurities in accordance with the ICH and FDA regulatory guidelines¹³ for peptide therapeutics.

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