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Applikationsbericht

Streamlining Impurity Analysis in Atorvastatin with the ACQUITY[™] QDa[™] II Mass Detector for Enhanced Detection and Quantification

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Abstract

- Demonstrate the practical application of the QDa II Mass Detector to complement UV chromatographic workflows for routine analysis of atorvastatin API impurities
- Assess the suitability of an analytical workflow for determining selected impurities at qualification thresholds of ≤0.15% as per ICH-Q3 guidelines
- · Underline the benefits of in-source fragmentation to gain deeper insights into the structural characteristics of unknown compounds
- Highlight the user-friendly features and advantages of the Empower[™] Chromatography Data System (CDS)
 for compliant impurity testing in a regulated pharmaceutical environment

Introduction

Streamlining Impurity Analysis in Atorvastatin with the ACQUITY[™] QDa[™] II Mass Detector for Enhanced Detection 1 and Quantification Atorvastatin is one of the most widely prescribed medications globally for managing conditions such as hypercholesterolemia and reducing the risk of cardiovascular events. Efficacy in lowering LDL cholesterol levels has made it a prominent drug supplied to millions of patients worldwide. However, the presence of impurities in the pharmaceutical formulation of the drug can pose a significant concern for patient safety. These impurities can originate during the synthesis process of the active substance, or as degradation products under certain environmental conditions, and may possess toxicological or pharmacological properties that could compromise the safety and efficacy of the final drug product.

Atorvastatin is typically administered as the calcium salt of the active hydroxyl acid, in doses of between 10–80 mg per day.¹ According to ICH-Q3 guidelines, drugs consumed with a maximum daily dosage of less than 2.0 g per day should have a qualification threshold for impurities of 0.15%.²

According to the European Pharmacopoeia monograph for atorvastatin as an active substance, the following limits are prescribed for its impurities: 0.3% for related impurity A and B; 0.15% for Impurity C and D; and 0.1% for other unspecified impurities.³

Traditional methods for impurity testing, namely chromatographic techniques coupled with UV detection, serve as the cornerstone for quality control (QC) in pharmaceutical analysis. Supplementing this workflow with mass detection enables confident and rapid identification, characterization, and quantification of impurities. Moreover, mass detection can provide a greater scope to analysis with additional molecular weight information that aids in impurity identification, even at very low limits of detection.

In this application, we aim to demonstrate the suitability of mass detection for fast, accurate, identification and quantification of atorvastatin API-related impurities. Additionally, we aim to characterize impurities through insource fragmentation, employing a pseudo-MS/MS approach, thereby presenting a comprehensive solution for impurity analysis.

Experimental

Commercially available samples of atorvastatin calcium salt and three related impurities (impurity A, C, and I), were purchased from Sigma Aldrich and stored at 4 °C until analysis. Samples were dissolved in methanol (MeOH), and serially diluted with a solvent mixture of 40:60 H₂O:MeCN for analysis.

LC-MS Experimental Conditions

LC system:	ACQUITY UPLC [™] H-Class System with FTN-R
	Sample Manager
Detection:	ACQUITY QDa II Mass Detector
UV system:	Photo diode array (PDA) Detector
Column:	ACQUITY UPLC CSH [™] Phenyl-Hexyl column, 1.7 µm, 2.1 mm x 100 mm (p/n: 186005407)
Column temperature:	30 °C
Sample temperature:	10 °C
Injection volume:	2 μL
Flow rate:	0.4 mL/min
Run time:	27.5 min
Mobile phase A:	10 mM Ammonium acetate in H2O
Mobile phase B:	Acetonitrile with 0.1% Formic acid
Vials:	Clear glass 12 x 32 mm screw neck vial (186000273)
Ionization:	Positive Electrospray (ES+)
Capillary voltage:	1.1 kV
Desolvation temperature:	600 °C

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LC Gradient

Time (min)	Flow rate (mL/min)	%A	%В	Curve
Initial	0.4	90	10	6
8.0	0.4	90	10	6
9.0	0.4	60	40	6
18.0	0.4	50	50	6
18.1	0.4	20	80	6
22	0.4	20	80	6
22.1	0.4	90	10	6
27.5	0.4	90	10	6

Software

Data acquisition, processing, and reporting:

Empower CDS 3.0

Results and Discussion

Method Development

Unlike the method prescribed by the European Pharmacopoeia monograph, the analytical method presented in this application was developed without the use of tetrahydrofuran (THF) as a part of the mobile phase, which was avoided due to its toxicity and instability.⁴ Instead, an ammonium acetate buffer was added to the aqueous mobile phase, and formic acid was added to the organic mobile phase, as a means to reduce pH across the sample run and improve peak shape. Several LC columns and mobile phase compositions were also tested, including optimization of the mobile phase to a pH of 4.0–4.2, which did not increase the resolution of chromatographic peaks when compared to the final method.⁵ Ultimately, the stated method is approximately 60 minutes shorter than the European Pharmacopoeia monograph method and uses less than 10% of the solvent

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volume per injection.

The method was optimised using a 1 μ g/mL mix of atorvastatin and selected impurities (A, C, and I). Cone voltage, capillary voltage, and desolvation temperature were all optimized. Despite most literature methods opting for a C₁₈ column, the CSH Phenyl-Hexyl column proved optimal for improving separation between the atorvastatin compound and related impurity C. A column temperature of 30 °C was optimal to reduce peak tailing of the API.

Initially, a 25 µg/mL sample of atorvastatin was spiked with the selected impurities to assess method performance, with concentrations ranging from 0.36–0.42% relative to the API. Three unknown impurities, labelled as Unknown Impurity 1, 2, and 3 in Figure 1, were also quantified in the PDA trace-at 0.11, 0.14, and 1.15% respectively-relative to the API.



Figure 1. PDA chromatogram of atorvastatin (25 µg/mL) spiked with related impurities (A, C, and I), annotated with mass information and (%) peak area relative to the API. Unknown impurities are also annotated. Inset shows more clearly the separation between the API peak and related impurity C.

Linearity

To assess linearity, a serial dilution was performed for both the API and an impurity mix standard (containing impurities A, C, and I). All calibration curves were acquired using the stated method, which had a total run time of 27.5 minutes.

Streamlining Impurity Analysis in Atorvastatin with the ACQUITY[™] QDa[™] II Mass Detector for Enhanced Detection 5 and Quantification A bracketed calibration plot of atorvastatin is shown in Figure 2.

- \cdot The compound was found to be linear (1/X weighted) from 0.01–5.0 µg/mL with an R² value of 0.999
- The percentage of the observed values (% residuals) for bracketed calibration curves were < 15.4% in all cases
- The peak of the lowest calibration point (0.01 μg/mL), equivalent to 20 pg on-column, had a signal-to-noise ratio of 136 (peak to peak)



Figure 2. Calibration curve for atorvastatin API analyzed over a concentration range of 0.01–5.0 μ g/mL.

Selected atorvastatin related impurities were also assessed for linearity and sensitivity (Table 1). R² values in all cases exceeded 0.998, with % residuals values of <16.3% in all cases. All calibration responses were linear (1/X weighted).

Impurity	Regression equation	R²	Linear range (µg/mL)	Residual error (% deviation from the line of fit)
Impurity A	y = 1.07e+007X + 1.70e+003	0.999	0.01–1.0	<11.7
Impurity C	y = 7.42e+006X + 5.38e+004	0.998	0.05-20.0	<6.5
Impurity I	y = 1.56e+007X + 3.17e+004	0.998	0.005-1.0	<16.3

Table 1. Summary of calibration results for impurities A, C, and I.

An example calibration plot for impurity A can be seen in Figure 3. The method showed no carryover in any of the subsequent blank injections across the analysis, despite the high concentrations of API injected. This was true for all injections of the atorvastatin API up to and including a concentration of 75 µg/mL.



Figure 3. Calibration curve for impurity A analyzed over a concentration range of 0.01–1.0 μ g/mL.

Quantification of Atorvastatin Impurities at 0.15% Threshold

After assessing the sensitivity of the QDa II Mass Detector for the related impurities, an evaluation was made to spike a 75 μ g/mL atorvastatin sample with 0.12% of an impurity mix (A, C, I) - with each impurity present at 0.09 μ g/mL. For simplicity, this sample will be referred to as 0.12% Threshold Sample from hereon. Compounds were

Streamlining Impurity Analysis in Atorvastatin with the ACQUITY[™] QDa[™] II Mass Detector for Enhanced Detection 7 and Quantification analysed using both UV and mass detection using scan mode. An example UV and associated MS TIC chromatogram for this sample are shown in Figure 4.



Figure 4. Stacked PDA chromatogram (A) and MS TIC chromatogram (B) of a 75 µg/mL atorvastatin API sample and related impurities (spiked at 0.12%) annotated with mass information and % area of each peak relative to the API. Insets show separation of the API and impurity C in more detail (A), and an overlay of a solvent blank MS TIC injection (B). Inset B shows smoothed data.

In the PDA trace in Figure 4A, three unknown peaks can be seen at 0.13%, 0.17%, and 1.16% respectively, relative concentrations comparable to those in the 25 μ g/mL sample shown in Figure 1. The corresponding MS TIC chromatogram in Figure 4B shows the presence of an additional unknown impurity, Unknown Impurity 4, with more clarity.

Streamlining Impurity Analysis in Atorvastatin with the ACQUITY[™] QDa[™] II Mass Detector for Enhanced Detection 8 and Quantification Seen also in Figure 4B is an overlay of a solvent blank sample acquired during the same analysis, demonstrating that unlabelled peaks in the MS TIC chromatogram of Figure 4B are not exclusive to the API sample.

To assess the repeatability of this method, 10 replicate injections of the 0.12% Threshold Sample were performed. The results of this can be seen in Table 2. The %RSD for the area of each impurity was < 9% in all cases, with the exception of related impurity A which had a %RSD of 13.4% over the 10 injections.

Repeatability of injections for impurities A, C, and I spiked at 0.12% level relative to 75 µg/mL atorvastatin							
Injection	% Impurity A	% Atorvastatin	% Impurity C	% Unknown impurity 1	% Unknown impurity 2	% Unknown impurity 3	% Impurity I
1	0.11	98.34	0.08	0.10	0.14	1.14	0.10
2	0.08	98.29	0.08	0.12	0.16	1.18	0.09
3	0.09	98.25	0.09	0.12	0.18	1.17	0.10
4	0.11	98.29	0.08	0.11	0.17	1.13	0.10
5	0.08	98.32	0.08	0.11	0.16	1.15	0.09
6	0.11	98.27	0.07	0.12	0.18	1.16	0.08
7	0.12	98.24	0.09	0.13	0.17	1.16	0.10
8	0.10	98.25	0.08	0.12	0.17	1.18	0.10
9	0.11	98.28	0.08	0.12	0.18	1.15	0.09
10	0.09	98.27	0.08	0.14	0.16	1.17	0.09
Mean	0.10	98.28	0.08	0.12	0.17	1.16	0.09
Std. dev	0.01	0.03	0.01	0.01	0.01	0.02	0.01
%RSD	13.4	0.03	6.6	8.8	7.1	1.4	7.1

Table 2. Relative %Area of quantified peaks in [254 nm, 4.8 mm] PDA trace of 75 μ g/mL atorvastatin sample, spiked with 0.12% of impurities A, C and I, over n=10 consecutive replicates.

Impurities A and C can be seen more clearly in the MS TIC chromatogram with the use of a divert valve, which acts to automatically divert flow to waste during a stated time period. An example of this feature in use can be seen in Figure 5, where the flow is diverted from 16.08 to 16.47 mins (RT interval of atorvastatin API elution).



Figure 5. MS TIC Chromatogram showing the use of the divert valve, directing flow to waste as the API elutes, allowing for easier quantification of impurity A [m/z 541.1] and impurity C [m/z 557.1].

Using the ACQUITY QDa II, mass spectra for each peak in the chromatogram can be easily obtained (Figure 6).



Figure 6. Mass spectra of atorvastatin, and atorvastatin related impurities at 1 µg/mL.

In-Source Fragmentation for Structural Information

Upon investigation, unknown impurity 1 and 4 were found to have a molecular weight of *m/z* 557, only 2 Da less than that of the API. Literature suggests this is likely a common photolytic degradant of the atorvastatin compound.⁶ However, unknown impurities two and three require further assessment to better understand their respective structures.

In-source fragmentation can provide valuable structural information about analyte molecules, allowing for the detection of characteristic fragment ions that can aid in the identification and characterization of the compound.

By increasing the cone voltage of the QDa II Mass Detector, one can perform pseudo-MS/MS of given compounds by achieving in-source fragmentation. An example of this can be seen in Figure 7, where unknown impurities two and three are investigated.

Applying a cone voltage of 35 V, the *m/z* 573.3 precursor mass of unknown impurity 3 fragments into *m/z* 454.

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An adduct ion $[M+K]^+$ at m/z 611 can be seen in this analysis, and further fragmentation at a cone voltage of 50V generated product ions with a m/z of 436, 394, 376, and 292-which align more closely with literature assessments of Atorvastatin Methyl Ester⁷. However, more analysis would be needed to be absolute in this statement.



Figure 7. Mass Spectrum of Unknown Impurity 2 (A) and Unknown Impurity 3 (B), using increased cone voltage to achieve In-Source Fragmentation as a means of pseudo-MS/MS, to better profile impurities based on fragmentation. Possible impurity profiles and fragmentation patterns are shown below each section of the figure.

A 50V cone voltage applied to the m/z 451.2 precursor ion of unknown impurity 2 generated a fragment at m/z 422 (Figure 7A). This fragmentation pattern matches both Desfluoro-Atorvastatin and Related Impurity H. The presence a subsequent m/z 380 fragment of m/z 422 indicates a higher likelihood of this compound being Related Impurity H.

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Mass Detection Detects Co-Eluting Impurities

Samples were stored at 10 °C for two weeks to monitor any impurity changes over time. A 75 μ g/mL atorvastatin sample spiked with 0.12% impurity mix was designated as a 'Degradation Test Sample'. Analysis shown in Figure 8 revealed intriguing differences between the mass spectrum of related impurity A in the Degradation Test Sample (Figure 8B) and a sample of similar concentration (0.10 μ g/mL impurity mix) without the API present (Figure 8A). A significant increase in *m*/*z* 557.2 was observed in the former, suggesting the presence of a photolytic degradant of the API co-eluting with related impurity A.





Upon re-quantification of the sample approximately 15 days after preparation, the relative amount of related impurity A compared to the API appeared to increase, from an average of 0.10% (see Table 2) to 0.16% as shown below in Figure 9. This difference is not attributed to a change in the concentration of related impurity A, but rather to a co-elution with another compound, which becomes more prominent as the API degrades. Consequently, this leads to an overestimation when quantifying using UV chromatography. The co-elution is illustrated in Figure 9, where an XIC for related impurity A (m/z 541.1) is overlayed against an XIC for the m/z 557.2 impurity, as well as the MS TIC scan. This highlights the importance of mass detection to ensure accurate quantification of impurities and proper management of co-elution.

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Figure 9. UV Chromatogram of 75 μ g/mL atorvastatin API sample spiked with 0.12% impurity A, C, and I, ~ 15 days after sample preparation. Relative concentration of impurity A (m/z 541) is overestimated, at 0.16%, due to co-elution with another impurity (m/z 557). Inset in the top left of the figure shows a MS TIC Scan (ESI+) overlayed against an XIC for both compounds, demonstrating the composition of the peak.

Empowering Data Reporting

This application extensively leveraged Empower CDS reporting functionality to facilitate the streamlined assessment of relative impurity concentrations within sample batches, as depicted in Figure 10. This robust feature allows users to swiftly make well-informed decisions, thereby ensuring both workflow compliance and data integrity.



Figure 10. Example of Empower CDS generated data report for an analysis of a 75 μ g/mL atorvastatin sample spiked with 0.12% impurity A, C, and I, and other unknown impurities.

Conclusion

This work demonstrates the significant benefits for end-users in adopting the ACQUITY QDa II Mass Detector into existing UV workflows for routine testing of impurities in APIs, by demonstrating the capability to:

- Effortlessly annotate peaks with molecular weight, for simple mass confirmation within an analysis, streamlining the confirmation process of known impurities
- Precisely quantify impurity levels, ensuring atorvastatin related impurities A, C, and I are maintained below a qualification threshold of 0.15% relative to the API
- Enhance structural insights with in-source fragmentation, to facilitate the confirmation of molecular weights for unknown impurities, offering deeper understanding into compound structures
- Safeguard against biases in reporting of known impurities, with valuable insights from mass detection enabling interference detection, preserving product quality, and expediting product release timelines

In conjunction with Empower CDS software, this analytical approach further elevates the impurity workflow, delivering simplicity, efficiency, and upholding compliance and data integrity standards.

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