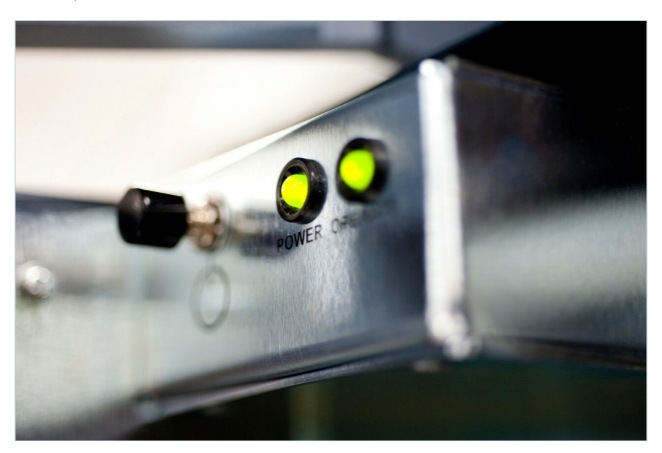
# Waters™

응용 자료

# Future-proofing the QC Laboratory for UPLC While Enabling the Faithful Analysis of Legacy HPLC Methods

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#### **Abstract**

This application note describes how the AstraZeneca QC department in Macclesfield has successfully transferred and run all registered QC methods on the Waters ACQUITY UPLC H-Class System with three high throughput products that were successfully developed and validated for UPLC.

#### **Benefits**

- The capability to faithfully and robustly run legacy HPLC methods while providing access to true UPLC technology when desired
- No modifications required to provide seamless development, transfer and implementation of UPLC methods
- Unprecedented efficiency savings along with solvent usage costs and corresponding solvent disposal costs

#### Introduction

In this application note, we describe how the AstraZeneca QC department in Macclesfield (a global center for developing new technologies for QC) has successfully transferred and run all registered QC methods on the Waters ACQUITY UPLC H-Class System with three high throughput products that were successfully developed and validated for UPLC. In efforts to update and modernize their labs, it was critical for Astra Zeneca to ensure new technology would be efficient, easy to adopt, and cost effective. AstraZeneca updated their LC platforms by implementing Waters UPLC in their pharmaceutical development department with the intention of developing all new products on this platform. While future-proofing the QC department to receive newer UPLC methods, it was critical to retain the ability to faithfully and robustly run legacy chromatography methods. The technology of choice was the ACQUITY UPLC H-Class System, which has now been deployed throughout the AstraZeneca QC department based at Macclesfield, UK.

Within this body of work, we will give an example of a high profile compound 'B' legacy HPLC method transferred from an Agilent 1100 to a Waters ACQUITY UPLC H-Class System along with the newly developed UPLC method validated on the same instrument.

# Experimental

The new UPLC method for compound B degradant products was created using the ACQUITY UPLC Columns Calculator, to simplify transfer and scale HPLC methodology quickly to UPLC conditions with equivalent performance (with significantly reduced runtimes and solvent savings) ensuring it satisfied the system suitability criteria stated in the legacy HPLC method.

Impurities 1 and 2 of compound 'B' were validated over a range of 50% to 200% of their respective specification limits in the presence of the main compound 'B'.

#### HPLC conditions (Agilent 1100 or Waters ACQUITY UPLC H-Class)

Column:	$C_8$ 4.6 mm $\times$ 250 mm, 5 $\mu$ m
Flow rate:	1.3 mL/min
Injection volume:	50 μL
Run time:	30 min
Detection:	UV

#### UPLC conditions (ACQUITY UPLC H-Class equipped with ACQUITY TUV Detector)

Column:	Waters ACQUITY UPLC BEH 2.1 mm $\times$ 100 mm,
	1.7 μm Column
Flow rate:	0.3 mL/min
Injection volume:	4.2 μL
Run time:	6.86 min

#### Data management

### Results and Discussion

For the ACQUITY UPLC H-Class to be a successful forward facing platform for the quality control environment, it must first be able to faithfully and robustly reproduce the chromatography generated on the laboratory's existing HPLC platform. Figure 1 shows the comparison of Compound B's system suitability sample (SST) run on the Agilent 1100 (top), the Waters ACQUITY UPLC H-Class System in HPLC mode (middle), and the ACQUITY UPLC H-Class System again using the newly developed UPLC method (bottom).

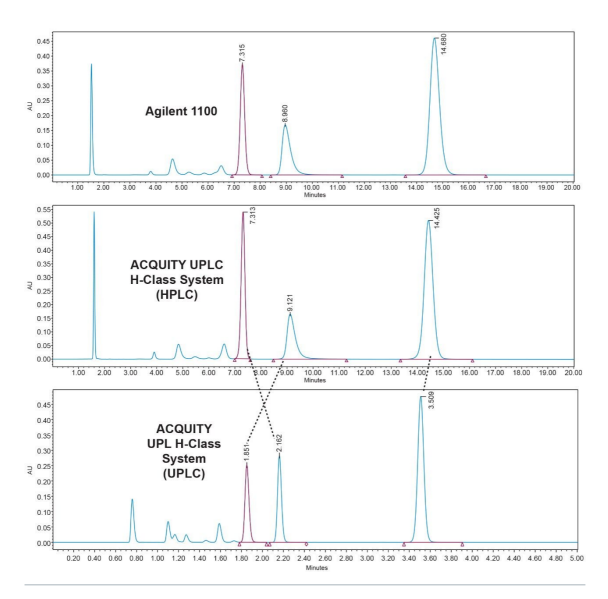


Figure 1. Comparison of Compound 'B' SST sample run on Agilent 1100 HPLC (top), Waters

ACQUITY UPLC H-Class System in HPLC mode (middle), and Waters ACQUITY UPLC H-Class in

UPLC mode (bottom).

The ACQUITY UPLC H-Class System has reliably replicated the chromatography from the Agilent 1100 and reproduced the relative retention times (RRT's) of impurities 1 and 2 with respect to the main peak, as shown in Table 1. In HPLC mode, the ACQUITY UPLC H-Class System also consistently reproduced the peak areas of Compound B and its related impurities compared to those obtained on the legacy LC system, as shown in Table 2.

Compound B	Main peak RT (mins)	Impurity 1 RT (mins)	Impurity RRT wrt main peak	Impurity 2 RT (mins)	Impurity RRT wrt main peak	
Agilent 1100	14.680	7.315	0.500	8.960	0.610	
ACQUITY UPLC H-Class System <b>HPLC</b> mode	14.425	7.313	0.510	9.121	0.630	
ACQUITY UPLC H-Class System <b>UPLC</b> mode	3.509	2.162	0.620	1.851	0.530	

Table 1. Retention times / relative retention times of compound 'B' and impurities 1 and 2 generated using the legacy method on the Agilent 1100 and the Waters ACQUITY UPLC H-Class System, along with the newly developed UPLC method results obtained from Waters ACQUITY UPLC H-Class System.

Compound B	ompound B Main peak area		Relative peak area wrt main peak	Impurity 2 peak area	Relative peak area wrt main peak	
Agilent 1100	11891513	4462703	0.38	3896619	0.33	
ACQUITY UPLC H-Class System <b>HPLC</b> mode	11094170	4436024	0.40	3477557	0.31	

Table 2. Relative peak areas with respect to the main peak of impurities 1 and 2. Results show consistent relative areas between the Agilent 1100 and the Waters ACQUITY UPLC H-Class System in HPLC mode.

The UPLC method had a runtime of under seven minutes compared to the legacy method runtime of 30 minutes. There is also a marked improvement in peak symmetry. Impurities 1 and 2 have switched elution order, although this has not compromised system suitability criteria as shown in Figure 1.

#### Validation

Once the newly developed UPLC method for Compound B degradants had satisfied system suitability criteria, the method was subject to a partial validation based on ICH Guidelines Q2<sup>1</sup> covering linearity, recovery, repeatability, and limits of detection and quantitation (LOD and LOQ respectively).

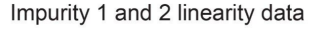
The range of the the validation covered 50% to 200% of the impurities respective specification limits (this exceeds the ICH Guideline's suggestion of 70% to 130% for added assurance of method robustness).<sup>2</sup>

Table 3 summarizes the validation data obtained.

Raw linearity data is presented in Figure 2 and Table 3, method precision data in Table 4, and impurity 1 and 2 recovery raw data is presented in Tables 5 and 6 respectively.

Conc Impurity 1 (µg/mL)	Conc Impurity 2 (μg/mL)	Peak area Impurity 1	Peak area Impurity 2
0.0822	0.1315	709	2114
0.1233	0.1972	1111	3205
0.1644	0.2630	1463	4295
0.2466	0.3944	2216	6418
0.3288	0.5260	2884	8520

Table 3. Linearity raw data.



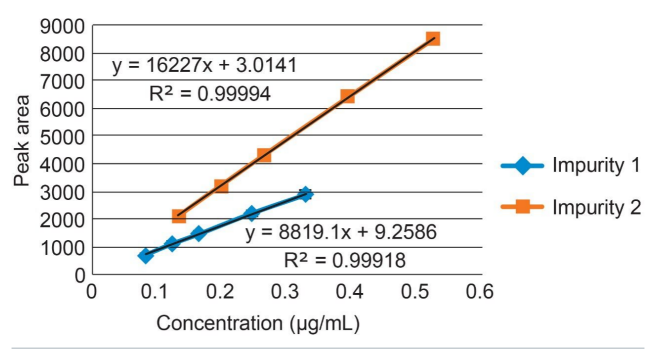


Figure 2. The UPLC method for Compound 'B' degradants comfortably satisfied the acceptance criteria with linearity. Linearity was performed over the range of 50% to 200% of respective specification limits of impurities 1 and 2.

Peak area Impurity 1	Peak area Impurity 2	
1446	4261	
1448	4271	
1453	4334	
1447	4286	
1493	4342	
1472	4276	
1460	4295	Mean
18.93	34.36	SD
1.30	0.80	%RSD

Table 4. Performed using six separate preparations of Compound 'B' standard spiked with impurities 1 and 2 at their respective impurity limits (figures adjusted for background impurity content in standard).

Recovery

# Impurity 1 recovery data

Mean % Recovery	% Recovery	Mean peak area 100% (n=6)	μg/mL	Peak area	% of Nominal
	98.219	1460	0.0822	717	50
	99.178	1460	0.0822	724	50
98.8	98.904	1460	0.0822	722	50
	100.411	1460	0.1644	1466	100
	103.630	1460	0.1644	1513	100
	99.178	1460	0.1644	1448	100
	98.973	1460	0.1644	1445	100
	100.479	1460	0.1644	1467	100
100.8	101.849	1460	0.1644	1487	100
	100.205	1460	0.3288	2926	200
	103.904	1460	0.3288	3034	200
101.6	100.822	1460	0.3288	2944	200
	100.500	Overall mean			
	1.800	SD			
	1.800	%RSD			

Table 5. Recovery data for impurity 1 covering 50% to 200% of the range of specification limit. The 50% and 200% levels were prepared in triplicate and the 100% level n=6 (100% data also used for precision).

#### Impurity 2 recovery data

Mean % Recovery	% Recovery	Mean peak area 100% (n=6)	μg/mL	Peak area	% of Nominal
	100.210	4295	0.1315	2152	50
	98.393	4295	0.1315	2113	50
98.4	96.717	4295	0.1315	2077	50
	99.208	4295	0.2630	4261	100
	99.441	4295	0.2630	4271	100
	100.908	4295	0.2630	4334	100
	99.790	4295	0.2630	4286	100
	101.094	4295	0.2630	4342	100
100.0	99.558	4295	0.2630	4276	100
	99.371	4295	0.5260	8536	200
	98.987	4295	0.5260	8503	200
99.2	100.210	4295	0.5260	8561	200
	99.4	Overall mean			
	1.2	SD			
	1.2	%RSD			

Table 6. Recovery data for impurity 2 covering 50% to 200% of the range of specification limit. The 50% and 200% levels were prepared in triplicate and the 100% level n=6 (100% data also used for precision).

	Runtime/month (hours)			Solvent volume/month (litres)			Solvent cost/month					
	HPLC	UPLC	% Save	Actual saving	HPLC	UPLC	% Save	Actual saving	HPLC	UPLC	% Save	Actual saving
Content	120.0	27.4	77.2	92.6	9.36	0.49	94.9	8.87	£30.51	£1.60	94.8	£28.91
Dissolution	81.7	14.9	81.8	66.8	6.37	0.72	88.7	5.65	£20.77	£2.34	88.7	£18.43
Assay/ related substances	140.0	32.0	77.1	108.0	10.92	0.58	94.7	10.30	£35.60	£1.89	94.7	£33.71
Total				267.4				24.82				£81.05

Table 7. Estimated workflow efficiency and solvent cost savings with the implementation of UPLC Technology for all Compound 'B' methods based on AstraZeneca batch throughput.

# Conclusion

The UPLC data detailed for Compound 'B' shows a time savings of between 77.1% to 81.8% equating to over 267 hours per month with solvent savings between 88.7% to 94.8% per month. This not only impacts solvents costs associated with purchase and disposal, but also reduces the need for large storage volume impacting space savings and health and safety.

The Waters ACQUITY UPLC H-Class System's success in transitioning legacy methods within the Quality Control environment of AstraZeneca exemplifies the instrument's ability to offer a seamless alternative to existing HPLC platforms while uniquely offering the option of true UPLC Technology when desired.

AstraZeneca have successfully run all registered QC methods on the ACQUITY UPLC H-Class System with three high throughput products transferred and validated successfully using UPLC Technology.

The success of the Waters ACQUITY UPLC H-Class System in the Quality Control department of AstraZeneca Macclesfield has led to a wider adoption globally of the ACQUITY UPLC H-Class System by AstraZeneca.

#### References

- 1. ICH Guidelines: Validation of analytical procedures: Text and methodology Q2(R1).
- 2. USP General Chapter, <621> Chromatography, USP36-NF31, The United States Pharmacopeia Convention, official December 1, 2013.

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