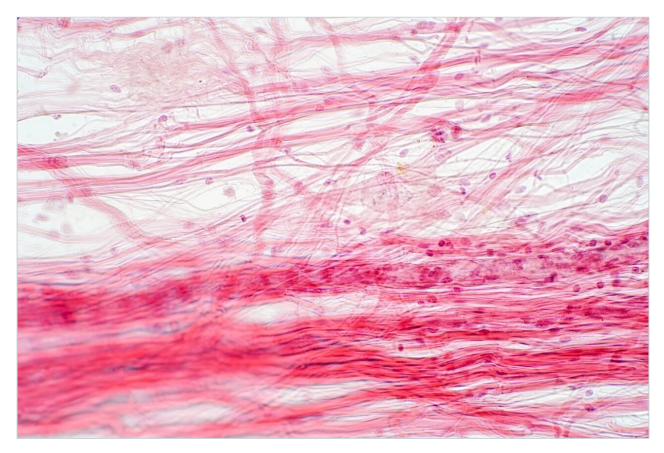
# Waters<sup>™</sup>

Application Note

# Rapid Analysis of Pharmaceuticals in Human Tissues Using the ACQUITY UPLC with 2D Technology

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For forensic toxicology use only.

#### Abstract

This application demonstrated the automated and fast method development capability of the ACQUITY UPLC with 2D Technology for the analysis of pharmaceuticals in human tissue samples.

#### Benefits

- · Fast extraction protocol (45 min)
- · Trace level detection (ppt)
- · 90 sec homogenization

### Introduction

According to the Scientific Working Group for Forensic Toxicology (SWGTOX), the field of forensic toxicology handles the analysis of drugs or chemicals in biological materials, and the interpretation of those results for medico-legal purposes.<sup>1</sup> In this field, forensic toxicologists often work with medical examiners to perform postmortem toxicological analyses on blood or biological tissues of deceased individuals in order to determine cause and manner of death.<sup>1</sup> Because these results are relied upon in a court of law, validity, reliability, accuracy and precision of the analytical techniques used to perform these analyses are essential.

The core focus of a forensic toxicology laboratory is the accurate identification and quantitation suspected drugs or chemicals in biological samples. The target matrix can vary between blood, plasma, urine, saliva, vitreous fluid, hair, nails, and organs such as brain, heart, lung, liver, kidney, spleen, and stomach contents. The Forensic Toxicology Research Team at the Federal Aviation Administration performs such analyses on samples from victims of fatal aircraft accidents to provide insight to the analysis of accident causation.<sup>2</sup> Aircraft accidents and crashes are often brutal enough to severely impair any human remains, which is why the toxicologists must rely on more complex biological tissues for analysis, i.e. brain, heart, lung, liver, kidney, spleen, etc. Additionally, they must have the ability to detect and measure many substances, from drugs and alcohol, to toxic gases and industrial chemicals.<sup>2</sup> Therefore, there is a need to develop multi-residue analyses and efficient sample preparation methods in order to analyze samples in a timely manner.

The analytical techniques currently available are divided into two categories, some platforms are used for screening methods (qualitative) and other solutions are used for confirmation methods (quantitative). Most

laboratories are usually equipped with gas chromatography (GC) or liquid chromatography (LC) hyphenated to a mass spectrometer (MS). For several decades, GC-MS was the tool of choice for bio-analysis. With the introduction of atmospheric pressure ionization technique, LC-MS is now the most popular technique in the field of forensic toxicology.

Detection and quantification of drugs in complex matrices is difficult to accomplish due to time-consuming extraction processes, and the difficulty to detect an analyte at trace levels. A robust extraction and clean up methodology, in which a homogenization step precedes, is a must in order to reach a target limit of detection (LOD) and to maintain instrument performance. The use of advanced hyphenated instrumentation platforms, such as UPLC-MS/MS has allowed analysts to detect trace levels of analytes. Traditional extraction techniques used in most laboratories are decades old and do not have the robustness to produce quality results. A micro extraction protocol combined with a multi-dimensional chromatography (2D LC-MS/MS) can decrease sample preparation time without sacrificing the quality seen with current single dimension chromatography techniques.<sup>3,4,5</sup>

## Experimental

Two MRM transitions (quantification and confirmation) for all drugs were selected and optimized. The MRM conditions are listed in Table 1. All human biological specimens used for this study were provided by the Federal Aviation Administration (FAA).

	lon Mode	Precursor ion	Cone	Product ion	CE
Zalaidam	ESI	200.2	40	235.3	35
Zolpidem	ESI+	308.3	40	263.3	25
Citolapram	ESI+	325.4	30	109.1	25
Citolapiani	LOIT	525.4	30	262.4	20
Norbuprenorphine	ESI+	414.6	30	101.2	45
Norbuprenorphille	LJIT	414.0	50	57.2	50
Oxycodone	ESI+	316.2	30	298.2	20
Oxycodone	LJIT	510.2	50	241.2	30
Normonoridino	ESI+	234.2	30	160.2	15
Normeperidine	ESIT	234.2	30	56.2	25
Dextrorphan	ESI+	258.3	30	157.3	35
Dextrorphan	LOIT	200.0	30	133.1	30
Doutromothorphon	ESI+	272.2	30	147.2	30
Dextromethorphan	ESIT	212.2	30	215.2	25
Diazanam	EQL	285.1	20	154.1	25
Diazapam	ESI+		30	105.1	25
Diltiorom	ESI	415 2	20	178.2	25
Diltiazem	ESI+	415.3	30	370.2	15
Quotionino	ESI+	384.5	30	253.3	25
Quetiapine	ESIT	304.0	30	221.3	30
Diphenydramine	ESI+	256.4	30	167.2	10
Dipitettyurainine	LOIT	200.4	30	152.2	30
Buprenorphine	ESI+	468.4	30	55.3	45
Duprenorphille	LOIT	400.4	30	84.3	40
Promethazine	ESI+	285.4	30	86.2	20
FIOITIettildzitte	LOIT	200.4	30	198.2	20
Dibudrocodoine	ESL	202.2	20	171.1	40
Dihydrocodeine	ESI+	302.3	30	199.2	35
Dovulomine	EQL	071.0	20	182.2	15
Doxylamine	ESI+	271.3	30	167.2	25
Floooloida	EQ.	415.0	20	98.1	25
Flecainide	ESI+	415.3	30	398.4	25
Uudeomoushaua	EQ1.	206.0	20	157.3	40
Hydromorphone	ESI+	286.2	30	185.1	30
Nordiaranam	ECL	071 1	20	140.2	25
Nordiazepam	ESI+	271.1	30	200.0	05

Loading conditions Loading: MilliQ Water (pH 7) Flow rate: 2 mL/min AT-column dilution: 5% (0.1 mL/min Loading pump and 2 mL/min Diluting pump)

## UPLC conditions

UPLC system:	ACQUITY UPLC with 2D Technology configured for "Trap and Elute" with AT-column dilution
Runtime:	10 min
Column:	ACQUITY UPLC BEH C <sub>18</sub> , 2.1 x 50 mm, 1.7 $\mu$ m
Column temp.:	60 °C
Mobile phase A:	Water + 0.5% formic acid
Mobile phase B:	Acetonitrile + 0.5 % formic acid
Elution:	5 minute linear gradient from 5% (B) to 95% (B)
Flow Rate:	0.500 mL/min (Elution pump)
Injection volume:	100 µL
MS conditions	

MS System:	Xevo Q-ST TQ-S

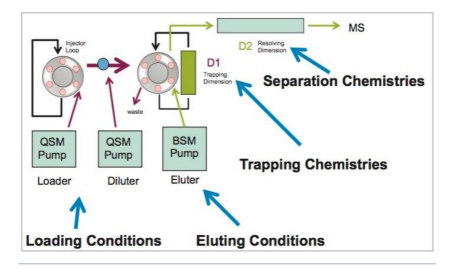
Ionization mode:	ESI Positive
Capillary voltage:	3.0 kV
Cone voltage:	90.0 V
Source temp.:	150 °C
Desolvation temp.:	550 °C
Desolvation gas:	1100 L/hr
Cone gas:	50 L/hr

# Results and Discussion

#### 2D LC method development

The analysis of started with the chromatography optimization of the 2D LC-MS/MS. The 2D LC-MS/MS is setup as depicted in Figure 3. This configuration was constructed with two quaternary pumps and one binary pump. The binary pump was set for gradient elution and the quaternary pumps were plumbed for "AT-column dilution" to create two distinct streams (loader and dilutor). The loader pump was set 0.1 mL/min for loading the extracts from the injection loop onto a 50  $\mu$ L mixer, while the dilutor pump was set at 2 mL/min flow rate for dilution following a re-focusing effect on the trap column. From the chemical structures of the target analytes, a high retention strength sorbent material (Oasis HLB, 40mg) was selected for the trap column, while a high XBridge Hybrid C<sub>18</sub> sorbent (BEH C<sub>18</sub>) was chosen for the analytical column. The next phase of the optimization was to select the trapping and elution conditions. As seen in previous publications, a 6x6 2D LC evaluation grid gives an excellent starting point to provide an overview of the chromatographic behavior for a target analyte. For this application, the 2D LC optimization process focused with methods 3, 6, 9, 12, 15, and 18. The results are tabulated in Table 2. The color coded chart was created to identify which analytical conditions give the best chromatographic profile with a quick visual survey. The green box depicts a Gaussian peak shape for quantification analysis. The yellow box was used to flag chromatography issues, such as peak split, tailing, shoulder or leading profiles. Finally, the red box indicates an absence of signal,

most likely due to breakthrough effect during loading phase on the trap column or poor elution from the trap onto the analytical column. Additional parameters can be adjusted to ensure proper mass transfer during loading and elution phase. One parameter in particular is the sorbent bed mass on the first dimension. Two sorbent bed masses (40 mg vs 80 mg) were evaluated for the retention and elution of the target analytes. As shown in Table 2, method 9 using an HLB 80 mg bed mass and method 6 using HLB 40 or 80 mg provided the best chromatography performance for all 21 target analytes.



*Figure 3. 2D LC configuration with AT-column dilution (3 pumps design).* 

	Meth 3	Meth 3	Meth 6	Meth 6	Meth 9	Meth 9	Meth 12	Meth 12	Meth 15	Meth 15	Meth 18	Meth 18
	40 mg	80 mg	40 mg	80 mg	40 mg	80 mg	40 mg	80 mg	40 mg	80 mg	40 mg	80 mg
	HLB	HLB	HLB	HLB	HLB	HLB	HLB	HLB	HLB	HLB	HLB	HLB
Normeperidine	pH 3	pH 3	pH 7	pH 7	pH 10	pH 10	pH 3	pH 3	pH7	pH 7	pH 10	pH 10
(water)	-67	e7	87	87	87	87	e6	e6 tail	87	e6 tail	e7	eî
Normeperidine (MeOH)	67	87	87	87	e7	e7	e6 tail	e6 tail	e7	e6 tail	e7	e7
Normeperidine (ACN)	e7	88	87	eð	e7	e6	26	e5 tail	e6 tail	e6 tail	86	26
Dextrorphan (water)	e7	e7	e7	e7	e7	e7	e6 tail	e6 tail	86	e6 tail	67	e6 tail
Dextrorphan(MeOH)	e7	e7	87	e7	e7	e7	e6 tail	e6 tail	87	e6 tail	87	e7 tail
Dextrorphan (ACN)	.e6	86	e7	eô	e7	86	eő	e5 tail	e6 tail	e6 tail	eô	e6 tail
Nordiazepam (water)	e7	86	e7	e6	e7	96	e7	87	87	67	87	e7
Nordiazepam (MeOH)	e7	86	87	e6	e7	86	e7	97	87	ę7	e7	e7
Nordiazepam (ACN)	97	86	e7	e6	e7	88	67	e7	87	e7	67	87
Dextromethorphan (water)	67	87	26	66	e7	67	26	e6 tail	86	mp e6 tail	88	26
Dextromethorphan (MeOH)	07	87	87	e7	87	-07	67	e7 tail	e7	e7 tail	e7	=1
Dextromethorphan (ACN)	67	e?	ef	e7	e7	67	e6	e6 tail	87	e6 tail	86	26
Diazepam (water)	mp.e7	mp e6	e7	86	e7	66	mp e7	mp e7	67	e?	67	el
Diazepam (MeOH)	mp e7	mp e6	mp e7	mp e6	mp e7	mp e6	mp e7	mpie7	e7	26	e7	e7
Diazepam (ACN)	mp e6	mp e6	e7	mp e6	e7	e6	mp e7	mp e7	87	86	87	87
Promethazine (water)	67	87	e7	e6	87	87	87	e6	88	86	e6	e6
Promethazine (MeOH)	e7	e7	e7	e7	e7	97	e7	-97	87	e7	67	et
Promethazine (ACN)	97	96	e7	86	e7	e6	26	e6	96	86	e6	mp e6
Oxazepam (water)	87	mple6	87	66	67	86	mp e7	mpel	67	mp e7	67	e7
Oxazepam (MeOH)	187	mp e6	87	66	67	66	mp e7	mp:e7	87	e7	e7	el
Oxazepam (ACN)	e7	mp e6	e7	66	e7	66	mp e7	mpie7	-87	67	e7	el
Termazepam (water)	87	e7	67	66	87	eð	87	87	e7	đ	67	e7
Termazepam (MeOH)	e7	mp.e6	e7	mpie6	e7	mp e6	mp.e7	mp a7	87	el.	67	e7
Termazepam (ACN)	e7	mp e6	e7	mp.e6	e7	mp:e6	mp e7	mpel	-87	67	07	e7
Flecainide (water)	- 07	e7	87	e7	<u>e7</u>		67	67	87	96	e7	87
Flecainide (MeOH)	87	87	87	87	87	- 97	87	87	87		67	e7
Flecainide (ACN)	e7	87	87	87	e7	197	67	86	47	e7	87	87
Diphenhydramine (water)	86	87	e6	e6	e6	e6	26	e6	89	26	66	65
Diphenhydramine (MeOH)	e7 e7	e7	87	el.	87	e7	8/	66	87	86	67	e/
Diphenhydramine (ACN)		e7	87	e7.	e7	87	86	66	87	e6	67	26
Hydromorphone (water)	mp e5	25	86	e6	en .	86	85	65	87	26	66	26
Hydromorphone (MeOH) Hydromorphone (ACN)	sp e5	20	26 26	06 06	e6 tail sp e5	86. 85	27	94	e6	86	86 86	e6 #5
Dihydrocodeine (water)	26	26	86	86	86	86	all	eß	67	86	67	26
Dihydrocodeine (MeOH)		e6	e6	66	87	e7	aß	26	87	63	e7	a7
Dihydrocodeine (ACN)	e4	mp e4	86	e6	86	86	E.	cu	e6		66	66
Zolpidem (water)	e7	e/	ef	e7	67	e7	e7 tail	97	97	ed.	97	87
Zolpidem (MeOH)	e7	e7	e7	e7	e7	- 19	07	e7	87	67	67	e7
Zolpidem (ACN)	117	87	e7	27	el	el	e7 tail	e7			67	01
N Desmethyl citolapram (water)	mp e7	mp e7	mp e6	mp e6	87	mp e7	mp e6 tail	mp e6	mp e6 tail	mp e6	mp e6 tail	mp e6 tail
N Desmethyl citolapram (MeOH)	mp.e7	mp e7	e7	e7	87	e7	mp e6 tail	mp e6	e7 tail	mp e6	e7 tail	mp e6 tail
N Desmethyl citolapram (ACN)	mp:e7	mp e7	e7	e7	e7	e7	mp e6 tail	mp e6	e7 tail	mp e6	mp e7 tail	mp e6 tail
Oxycodone (water)	e6	mp e6	efi	e6	87	mp e7	mp.a6	mp e6 tail	e6	mp e6	mp.e7	mp e7
Oxycodone (MeOH)	e6	mp e6	eű	e6	e7		mp e6	mp e6 tail		69	mp e7	mp-e7
Oxycodone (ACN)	mp e5	mp e6	e6	66	e7	mp e7	mp e6	mp e6 lead	e6	26	mpie7	mp e7
Citolapram (water)	mp.e7	mp e7	eő	mp.e6	e7	e7	mp.e7	mp.e6	e6	mp.e6	07	67
Citolapram (MeOH)	mp e7	mp e7	e7	e7	e7	87	mp e7	mpie7	87	e7	67	27
Citolapram (ACN)	mp.e7	mp e7	e7	e7	e7	-87	mg e/	mpie7	đ	e7	e7	e7
Quetiapine (water)	e7	e7	e7	eß	87	e7	e7	e7	ø	ø	e7	e7
Quetiapine (MeOH)	07	e6	67	86	87	e7	e7	e7	e7	e7	e7	e7
Quetiapine (ACN)	e6	86	e6	86	87	86	e6	e6	87	86	67	26
Norbuprenorpine (water)	mp eS	mp.e6	mp e5	mp.e5	mpleS	mp.e5	mp e5	mp e5	mple5	mpie4	mp-e5	mp e5
Norbuprenorpine (MeOH)	mp e5	mp e5	mp e5	mp-e5	mp.e5	mp e6	mp e5	mp e5	mp e5	mp e5	mp e5	mp eā
Norbuprenorpine (ACN)	mp e5	mp e5	mp e5	mp e5	mp.e5	mp e5	mp e5	mp e5	mp e5	mp e5	mp e5	mp e5
Diltiazem (water)	67	87	e7	ଶ	67	e7	el	e7	87	e7	e7	87
Diltiazem (MeOH)	89	e7	88	87	68	67	67	e7	66	67	e7	e7
Diltiazem (ACN)	e7	e7	87	e7	89	67	e7	e7	e7	67	e7	67
Buprenorphine (water)	e6	86	e6	85	e6	e6	e6	e6	85	85	eá	e6
Buprenorphine (MeOH)	e6	e6	86	e6	96 86	-06	efi	e6	e6	eő	85	e6
Buprenorphine (ACN)	66	86	e6	66		86	96			66	26	

#### Table 2. 6x6 grid results.

The rationale behind the selection of Method 6 related to the fact that the loading conditions for the target analytes on the trap column can be done at pH 7, while Method 9 utilize a high pH additive ( $NH_4OH$ ). Therefore, as cost saving measures, the final protocol will use a pH 7 loading onto an 40 mg HLB on the first dimension, followed by an elution with acetonitrile at pH 3 onto a BEH C<sub>18</sub> analytical column (See Figure 4).

The final separation showed excellent Gaussian peak shapes for all analytes. However water spikes exhibited lower intensities, which is expected due to secondary interactions with the active sites, most likely due to ion exchange retention with the glass vial surface. The ionic interaction can be eliminated by simply changing the diluent. In this case, methanol and acetonitrile diluents both gave higher intensities (See Figure 5).

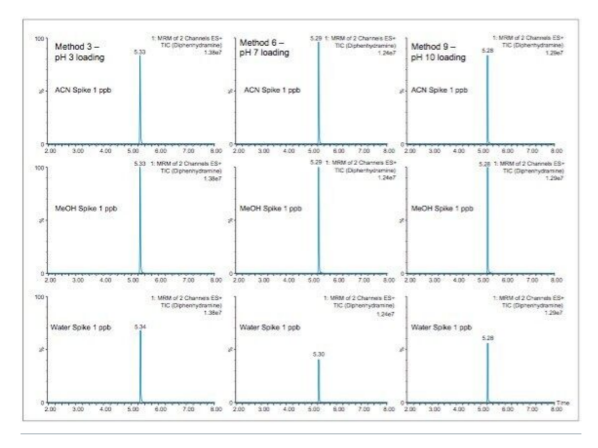


Figure 4. Method 6 chromatogram at 1 ppb in acetonitrile.

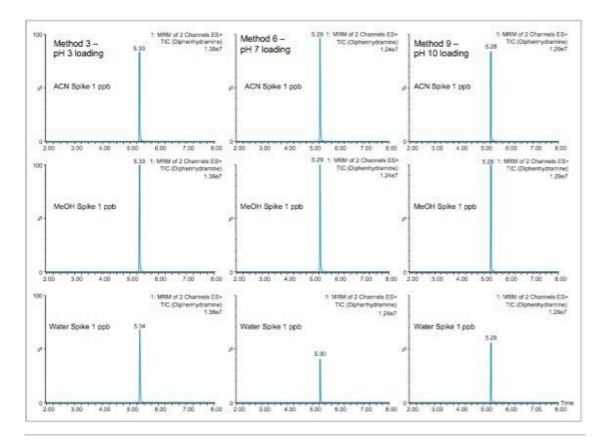


Figure 5. Results for method 3, 6, and 9 with 40 mg HLB bed mass for Diphenhydramine.

#### SPE extraction evaluation

After selecting the optimum 2D LC conditions, the work focused on the extraction optimization. The first step of the process targeted the choice of the sorbent. In this scenario, a mixed mode sorbent (Reversed Phased/Cation Exchange, Oasis MCX) was selected since all target analytes contain an amine functionality in their chemical structures. Hence, the evaluation started with two sorbent masses (60 and 150 mg) as presented in Table 3. The workflow started by loading a 2 mL water spike at 1 ppb and proceeded with a pH 3 water wash to ionize the basic compounds so they are captured onto the cation exchange portion of the sorbent. With target basic analyte secured, the reversed phase portion of the sorbent was eluted with a pH 3 high organic solvent wash. In this instance, a 100% Methanol with 2% formic acid was used for the secondwash. The elution of the basic analyte was performed with 100% acetonitrile with 2% ammonium hydroxide. The high pH value neutralizes the amine functionality, thus releasing all basic analytes from the cation exchange sorbent. The last wash and the final elution were collected to monitor if all analytes were in fact retained as predicted. As seen in table 2, the 60 mg sorbent bed showed signs of breakthrough for oxazepam, temazepam and N Desmethylcitolapram. When compared to a 150 mg sorbent bed, oxazepam and temazepam exhibited no breakthrough during the Methanol wash. However at this point in the

evaluation, it was clear that several analytes exhibited poor recovery with the MCX cartridge. These issues were resolved by selecting a mixed mode sorbent with a reversed phased portion and a weak cation exchange portion (Oasis WCX). The methodology is similar, but the wash step and elution are governed by pH to ionize or neutralize the weak cation exchanger on the sorbent, as opposed to the analyte itself. The side by side comparison between MCX and WCX is presented in Table 4. As shown, Normeperidine, Dextrorphan, Dextromethorphan, N-Desmethylcitolapram and Norbuprenorphine, all show poor recoveries when using a strong cation exchanger. For two analytes, the results show a 10x signal difference between MCX and WCX. For those problem analytes, a dual methodology was crafted and two target analytes were used as markers (Citolapram and Diphenhydramine) for recovery evaluation purpose.

	MCX 60 mg		MCX 60 mg		MCX 150 mg		MCX 150 mg	
1.7	Wash MeOH 2% FA	Mean	Elution ACN 2% NH <sub>4</sub> OH	Mean	Wash MeOH 2% FA	Mean	Elution ACN 2% NH_OH	Mean
	Area	mean	Acrea	Weall	Area	wiean	Acrea	wear
Normeperidine	5441		282621		3217		114004	
	4525		352711		4399		142777	
	5545	5170	348768	328033	2475	3364	142185	132989
		1.57%	10000000			2.47%		low Re
Dextrorphan	1922		228514 280310		4663		17365 21185	
	1880	1926	284213	264346	4103	4696	21618	20056
	1000	0.72%	CONCIO	A CONTRACTOR	1100	18.97%	21010	low Re
Nordiazepam	453		203376		627		227462	
	594		252200		452		246626	
	477	508	255220	236932	604	561	258576	24422
Dovalamina	5196	0.20%	660525	0-01211-02000	5848	0.23%	654529	
Doxylamine	6600		668535 773747		5998		765944	
	6939	6245	762365	734882	4881	5576	777176	73255
		0.84%				0.76%		
Dextromethorphan	5994	10/251/00/00	406573		5046	10000000	295028	
	5225		551406		4873		395256	
	4944	5388	561553	506511	5144	5021	419907	37006
Diazonam	1978	1.06%	177922		1325	1.35%	203456	low Re
Diazepam	1539		223835		1871	200000	203456	
	1476	1664	227167	209641	1672	1623	228516	21880
Section Street Contractor	1000000	0.79%	100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100 - 100		Show they	0.74%	0.00.0003512-0	
Promethazine	7934	2.531025007680	451453		8713	8700001002	461876	
	7232	7770	642717	500500	9263	0105	642419	an second
	8153	7773	677627	590599	9520	9165 1.53%	688151	09148
Oxazepam	51897	1.3170	122881		4089	1.3370	264526	
2011 10 10 10 10 10 10 10 10 10 10 10 10	40530		121777		3542		312994	
	49072	47166	124248	122969	4709	4113	327211	30157
	Breakthrough	27.28%		low Rec		1,35%		
Temazepam	56319		74337		3587		510018	
	58663 53763	56248	74699 75706	74914	3263 3212	3354	551183 548403	63653
	Breakthrough	42.88%	13/00	low Rec	3212	5.88%	348403	200.20
Flecaninide	3530	1210010	564321	10111100	3978	0.0070	593246	
	4794		735545		3331		812857	
	3264	3863	767391	689086	3855	3721	852158	75275-
O'		0.53%		- and the second second	10000200	0.49%		
Diphenhydramine	1347		130365		6193		126349 166090	
	1174 1626	1382	165624 169623	155204	7741 6314	6749	166090	15.517
	1020	0.89%	103023	155204	0314	4.17%	1/30/3	15050771
Hydromorphone	347	107702021.040V	42595		150	0004950250	78233	
	210		44585		132		80272	
	154	237	45313	44164	164	149	80441	79649
Dihudaa adalara	151	0.53%	100.000			0.19%	101024	
Dihydrocodeine	451 417		193420 202722		446 415		161234 172063	
	314	394	209123	201755	196	352	171609	16830
3020000000		0.19%	200.20			0.20%		low Re
Zolpidem	12152		1299467		2123		1267340	
20.	13871		1389542		2869		1366250	
	10986	12336	1419628	1369546	2672	2555	1380714	133810
DesmethylCitolapram	67930	0.89%	261488		45163	0.19%	25522	
beamethylontolapram	75121		356963		49196		35525	
	72058	71703	388218	335556	47818	47392	35001	32016
	Breakthrough	17.61%	KibridantoVi - 1		Breakthrough	59.69%	2010220	low Re
Oxycodone	539		397220		1316		370819	
C Sheri wa sh	534		420253		1238	100.	398264	2020
	530	534	427808	415094	1629	1394	402835	39063
Citolapram	5931	0.13%	550657		4589	0.36%	521151	
onolapiani	6539		731095		5966		702231	
	6803	6424	773777	685176	6610	5722	734192	65252
		0.93%				0.87%		
Quetiapine	9066		516308		12078		450202	
-1099/ALTE 154:200/6032	10734	11105	617433	507657	11410	10016	533633	T D D D D D D D D D D D D D D D D D D D
	13784	11195 1.84%	659231	597657	15261	12916 2.48%	540583	508 3
Norbuprenorphine	253	1.04%	13998		152	2.4070	8774	
Norbaprenorphine	203		16520		188		11343	
	274	248	17578	16032	150	163	11129	10415
		1.52%				1.54%		low Re
Diltiazem	19983		1701151		19237		1692454	
	19711		2107899		20532		2087306	
	19746	19813	2134585	1981212	20085	19951	2127205	196898
Bupreporchine	1310	0.99%	48300		105	1.00%	12476	
Buprenorphine	1310		46300 61701		195		43476 58034	
	1338 1491	1380	62986	56996	144	175	58454	54901

Table 3. Recovery values for MCX 60 mg versus MCX 150 mg cartridge.

	WCX 100 mL extracts	WCX 100 mL extracts	MCX 100 mL extracts	MCX 100 mL extracts	
	wash - MeOH + FA	elution - ACN + FA	wash - MeOH + FA	elution - ACN + NH <sub>4</sub> OH	
Normeperidine	10043	472271	1085	143796	
Dextrorphan	72	256835	178	12510	
Nordiazepam	129466	15397	204	196527	
Doxylamine	1049	129629	56	298312	
Dextromethorphan	149	348720	729	278889	
Diazepam	124873	25360	212	336645	
Promethazine	610	539304	1817	718461	
Oxazepam	134574	12057	522	113356	
Temazepam	11833	32516	0	525964	
Flecainide	1020	444358	598	549411	
Diphenhydramine*	0	377237	4084	458358	
Hydromorphone	0	4882	0	6504	
Dihydrocodeine	0	21500	0	23872	
Zolpidem	31687	692233	135	1291959	
Ndesmethylcitalopram	0	409467	0	60843	
Oxycodone	0	172149	0	181488	
Citalopram*	1492	796273	4150	1447050	
Quetiapine	652	430552	0	497057	
Norbuprenorphine	134	11497	0	8789	
Dilitazem	51	2726724	27332	3279491	
Buprenorphine	1430	18819	933	41542	

Table 4. Recovery values for WCX 150 mg vs MCX 150 mg cartridge.

The next phase of the application was to optimize the solid-liquid extraction of the solid sample (tissue) and evaluate the proper loading condition onto the mixed mode SPE sorbent. Store-bought calf liver was used for the sample preparation optimization, in order to preserve the human tissue specimens. When analyzing tissue samples, the homogenization process is typically performed with a common kitchen blender or a hand-held homogenizer (ex: Polytron). Those techniques can be cumbersome and are difficult to apply to small mass samples. In recent years, novel developments with ceramic or stainless steel ball bearings in combination with high speed orbital shakers have shown the ability to reach complete cell membrane breakdown in less than 60 seconds. With variable cycle speed, this novel homogenization protocol can process sample sizes from 0.1 to 5 grams. In this application, the mass range of tissue sample was set at 1.0 grams with to 4 mL extraction solvent ratio. In Figure 6, various organic solvents (acetonitrile, methanol, acetone) and pH range (2,7 and 10) were evaluated to measure which extraction conditions give maximum recovery percentage. In this application, the extraction of tissue with acetonitrile with no additives gave the highest signal.

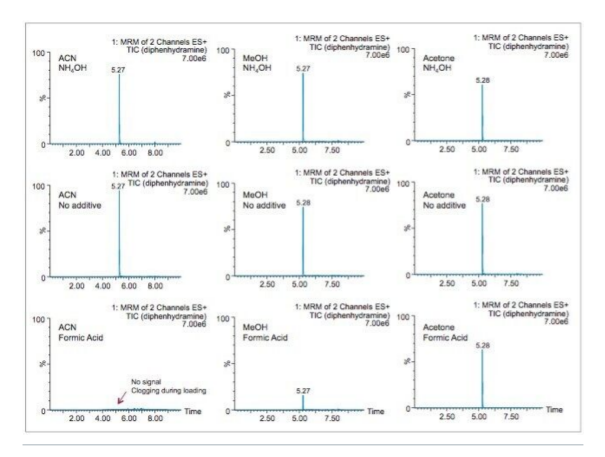


Figure 6. Optimization of the solid-liquid homogenization process.

Once the tissue sample was completely homogenized, it was centrifuged which produced a solid pellet on the bottom of the tube with the organic supernatant above. The organic supernatant was then filtered and decanted. Depending on the extraction conditions (pH and polarity), the target analyte is expected to be in solution and un-bound in the extraction solvent. In some applications, this crude extract can be used directly for quantification, however there is a high risk the raw sample extract will seriously reduce the robustness of the LC-MS/MS performance after a few injections. In traditional SPE protocols, when the target analyte is dissolved in a high percentage of organic solvent, the supernatant is usually evaporated to dryness and reconstituted in an aqueous diluent for further clean up. In instances where an evaporation-to-dryness step is needed, there is a risk of evaporative loss or possible re-dissolution issues. An effective way to avoid this lengthy step is to simply dilute the organic supernatant in a large aqueous volume at an organic/water ratio of less than 5%. A water volume between 100 and 200 mL is more than adequate to reach low organic ratio without any risk of breakthrough on the trapping column during loading phase. It may be perceived as a drawback, since the loading volume is quite large. However, with a loading flow rate at 10 mL/min using a large bore SPE barrel (6 cc with 150 mg bed mass), a 100 mL sample can be concentrated in 10 min, while evaporating to dryness can take several hours to complete.

The chromatograms in Figure 7 show the chromatography profile for an acetonitrile standard, water extracted standard and a spiked liver sample at 1 ppb level using the finalized extraction protocol. It is worth mentioning the stable baseline in both the water and liver extract, which is an indication that the extraction protocol, completed in 30 minutes, is producing a very clean extract. Table 5 depicts the overall recovery ratio for a liver tissue sample. Results demonstrated that 18 analytes have recovery values, measured against a post spiked deuterated internal standard (liver ion ratio recovery), within an acceptable range of 75% to 110%. The other analytes still show recovery ratio above 50%. The overall performance of both extraction methods gave an excellent linearity range as shown in Table 6. The R<sup>2</sup> values for all analytes ranged from 0.995 to 0.999 values. The limit of detection (LOD) for all analytes was set at 0.001 ng/mL (3x Sigma value).

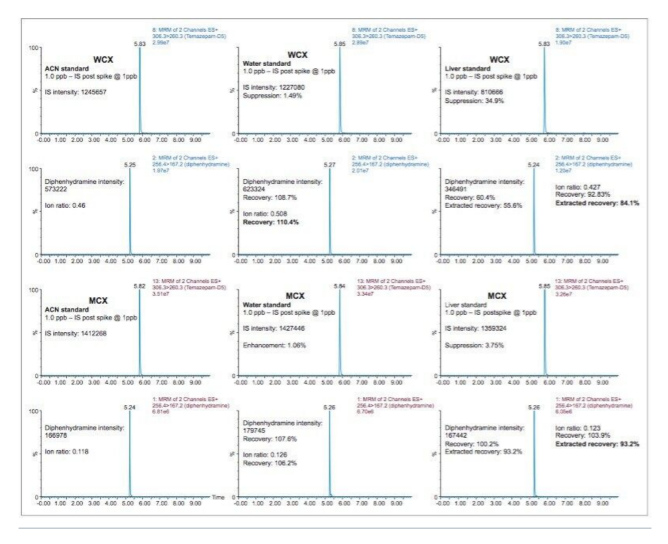


Figure 7. MCX vs WCX Chromatogram for MeOH std, water extracted std water, and matrix match extracted std for Diphenhydramine.

Compound	Water std	Water ion ratio	Liver std	Liver ion ratio
MCX method	recovery %	recovery %	recovery %	recovery %
Nordiazepam	90.9	94.7	90.6	103.8
Doxylamine	117.4	100.5	80.8	67.5
Diazepam	91.8	95.6	70.6	80.9
Promethazine	62.7	62.0	80.3	84.3
Oxazepam	57.1	56.5	74.9	78.6
Temazepam	82.4	81.5	76.5	80.3
Flecainide	82.3	70.5	92.6	77.3
Diphenhydramine	107.6	106.2	93.2	97.9
Hydromorphone	54.1	79.4	57.6	63.4
Dihydrocodeine	63.8	78.9	64.9	59.4
Zolpidem	98.6	97.6	88.0	92.4
Oxycodone	70.6	69.8	67.9	71.3
Citalopram	84.9	84.0	75.5	79.2
Quetiapine	69.1	68.4	104.1	109.2
Diltiazem	94.9	93.9	90.8	95.3
Buprenorphine	87.6	108.5	81.1	74.2
WCX method				
Normeperidine	100.2	101.9	70.6	106.9
Dextrorphan	61.7	67.7	31.0	47.0
Dextromethorphan	65.8	66.8	49.1	74.3
Diphenhydramine	108.7	110.4	55.6	84.1
Ndesmethylcitalopram	67.2	75.5	52.5	68.7
Citalopram	66.3	67.3	55.2	83.6
Norbuprenorphine	73.5	82.6	50.8	66.6

Table 5. Recovery values for water extract vs calf liver extract.

Compound - MCX	IS	Linearity	Weighting	range (ng/mL)	R <sup>2</sup>	LOD (ng/mL)
Nordiazepam	nordiazepam-d5	quadratic	1/x	0.025 - 10	0.997	0.001
Doxylamine	doxylamine-d5	quadratic	1/x	0.025 - 10	0.996	0.001
Diazepam	nordiazepam-d5	quadratic	1/x	0.025 - 10	0.999	0.001
Promethazine	temazepam-d5	quadratic	1/x	0.05 - 10	0.997	0.001
Oxazepam	temazepam-d5	quadratic	1/x	0.05 - 10	0.995	0.001
Temazepam	temazepam-d5	quadratic	1/x	0.05 - 10	0.998	0.001
Flecainide	doxylamine-d5	quadratic	1/x	0.050 - 10	0.998	0.001
Diphenhydramine	temazepam-d5	quadratic	1/x	0.025 - 10	0.996	0.001
Hydromorphone	hydromorphone-d5	quadratic	1/x	0.1 - 10	0.997	0.001
Dihydrocodeine	dihydrocodeiene-d6	quadratic	1/x	0.025 - 10	0.995	0.001
Zolpidem	temazepam-d5	quadratic	1/x	0.050 - 10	0.998	0.001
Oxycodone	temazepam-d5	quadratic	1/x	0.05 - 10	0.999	0.001
Citalopram	temazepam-d5	quadratic	1/x	0.050 - 10	0.997	0.001
Quetiapine	temazepam-d5	quadratic	1/x	0.050 - 10	0.996	0.001
Diltiazem	temazepam-d5	quadratic	1/x	0.025 - 1	0.996	0.001
Buprenorphine	dihydrocodeiene-d6	quadratic	1/x	0.05 - 10	0.996	0.001
Compound - WCX						
Normeperidine	temazepam-d5	quadratic	1/x	0.05 - 10	0.999	0.001
Dextrorphan	temazepam-d5	quadratic	1/x	0.05 - 10	0.996	0.001
Dextromethorphan	temazepam-d5	quadratic	1/x	0.010 - 10	0.999	0.001
Diphenhydramine	temazepam-d5	quadratic	1/x	0.05 - 10	0.999	0.001
Ndesmethylcitalopram	ndesmethylcitalopram-d5	quadratic	1/x	0.025-10	0.999	0.001
Citalopram	temazepam-d5	quadratic	1/x	0.05 - 10	0.998	0.001
Norbuprenorphine	ndesmethylcitalopram-d5	quadratic	1/x	0.05 - 10	0.999	0.001

Table 6. Linear range and detection limits.

#### Sample quantification

When analyzing highly complex sample types (class C matrix or solid samples), extraction recoveries are most often overwhelmed by matrix effects, which can lead to either suppression or enhancement in the MS detector. These effects are related to the inability of the sample clean up protocol to fully remove interferences from the raw sample.

In this work, the extraction protocol relied heavily on the use of a mixed mode sorbent using two trapping mechanisms. In this application, the Oasis MCX and WCX both have a reverse phase and cation exchange ligands to fractionate target basic analytes from neutral and acidic interferences. As seen in Figures 8 and 9, the MCX and WCX extracts for citolapram in various human tissue sample showed outstanding clean chromatograms at concentration values between 1.0 and 0.05 ppb.

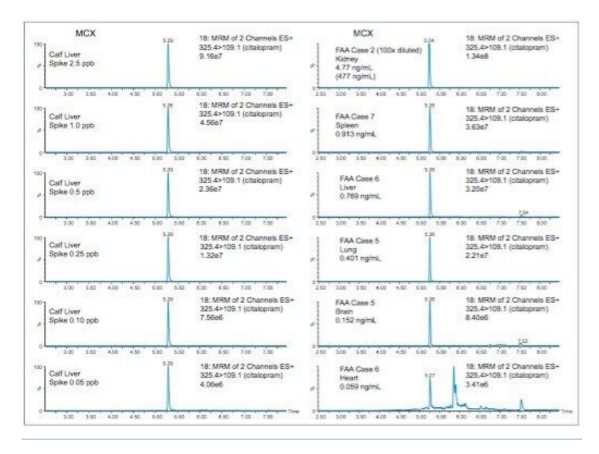


Figure 8. MCX chromatograms for tissue samples.

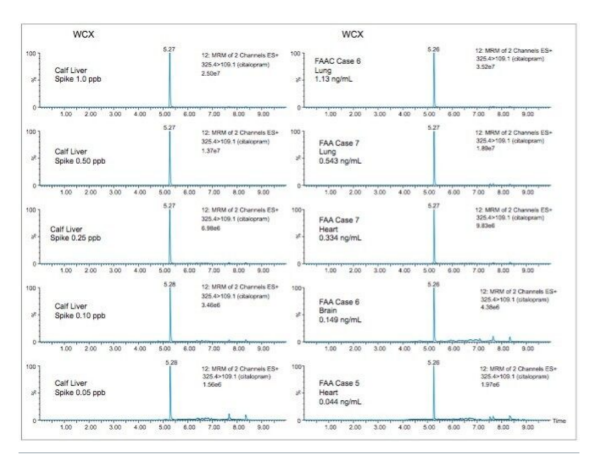


Figure 9. WCX chromatograms for tissue samples.

The results for the analyses of biological specimens (heart, brain, lung, liver, kidney, and spleen) are presented in Table 7. The extracts were quantified against a matrix match standard (calf liver) with a corresponding deuterated internal standard. The results were quantified within the linear range of 0.01 to 10 ng/mL. Therefore, some tissue extracts were subjected to a 100:1 dilution step before injection, to avoid flat top peak shape due to detector saturation. From the case studies in this application, case 7 tested positive for dextromethorphan (cough suppressant) and case 5 tested positive for flecainide (antiarrythmic agent). Also, case 2 tested positive for citolapram (antidepressant). As seen, since citolapram was selected as an efficiency marker for the MCX and WCX protocols, the results show comparable and precise performances for a variety tissue samples. The column chemistries used for this application gave an excellent performance analyzing well over 1000 sample injections.

Quantifcation (ng/mL)	WCX extracts					MCX EXTRACTS												
	A	В	С	D	E	F	G	н	1	J	к	L	М	N	0	P	Q	R
Case 2 Heart	ND	0.08	1.23	493.0	669.0	ND	TLD	ND	0.03	5.29	0.01	0.73	9.15	690.0	96.7	ND	0.32	ND
Case 2 Lung	ND	0.02	0.36	973.0	740.0	ND	0.05	ND	0.03	2.82	0.02	0.76	7.25	966.0	76.0	ND	0.37	ND
Case 2 Liver	ND	0.02	0.18	1690.0	1070.0	ND	0.03	ND	0.16	1.70	0.02	3.10	9.90	1012.0	327.0	ND	0.56	ND
Case 2 Kidney	ND	TLD	2.15	544.0	504.0	ND	TLD	ND	0.40	10.8	0.05	1.33	6.05	477.0	221.0	ND	0.18	ND
Case 2 Spleen	ND	0.00	1.69	445.0	643.0	ND	TLD	ND	TLD	8.83	0.02	1.54	4.40	616.0	90.0	ND	0.22	ND
Case 2 Brain	0.01	0.05	2.72	115.0	525.0	ND	TLD	ND	TLD	8.78	0.01	0.60	3.65	587.0	82.90	ND	0.21	ND
Case 5 Heart	ND	0.17	TLD	0.03	0.04	ND	ND	ND	830.0	ND	ND	ND	ND	0.18	TLD	44.6	0.57	ND
Case 5 Lung	ND	0.23	TLD	0.02	0.02	ND	ND	ND	500.0	ND	ND	ND	ND	0.40	0.013	20.9	0.34	ND
Case 5 Liver	ND	0.09	0.01	0.11	0.28	ND	ND	ND	970.0	ND	ND	ND	ND	0.07	TLD	117.0	0.97	ND
Case 5 Kidney	ND	0.07	TLD	0.13	0.17	ND	ND	ND	588.0	ND	ND	ND	ND	0.07	TLD	35.3	0.45	ND
Case 5 Spleen	ND	0.06	TLD	0.51	1.00	ND	ND	ND	423.0	ND	ND	ND	ND	0.03	TLD	48.0	0.68	ND
Case 5 Brain	ND	0.01	TLD	0.40	0.60	ND	ND	ND	138.0	ND	ND	ND	ND	0.15	TLD	10.4	0.14	ND
Case 6 Heart	ND	0.06	0.03	0.20	0.43	0.04	TLD	ND	0.23	TLD	ND	ND	ND	0.06	0.03	ND	ND	ND
Case 6 Lung	ND	0.40	0.10	0.61	1.13	0.05	TLD	ND	0.04	0.08	ND	ND	ND	1.17	0.16	ND	ND	ND
Case 6 Liver	ND	0.21	0.03	0.31	0.43	0.08	TLD	0.008	TLD	0.09	ND	ND	ND	0.77	0.11	ND	ND	ND
Case 6 Kidney	0.81	0.06	0.16	0.38	0.40	0.24	0.04	0.053	TLD	0.35	ND	ND	ND	3.63	0.12	ND	ND	0.04
Case 6 Spleen	ND	0.35	0.04	0.75	1.45	0.14	TLD	0.032	TLD	0.08	ND	ND	ND	0.43	TLD	ND	ND	ND
Case 6 Brain	ND	0.14	0.02	0.06	0.15	0.02	TLD	ND	TLD	0.01	ND	ND	ND	0.56	0.023	ND	ND	ND
Case 7 Heart	0.38	448.0	0.38	0.13	0.33	ND	0.26	ND	ND	0.62	ND	ND	ND	0.41	TLD	ND	ND	ND
Case 7 Lung	2.23	1684.0	4.26	0.25	0.54	ND	3.06	ND	ND	5.30	ND	ND	ND	0.84	0.16	ND	ND	ND
Case 7 Liver	1.18	1258.0	1.61	0.32	0.57	ND	0.81	ND	ND	2.90	ND	ND	ND	0.13	TLD	ND	ND	ND
Case 7 Kidney	2.38	365.0	0.67	0.13	0.17	ND	0.41	ND	ND	0.98	ND	ND	ND	0.62	0.05	ND	ND	ND
Case 7 Spleen	0.73	554.0	0.47	0.25	0.59	ND	0.48	ND	ND	1.07	ND	ND	ND	0.91	0.07	ND	ND	ND
Case 7 Brain	0.25	391.0	0.31	0.27	0.40	ND	0.15	ND	ND	0.46	ND	ND	ND	0.73	0.16	ND	ND	ND
WCX extracts (Detected)	MCX ex	tracts (De	etected	WCX e	extracts (	Un-De	tected)	МСХ	extracts	(Un-D	etected)		Trace level Detection (TLD) <0.01 ng/mL					
A: Dextrorphan	F: Nord	liazepam		Norm	eperidine	9	Promethazine						Not Detected (ND)					
B: Dextromethorphan	G: Doxy	lamine						Oxaz	epam									
C: Diphenhydramine	H: Diaz	epam						Tema	zepam									
D: ndesmethylcitalopram	I: Fleca	inide																
E: citalopram	J: Diphe	enhydrami	ine															
	K: Hydr	omorphor	ne															
	L: Dihy	drocodein	e															
	M: Zolp	idem																
	N: Cital	opram																
	O: Quet	tiapine																
	P: Diltia	zem																

Table 7. Quantification values for human tissue samples.

Q: Buprenorphine R: Oxycodone

# Conclusion

This application demonstrated the automated and fast method development capability of the ACQUITY UPLC with 2D Technology for the analysis of pharmaceuticals in human tissue samples. The quantification limit was set at 10 ppt using a 1.0 g of sample. The micro extraction protocol offered the option to evaluate several elution parameters in a short time period. The elution optimization was completed within a 4 hrs hands-on work and the 2D LC results were analyzed using an over-night run using a multi-methods sample list (18 hrs). With the extraction protocol optimized, the final protocol produced a clean extract in 30 minutes without any evaporation to dryness and reconstitution into initial mobile phase conditions.

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