

Extraction of Illicit and Prescribed Drugs from Enzyme-Hydrolyzed Urine Using ISOLUTE® HYDRO DME+ Prior to UPLC-MS/MS Analysis

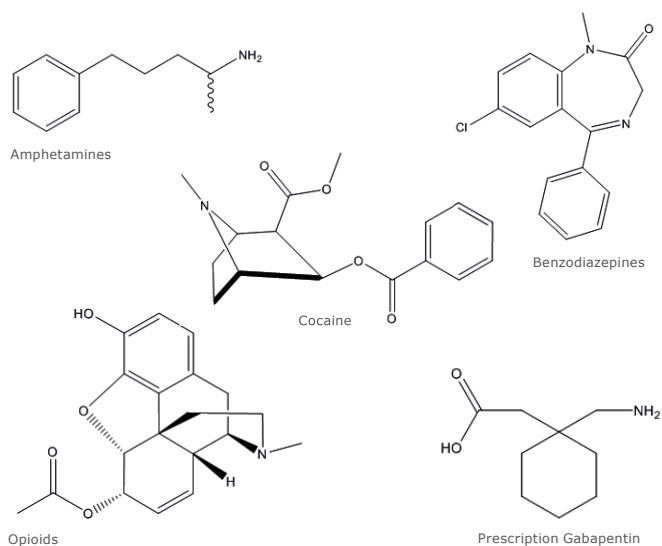


Figure 1. Example analyte structures by class.

Introduction

This application note describes the extraction of 25 illicit and prescribed drugs from hydrolyzed urine using ISOLUTE® HYDRO DME+ Dual Mode Extraction plates prior to UPLC-MS/MS analysis.

ISOLUTE HYDRO DME+ products provide extremely efficient removal of matrix components and hydrolysis enzyme from urine samples, using a simple pass through workflow.

Because of the enhanced sample clean up delivered by ISOLUTE HYDRO DME+ products, analyte sensitivity is significantly increased compared to traditional dilute and shoot (D&S) techniques, resulting in reduced limits of quantitation (LOQ). ISOLUTE® HYDRO DME+ plates and columns are ideal for urinary drugs of abuse and pain management applications because the inclusion of Biotage® HYDRO frit technology means that urine samples can be hydrolyzed in-situ in the column, eliminating the need for post hydrolysis sample transfer steps.

The simple sample preparation procedure described delivers clean extracts and recoveries above 65% for the majority of analytes. The limits of quantitation all meet and exceed the sensitivity requirements set by SAMHSA and EWDTs for workplace testing applications.

Analytes

Ecgonine methyl ester, pregabalin, morphine, oxycodone, amphetamine, gabapentin, codeine, 6-monoacetylmorphine, MDMA, hydrocodone, mephedrone, benzoylecgonine, ketamine, 7-aminoclonazepam, cocaine, norbuprenorphine, 7-aminoflunitrazepam, buprenorphine, PCP, EDDP, oxazepam, methadone, Zaleplon, flunitrazepam and ritalinic acid.

Sample Preparation Procedure

Format

ISOLUTE® HYDRO DME+ 400 mg Fixed Well Plate, part number: 970-0400-PZ01.

Sample Pre-treatment

(hydrolysis using β -Glucuronidase enzyme (*Helix pomatia*))

To 500 μ L of urine, add 25 μ L of internal standard mix at concentration 1 ng/ μ L and allow equilibration to take place at room temperature for 1 hour.

Apply 25 μ L of enzyme solution to 450 μ L of ammonium acetate (50 mM pH 5.0) and vortex briefly. Add this mix to the urine spiked with internal standard (as above) and vortex briefly.

Apply a 100 μ L aliquot of this sample (matrix/IS/enzyme/buffer mix) to the ISOLUTE HYDRO DME+ product and incubate for 2 hours at 60 °C.

Extraction Procedure and Post-Extraction

Allow the sample to cool to room temperature and position a 96-well collection plate under the extraction plate. Add acetonitrile (600 μ L) onto the hydrolyzed urine sample. Perform 5x aspirate/dispense steps with an electronic 8-channel pipette to ensure sufficient mixing.

Using a Biotage® Pressure+ 96 Positive Pressure Manifold, apply approximately 5 PSI of positive pressure to elute the acetonitrile. The samples may be analysed by UPLC-MS/MS without an evaporation step*. Simply cover the collection plate with a sealing mat prior to transfer to the autosampler.

*If increased analyte sensitivity is required, the samples may be evaporated using a Biotage® SPE Dry 96 at 40L/min at 40 deg C and reconstituted in a low solvent volume prior to UPLC-MS/MS analysis. If so, a 100 μ L volume of methanolic hydrochloric acid (50mM) should be added to each well prior to evaporation to prevent the loss of more volatile analytes such as amphetamine.

UHPLC Conditions

Instrument

Waters ACQUITY UPLC with 20 μ L loop

Column

Restek Raptor™ Biphenyl 2.7 μ m (100 x 2.1 mm id) with Raptor™ Biphenyl EXP guard cartridge

Mobile Phase

A: 2 mM ammonium formate (aq), 0.1 % formic acid

B: 2 mM ammonium formate (methanol), 0.1 % formic acid

Flow Rate

0.4 mL min

Gradient Details

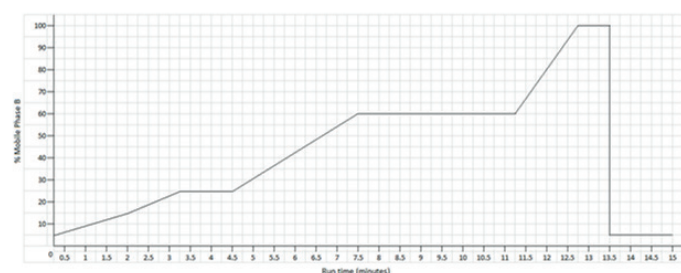


Figure 2. Graphical representation of LC conditions

Table 1. Gradient Conditions. Curve 6: Linear Gradient.

Time (min)	%A	%B	Curve
0.00	95	5	6
2.00	85	15	6
3.25	75	25	6
4.50	75	25	6
7.50	40	60	6
11.25	40	60	6
12.75	0	100	6
13.50	0	100	6
13.51	95	5	6
15.00	95	5	6

Column Temperature

40 °C

Injection Volume

1 μ L (partial loop without overfill)

Sample Temperature

20 °C

MS/MS Conditions

Instrument

Waters Premier XE triple quadrupole mass spectrometer equipped with an electrospray interface for mass analysis.

Source Temperature

150 °C

Desolvation Temperature

450 °C

Positive ions acquired in the multiple reaction monitoring (MRM) mode are described in Table 2:

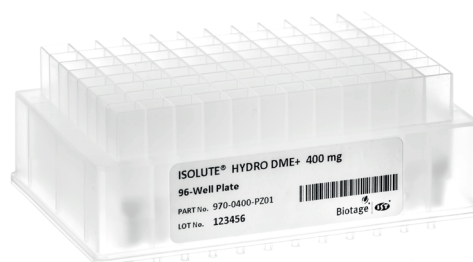
Table 2. MRM Conditions.

Compound	MRM Transition	Cone Voltage (V)	Collision Energy (eV)
Ecgonine Methyl Ester (EME)	182.2 > 82.0	50	15
Pregabalin	160.2 > 55.2	18	25
Morphine	286.2 > 201.0	42	25
Oxymorphone	302.2 > 198.1	34	37
Amphetamine	136.0 > 118.9	16	9
Gabapentin	172.3 > 137.1	23	15
Codeine	300.3 > 215.1	42	25
6-MAM	328.2 > 165.1	44	33
MDMA	194.1 > 163.0	20	13
Hydrocodone	300.2 > 199.1	46	33
Mephedrone	178.1 > 160.0	35	12
Ritalinic Acid	220.2 > 84.1	24	21
Benzoylcegonine (BZE)	290.1 > 168.0	30	18
Ketamine	238.1 > 124.9	25	27
7-amino-clonazepam	286.2 > 121.0	40	30
Cocaine	304.2 > 182.0	30	20
Norbuprenorphine	414.3 > 101.0	55	42
7-amino-flunitrazepam	284.2 > 135.0	40	27
Buprenorphine	468.3 > 468.3	55	5
PCP	244.2 > 158.9	20	15
EDDP	278.2 > 234.2	26	30
Oxazepam	287.2 > 241.0	30	21
Methadone	310.2 > 265.2	26	15
Zaleplon	306.2 > 264.2	40	22
Flunitrazepam	314.2 > 268.2	40	25

Results

Analyte recovery, reproducibility, linearity and cleanliness studies were performed using intact urine from healthy volunteers.

Recovery data shown in Figure 3 demonstrates that this protocol provides extraction recovery of 65% or greater for the majority of analytes while simultaneously removing common urinary components. Where the recovery value was less than 65%, the sensitivity was more than sufficient to meet established cut off limits for drugs of abuse testing and prescription drug monitoring. RSD values were below 10%.



Note: The direct inject approach allows maximum sample throughput, however if greater signal is required, the samples may be evaporated and reconstituted in a low solvent volume prior to UPLC-MS/MS analysis (see page 1 for details).

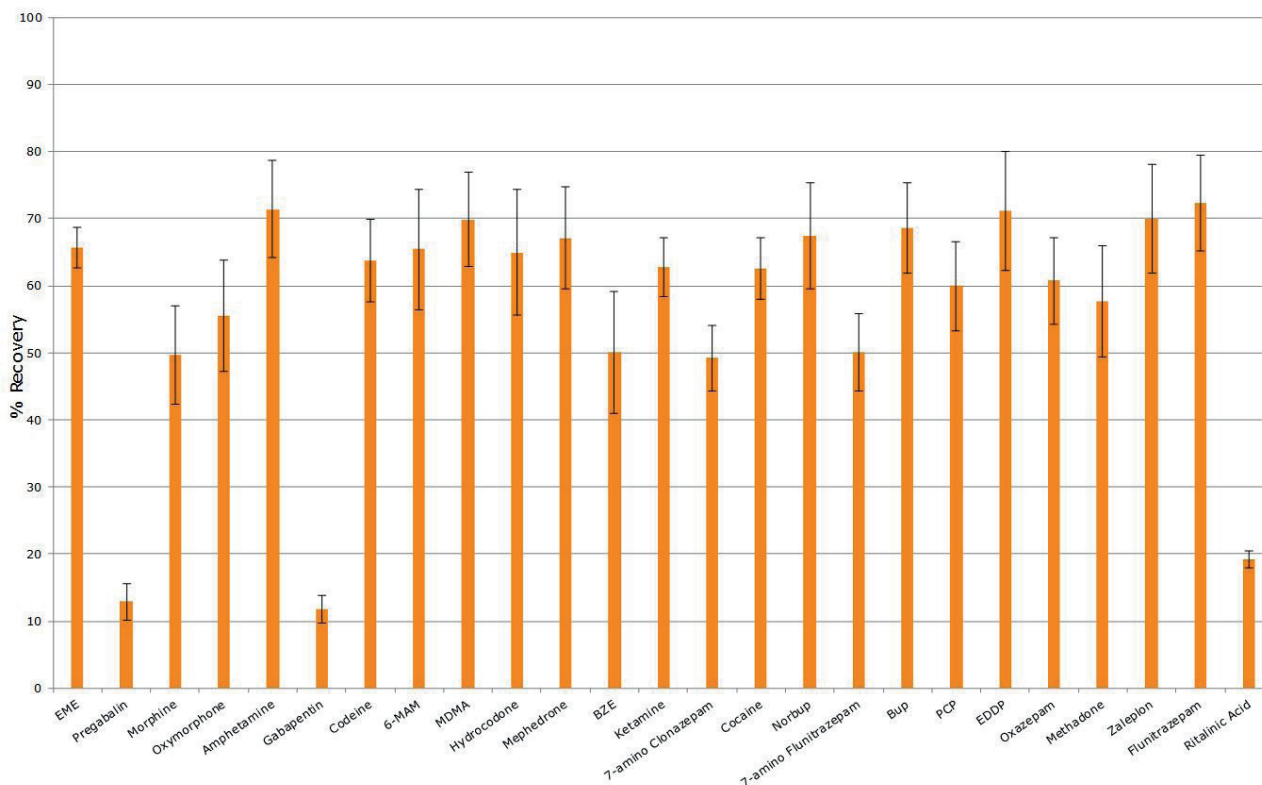


Figure 3. Typical analyte recoveries and RSD (n=7, shown as error bars) for hydrolyzed urine following ISOLUTE® HYDRO DME+ processing and direct UPLC-MS/MS injection.

Following recovery determination, analytes were extracted from urine spiked before hydrolysis at levels 10, 20, 50, 100, 200 and 400 ng/mL to construct calibration curves. Representative curves are shown in Figure 4.

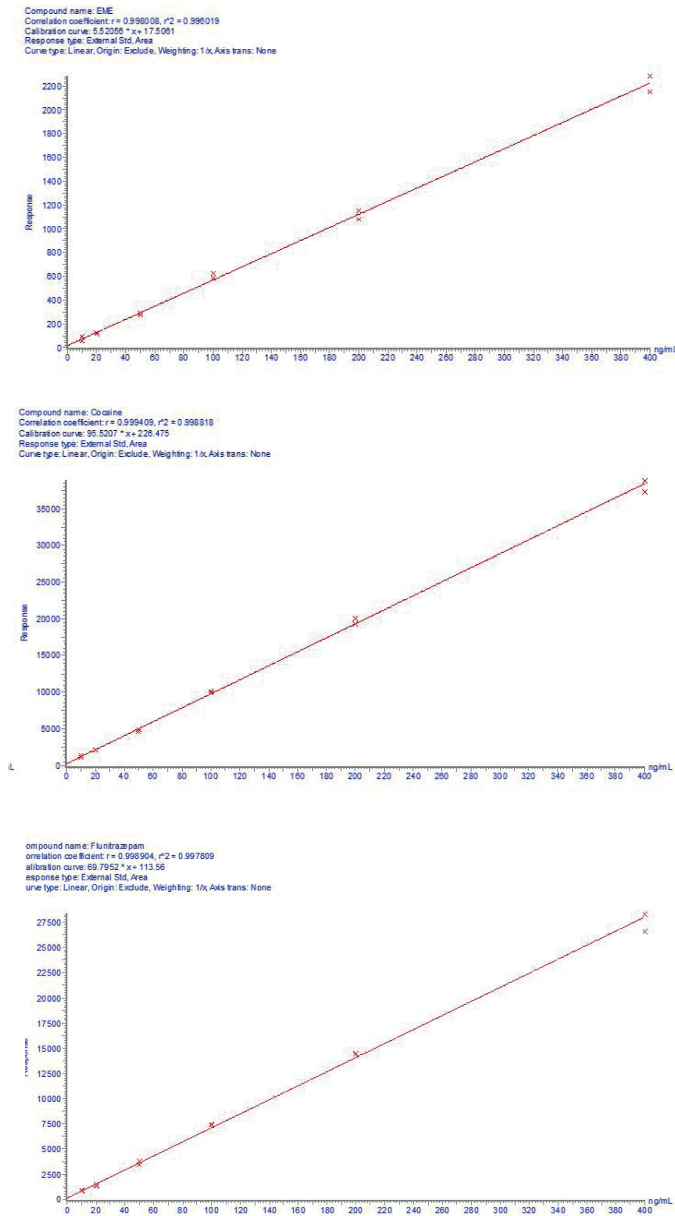


Figure 4. Calibration curves of application analytes EME, cocaine and flunitrazepam constructed following extraction of hydrolyzed urine using ISOLUTE® HYDRO DME+. Analyte concentrations are 10, 20, 50, 100, 200 and 400 ng/mL showing r^2 values of greater than 0.99. Internal standard concentrations are at 50 ng/mL.

Matrix Component Removal

Urea and creatinine, along with other urinary matrix components and hydrolysis enzyme, can have a detrimental effect on quantitation of desired analytes. Figures 5 and 6 illustrate the extent of removal of urea and creatinine from urine using ISOLUTE® HYDRO DME+ products, compared to the levels present in non-purified urine (as used in dilute and shoot (D&S) experiments).

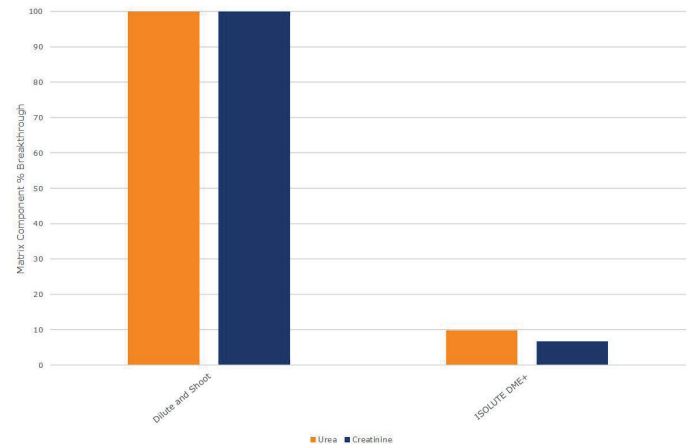


Figure 5. Chart demonstrating urea and creatinine % breakthrough into sample extract with and without ISOLUTE® HYDRO DME+ clean up.

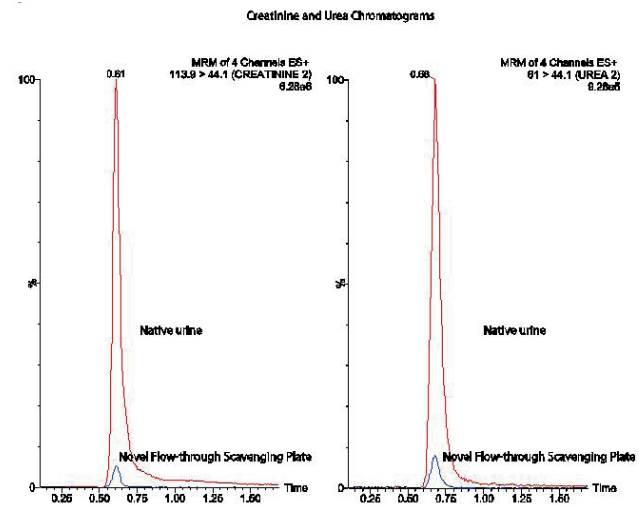


Figure 6. MRM chromatograms illustrating relative Creatinine (left) and Urea (right) content in hydrolyzed urine: (red) following ACN crash, (blue) post processing through ISOLUTE® HYDRO DME+.

Chemicals and Reagents

- » Reference standards (including deuterated internal standards), ammonium acetate (reagent grade $\geq 98\%$), ammonium formate (LC-MS grade), formic acid (LC-MS grade) and β -Glucuronidase enzyme (Type HP-2, aqueous solution $\geq 100,000$ units/mL) were purchased from Sigma- Aldrich Company Ltd. (Gillingham, UK).
- » HPLC-grade solvents (acetonitrile, methanol) were purchased from Honeywell Research Chemicals (Bucharest, Romania).
- » Water used was 18.2 MOhm-cm, drawn daily from a Direct-Q5 water purifier.
- » Ammonium acetate (50 mM aq pH5) was prepared by adding 3.854 g of ammonium acetate to 1 L of deionized water. The pH was adjusted using formic acid (as above).
- » Mobile phase A (2 mM ammonium formate (aq), 0.1 % formic acid) was prepared by adding 126 mg of ammonium formate to 500 mL of purified water, adding 1mL of concentrated formic acid and making up to 1 L with purified water
- » Mobile phase B (2 mM ammonium formate (methanol), 0.1 % formic acid) was prepared by adding 126 mg of ammonium formate to 500 mL of HPLC grade methanol, adding 1mL of concentrated formic acid and making up to 1 L with HPLC grade methanol
- » 50 mM hydrochloric acid in methanol was prepared daily by adding 100 μ L of 12M concentrated hydrochloric acid to 23.9 mL of HPLC-grade methanol.

Ordering Information

Part Number	Description	Quantity
970-0400-PZ01	ISOLUTE® HYDRO DME+ 400 mg Plate	1
PPM-96	Biotage® PRESSURE+ 96 Positive Pressure Manifold	1
SD-9600-DHS-EU	Biotage® SPE Dry 96 Sample Evaporator 220/240 V	1
SD-9600-DHS-NA	Biotage® SPE Dry 96 Sample Evaporator 100/120 V	1
121-5203	Collection Plate, 2 mL Square	50
121-5204	Pierceable Sealing Cap	50

Additional information

All data shown in this application note was generated using real, intact matrix, obtained from human volunteers.

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