# Small Molecules Automated Extraction from Human Breast Milk Using the Extrahera and the **EVOLUTE EXPRESS CX Prior to LC-MS/MS Analysis**

500

20.0

20.0

25.0

100.0

10.0 20.0

5.0 10.0

Time

(min)

1-2

1-2

1-2

1-2

1-2

5

2-3

5-10

2-3

5-10

Pressure

(psi)

0.5-3

0.5-3

0.5-3

0.5-3

0.5-3

20

0.5-3

N/A

0.5-3

N/A

Mohamed Youssef<sup>1</sup>, Mario Merida III<sup>1</sup>, Jeremy Smith<sup>1</sup>, Stephanie Marin<sup>1</sup>, Jillian Neifeld<sup>1</sup>, Elena Gairloch<sup>1</sup> <sup>1</sup>Biotage, 10430 Harris Oaks Blvd., Suite C, Charlotte North Carolina 28269, USA

Sample Preparation

limit of detection for THCOOH.

5.0 10.0

1.3 2.5

125 250

5.0 10.0

5.0 10.0

2.5

Table 1 Spiked Concentrations In Cals and OCs

Volume (uL)

1000

1000

300

300

300

N/A

600

N/A

600

N/A

Table 2. EXTRAHERA Extraction protocol

**Chromatography Parameters** 

Table 3. Shimadzu Nexera X2 UPLC setup

Mass Spectrometry Parameters

The extraction method is shown in (Table 2).

Phencyclidine(PCP) 2.5 5.0

and analyzed via LC/MS-MS

5.0

EVOLUTE® EXPRESS CX SPE/EXTRAHERA Procedure

Extracts were then reconstituted with 200 uL of 5% MeOH in water

Solvent/Equipmen

Methano

4 % Formic Acid

Sample

4 % Formic Acia

70:30 Hexane/EtOAc

N/A

2% Acetic Acid in ACN

Using SPEDry @ 40°

DCM/IPA/NH4OH[78:20:2]

Using SPEDry @ 40°

Instrument: A SCIEX 5500 triple quadrupole Mass Spectrometer with

Turbo Ionspray® Ion interface (Foster City, CA) was used. Optimized

source parameters are shown in table 5 (MRM transition parameters

not shown, but available upon request). Retention window for MRM

was set at 60 seconds with target scan time at 2.5 seconds.

Parameters

Restek Raptor Biphenyl 2.7 µm, 50 x 3.0mm

0.1 % Formic acid in H2O

0.1 % Formic acid in MeOH

0.4 ml/min

40°C

10°C

20 uL

Benzoylecognine

Caffeine

d-Amphetamine

Methadone

Morphine

Oxazepam

Oxycodone

Secobarbital

тнсоон

Step

Condition

Equilibration

Sample Load

Wash #1

Wash #2

Drv

Elution 1

Evaporate

Elution 2

Evaporate

HPLC

Column

ΜΡΔ

MPB

Flow Rate

Column Temp

Sample Temp

Injection Vol.

d-Methamphetamine 5.0

Breast Milk Sample Pre-treatment and Preparation

Calibrators were prepared by spiking the target analytes into drug-

free human breast milk. Serial dilutions were used to achieve the

remaining standard calibration concentrations (Table 1). In 1.5 mL

micro-centrifuge tubes, 200 µL of methanol was added to 200 µL of

each calibrator and control and vortexed for 1 minute then

centrifuged for 10 minutes at 13,000 rpm. 300 µL of the resulting

supernatant was then loaded onto the EVOLUTE® EXPRESS CX 30 mg

plate, and went through extraction and elution (Table 2).

Conditioning and equilibration steps are optional to reach a lower

Analyte conc. In ng/mL Cal 1 Cal 2 Cal 3 Cal 4 Cal 5 Cal 6 QC 1 QC 2

 15.0
 20.0
 30.0
 40.0
 10.0
 20.0

 3.75
 5.0
 7.5
 10.0
 2.5
 5.0

375 500 750 1000 250

15.0 20.0 30.0 40.0 10.0

 5.0
 10.0
 15.0
 20.0
 30.0
 40.0
 10.0

 6.3
 12.5
 18.75
 25.0
 37.5
 50.0
 12.5

25.0 50.0 75.0 100.0 150.0 200.0 50.0

15.0 20.0 30.0 40.0

7.5 10.0 15.0 20.0

 2.5
 5.0
 7.5
 10.0
 15.0
 20.0
 5.0
 10.0

 10.0
 20.0
 30.0
 40.0
 60.0
 80.0
 20.0
 40.0

 2.5
 5.0
 7.5
 10.0
 15.0
 20.0
 5.0
 10.0

# Introduction

Breast feeding is beneficial in meeting the nutritional and immunological needs of infants. Using illegal drugs while breast feeding can have severe consequences for both infant and mother. At Biotage, we developed a new extraction protocol for 12 common drugs of abuse (DOA) to be detected in breast milk (Figure 1) using mixed-mode polymeric cation exchange solid phase extraction (SPE). The EVOLUTE® EXPRESS CX (figure 3) extraction allowed for highly-sensitive detection of analytes including Benzoylecgonine, (-)-Cotinine, Caffeine, d-Amphetamine, d-Methamphetamine, Methadone, Morphine, Oxazepam, Oxycodone, Phencyclidine (PCP), Secobarbital, and (l)-9-Carboxy-11-nor-Delta-9-THC (THC-COOH).

Extrahera (figure 2), a bench top automated extraction system. provided minimal sample intervention and high throughput for the analysis of these DOA. Using the combination of reliable automation and SPE sample preparation techniques, a method was developed demonstrating the precision, accuracy, linearity, and sensitivity necessary for a robust guantitative workflow.



Figure 1. Breast Milk Composition Figure 2. EXTRAHERA Automation System



# Figure 3. EVOLUTE EXPRESS CX Interferences Removal

# Experimental

# **Reagents and Materials**

# Standards

All standards were purchased from Cerilliant (Round Rock, TX), HPLC grade water and methanol (MeOH) were purchased from Sigma Aldrich (St. Louis, MO) in addition to reagent grade dichloromethane (DCM), formic acid, acetic acid (CH<sub>3</sub>COOH), Acetonitrile (ACN), ammonium hydroxide (NH4OH) Hexane (C6H14), ethyl acetate (EtOAc). EVOLUTE® EXPRESS CX (30 mg bed) extraction plate (601-0030-PX01), and Biotage® SPE Dry 96 (SD-9300-DHS-NA), Extrahera (414001) were supplied by Biotage. The LC column was provided by Restek Corp. QC materials were generously donated from UTAK labs (product # 21801-4).

Ionization Spray Voltage	+1500(V)	CAD	Medium
Source Temp	600 °C	GS1	50
Curtain	30 (V)	GS2	50

Pos & Neg mode switching Table 4. SCIEX 5500 Triple Quadrupole ESI (+/-) Turbo Ionspray® Source Parameters.

# Results

Analytes were extracted from the calibrators and OC material by protein precipitation followed by SPE. Samples were then analyzed via an LC/MS-MS system and a biphenyl column using a 5-minute gradient. The DOA standard curves in breast milk had excellent linearity within the measured range (Table 1) with R<sup>2</sup> values greater than 0.995 (figure 4). When an excess of organic solvent is injected on the LC-MS/MS, there is an increased chance of peak broadening and peak tailing. Using the EVOLUTE® EXPRESS CX method, the peak shape was investigated to determine if it was still acceptable (Figure



Figure 4. Calibration curve for benzovlecaonine (BE



# Figure 5. Peak Shape at lowest calibration for BE at 5 ng/mL

Carryover was assessed by analyzing the area count of the blank calculated as percentage of the mean peak area of the lowest calibrator concentration. No significant carryover (0%) was determined for all analytes. Matrix effects were also investigated for all analytes by comparing the AUCs of standard curves prepared in 100% water to those in drug-free human breast milk. No significant matrix effects were observed (<10%). Washing the plate with a Hexane/EtOAc mixture eliminated interferences from the matrix and eliminated lipids and proteins left in the sample

The OC samples were analyzed to obtain inter- and intra-day precision and accuracy values. Accuracies determined were within (10%) and coefficient of variation values were all (<10%) for the concentrations within the measured range (Table 5).

All analytes had excellent process efficiency within ± 10% indicating efficiency in sample clean up to eliminate the lipids and the proteins. Using two different elution solvents helped increase recovery for analytes from different classes

### Interday % Accuracy (n=3) Intraday % Precision (n=3) QC Conc In ng/mL Intraday % Accuracy (n=3) Analyte Benzolyacov 100.3 7.6 95.8 6.3 10 91.9 20 3.4 92 3.3 Cotinine 2.5 93.6 90.6 1.9 8.7 5 94.5 2.9 62.5 2.4 Caffeine 100.3 250 82 93.8 53 500 97.2 3.6 95.4 3.4 d-Amphetamine 98.6 96.8 5.6 20 90.7 4.6 93 3.7 d-Methamphetan 93.6 90.6 2.2 10 87 20 94.5 2.9 92.5 2.9 Methadone 12.5 98.2 92.6 7.9 5.4 25 97.2 3.9 94.4 3.5 Morphine 93.5 95.3 5.7 10 7.9 91.7 4.8 20 5.4 92.1 Oxazepam 94.9 92.1 2.8 50 4.6 100 93.5 3.5 94.8 2.9 Oxycodone 96.2 92.6 4.9 92.1 3.9 93.5 3.9 hencyclidine (PC 92.3 0.1 95.8 10 97.1 5.6 91.2 4.8 Secobarbital 95.1 4.6 91.3 2.9 92.4 3.5 93.5 3.1 40 THCOOH 92.6 88.1 4.3 5.1 10 93.1 3.8 93.5 4.2

# Table 5. Inter & intra-day precision and accuracy

# Conclusions

- » Drugs of abuse in human breast milk can be automated and guantified accurately with this method. This analytical method employs protein precipitation followed by EVOLUTE® EXPRESS CX. Interferences including proteins, sugars, non-ionisable molecules and phospholipids are removed during the elution step, ensuring extremely clean extracts and enabling analysis of 12 DOA from different classes.
- » Using commercial quality controls ensured the integrity of making in-house calibrators
- » Using two different elution solvents allowed for recovery of analytes from different classes that were detected using positive and negative mode on the mass spec.
- » Automating this extraction protocol can provide savings in analysis time, solvent consumption, and minimizes manual sample intervention when compared to traditional sample preparation techniques.

# References

Marchei, E., Escuder, D., Pallas, C., Garcia-Algar, O., Gómez, A., Friguls, B., Pellegrini, M. and Pichini, S. (2019). Simultaneous analysis of frequently used licit and illicit psychoactive drugs in breast milk by liquid chromatography tandem mass spectrometry.

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