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Extraction of gamma-Hydroxybutyric Acid (GHB) from Urine Using ISOLUTE® SLE+ Prior to GC/MS Analysis

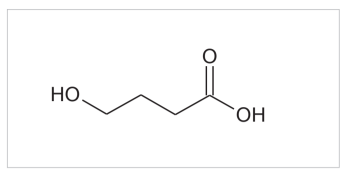


Figure 1. Structure of gamma-Hydroxybutyric acid (GHB)

Introduction

This application note describes the extraction of GHB from urine using supported liquid extraction and subsequent analysis by GC/MS.

ISOLUTE® SLE+ Supported Liquid Extraction plates and columns offer an efficient alternative to traditional liquid-liquid extraction (LLE) for bioanalytical sample preparation, providing high analyte recoveries, no emulsion formation, and significantly reduced sample preparation.

Analytes

Gamma-Hydroxybutyric acid (GHB) & GHB-D6

Sample Preparation Procedure

Sample Pre-treatment: Dilute pre-treated urine (0.2 mL) with 0.2% formic acid (aq) (0.2 mL). Spike GHB-D6 internal

standard and vortex mix thoroughly.

Format: ISOLUTE° SLE+ 200 µL Fixed Well Plate, part number 820-0200-P01

Sample Loading: Load the pre-treated urine (200 µL total volume) onto each well and apply a pulse of vacuum

or positive pressure (3–5 seconds) to initiate flow. Allow the sample to absorb for 5 minutes.

Analyte Extraction: Apply ethyl acetate (1 mL) and allow to flow under gravity for 5 minutes. Apply vacuum or

positive pressure to pull through any remaining extraction solvent (5-10 seconds).

Format: ISOLUTE° SLE+ 400 µL Sample Capacity columns, part number 820-0055-B

Sample Loading: Load the pre-treated urine (200 µL total volume*) onto the column and apply a pulse of

vacuum or positive pressure (3-5 seconds) to initiate flow. Allow the sample to absorb for

5 minutes.

*Note: Column is underloaded

Analyte Extraction: Apply ethyl acetate (1 mL) and allow to flow under gravity for 5 minutes. Apply a further

aliquot of ethyl acetate (1 mL) and allow to flow for another 5 minutes under gravity. Apply vacuum or positive pressure to pull through any remaining extraction solvent (5–10 seconds).

Post Elution, Reconstitution and Derivatisation:

Dry the extract in a stream of air or nitrogen at ambient temperature using a SPE Dry

(20 to 40 L/min) or TurboVap (1.0 bar) for 30 mins.

Upon dryness, reconstitute with ethyl acetate (50 µL) and BSTFA/1% TMCS (50 µL) and vortex

for 20 seconds. Transfer to a high recovery glass vial.



GC Conditions

Instrument: Agilent 7890A with QuickSwap

Column: Phenomenex Zebron ZB-35, 30 m x 0.25 mm ID x 0.25 μm

Carrier Helium 1.2 mL/min (constant flow)

Inlet: 250 °C, Split (ratio 10:1), 12 mL/min, septum purge flow: 3 mL/min

Injection: 1 µL

Wash Solvent: Ethyl acetate

Oven: Initial temperature 70 °C

Ramp 10 °C/min to 140 °C

Ramp 120 °C/min to 350 °C

Post Run: Back-flush for 1.6 minutes (2 void volumes)

Transfer Line: 280 °C

MS Conditions

Instrument: Agilent 5975C

Source: 230 °C

Quadrupole: 150 °C

MSD mode: SIM

SIM Parameters

Table 1. Ions acquired in the selected Ion Monitoring (SIM) mode

SIM Group	Analyte	Quantifier Ion	Quantifier Ion
1	GHB-D6	239	240
1	GHB	233	117

Results

The optimized protocols for ISOLUTE SLE+ plates and columns demonstrated reproducible recoveries between three unique donors, as shown in Figure 2. Percentage recovery was 52-69%. RSDs for GHB were below 9% on the 200 µL capacity plate format and below 4% on 400 µL capacity column format. **Figure 3**. shows linearity data for plate and column formats ($r^2 > 0.999$ for both formats). This application is unusual because of the relatively high concentration cut-off requirement ($10 \mu g/mL$) for illicit GHB. The low matrix volume used and relatively low recoveries do not adversely affect the required assay performance and sufficient sensitivity was achieved.



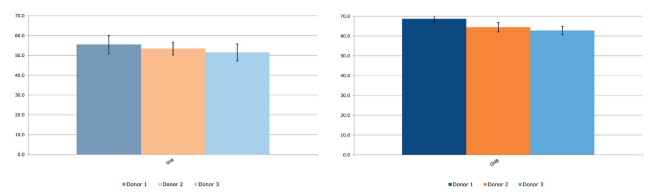


Figure 2. Typical analyte % extraction recoveries (n=7) using the ISOLUTE $^{\circ}$ SLE+ protocol using 200 μ L capacity plate format (left) and 400 μ L capacity column format (right)

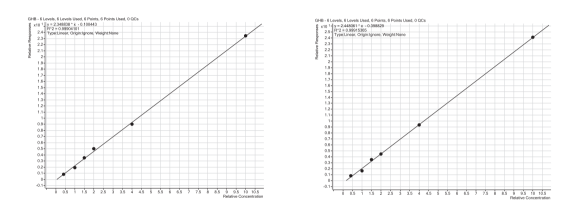


Figure 3. Calibration curves for extracted levels of spiked urine, using ISOLUTE* SLE+ protocol on 200 μL capacity plate format (left) and 400 μL capacity column format (right). Concentrations are 2.5, 5, 7.5, 10, 20 and 50 μg/mL. r² values are greater than 0.999.

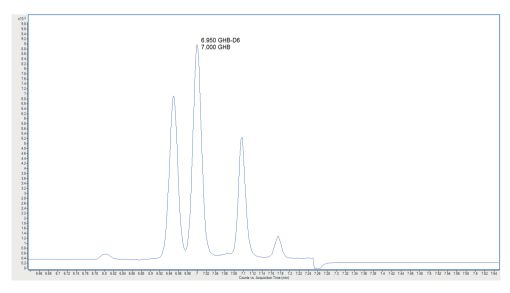


Figure 4. GC/MS chromatography for urine (Cal 1) spiked at 2.5 ng/mL. Compared to other more traditional solid phase extraction procedures, this ISOLUTE* SLE+ method results in a baseline with reduced noise and thus offers greater confidence when qualifying GHB.



Additional Information

0.2% formic acid (aq) is prepared by adding $100~\mu L$ concentrated formic acid (commercially available 98%) to 49.9~mL HPLC grade water

Ordering Information

Part Number	Description	Quantity
820-0200-P01	ISOLUTE® SLE+ 200 μL Supported Liquid Extraction Plate	1
820-0055-B	ISOLUTE® SLE+ 400 μL Sample Volume Columns	50
PPM-48	Biotage® PRESSURE+ 48 Positive Pressure Manifold	1
SD-9600-DHS-EU	Biotage® SPE Dry Sample Concentrator System 220/240 V	1
SD-9600-DHS-NA	Biotage® SPE Dry Sample Concentrator System 100/120 V	1
C103198	TurboVap° LV, 100/120V	1
C103199	TurboVap* LV, 220/240V	1

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