Automated Extraction of 11-nor-9-Carboxy-Δ⁹-THC from Hydrolyzed Urine Using ISOLUTE[®] SLE+ Prior to GC/MS Analysis

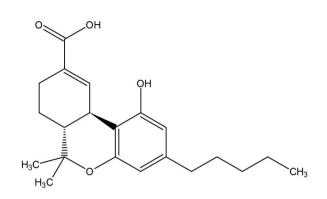


Figure 1. Structure of 11-nor-9-carboxy- Δ^9 -THC.

Introduction

This application note describes the fully automated extraction of carboxy-THC from urine, following base hydrolysis prior to GC/MS analysis. The method was automated using Biotage[®] Extrahera[™], configured for use with ISOLUTE SLE+ columns.

This application note describes an effective and efficient ISOLUTE SLE+ protocol optimized for extraction of 1 mL of pre-hydrolyzed urine. The simple sample preparation procedure delivers clean extracts and analyte recoveries greater than 80% with RSDs lower than 10% for carboxy-THC and its deuterated internal standard. Using Biotage Extrahera, 24 samples are extracted in approximately 35 minutes. Limit of quantitation is below the SAMHSA/EWDTS confirmation cut off of 15 ng/mL for workplace testing applications.

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

Carboxy-THC and carboxy-THC-D₉ as internal standard

Sample Preparation Procedure

Format

ISOLUTE[®] SLE+ 1 mL Sample Volume columns (Tabless), part number 820-0140-CG.

Sample Pre-treatment

Apply 20 μ L of a 1 ng/ μ L aqueous internal standard solution to 1 mL of urine and allow to equilibrate for 1 hr at room temperature. Add 50 μ L sodium hydroxide (10N) to this urine sample and heat at 60 °C for 20 minutes. Allow to cool and add 60 μ L glacial acetic acid.

Sample Loading

Load 1 mL of the hydrolyzed sample 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.

Analyte Extraction

Apply hexane/ethyl acetate (2.5 mL, 50/50, v/v) and allow to flow under gravity for 5 minutes. Apply a further aliquot of hexane/ethyl acetate (2.5 mL, 50/50, v/v) 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 and Derivatization

Dry the extract in a stream of air or nitrogen using a Biotage[®] TurboVap[®] (1.2 L/min at 40 °C for 25 mins).

Reconstitute the extracts with 250 μ L ethyl acetate and vortex for 10 seconds before transferring to high recovery GC vials. Dry the extract in a stream of air or nitrogen using a TurboVap (1.2 L/min at 40 °C for 10 mins).

Upon dryness, reconstitute with 20 μL ethyl acetate and 20 μL BSTFA:TMCS 99:1 and vortex for 20 seconds. Place in a heating block set to 70 °C, for 25 minutes. Remove vial from the block and allow to cool.



GC Conditions

Instrument

Agilent 7890A with QuickSwap

Column

Restek Rxi-5ms, 30 m x 0.25 mm ID x 0.25 µm

Carrier

Helium 1.2 mL/min (constant flow)

Inlet

280 °C, Splitless, purge flow: 50 mL/min at 1.0 min

Injection Volume

2 µL

Wash Solvents

Methanol and ethyl acetate

Oven

Initial temperature 125 °C

Ramp 50 °C/min to 300 °C, hold for 2.5 minutes

Ramp 50 °C/min to 330 °C, hold for 1.4 minutes

Post Run

Backflush 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

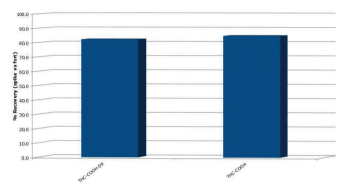
SIM Parameters

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

SIM Group	Analyte	Target (Quant) Ion	1st Qual Ion	2nd Qual Ion
1	THC-COOH-D ₉	380	479	
1	THC-COOH	371	488	473

Results

This optimized SLE+ protocol demonstrated analyte recoveries of 82% and 85% from urine for the carboxy-THC-D₉ and carboxy-THC respectively, as shown in Figure 2. RSDs were lower than 10% (n=7).





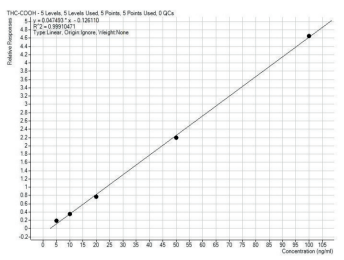


Figure 3. A calibration curve of carboxy-THC constructed following extractions using ISOLUTE SLE+ with the optimal protocol. Analyte concentrations shown here are 5, 10, 20, 50 and 100 ng/mL with r² values of greater than 0.999. The carboxy-D₉ deuterated internal standard concentrations is at 20 ng/mL.

The analyte signal allows an approximate inferred limit of quantitation of between 5 and 10 $\rm ng/mL.$





Additional information

- **Sodium Hydroxide 10N** is prepared with 40 g of pellets » in 100 mL deionized water. Prepare this solution with extreme care. As an added precaution, place the beaker on ice prior to gradual pellet addition.
- » Glacial Acetic Acid was purchased from Sigma-Aldrich at ≥ 99.85%
- » The use of MTBE is an acceptable alternative to hexane/ethyl acetate (50/50, v/v)
- » For manual extraction protocols, ISOLUTE® SLE+ 1 mL Sample Volume columns are available in tabbed format (p/n 820-0140-C) and can be used as an alternative to the tabless version used in this Biotage[®] Extrahera[™] automated protocol.

Ordering Information

Part Number	Description	Quantity
820-0140-CG	ISOLUTE® SLE+ 1 mL Sample Volume Column (Tabless)	30
414001	Biotage [®] Extrahera [™]	1
415041	Configuration Kit 24 Positions, Dual Flow	1
415000	TurboVap® LV Evaporator	1

EUROPE

Main Office: +46 18 565900 Toll Free: +800 18 565710 Fax: +46 18 591922 Order Tel: +46 18 565710 Order Fax: +46 18 565705 order@biotage.com Support Tel: +46 18 56 59 11 Support Fax: + 46 18 56 57 11 Outside US: +1 704 654 4900

NORTH & LATIN AMERICA

Main Office: +1 704 654 4900 Tel: +81 3 5627 3123 Toll Free: +1 800 446 4752 Fax: +1 704 654 4917 Order Tel: +1 704 654 4900 Order Fax: +1 434 296 8217 ordermailbox@biotage.com Support Tel: +1 800 446 4752 eu-1-pointsupport@biotage.com us-1-pointsupport@biotage.com

JAPAN

Fax: +81 3 5627 3121 jp_order@biotage.com

CHINA

Tel: +86 21 68162810 Fax: +86 21 68162829 cn_order@biotage.com

KOREA

Tel: + 82 31 706 8500 Fax:+ 82 31 706 8510 korea_info@biotage.com jp-1-pointsupport@biotage.com cn-1-pointsupport@biotage.com kr-1-pointsupport@biotage.com

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