# APPLICATIONS



# A Sensitive and Fast LC-MS/MS Method for Measurement of Nicotine and Metabolites in Human Urine

Shuguang Li<sup>1</sup>, Seyed Sadjadi<sup>1</sup>, Carrie J. Haglock<sup>2</sup>, Simon Lomas<sup>1</sup> and Jeff Layne<sup>1</sup> <sup>1</sup>Phenomenex, Inc., 411 Madrid Ave., Torrance, CA 90501 USA <sup>2</sup>ARUP Laboratories, 500 Chipeta Way, Salt Lake City, UT 84108 USA

A method is illustrated for simultaneous analysis of nicotine, two of its major metabolites, cotinine and 3-hydroxycotinine, as well as nornicotine and anabasine from human urine samples. The method described uses Strata<sup>®</sup>-X-C solid phase extraction (SPE) cartridges for sample clean-up and concentration, followed by fast (<6 min) LC/MS/MS analysis using a Gemini<sup>®</sup> NX-C18 column.

### Introduction

0

Revision:

PHEN-RUO-00029

TN-1161

Nicotine is the most abundant alkaloid present in all tobacco products along with being a major tobacco-specific component in both mainstream tobacco smoke and environmental tobacco smoke (ETS). Determination of nicotine metabolism / pharmacokinetics provides a useful tool for estimating uptake of nicotine and tobacco-related toxicants, understanding the pharmacologic effects of nicotine and nicotine addiction. (Xu et al 2004).

In addition to nicotine and its metabolites, tobacco products also contain other alkaloids that can serve as unique markers of tobacco use. Two such examples are anabasine and nornicotine, which are present in tobacco products but not in nicotine replacement therapies.

Our goal was to develop a sensitive, specific, accurate and fast analytical method to simultaneously quantify nicotine and metabolites in human urine using SPE for sample cleanup and concentration, and fast LC/MS/MS analysis using a Gemini NX-C18 column.

### **Materials and Methods**

All reagents and solvents were HPLC or analytical grade. HPLC grade methanol and acetonitrile was purchased from Honeywell, Burdick & Jackson (Muskegon, MI). Milli-Q Water was used for reagents preparation, SPE, sample preparation and to prepare the LC mobile phase. Anabasine, Cotinine, Nicotine, Nornicotine, and Ammonium Bicarbonate was purchased from J.T. Baker, Inc., (Phillipsburg, NJ). Ammonium hydroxide was purchased from Sigma-Aldrich. Trans-3'-Hydroxycotinine-d3 was purchased from Toronto Research Chemicals, Inc. (North York, ON, Canada). Nicotine-d4 was purchased from C/D/N Isotopes, (Quebec, Canada). Cotinine-d4 was purchased from C/D/N Isotopes, (Quebec, Canada). Nornicotine-d4 was purchased from C/D/N Isotopes, (Quebec, Canada).

An Agilent 1200 Series HPLC (Agilent<sup>®</sup> Technologies Inc., Santa Clara, CA USA) was interfaced with API 4000<sup>™</sup> MS/MS with ESI TurbolonSpray<sup>®</sup> (AB SCIEX Foster City, USA) operated in positive ionization mode (ESI+).

### **Sample Preparation**

The individual deuterated internal standard (IS) stock solutions (1000 ng/mL) were prepared in acetonitrile and stored at 4 °C until use. A working solution containing all IS at a final concentration of 100 ng/mL was prepared in acetonitrile. 100  $\mu$ L of the IS working solution was then added to 1 mL of diluted human urine specimen, yielding a final urine IS concentration of 100 ng/mL.

For calibrators, a stock solution of 1 mg/mL of each analyte was prepared in acetonitrile and stored at 4°C until use. The working solution of 25 µg/mL was prepared by dilution with methanol. A standard calibration curve was generated by spiking different aliquots of the working stock solution into blank human urine specimen; yielding a nine-point calibration curve (1, 2.5, 5, 10, 25, 50, 100, 250, and 500 ng/mL).

Three human urine quality control (QC) samples were prepared in triplicates with a different lot of working standard solution to yield QC concentration of 4, 40 and 400 ng/mL, respectively. The samples were prepared for analysis using the SPE procedure.

The urine sample was prepared by diluting  $0.5 \,\text{mL}$  urine samples with  $0.5 \,\text{mL}$  of 20 mM ammonium acetate, pH 4, and adding 100  $\mu$ L internal standards.

### Solid Phase Extraction (SPE)

	=,
Cartridge:	Strata-X-C (60 mg/3 mL)
Part No.:	8B-S029-UBJ
Condition:	2 mL Methanol (1-2 mL/min)
Equilibrate:	2 mL Ammonium acetate buffer
Load:	0.5 mL Diluted urine sample
Wash 1:	2 mL Ammonium acetate buffer
Wash 2:	2 mL 30 % Methanol in water
Dry:	> 10" Hg for 5 min to remove residual water
Elute:	2 x 2 mL 1.5 % Ammonium hydroxide in methanol
Dry down:	Nitrogen gas at 55 °C
Reconstitute:	500 µL of Acetonitrile/20 mM Ammonium bicarbonate (10:90)

### LC/MS/MS

Column:	Gemini 3 µm N	(-C18					
Dimensions:	50 x 2.0 mm						
Part No.:	00B-4453-B0						
Mobile Phase:	A: 20 mM Amm B: Acetonitrile	onium bicarbonate					
Gradient:	Time (min)	B (%)					
	0	10					
	3	75					
	3.1	10					
	5	10					
Flow Rate:	0.5 mL/min						
Temperature:	25 °C						
Injection:	10 µL						
Detection:	AB SCIEX API 4	000 <sup>™</sup> MS/MS (ESI+)					

### 

# APPLICATIONS

### MS/MS Conditions

IOIIIZuuoiii	LUI
Polarity:	Positive
Scan Type:	MRM
Curtain Gas (CUR):	50
Gas 1 (GS1):	50
Gas 2 (GS2):	50
IS:	5500
Collision Gas (CAD):	5
Interface Heater (Ihe):	0n
Temperature (TEM):	550 °C
Entrance Potential (EP):	10

### Table 1.

Mass Dependant Parameters

Compounds	Q1 Mass (amu)	Q3 Mass (amu)	Dwell Time (msec)	DP	CE	СХР
Nicotine	163.1	132.1	50	56	23	14
Nicotine-d4	167.1	136.0	50	56	21	14
Cotinine	177.1	80.1	50	70	31	8
Cotinine-d3	180.1	80.1	50	70	31	8
Nornicotine	149.1	80.1	50	71	29	8
Nornicotine-d4	153.1	84.1	50	70	29	8
3-0H-cotinine	193.1	80.1	50	70	31	8
3-0H-cotinine-d3	196.1	80.1	50	66	35	8
Anabasine	163.1	120.1	50	70	31	8

### **Results and Discussion**

The use of the rugged pH stable Gemini® NX-C18 column allowed for fast elution of nicotine and its metabolites in less than 3 minutes (Figure 1). This fast separation allows for multiplexing techniques to handle the analysis of large numbers of samples. In ESI positive mode, nicotine and its metabolites were detected by monitoring the mass transitions and their deuterium-labeled internal standards listed in Table 1. The most abundant transition for each analyte was used for quantification. The second mass transition (not listed) served as a confirmation for each analyte. Figure 2 shows the extracted ion chromatograms for nicotine, cotinine, 3-hydroxycotinine, nornicotine and anabasine at concentrations of 10 ng/mL in extracted human urine. Note that anabasine and nicotine have the same parent mass, accounting for the 3rd peak in the anabasine XIC (Figure 2). Since the deuterium-labeled internal standard was not available for anabasine, cotinine-d3 served as its internal standard because it has the nearest retention time (Figure 2). The pH of the mobile phase was adjusted so that anabasine and cotinine elute as closely as possible. Anabasine can be separated at a slightly higher pH (pH 9) if the deuterated internal standard were to become available.

### Figure 1.

Nicotine, Cotinine, 3-OH-cotinine, Nornicotine and Anabasine analysis (10 ng/mL urine extracted standard)



### Figure 2.

Extracted Ion Chromatograms for Nicotine, Cotinine, 3-OH-cotinine, Nornicotine and Anabasine at a concentration of 10 ng/mL in human urine



## APPLICATIONS



### **Results and Discussion (con't)**

Standard calibration curves were generated over the concentration range of 1.0 ng/mL to 500 ng/mL by plotting the relative response (peak area of nicotine and its metabolites / peak area of internal standards) versus concentration. The standard calibration curves were linear over the calibration ranges with R<sup>2</sup> values of 0.9997 for nicotine, 0.9979 for cotinine, 0.9997 for nornicotine, 0.9973 for 3-OH-cotinine and 0.9995 for anabasine, separately (**Figure 4**).

Sensitivity of the method was evaluated by determining the lowest level concentration with a signal to noise of at least 10:1 for limit of quantification (LOQ). At the lowest level standard concentration (1.0 ng/mL) the signal-to-noise ratios were 46.1 for nicotine, 141.5 for cotinine, 16.1 for nornicotine, 50.7 for 3-OH-cotinine and 24.5 for anabasine, separately (**Figure 3** and **Table 2**). Therefore, the LOQs were estimated to be < 1.0 ng/mL for nicotine, cotinine, nornicotine, 3-OH-cotinine and anabasine (**Table 2**). Furthermore, the current method achieved additional sensitivity by reducing specimen size from 1000 µL to 500 µL urine. If greater sensitivity is required or if the intended detector cannot meet the 1 ng/mL sensitivity, reducing the final elution volume by half is an easy approach. The upper limit of quantification was 500 ng/mL for all the analytes (**Table 2**).

#### Table 2.

### Statistical Data of Nicotine and Metabolites in Urine by LC/MS/MS

Analyte	1.00	111.00	Y-intercept	R <sup>2</sup>	Intra Assay Precision % (N = 3)					S/N	RT	
	LUQ	ULUQ			4 ng/mL		40 ng/mL		400 ng/mL		1 ng/	min
	ng/mL	ng/mL			Mean	% CV	Mean % CV		Mean	% CV mL		
Nicotine	1	500	0.0495x +0.0503	0.9997	96.13	3.72	105.67	2023	102.00	0.80	46.10	2.31
Cotinine	1	500	0.0386x +0.0372	0.9979	94.84	3.06	100.00	0.99	101.37	3.98	141.50	1.73
3-0H-cotinine	1	500	0.0398x +0.044	0.9973	103.10	5.85	98.20	1.92	107.33	4.95	50.70	1.16
Nornicotine	1	500	0.0285x +0.0233	0.9997	101.53	5.94	100.67	1.66	105.43	6.09	16.10	1.08
Anabasine	1	500	0.0132x +0.0311	0.9995	97.03	2.77	107.00	2.64	95.53	4.67	24.50	1.71

LOQ = Limit of quantification

ULOQ = Upper limit of quantification

**CV** = Coefficient of Variation (Standard Deviation / Mean)

S/N = Signal / Noise

RT = Retention time



### **TN-1161**



**APPLICATIONS** 



Nicotine and metabolites analysis (1 ng/mL extracted standard)





### TN-1161

**APPLICATIONS** 

### Figure 4.

?

Standard curves from 1 ng/mL to 500 ng/mL for Nicotine, Cotinine, Nornicotine, 3-OH-cotinine and Anabasine





Ophenomenex

...breaking with tradition<sup>™</sup>

## APPLICATIONS



### **Results and Discussion (con't)**

Three levels of QC samples were prepared at 4, 40 and 400 ng/mL. These concentrations were selected to represent low, medium, and high concentration across the calibration range for each analyte. The three level QC samples were extracted in the same way as the actual sample described above and analyzed in triplicates to assess reproducibility. The mean expected recovery of the lowest level QC samples at 4 ng/mL was 96.13 % for nicotine, 94.80 % for cotinine, 103.10 % for 3-OH-cotinine, 101.53 % for nornicotine and 97.03 % for anabasine (**Table 2**).

The percentage Coefficient of Variation (%CV) for the intra assay precision were from 2.77% to 5.85% for levels of 4 ng/mL, 0.99% to 2.64% for levels of 40 ng/mL and 0.8% to 6.09% for levels of 400 ng/mL, respectively (**Table 2**). No endogenous signal was found in 3 nonsmoker urine specimens, demonstrating the selectivity of the method. There were no carryovers observed by injecting blank urine samples after the highest calibrator (500 ng/mL).

The chromatogram in Figure 5 illustrates a urine/matrix suppression study when using an ESI source and represents a post-column infusion of high concentration standards of nicotine, cotinine, 3-OH-cotinine, nornicotine and anabasine, while a low level urine extract is injected on column. The top trace contains the MRM transitions for all five analytes and the bottom trace is the four internal standards channel (nornicotine-d4, 3-OH-cotinined3, cotinine-d3 and nicotine-d4). As expected, this section of the chromatogram contains highly polar components of the extract and is virtually un-retained. Nicotine, its metabolites and their internal standards elute between 1 to 2.5 min, in the region of stability between 0.8 - 2.8 min. The region where the mobile phase is mostly organic, 2.9 – 4.5 min, produces the most dramatic signal increase. This is to be expected as the ionization source becomes more productive with a lower viscosity and lower boiling point solvent.

#### Figure 5.



Urine matrix effect on Nicotine, its metabolites and their IS's response using an ESI source

We optimized the SPE extraction procedure to allow for the best recovery of the urinary nicotine and its metabolites and to have the greatest potential for high-throughput sample preparation and automation. The Strata®-X-C sorbent represented the most durable and selective material.

### Conclusion

A rapid analysis of urinary nicotine and its metabolites was developed. The application of the Gemini<sup>®</sup> NX-C18 column in this method results in a shorter chromatographic analysis time, providing a productivity benefit for clinical testing laboratories with a dramatic increase in efficiency while simultaneously reducing costs due to solvent consumption. Sample preparation using Strata-X-C SPE concentrates the nicotine and metabolites and removes potential sample matrix interferences, which coupled with high efficiency Gemini NX-C18 column provides for the low level detection and quantitation of nicotine and its metabolites in human urine.

#### References

- R.A. Davis, M. Curvall. Determination of nicotine and its metabolites in biological fluids: In vivo studies. In: J.W. Gorrod P. Jocob, eds. Analytical determination of nicotine and related compounds and their metabolites. Amsterdam: Elsevier Science, **1999**, 583-643.
- 2. Kyerematen GA, Vesell ES. Drug Metab. Rev., 1991, 23, 3.
- 3. Kakajima M, Drug Metab. Dispos., 1996, 24, 1212.
- 4. Matt GE. Biomarkers, 2006, 11, 507-523.
- 5. T. Tuomi, T. Johnsson, K. Reijula. Clin. Chem. 1999, 723, 185-94.
- 6. X Xu, M.M. Iba, C. P. Weisel. Clin. Chem., 2004, 50, 2323-2330.

Ophenomenex ...breaking with tradition™

### **Ordering Information**

PLICATIONS

### **Gemini® HPLC Columns**

3 µm Micro	bore, Minibore and	d Narrow Bore Co	olumns (mm)					Security	Guard™ Cartridges (mm)
Phases	20 x 2.0	30 x 2.0	50 x 2.0	100 x 2.0	150 x 2.0	50 x 3.0	100 x 3	8.0 150 x 3	3.0 4 x 2.0*
NX-C18	00M-4453-B0	00A-4453-B0	00B-4453-B0	00D-4453-B0 0	0F-4453-B0	00B-4453-	Y0 00D-4453	3-Y0 00F-445	3-Y0 AJ0-8367
									for ID: 2.0-3.0 mm
3 µm Analy	tical Columns (mn	n)		Security	Guard Cartrid	lges (mm)			
Phases	50 x 4.6	100 x 4.6	150 x 4.6	250 x 4.6	4 x 3	.0*			
NX-C18	00B-4453-E0	00D-4453-E0	00F-4453-E0	00G-4453-E0	AJ0-8	368			
					for ID: 3.2-	-8.0 mm			
5 µm Minib	ore and Narrow Bo	ore Columns (mn	1)					Security	Guard Cartridges (mm)
Phases	30 x 2.0	50 x 2.0	150 x 2.0	50 x 3.0	100 >	c 3.0	150 x 3.0	250 x 3.0	4 x 2.0*
NX-C18	00A-4454-B0	00B-4454-B0	) 00F-4454-B	00B-4454-Y	′0 00D-44	54-Y0 (	0F-4454-Y0	00G-4454-Y0	AJ0-8368
									for ID: 2.0-3.0 mm
5 µm Analy	rtical Columns (mm	n)		SecurityGuard C	artridges (mn	n)		For Gerr	nini Capillary or Prepar
Phases	50 x 4.6	100 x 4.6	150 x 4.6	250 x 4.6	4 x 3.0*			Columns	s, Guards, and Adapte
NX-C18	00B-4454-E0	00D-4454-E0	00F-4454-E0 0	00G-4454-E0	AJ0-8368			Phenom	enex technical consul
				for	ID: 3 2-8 0 mn	n		distribut	or.

for ID: 3.2-8.0 mm

### **Ordering Information**

Strata <sup>®</sup> -X-C Solid Phase Extraction (SPE)									
Format	Sorbent Mass	Part Number	Unit						
Microelution 96-V	Vell								
	2 mg	8M-S029-4GA	1 Plate/Box						
96-Well Plate									
	10 mg	8E-S029-AGB	2 Plates/Box						
145	30 mg	8E-S029-TGB	2 Plates/Box						
1	60 mg	8E-S029-UGB	2 Plates/Box						
Tube									
	30 mg	8B-S029-TAK*	1 mL(100/box)						
	30 mg	8B-S029-TBJ	3 mL (50/box)						
	60 mg	8B-S029-UBJ*	3 mL (50/box)						
	100 mg	8B-S029-EBJ	3 mL (50/box)						
	100 mg	8B-S029-ECH	6 mL (30/box)						
	200 mg	8B-S029-FBJ	3 mL (50/box)						
	200 mg	8B-S029-FCH	6 mL (30/box)						
	500 mg	8B-S029-HBJ	3 mL (50/box)						
	500 mg	8B-S029-HCH	6 mL (30/box)						

\*Tab-less tubes available. Contact Phenomenex for details.



If Phenomenex products in this technical note do not provide at least an equivalent separation as compared to other products of the same phase and dimensions, return the product with comparative data within 45 days for a FULL REFUND



## PI ICATIONS



**Australia** t: +61 (0)2-9428-6444

auinfo@phenomenex.com

Austria t: +43 (0)1-319-1301 anfrage@phenomenex.com

Belaium t: +32 (0)2 503 4015 (French) t: +32 (0)2 511 8666 (Dutch) beinfo@phenomenex.com

Canada t: +1 (800) 543-3681 info@phenomenex.com

**China** t: +86 400-606-8099 cninfo@phenomenex.com

**Denmark** t: +45 4824 8048 nordicinfo@phenomenex.com

Finland t: +358 (0)9 4789 0063 nordicinfo@phenomenex.com

France t: +33 (0)1 30 09 21 10 franceinfo@phenomenex.com

**Germany** t: +49 (0)6021-58830-0 anfrage@phenomenex.com

India t: +91 (0)40-3012 2400 indiainfo@phenomenex.com

Ireland t: +353 (0)1 247 5405 eireinfo@phenomenex.com

Italy t: +39 051 6327511 italiainfo@phenomenex.com

Luxembourg t: +31 (0)30-2418700

### nlinfo@phenomenex.com

www.phenomenex.com

Phenomenex products are available worldwide. For the distributor in your country, contact Phenomenex USA, International Department at international@phenomenex.com

#### Mexico

t: 01-800-844-5226 tecnicomx@phenomenex.com

The Netherlands

t: +31 (0)30-2418700 nlinfo@phenomenex.com

**New Zealand** t: +64 (0)9-4780951 nzinfo@phenomenex.com

Norway t: +47 810 02 005

- nordicinfo@phenomenex.com
- Portugal
- t: +351 221 450 488 ptinfo@phenomenex.com

### Singapore

- t: +65 800-852-3944 sginfo@phenomenex.com
- **Spain** t: +34 91-413-8613
- espinfo@phenomenex.com
- **Sweden** t: +46 (0)8 611 6950
- nordicinfo@phenomenex.com

#### Switzerland t: +41 61 692 20 20

swissinfo@phenomenex.com

### United Kingdom

- t: +44 (0)1625-501367 ukinfo@phenomenex.com
- USA t: +1 (310) 212-0555 info@phenomenex.com

### All other countries Corporate Office USA

- info@phenomenex.com

### Terms and Conditions

Subject to Phenomenex Standard Terms and Conditions, which may be viewed at http://www.phenomenex.com/TermsAndConditions.

### Trademarks

Gemini and Phenomenex are registered trademarks and SecurityGuard, Strata-X, and Giga are trademarks of Phenomenex. Agilent is a registered trademark of Agilent Technologies. TurbolonSpray is a registered trademark and API 4000 is a trademark of AB SCIEX.

Disclaimer

Comparative separations may not be representative of all applications. Phenomenex is not affiliated with Agilent Technologies.

Gemini is patented by Phenomenex, U.S. Patent No. 7.563.367

Strata-X is patented by Phenomenex. U.S. Patent No. 7,119,145

FOR RESEARCH USE ONLY. Not for use in clinical diagnostic procedures. © 2018 Phenomenex, Inc. All rights reserved.