

## STYRE SCREEN®

POLYMERIC SORBENT SPE



INNOVATION THROUGH CHEMISTRY

# STYRE SCREEN® Extraction Columns

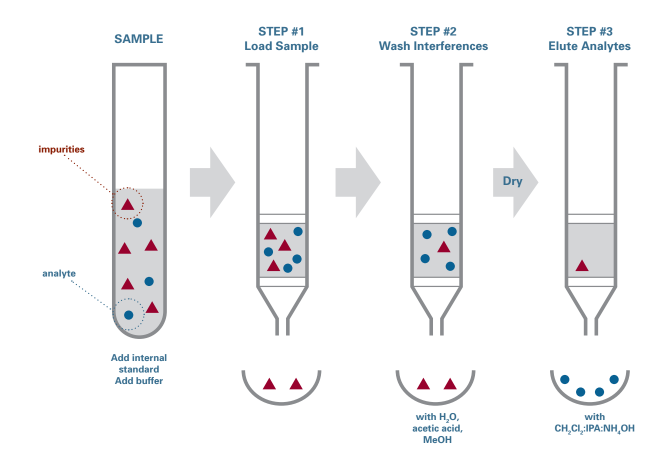
Styre Screen® extraction sorbents are formulated with an ultra clean, highly cross-linked styrene and divinylbenzene copolymer sorbent. The sorbent can be functionalized with any of the same phases as our silica based sorbents. Possibilities include standard hydrophilic, hydrophobic, or ion exchange functionalities as well as copolymeric phases such as the DBX phase. Styre Screen® particles have an average particle size of 30 microns. This polymeric sorbent has a very high analyte capacity, ideal for standard solid phase extraction applications. This higher capacity translates into a lower bed mass requirement in order to retain the same analyte quantity as a traditional silica particle. Lower bed mass also means extractions can be run at faster flow rates and with less solvent usage. The Styre Screen® sorbent also eliminates the need for an initial column conditioning step. All these attributes ultimately result in excellent cost benefit.

## **Advantages:**

- No conditioning step
- High and reproducible recoveries
- highly cross-linked sorbent minimizes bead swelling
- Reduced sorbent mass
- Improved flow rates
- pH stable from 1 14
- Reduced solvent use
- High sorbent capacity
- Methods for NIDA/SAMHSA 5 Drugs



## **STYRE SCREEN® General Application**



## STYRE SCREEN® DVB - Polystyrene Divinylbenzene

**Application:** Retention of neutral and aromatic compounds, useful for screening applications where a broad range of analytes is to be extracted

### Structure:



COLUMNS						
Tube Volume (mL		Sorbent Amount (mg)		Units per Pack	Part Number	
1	1	0		100	SSDVB0X1	
1	3	0		100	SSDVB031	
1	1	00		100	SSDVB111	
3	3	0	)		SSDVB033	
6	5	50		50	SSDVB056	
6	2	200		30	SSDVB206	
6	5	00		30	SSDVB506	
10	1	00		50	SSDVB11Z	
		WEL	L PLA	ГЕ		
Number of wells	Sorbent Amount (mg)	Units per	pack	Extended Drip Tip	Part Number	
48	60	1		NO	WSH48DVB406	
96	30	1		NO	WSHDVB403	
96	50	1		NO	WSHDVB405	
96	60	1		NO	WSHDVB406	

## STYRE SCREEN® DBX – Octadecyl (C18) and Benzenesulfonic Acid – Mixed Mode

Application: Retention of weakly basic and hydrophobic compounds



$$\begin{array}{c} \left( \begin{array}{c} PS/DVB \end{array} \right) & O \\ & \parallel \\ -S - O^{\odot} \\ \end{array} \\ \begin{array}{c} \\ O \end{array}$$

COLUMNS						
Tube Volume (mL)	)	Sorbe Amount			Units per Pack	Part Number
1		30			100	SSDBX031
3		30			50	SSDBX033
3		30			500	SSDBX033-D
3		60			50	SSDBX063
6		50			50	SSDBX056
6		50			500	SSDBX056-D
6		150	150		50	SSDBX(150)06
6		200	)		50	SSDBX206
10		50	50 50 SSDBX05Z		SSDBX05Z	
WELL PLATE						
Number of wells	Sorbent A	Amount (mg)	Units per	pack	Extended Drip Tip	Part Number
96	;	30	1		NO	WSHDBX403

### Structure:

## STYRE SCREEN® BCX – Benzensulfonic Acid – Cation Exchange

**Application:** Retention of weakly basic compounds

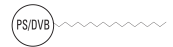
PS/DVB	0   
	U

COLUMNS						
Tube Volume (mL)	Sorbent Amount (mg)	Units per Pack	Part Number			
1	30	100	SSBCX031			
3	30	50	SSBCX033			
3	60	50	SSBCX063			
6	50	50	SSBCX056			

### Structure:

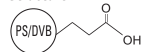
### **STYRE SCREEN® C18 – Reverse Phase**

**Application**: Retention of hydrophobic compounds



COLUMNS						
Tube Volume (mL)	Sorbent Amount (mg)	Units per Pack	Part Number			
1	30	100	SSC18031			
3	30	50	SSC18033			
6	50	50	SSC18056			
6	200	50	SSC18206			
6	300	50	SSC18306			
6	500	50	SSC18506			
75	5000	10	SSC1815M75			

### Structure:



## **STYRE SCREEN® CCX – Carboxylic Acid – Cation Exchange**

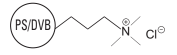
Application: Retention of basic compounds, particulary strong bases

	COLUMNS						
Vol	Tube ume (mL)	Sorbent Amount (mg)	Units per Pack	Part Number			
	1	30	100	SSCCX031			
	3	30	50	SSCCX033			
	3	50	50	SSCCX053			
	3	60	50	SSCCX063			
6		50	50	SSCCX056			
WELL PLATE							
Number of Wells	Sorbent Amount (mg)	Units per Pack	Extended Drip Tip	Part Number			
96	30	1	NO	WSHSSCCX103			

### Structure:

## **STYRE SCREEN® QAX – Quaternary Amine – Anion Exchange**

Application: Retention of weakly acidic compounds



**Structure: Proprietary** 

COLUMNS						
Tube Volume (mL)	Sorbent Amount (mg)	Units per Pack	Part Number			
1	30	100	SSQAX031			
3	30	50	SSQAX033			
6	50	50	SSQAX056			
6	150	50	SSQAX(150)06			

### STYRE SCREEN® THC

Application: Retention of THC and THC metabolites (THC-delta-9,

THC-hydroxy metabolite and THC-carboxy metabolite)

COLUMNS					
Tube Volume (mL)	Sorbent Amount (mg)	Units per Pack	Part Number		
1	30	100	SSTHC031		
3	60	50	SSTHC063		
6	60	50	SSTHC066		
10	60	50	SSTHC06Z		
6	100	50	SSTHC116		
10	100	50	SSTHC11Z		

## **COCAINE AND BENZOYLECGONINE IN URINE**

### Part #

SSDBX033 – STYRE SCREEN® DBX 30 mg, 3 mL Tube SLDA50ID21-5UM – SELECTRA® DA HPLC Column 50 x 2.1 mm, 5 µm

#### 1. PREPARE SAMPLE:

To 1 mL of urine add internal standard(s) and 300  $\mu$ L 100mM HCl. Mix/Vortex.

### 2. APPLY SAMPLE:

Load at 1 to 2 mL/minute

### 3. WASH COLUMN:

 $1 \times 1 \text{ mL D.I. H}_2\text{O}$ 

1 x 1 mL 100 mM HCI

1 x 1 mL CH<sub>3</sub>OH

Dry column (10 minutes at full vacuum or pressure)

### 4. ELUTE COCAINE/METABOLITE:

 $2 \times 0.5 \text{ mL CH}_2\text{Cl}_2/\text{ IPA /NH}_4\text{OH (78:20:2)}$ 

Collect eluate at 1 to 2 mL/minute

NOTE: Prepare elution solvent daily. Add IPA /NH $_4$ OH,

mix, then add  $CH_2Cl_2$  (pH 11-12)

### **5. DRY ELUATE:**

Evaporate to dryness at < 40 °C

### **6. RECONSTITUTE**

Reconstitute sample in 100 µL of mobile phase

### **PARAMETERS**

Instrument: API 3200 Qtrap MS/MS with Shimadzu Prominence UFLC HPLC Column: SELECTRA® DA HPLC Column 50 x 2.1mm, 5 µm

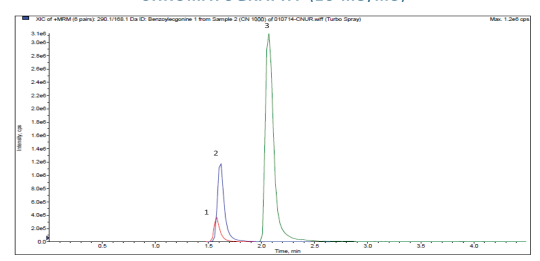
Polarity: Positive Injection Volume: 10 μL Flow Rate: 0.7 mL/minute

Mobile Phase A: 0.1% formic acid in water Mobile Phase B: 0.1% formic acid in MeOH

**Gradient:** 

Time	%A	%B	
0.00	75	25	
3.00	50	50	
3.01	10	90	
4.00	75	25	
5.50	STOP		

### CHROMATOGRAPHY (LC-MS/MS)



Analyte		MRM Tra	ansitions	Relative
		Q1	<b>O</b> 3	Retention Time (minutes)
1.	Benzolyecgonine D <sub>8</sub>	298.1	171.1	1.58
2.	Benzolyecgonine	290.1	168.1	1.60
3.	Cocaine	304.1	182.1	2.0

# THC, THC-OH, AND THC-COOH CONFIRMATIONS IN WHOLE BLOOD BY LC-MS/MS OR GC-MS USING 100 MG STYRE SCREEN® SSTHC

### Part #

SSTHC116 – STYRE SCREEN® THC 100 mg, 6 mL Tube SLDA100ID21 – 5UM – SELECTRA® DA HPLC Column 100 x 2.1mm, 5 µm

### 1. PREPARE SAMPLE:

To 1-2 mL whole blood add appropriate internal standards prepared in alcohol

Add drop-wise 2.5 mL Ice Cold acetonitrile

Mix thoroughly and centrifuge

Decant acetonitrile into a clean tube.

Evaporate acetonitrile under a stream or air or nitrogen to  $\sim 200 \text{ uL}$ 

Add 2 mL D.I.  $H_2O$  (pH of  $H_2O$  must be ~ 6.0-7.0)

### 2. APPLY SAMPLE:

Load at 1 to 2 mL/minute.

### 4. WASH COLUMN:

Wash with 2 mL (84: 15: 1) D.I.  $\rm H_2O$ : Acetonitrile:  $\rm NH_4OH$  (made fresh daily) Dry column under full vacuum or pressure for 10-15 minutes

### 5. ELUTE THC & metabolites:

1 x 3 mL Hexane/ Ethyl Acetate/ Glacial Acetic Acid (49: 49:2) Collect at 1-2 mL/ minute.

### 6. DRY ELUATE:

Evaporate to dryness at < 40° C.

#### 7. RECONSTITUTE / DERIVATIZE:

• LC-MS/MS: Reconstitute sample in 100  $\mu$ L of mobile phase Inject 5  $\mu$ L.

### **PARAMETERS**

Instrument: API 4000 Qtrap MS/MS with Agilent 1200 Binary Pump SL HPLC Column: SELECTRA® DA HPLC Column  $100 \times 2.1 mm$ ,  $5 \mu m$ 

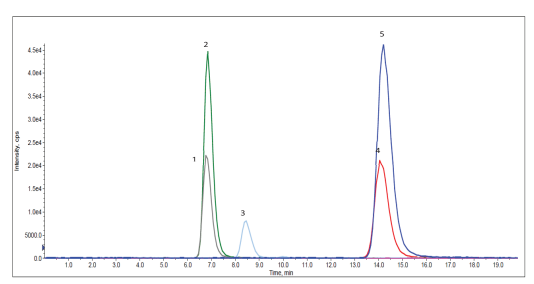
Polarity: Negative/ Positive Injection Volume: 5 µL Flow Rate: 0.5 mL/minute Reconstitute: 100 µL

Mobile Phase A: 0.1% formic acid in water Mobile Phase B: 0.1% formic acid in MeOH

**Gradient:** 

Time	%A	%B		
0.00	25	75		
20.00	STOP			

### CHROMATOGRAPHY (LC-MS/MS)



Analyte		MRM Tra	nsitions	Relative
		Q1	<b>O</b> 3	Retention Time (minutes)
1.	HYDROXY DELTA 9-THC D <sub>3</sub>	334.0	316.2	6.80
2.	HYDROXY DELTA 9-THC	330.9	313.2	6.88
3.	CARBOXY DELTA 9-THC	334.0	316.2	8.47
	CARBOXY DELTA 9-THC D <sub>3</sub>	348.3	303.0	-
4.	DELTA 9-THC D <sub>3</sub>	318.2	196.2	14.20
5.	DELTA 9-THC	315.2	196.2	14.31

# OPIATES IN BLOOD, PLASMA/SERUM, URINE, OR TISSUE

### Part #

SSDBX033 – STYRE SCREEN® DBX 30 mg, 3 mL Tube BETA-GLUC-10 – SELECTRAZYME® Beta-glucuronidase SLDA50ID21-5UM – SELECTRA® DA HPLC Column, 50 x 2.1 mm, 5 µm SPHACE4501-5 – Select pH Buffer Pouches 100mM Acetate pH 4.5

### 1. PREPARE SAMPLE:

**Blood:** To 1 mL of 100 mM phosphate buffer (pH 6.0) add internal standards.

Add 1-2 mL of blood, plasma/ serum, or 1 g (1:4) tissue homogenate.

Mix/vortex and let stand for 5 minutes

Add 2 mL of 100 mM phosphate buffer (pH 6.0). Mix/vortex

Sample pH should be  $6.0 \pm 0.5$ .

Centrifuge for 10 minutes at 2000 rpm and discard pellet

### Urine: PREPARE SAMPLE FOR ENZYME HYDROLYSIS OF GLUCURONIDES:

To 1-2 mL of urine sample, add 1 mL of acetate buffer (pH 5.0) containing

5,000 units/mL of Selectrazyme® ß-glucuronidase.

Optionally, add 1 mL of acetate buffer and 25-50 µL of concentrated

ß-glucuronidase.

Vortex and heat for 1-2 hours at 65 °C.

Allow sample to cool

Do not adjust pH~ sample is ready to be added to the extraction column.

### 2. APPLY SAMPLE:

Load at 1 to 2 mL/minute.

### 3. WASH COLUMN:

1 x 1 mL D.I. H<sub>2</sub>O.

1 x 1 mL 100 mM acetate buffer (pH 4.5).

1 x 1 mL CH<sub>3</sub>OH.

Dry column (5 minutes at full vacuum or pressure).

### 4. ELUTE OPIATES:

2 x 0.5 mL CH<sub>2</sub>Cl<sub>2</sub>/ IPA/ NH<sub>4</sub>OH (78:20:2)

Collect eluate at 1 to 2 mL/minute.

NOTE: Prepare elution solvent daily. Add IPA/ NH<sub>4</sub>OH, mix, then add CH<sub>2</sub>Cl<sub>2</sub> (pH 11-12).

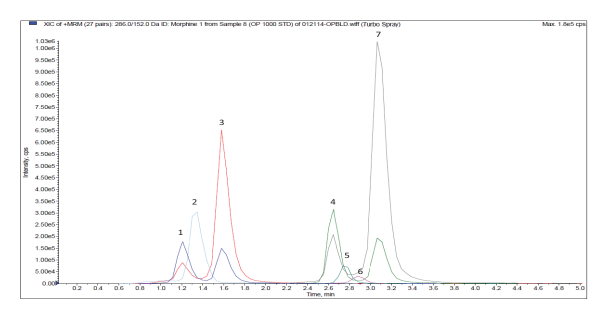
### 5. DRY ELUATE:

Evaporate to dryness at < 40 °C.

### 6. RECONSTITUTE:

Reconstitute sample in 100 µL of mobile phase

### CHROMATOGRAPHY (LC-MS/MS)



Analyte		MRM Transitions		Relative
		Q1	<b>O</b> 3	Retention Time (minutes)
1.	Morphine	286.0	152.0	1.21
2.	Oxymorphone	302.0	227.0	1.30
3.	Hydromorphone	286.0	185.0	1.60
4.	Codeine	300.0	152.0	2.65
5.	6-MAM	328.0	165.1	2.75
6.	Oxycodone	316.0	240.0	2.85
7.	Hydrocodone	300.0	199.0	3.10

### **PARAMETERS**

Instrument: API 3200 Qtrap MS/MS with Shimadzu Prominence UFLC HPLC Column: SELECTRA® DA HPLC Column 50 x 2.1mm, 5  $\mu$ m

Polarity: Positive

Injection Volume: 10  $\mu$ L Flow Rate: 0.6 mL/minute

Mobile Phase A: 0.1% formic acid in water Mobile Phase B: 0.1% formic acid in MeOH

Gradient:

Time	%A	%B
0.00	85	15
7.00	40	60
7.01	20	80
8.00	85	15
9.00	STOP	

### Note:

Hydroxylamine can be added to sample within method if oxime derivative is preferred.

Following hydrolysis, add 200 µL 10% Hydroxylamine solution.

Heat to 60 °C for 30 min in a heating block.

Allow sample to cool then adjust pH back to 5 with 1.0 M NaOH.

Centrifuge for 10 minutes at 2000 rpm and discard pellet

Sample is now ready to be added to the extraction column

## BASIC ANALYTES IN URINE BY LC-MS/MS USING STYRE SCREEN® BCX

### Part #:

SSBCX056 – STYRE SCREEN® BCX SPE Cartridge, 50 mg / 6 mL BETA-GLUC-10 – SELECTRAZYME® Beta-glucuronidase SLDA50ID21-3UM – SELECTRA® DA HPLC Column, 50 x 2.1mm, 3µm

### 1. PREPARE SAMPLE

Hydrolysis: To 1 mL of urine sample, add 1 mL of acetate buffer (pH=5) and 50 μL of concentrated  $\beta$ -glucuronidase. Vortex and heat for 1-2 hours at 65 °C. Do not adjust pH $\sim$  sample is ready to be added to the extraction plate.

### 2. APPLY SAMPLE

Load sample directly to column without any preconditioning. Pull sample through at a rate of 1-2 mL/ minute.

### 3. WASH

1 x 1 mL 100mM Acetic Acid 1 x 1 mL MeOH. Dry column (5 mins at >10 inches Hg).

### 4. ELUTION

2 x 0.5 mL MeOH/NH $_4$ OH (98/2), collect eluate at 1 to 2 mL/min.

NOTE: Prepare elution solvent daily.

### 5. DRY ELUTE

Evaporate fraction to complete dryness under stream of dry air or Nitrogen at  $\sim$  35  $^{\circ}\text{C}.$ 

### 6. RECONSTITUTE

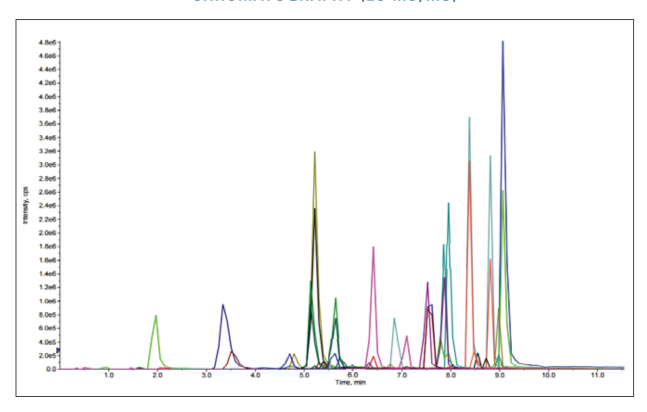
Reconstitute sample in 100 µL of mobile phase

### **ANALYTES TARGETED WITHIN METHOD**

Analyte
Morphine
Hydromorphone
Codeine
Hydrocodone
6-MAM
Bezoylecgonine
Cocaethylene
Cocaine
Ketamine
PCP
Norketamine
Amp
Methamp
MDA
MDMA
Buprenorphine

Analyte
EDDP
Methadone
Pyrovalerone
3,4MDPV
Mephedrone
Ethylone
Butylone
Fentanyl
Naltrexone
Naloxone
Tramadol
Norfentanyl
Oxymorphone
Oxycodone
Norbuprenorphine

## **CHROMATOGRAPHY (LC-MS/MS)**



### **PARAMETERS**

Instrument: Agilent 1200 Binary Pump SL

Detector: API 4000 Otrap MS/MS

**HPLC Column:** SELECTRA® DA HPLC Column 50 x 2.1mm, 3  $\mu m$ 

**Polarity:** Positive **Injection Volume:** 10 μL **Flow Rate:** 0.4 mL/minute

Mobile Phase A: 0.1% formic acid in water Mobile Phase B: 0.1% formic acid in MeOH

**Gradient:** 

Time	%A	%В		
0.00	90	10		
0.50	90	100		
4.00	60	40		
7.50	15	85		
8.50	0	100		
12.00	0	100		
12.20	90	10		
15.00	STOP			

### **PRICES AND TERMS**

Our prices are subject to change without notice. The price in effect when we receive your order will apply. All prices are in US Dollars and are F.O.B. Terms of payment are net 30 days.

### MINIMUM ORDERS

We welcome all orders, therefore, we do not have a minimum order requirement. When ordering, please include your purchase order number, complete "Ship To" and "Bill To" address, catalog number, quantity, and description of product(s). Also include your name and a phone number where you can be reached should we have any questions concerning your order.

### **SHIPMENTS**

Normal processing is within 24 hours after receipt of an order. Unless special shipping requests have been made, our trained staff will send all orders by UPS Ground service. The appropriate shipping charges (freight & insurance costs) will be added to the invoice, unless otherwise instructed by the customer.

### **SPECIAL PRICING**

We offer special pricing for volume purchases and standing orders. These discounts apply to bonded phase extraction column purchases only. Please call a sales representative for more information on special pricing qualifications.

### **RETURN POLICY**

Our Quality Manager will handle all returns. Before returning merchandise, please call to obtain a return authorization number from the quality manager. We will need to know the reason for the return, date of purchase, purchase order number and invoice number in order to issue a return authorization number. Return merchandise must be received before a credit can be issued. Returns will not be accepted after 90 days. A restocking fee of 25% of the price paid, or a minimum of \$25.00 (whichever is greater) will be charged on all returns.

### **WARRANTY**

All products manufactured by UCT are guaranteed against defects in materials and workmanship for a period of 90 days after shipment. UCT will replace any items that prove to be defective during this time period.

The exclusive remedy requires the end user to first advise UCT of the defective product by phone or in writing. Secondly, the defective product must be returned within 30 days after proper approval from our Quality Manager. All returns must indicate the purchase order number, the lot number and the shipping date. UCT's total liability is limited to the replacement cost of UCT products.

This warranty does not apply to damage resulting from misuse.

### **Contact Us**

Fax:

Phone: 215.781.9255

800.385.3153

215.785.1226

UCT, Inc.

2731 Bartram Rd. Bristol, PA 19007

Email: info@unitedchem.com









