MACHEREY-NAGEL

Modern polymeric SPE phases



- Well-defined portfolio of polymeric SPE phases
- Broad application range
- High performance adsorbents





Do you want to squeeze the best out of your samples?



CHROMABOND® HLB	Hydrophilic-lipophilic balance NVP/DVB copolymer	page 04-07
CHROMABOND® HR-X	Hydrophobic PS/DVB copolymer	page 08-09
CHROMABOND® HR-XC	Strong mixed-mode cation exchanger on PS/DVB copolymer basis	page 10-11
CHROMABOND® HR-XA	Strong mixed-mode anion exchanger on PS/DVB copolymer basis	page 12-13
CHROMABOND® HR-XCW	Weak mixed-mode cation exchanger on PS/DVB copolymer basis	page 14-15
CHROMABOND® HR-XAW	Weak mixed-mode anion exchanger on PS/DVB copolymer basis	page 16-17

Characteristics

- State-of-the-art spherical polymers with different particle sizes to suit sample volume and matrix
- Optimized pore structure and high specific surface
- High purity adsorber material
- Extremely low blind values
- High specific surface
- pH stability of 1–14

Benefits for you

Save time and reduce costs

- Well-defined portfolio of polymer phases to suit your application
- Excellent enrichment of neutral, acidic and basic compounds
- Outstanding price / performance ratio

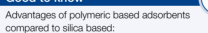
Robust methodology and less pain during method development

- Good reproducibility
- COMPETITIVE ADVANTAGE Cleaner samples and protection of your HPLC and GC instruments
- High loadability and outstanding performance
- Ideal flow properties
- Consistent recoveries

No risk

• Test samples available on request.

Good to know



- Higher capacity of up to 30 wt % (silica gel 3-5 wt %)
- pH stability of 1–14 (silica gel ~ 2–8)

Modern polymeric CHROMABOND® SPE phases

HR-X

Selection guide

Adsorbent

The continuous strive to improve SPE methods led to the development of our portfolio of CHROMABOND® polymer phases.

Stationary phase selection Matrix Aqueous (water, serum, blood, plasma, aqueous extracts) SPE mode Reversed phase Ion exchange Analyte Non polar to Strong bases Acids Strong acids Bases pKa 2-8 characteristic moderate polar pKa >10 pKa < 1 pKa 2-10

HR-XCW

HR-XA

HR-XAW

HR-XC



CHROMABOND® HI B

Technical data

Hydrophilic-lipophilic balanced N-vinylpyrrolidone-divinylbenzene copolymer (NVP/DVB)

SPE mode: Reversed phase Interactions: Hydrophobic and polar

Particle shape: Spherical pH stability:

Particle size: 60 um and 30 um

Pore size: 65 Å Specific surface: 750 m²/g

Special characteristics

- Applicable for a wide range of analyte polarities
- High loadability and outstanding performance
- Water wettable even if bed runs dry, SPE can be continued

Recommended application

- Medium polar organic molecules from polar matrices
- Drugs and pharmaceuticals from urine, blood, serum and plasma
- Tetracyclines and alkaloids from serum
- Pesticides from water

Standard SPE procedure for CHROMABOND® HLB (subsequent HPLC analysis)

MN Appl. No. 306300



CHROMABOND® HLB/3 mL/200 mg, REF 730924

Sample pretreatment:

Individual sample preparation in reference to the compounds and matrix. (Adjust pH value if necessary)

5 mL methanol, then 5 mL dist. water Conditioning: Sample application: Slowly aspirate sample through column

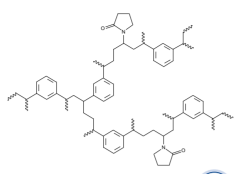
Washing: 5 mL dist. water

Drying: 10 min with applied vacuum

Elution: 8 mL methanol Evaporation: Under nitrogen

Reconstitution: In 1 mL dist. water + 0.1 % formic acid





Good to know



- Oasis® HLB
- Strata[™]-X
- SupelTM-Select HLB
- Supra-Poly® HLB
- Isolute[®] ENV+

Standard SPE procedure for CHROMABOND® HLB (subsequent GC analysis)

MN Appl. No. 306310

Column type:

CHROMABOND® HLB/3 mL/200 mg, REF 730924

Sample pretreatment:

Individual sample preparation in reference to the compounds and matrix. (Adjust pH value if necessary)

5 mL solvent (e.g., ethyl acetate), Conditioning:

5 mL methanol, 5 mL dist. water

Sample application: Slowly aspirate sample through column

Washing: 5 mL dist. water

Drying: 10 min with applied vacuum

Elution: Solvent¹⁾ (typical solvents: ethyl acetate, MTBE,

methylene chloride)

Under nitrogen, dry with sodium sulfate2), adjust Evaporation:

to final volume

1) usually nonpolar, therefore often 10 % methanol are added

2) e.g., with CHROMAFIX® Dry

Modern polymeric CHROMABOND® SPE phases

Applications

Tetracyclines and alkaloids from serum at pH 5

MN Appl. No. 306380

Chromatographic conditions

CHROMABOND® HLB/1 mL/30 mg

Oasis® HLB/1 mL/30 mg

MN REF:

Conditioning: 1 mL methanol, then 1 mL dist. water

Application: 1 mL serum pH 5, adjusted with formic acid

(spiked with 20 µg/mL of each analyte)

Washing: 1 mL dist. water

Drvina: 10 min with applied vacuum

Elution: 2 mL methanol Evaporation: Under nitrogen, 40 °C

Reconstitution: In 1 mL dist. water + 0.1 % formic acid

Recovery rates \pm RSD [%], n = 4

Compound	CHROMABOND® HLB	Oasis® HLB
Berberine	85.4 ± 0.3	82.5 ± 0.6
Chlortetracycline	72.1 ± 1.4	66.3 ± 2.8
Hydrastine	88.9 ± 2.6	99.3 ± 5.7
Oxytetracycline	82.3 ± 1.4	78.7 ± 1.4
Tetracycline	78.1 ± 1.4	70.7 ± 2.6

Further analysis: HPLC, according to MN Appl. No. 128180

EC 50/2 NUCLEOSHELL® RP 18plus, 2.7 µm Column:

MN REF: 763232.20

Eluent: A: dist. water + 0.1 % formic acid

B: acetonitrile + 0.1 % formic acid

Gradient: 2-60 % B in 4 min, 60 % B for 1 min, 60-2 % B in 0.5 min, 2 % B for 3 min

0.75 mL/min Flowrate:

Temperature: 22 °C

UV, 330 nm Detection:

Injection: 5 uL

Mycotoxins in wheat flour MN Appl. No. 306740

Chromatographic conditions CHROMABOND® HLB/60 µm/3 mL/200 mg Columns:

MN REF: 730924

Extraction:

• Weigh 4 g homogenized sample in an empty 50 mL centrifuge

 Add 8 μL mycotoxin standard mixture (β = 10 μg/mL each analyte in acetonitrile)

 Add 10 mL of water / acetonitrile mixture (20:80, v/v), shake vigorously and wait 10 min

 Add CHROMABOND® QuEChERS extraction Mix XII (REF 730648), shake vigorously for 1 min and cool the mixture down

Centrifuge at 4500 rpm for 20 min at 20 °C

Take organic phase for clean-up procedure

Conditioning: 6 mL acetonitrile

1 mL sample extract was aspirated with low vacuum into a vial Elution: 4 mL acetonitrile were aspirated with low vacuum into a vial Evaporation: Combine cleaned sample extract and acetonitrile eluate

and evaporate to dryness under nitrogen, 60 °C

Reconstitution: In 1 mL acetonitrile

Analyte	Recovery rate [%]	RSD [%], n = 5
Aflatoxin B1	88	2.6
Aflatoxin B2	91	5.0
Aflatoxin G1	85	2.6
Aflatoxin G2	88	4.5
HT-2 toxin	115	5.7
T-2 toxin	106	5.1
Zearalenone	49	3.4







Applications

Sulfa drugs from serum

MN Appl. No. 306340

Columns*: CHROMABOND® HLB/60 µm/1 mL/30 mg

Oasis® HLB/60 µm/1 mL/30 mg

MN REF:

Conditioning: 1 mL methanol, 1 mL dist. water

1 mL serum (spiked with 10 µg/mL of each Application:

analyte)

Washing: 1 mL dist. water

10 min with applied vacuum Drying:

Elution: 2 mL methanol

Evaporation: Under nitrogen, 40 °C

Reconstitution: In 1 mL dist. water + 0.1 % formic acid

Equivalence to Oasis® HLB

CHROMABOND® HLB shows equivalent recovery rates to

Oasis® HLB for the three tested sulfa drugs.

Further analysis: HPLC, according to MN Appl. No. 128130

EC 150/2 NUCLEODUR® C18 Pyramid, 3 µm Column:

MN REF: 760261.20

Dist. water + 0.1 % formic acid/methanol + 0.1 % Eluent:

formic acid (85:15, v/v), 5 min

Flow rate: 0.6 mL/min Temperature: 25 °C UV, 254 nm Detection 5 μL Injection:

Recovery rates ± RSD [%], n = 5

Compound	CHROMABOND® HLB	Oasis® HLB
Sulfadiazine	97.3 ± 2.9	92.0 ± 3.8
Sulfamerazine	94.4 ± 1.8	92.8 ± 1.6
Sulfathiazole	90.3 ± 2.9	89.6 ± 1.5

Chloramphenicol from honey

MN Appl. No. 306350

CHROMABOND® HLB/60 µm/3 mL, 200 mg Columns*:

Oasis® HLB, 3 mL, 200 mg

MN REF: 730924 Sample pretreatment:

Weigh out 5 g of honey. Add 4 mL water and shake rigorously for 30 sec. Spike with 1 mL standard solution (c = 5 ng/mL in methanol) and shake rigorously for 30 sec. Add 15 mL ethyl acetate and shake rigorously for 30 sec. Centrifuge at 3000 rpm for 10 min. Take 12 mL of supernantant for eluent exchange. Evaporate extracts to dryness at 40 °C under a stream of nitrogen.

Redissolve residue in 10 mL water.

Conditioning: 3 mL methanol (dispensing speed 1 mL/min), 5 mL

dist. water (disp. speed 1 mL/min)

9 mL water sample (disp. speed 3 mL/min over Application:

sample loop)

Washing: 10 mL dist. water (disp. speed 3 mL/min) 100 mL air (disp. speed 100 mL/min) Drying: 5 mL ethyl acetate /!methanol (80:20, v/v) Elution:

Drying: 100 mL air (disp. speed 100 mL/min)

under nitrogen, 40 °C Evaporation:

Reconstitution: in 1 mL dist. water/acetonitrile (95:5, v/v)

The SPE application was performed with a FREESTYLE® SPE automation system.

Further analysis: LC-MS/MS, according to MN Appl. No. 128140

EC 150/2 NUCLEODUR® π², 5 μm Column:

MN REF: 760624.20 Eluent: A: dist_water

5-95 % B in 7.5 min, 95 % B for 1 min, 95-5 % B

in 1 min. 5 % B for 5 min

Flow rate: Temperature: 35 °C

MS, Selected Reaction Monitoring (SRM) Detection:

Injection:

Recovery rates \pm RSD [%], n = 5

Compound	CHROMABOND® HLB	Oasis® HLB
Chloramphenicol-d5	90.9 ± 5.4	90.0 ± 9.3

Good to know

Antibiotics and pesticides contamination of agricultural products such as honey has been an issue in the recent years and resulted in stricter guidelines in food safety control.



^{*}Same conditions for all used columns. Due to a better comparability CHROMABOND® HLB and Oasis® HLB adsorbents (60 µm) were packed into equal column hardware. The shown chromatograms may not be representative of other applications

Modern polymeric CHROMABOND® SPE phases

Applications

Pesticides from tap water

MN Appl. No. 306360

Columns*:

CHROMABOND® HLB/60 µm/3 mL/200 mg

Oasis® HLB/60 µm/3 mL/200 mg

MN REF:

Conditioning: 5 mL methanol, 5 mL dist. water

1000 mL tap water (spiked with 50 ng of each Application:

analyte)

Washing: 10 mL dist. water

5 min with applied vacuum (-15 psi) Drying:

Flution: 6 mL acetonitrile Evaporation: Under nitrogen, 40 °C

Reconstitution: In 1 mL dist. water/acetonitrile (95:5, v/v)

Recovery rates \pm RSD [%], n = 5

Compound	CHROMABOND® HLB	Oasis® HLB
Acetamiprid	73.3 ± 5.0	112.1 ± 9.9
Atrazine	110.3 ± 17.8	114.0 ± 11.6
Azoxystrobin	74.7 ± 5.4	98.1 ± 10.8
Carbaryl	65.7 ± 5.4	69.1 ± 7.1
Chlorotoluron	82.7 ± 5.7	101.2 ± 3.8
Chlorpyrifos	50.3 ± 5.4	47.0 ± 3.7
Clofentezine	27.8 ± 2.7	21.4 ± 3.7
Clothianidin	69.4 ± 6.5	52.9 ± 2.9
Coumaphos	69.8 ± 4.8	82.3 ± 5.2
Cyanazine	99.8 ± 9.3	85.1 ± 7.2
Desethylatrazine	94.8 ± 15.1	87.4 ± 11.4
Desisopropylatrazine	92.5 ± 7.6	N/A
Diazinon	71.5 ± 7.9	73.3 ± 4.7
Difenoconazole	83.9 ± 6.5	28.8 ± 5.0
Diuron	70.0 ± 4.8	80.1 ± 8.4
Ethoprophos	72.4 ± 9.3	85.4 ± 7.2
Hexazinone	88.4 ± 7.7	104.3 ± 7.4
Imazalil	27.3 ± 15.7	N/A
Imidacloprid	93.4 ± 5.1	40.3 ± 5.2
Isoproturon	100.2 ± 4.2	102.8 ± 13.0
Linuron	84.5 ± 7.6	88.3 ± 9.5

Further analysis: LC-MS/MS, according to MN Appl. No. 128150

EC 50/2 NUCLEOSHELL® PFP, 2.7 µm Column:

763532.20 MN REF:

Eluent: A: dist. water + 0.1 % formic acid

B: acetonitrile + 0.1 % formic acid

5–95 % B in 15 min. 95 % B for 5 min. 95–5 % B in

1 min, 5 % B for 9 min

0.3 mL/min Flow rate:

Temperature: 40 °C

Detection: MS, Selected Reaction Monitoring (SRM)

Injection:

Compound	CHROMABOND® HLB	Oasis® HLB
Methabenzthiazuron	72.5 ± 5.3	48.0 ± 3.7
Methomyl	78.8 ± 5.4	83.6 ± 5.6
Metobromuron	73.8 ± 5.6	85.6 ± 9.3
Metolachlor	79.0 ± 5.2	89.2 ± 5.0
Monolinuron	75.4 ± 6.2	97.9 ± 7.2
Myclobutanil	101.8 ± 11.4	88.7 ± 14.5
Phosalone	63.8 ± 7.7	74.0 ± 4.0
Piperonylbutoxide	101.4 ± 8.6	99.7 ± 7.9
Propazine	102.1 ± 13.6	90.9 ± 9.4
Propyzamide	84.8 ± 7.1	86.4 ± 10.6
Terbuthylazine	107.9 ± 13.3	100.0 ± 13.6
Thiacloprid	74.1 ± 6.3	86.5 ± 10.8





CHROMABOND® HR-X

Technical data

Hydrophobic polystyrene-divinylbenzene copolymer (PS/DVB)

SPE mode: Reversed phase Interactions: Hydrophobic and π - π

Particle shape: Spherical pH stability:

Particle size: 85 um and 45 um Pore size: 55-60 Å Specific surface: 1000 m²/g

RP capacity: 390 mg/g (caffeine in water)

Recommended application

- Pharmaceuticals / active ingredients from tablets, creams and water
- Drugs and pharmaceuticals from urine, blood, serum and plasma
- Trace analysis of pesticides, herbicides, phenols, PAH and PCBs from water

Standard protocol for CHROMABOND® HR-X MN Appl. No. 304310

Column type:

CHROMABOND® HR-X/3 mL/200 mg, REF 730931

Sample pretreatment:

Individual sample preparation in reference to the compounds and

matrix (adjust pH value if necessary). Conditioning: 5 mL methanol, then 5 mL water

(do not let run the column dry!)

Sample aspiration: The prepared sample is passed through the column by vacuum or pressure (max. 1000 mL sample volume)

5 mL water/methanol (95:5, v/v)

Washing:

With nitrogen or air Drying: Elution: 3 x 2 mL methanol

Further analysis:

Evaporation and reconstitution (if necessary); HPLC or GC

These conditions are a starting point for SPE method development. Further optimization may be required to improve results.

Good to know



- Nexus

- StrataTM-X

A possible replacement for: ■ ENVI-Chrom P ■ Bakerbond H₂O-phobic DVB

Modern polymeric CHROMABOND® SPE phases

Applications

Determination of pyrrolizidine alkaloids

MN Appl. No. 306620

Chromatographic conditions

Columns: CHROMABOND® HR-X/85 µm/3 mL/200 mg

MN REF: 730921

Pretreatment: The following analysis were performed with standard

Conditioning: 5 mL methanol, 5 mL water

Application: 10 mL neutralized standard solution with a flow rate

of 3 mL/min

2 x 5 mL of water with a flow rate of 3 mL/min Washing:

5-10 min with vacuum Drying: 5 mL methanol Elution:

Eluent exchange: Add 1.0 mL water as keeper. Evaporate eluate to a volume of 0.5 mL at 40 °C under a stream of nitrogen and fill up

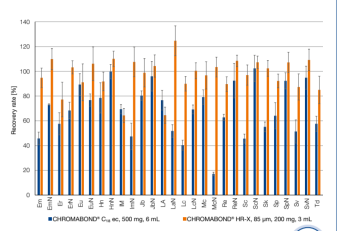
to 1.0 mL with water/methanol (95:5, v/v).

Further analysis:

HPLC determination of recovery rates with EC 150/2

NUCLEOSHELL® RP 18plus, 2.7 µm (REF 763236.20) in reference

to MN Appl. No. 127480



Superior to silica based RP phase

CHROMABOND® HR-X shows higher recovery rates for most tested pyrrolizidine alkaloids than CHROMABOND® C18 ec under the given conditions.

Enrichment of opiates

MN Appl. No. 306710

Chromatographic conditions

CHROMABOND® HR-X/45 µm/3 mL/60 mg Columns:

730936P45

Pretreatment: 400 µL methanolic standard solution were diluted

with 50 mmol/L phosphate buffer pH 7.0 to 20 mL 2.5 mL of this solution are equal to 5 ng of each

Conditioning: 3x1 mL methanol, 3x1 mL water, then 3x1 mL

50 mmol/L phosphate buffer pH 7.0

Aspiration: 2.5 mL of pretreated sample solution is passed

through the column at a flow of 1-2 mL/min

Washing: 3 x 1 mL 50 mmol/L phosphate buffer pH 7.0,

3 x 1 mL water

Drying: 5 mL air by pushing with a syringe Elution: 3 x 1 mL 0.1 % formic acid in methanol

Solvent change: Eluate is evaporated to dryness at 30 °C under a stream of nitrogen and then redissolved in organic solvent suited

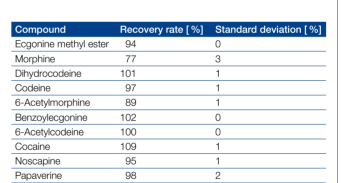
for the subsequent analysis.

Further analysis:

HPLC determination of recovery rates with EC 100/2

NUCLEOSHELL® Biphenyl, 2.7 µm (REF 763634,20) in reference

to MN Appl. No. 128880





CHROMABOND® HR-XC

Technical data

Strong cation exchanger based on polystyrene-divinylbenzene copolymer (PS/DVB)

SPE mode: Ion exchange and reversed phase (mixed-mode)

Interactions: lonic, hydrophobic and π – π

Particle shape: Spherical pH stability: 1-14

Particle size: $85 \mu m$ and $45 \mu m$

Pore size: 65-75 Å Specific surface: 800 m²/q

RP capacity: 300 mg/g (caffeine in water) Exchange capacity: 1.0 meq/g, pKa < 1

Recommended application

- Basic active ingredients from heavily matrix-contaminated samples, e.g., urine, plasma, serum
- Fungicides from food
- Basic analytes, e.g., amines
- Bases with pKa 2–10

Standard protocol for CHROMABOND® HR-XC MN Appl. No. 304790

Column type:

CHROMABOND® HR-XC/3 mL/200 mg, REF 730952

Sample pretreatment:

Individual sample preparation in reference to the compounds and matrix (adjust pH value if necessary).

5 mL methanol, then 5 mL water Conditioning:

(do not let run the column dry!)

Sample aspiration: The prepared sample is passed through the

column by vacuum or pressure

Washing 1: 2 mL 0.1 M HCl in water

Washing 2: / Elution 1: 2 mL methanol

(elution of neutral and acidic compounds)

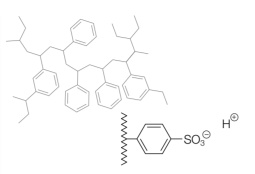
With nitrogen or air Drying: 5 mL methanol/5 % NH₃ Elution 2: (elution of basic compounds)

Further analysis:

Evaporation and reconstitution (if necessary); HPLC or GC

These conditions are a starting point for SPE method development.

Further optimization may be required to improve results.



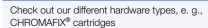
Good to know



A possible replacement for:

- Oasis® MCX
- Strata™-X-C
- StyreScreen® DBX
- HvperSep™ Retain CX

SPE hardware formats





Modern polymeric CHROMABOND® SPE phases

Applications

Enrichment of benzodiazepines

MN Appl. No. 306720

Chromatographic conditions

CHROMABOND® HR-XC 45 µm/3 mL/60 mg Columns:

MN REF: 730956P45

Pretreatment: 400 µL methanolic standard solution were diluted

with phosphate buffer pH 6.0 to 20 mL 2.5 mL of this solution are equal to 5 ng of each

analyte

Conditioning: 2 mL methanol, 2 mL phosphate buffer pH 6.0

Aspiration: 2.5 mL of pretreated sample solution is passed

through the column at a flow of 1-2 mL/min.

2 mL phosphate buffer pH 6.0, 2 mL methanol/ Washing:

> water (30:70, v/v), 3 mL 0.1 mol/L hydrochloric acid, 2 mL methanol/water (30:70, v/v), 0.1 mL methanol followed by 1 min drying, 2 mL methanol/water

(30:70, v/v)

5 min with a slight nitrogen stream Drying:

Elution: 2 x 1.5 mL 25 % aqueous ammonia solution/

ethylacetate (2:100, v/v)

Solvent change: Eluate is evaporated to dryness at 30 °C under a stream of nitrogen and then redissolved in organic solvent suited for the subsequent analysis.

Further analysis:

HPLC determination of recovery rates with EC 150/2 NUCLEOSHELL® Bluebird RP 18, 2.7 µm (REF 763436.20) in reference to MN Appl. No. 128890

Compound	Recovery rate [%]
Nortetrazepam	85
Tetrazepam	85
α-Hydroxytriazolam	87
Zaleplon	84
Nitrazepam	92
Oxazepam	104
Nordiazepam	83
N-Desmethylflunitrazepam	90
Lorazepam	89
Clonazepam	88
Desalkylflurazepam	102
Temazepam	103
Flunitrazepam	89
Lormetazepam	109
Clobazam	90
Diazepam	98





CHROMABOND® HR-XA

Technical data

Strong anion exchanger based on polystyrene-divinylbenzene copolymer (PS/DVB)

SPE mode: Ion exchange and reversed phase (mixed-mode)

Interactions: lonic, hydrophobic and π - π

Particle shape: Spherical pH stability: 1-14

Particle size: $85 \mu m$ and $45 \mu m$ 55-65 Å Pore size:

RP capacity: 350 mg/g (caffeine in water) Exchange capacity: 0.25 meg/g, pKa ~ 18

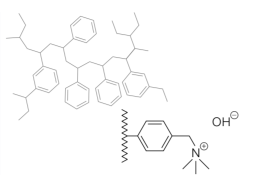
850 m²/q

Recommended application

- Acidic active ingredients from heavily matrix-contaminated samples, e.g., urine, plasma, serum
- Phenolic acids
- Acidic herbicides

Specific surface:

• Weak/medium-strength acids with pKa 2-8



Good to know



A possible replacement for:

- Oasis® MAX
- Strata™-X-A
- HyperSep™ Retain AX
- StyreScreen® QAX

Standard protocol for CHROMABOND® HR-XA MN Appl. No. 304970

Column type:

CHROMABOND® HR-XA/3 mL/200 mg/REF 730951

Sample pretreatment:

Individual sample preparation in reference to the compounds and

matrix (adjust a basic pH value).

Conditioning: 5 mL methanol, then 5 mL water

(do not let run the column dry!)

Sample aspiration: The basic sample is passed through the column

by vacuum or pressure (max. 1000 mL sample

2 mL 0.1 M NaOH in water Washing 1:

Washing 2: / Elution 1: 2 mL methanol

(elution of neutral and basic compounds)

Drying: With nitrogen or air

Elution 2: 5 mL methanol / 1-10 % formic acid

Further analysis:

Evaporation and reconstitution (if necessary); HPLC or GC

These conditions are a starting point for SPE method development.

(elution of acidic compounds)

Further optimization may be required to improve results.

Successful filtration

We recommend to use CHROMAFIL® Xtra syringe filters in combination with our SPE columns. For further information, please visit www.mn-net.com/chromafil.

Modern polymeric CHROMABOND® SPE phases

Applications

Fractions of acidic and basic analytes from serum

MN Appl. No. 305020

Chromatographic conditions

Column: CHROMABOND® HR-XA/85 µm/3 mL/200 mg

MN REF: 730951

Pretreatment: 1 µg/mL analytes in serum, adjusted on basic pH

with 1 N NaOH

Conditioning: 5 mL methanol, then 5 mL water (Do not let run the

column dry!)

The prepared sample is passed through the column Aspiration:

Washing: With 2.5 mL water impurities are removed

With nitrogen or air Drying:

Elution: Fraction A (basic analytes) is eluted with 5.0 mL

methanol

Fraction B (acidic analytes) with 5.0 mL methanol/

Evaporation and reconstitution with 1 mL of mobile phase from

subsequent HPLC.

Washing: 1.6 mL acetonitrile, 20 µL/s

Subsequent analysis:

Fraction A: HPLC determination on EC 125/4 NUCLEODUR® C8 Gravity,

5 μm (REF 760751.40) in reference to MN Appl. No. 118520

Fraction B: HPLC determination on EC 125/4 NUCLEODUR® C18 Gravity. 5 μm (REF 760100.40) in reference to MN Appl. No. 122230

Recovery rates:

Fraction A	Recovery [%]	Fraction B	Recovery [%]
Protriptyline	75	Suprofen	96
Nortriptyline	69	Naproxen	86
Doxepine	72	Tolmetin	85
Imipramine	80		
Amitriptyline	78		
Trimipramine	73		

Acidic pharmaceuticals from serum

MN Appl. No. 305000

Chromatographic conditions

CHROMABOND® HR-XA/85 µm/3 mL/200 mg Column:

MN REF: 730951

Pretreatment: 1 µg/mL pharmaceuticals in serum, adjusted on

basic pH with 1 N NaOH

Conditioning: 5 mL methanol, then 5 mL water (Do not let run the

Aspiration: The prepared sample is passed through the column

by vacuum

With the following washing mixtures impurities are Washing:

removed: a) 2.5 mL water · b) 2.5 mL 0.1 N NaOH ·

c) 5.0 mL methanol

Drying: With nitrogen or air

Elution: Analytes are eluted with 5 mL methanol/1% formic acid

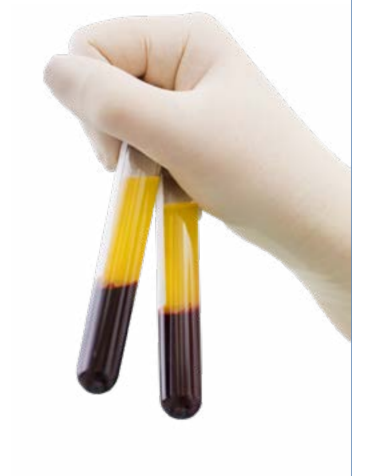
Evaporation to dryness and reconstitution with 1 mL of mobile

phase from subsequent HPLC.

HPLC determination of recovery rates with EC 125/4 NUCLEODUR® C18 Gravity, 5 µm (REF 760100.40) in reference to MN Appl. No. 122840

Recovery rates:

Compound	HR-XA [%]	Oasis® MAX [%]
Ketoprofen	90	85
Fenoprop	104	123
Fenoprofen	98	69
Flurbiprofen	106	98
Ibuprofen	88	58
Carprofen	69	89
Diclofenac	95	94
Meclofenamic acid	92	93



CHROMABOND® HR-XCW

Technical data

Weak cation exchanger based on polystyrene-divinylbenzene copolymer (PS/DVB)

SPE mode: Ion exchange and reversed phase (mixed-mode)

Interactions: lonic, hydrophobic and π - π

Particle shape: Spherical pH stability:

Particle size: 85 μm and 45 μm Pore size: 50-60 Å

Specific surface: 850 m²/q

RP capacity: 350 mg/g (caffeine in water) Exchange capacity: > 0.7 meg/g, pKa ~ 5

Recommended application

- Basic compounds like quaternary amines
- Active ingredients from heavily matrix-contaminated samples, e.g., urine, plasma, serum
- Strong bases with pKa > 10

Standard protocol for CHROMABOND® HR-XCW MN Appl. No. 305300

Column type:

CHROMABOND® HR-XCW/3 mL/200 mg, REF 730739

Sample pretreatment:

Individual sample preparation in reference to the compounds and matrix.

Conditioning: 5 mL methanol, then 5 mL water

(do not let run the column dry!)

Sample aspiration: The sample is passed through the column by

vacuum or pressure (max. 1000 mL sample

volume)

2 mL 5 % ag. NH₄OH solution Washing 1:

Washing 2: / Elution 1: 2 mL methanol

(elution of neutral and acidic compounds)

Drying: With nitrogen or air

Elution 2: 2 x 2 mL 1-5 % formic acid in methanol

(elution of strongly basic compounds)

Basic methanol (NH₃) can be used alternatively for elution 2 (e.g., for primary to tertiary amines). Here an interruption of the interactions with the cation exchanger results by a deprotonation of the analyte.

Further analysis:

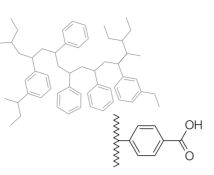
Evaporation and reconstitution (if necessary); HPLC or GC

These conditions are a starting point for SPE method development.

Further optimisation may be required to improve results.

HPLC columns

Are you looking for HPLC columns for subsequent analysis? Find an overview of our HPLC columns under the following link www.mn-net.com/hplc.



Good to know

A possible replacement for:

- Oasis® WCX
- StrataTM-X-CW

Modern polymeric CHROMABOND® SPE phases

Applications

Tricyclic Antidepressants

MN Appl. No. 305340

Column type:

CHROMABOND® HR-XCW/85 µm/3 mL/60 mg

MN REF: 730735

Pretreatment: 250 µL spiked serum, diluted with 1 mL 10 % formic

acid in water

Conditioning: 3 mL MeOH Equilibration: 3 mL water

Application: Slowly aspirate sample through the column 1 mL 5 % formic acid in water, then 1 mL MeOH After drying by vaccum (15 min) 3 mL 5 % formic

Further analysis:

Evaporate and redissolve in a suitable solvent for HPLC on NUCLEODUR® C8 Gravity, see MN Appl. No. 118520

Recovery rates:

Compound	HR-XCW	HR-XC*	PCA**	Oasis® WCX
Doxepine	79	5	11	41
Imipramine	79	9	20	67
Amitriptyline	91	9	14	46
Trimipramine	98	7	14	27

- * HR-XC: Basic analytes can not be eluted with slightly acidic organic conditions from the strong cation exchanger CHROMABOND® HR-XC, because the eluting power is not sufficient to dissociate the interaction with the ion exchanger. However, with the usage of basic methanol a complete elution can be achieved (please see also MN Appl. No. 304780).
- ** PCA: Due to the missing RP interactions of silica based weak cation exchanger, CHROMABOND® PCA gives only a small enrichment elution of the analytes







CHROMABOND® HR-XAW

Technical data

Particle size:

Weak anion exchanger based on polystyrene-divinylbenzene copolymer (PS/DVB)

SPE mode: Ion exchange and reversed phase (mixed-mode)

Interactions: lonic, hydrophobic and π – π

85 µm and 45 µm

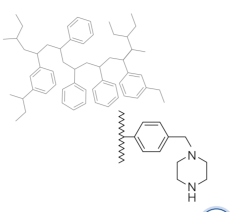
Particle shape: Spherical pH stability: 1-14

55-65 Å Pore size: Specific surface: 850 m²/g

RP capacity: 350 mg/g (caffeine in water) Exchange capacity: > 0.5 meg/g, pKa ~ 6

Recommended application

- Perfluorinated surfactants
- Acidic compounds like sulfonates
- Active ingredients from heavily matrix-contaminated samples, e.g., urine, plasma,
- Strong acids with pKa < 1



Good to know

A possible replacement for:

- Oasis® WAX
- StrataTM-X-AW

Standard protocol for CHROMABOND® HR-XAW MN Appl. No. 305200

Column type:

CHROMABOND® HR-XAW/3 mL/200 mg, REF 730748

Sample pretreatment:

Individual sample preparation in reference to the compounds and matrix.

5 mL methanol, then 5 mL water Conditioning:

(do not let the column run dry!)

The sample is passed through the column by Sample aspiration:

vacuum or pressure (max. 1000 mL sample

25 mM ammonium acetate in water Washing 1:

Washing 2: / Elution 1: 2 mL methanol

(elution of neutral and basic compounds)

Drying: With nitrogen or air

Elution 2: 2 x 2 mL 1-5 % ammonia in methanol

(elution of strongly acidic compounds)

Acidic methanol (formic acid) can be used alternatively for elution 2. Here an interruption of the interactions with the anion exchanger results

by a protonation of the analyte.

Further analysis:

Evaporation and reconstitution (if necessary); HPLC or GC

These conditions are a starting point for SPE method development.

Further optimisation may be required to improve results.

GC columns

For more information on our high performance GC capillary columns, please visit www.mn-net.com/optima

Modern polymeric CHROMABOND® SPE phases

Applications

Polyfluorinated compounds (PFCs) from fresh and sea water

Chromatographic conditions

MN Appl. No. 306730

Columns: CHROMABOND® HR-XAW/85 µm/3 mL/60 mg

MN REF: 730747

Pretreatment: 50 mL water sample spiked with PFC standard

mixture ($\beta = 0.5$ ng for each analyt in 50 mL water),

adjusted to pH value 7-8

Conditioning: 2 mL 0.1 % ammonium hydroxide in methanol, 2 mL

methanol, 2 mL water

Aspiration: Pretreated sample solution is passed through the

column at a flow of 5-10 mL/min

2 mL water, 2 mL 1.0 % formic acid in acetone/ Washing:

acetonitrile (50:50, v/v), 2 mL methanol

Drying:

2.4 mL 0.1 % ammonium hydroxide in methanol Elution:

Solvent change: Evaporate eluate to dryness at 40 °C under a stream of nitrogen and reconstitute in 0.5 mL water/methanol (40:60, v/v)

Did you know?

Properties of PFCs:

- Persistent in the environment
- Water-, dirt- and fat-repellent; resistant against aggressive chemicals
- Often toxic; many PFCs are bioaccumulative
- Thermally and chemically stable

Daily use of PFCs:

- Fire-fighting foam
- Paper finishing
- Fibre coating
- Textile coating, e.g., seat covers, carpets, outdoor clothing
- Food packaging, e.g., pizza cartons, paper cups
- Building material, e.g., water resistant lacquer

Recovery rates:

Matrix	Water		Seawater	
Analyte	Recovery	RSD	Recovery	RSD
	[%]	[%, n=3]	[%]	[%, n=3]
PFPeA	98	2.9	84	1.6
PFHxA	96	1.7	91	1.3
PFHpA	106	2.9	82	2.4
PFOA	99	2.3	86	2.5
PFNA	114	2.7	93	2.0
PFDA	110	2.6	90	2.3
PFUdA	96	5.3	85	3.5
PFDoA	84	1.6	76	2.1
PFTrDA	75	2.9	70	2.6
PFTeDA	66	4.3	74	4.0
L-PFBS	96	1.6	91	0.7
PFHxS	100	1.6	84	0.8
L-PFHpS	104	1.8	90	3.2
PFOS	103	2.0	84	2.3
L-PFDS	72	4.8	75	3.4
FOSA*	0	_	0	_
N-MeFOSAA*	3	_	0	_
N-EtFOSAA*	2	_	0	_
4:2 FTS	96	1.3	46	2.0
6:2 FTS	108	2.4	53	0.8
8:2 FTS	105	5.2	63	4.5
PFBA**	356	3.6	65	1.8
M ₄ -PFBA**	139	4.0	64	1.4
M ₄ -PFOA	101	3.7	89	2.8
M ₂ -PFHxA	95	2.2	84	0.5
M ₄ -PFHxS	96	2.2	84	1.7
M ₅ -PFNA	107	3.5	90	1.8
M ₄ -PFOS	101	2.4	82	1.2
M ₂ -PFDA	103	3.6	87	3.3
M ₂ -PFDoA	79	3.3	75	2.1
M ₂ -PFUdA	90	3.3	82	2.3
* Due to the organic	washing steps, the	ese analytes were	eluted into waste.	

** In accordance to the properties of the analyte molecules, a not satisfying S/N ratio is received resulting in an improper integration for calculating the recovery rate

Note: An LC-MS/MS method for determination of polyfluorinated compounds is shown in MN Appl. No 128900









Ordering information

CHROMABOND® HLB

	Volume	Adsorbent weight							Pack of
		30 mg	60 mg	100 mg	150 mg	200 mg	500 mg	1 g	
	CHROMABOND® HLB polypropylene columns (60 μm)								
	1 mL	730921		730922					30
	3 mL		730923			730924	730925		30
	6 mL				730944	730926	730927		30
Ш	15 mL						730928	730929	20
	CHROMABOND® H	LB polypropylene col	umns (60 µm)	· BIGpacks					
	3 mL		730923.250			730924.250			250
	6 mL					730926.250	730927.250		250
	CHROMABOND® H	LB polypropylene col	umns (30 µm)						
	1 mL	730921P30		730922P30					30
	3 mL		730923P30			730924P30			30
	6 mL				730944P30				30
TT	CHROMABOND® L	V-HLB (30 μm)							
	15 mL	732140	732141						30

	Size Minimum adsorbent weight	S 50 mg	M 120 mg	L 350 mg	Pack of
7.	CHROMAFIX® HLB cartridges (60 µm)				
		731921	731922	731923	50
	Adsorbent weight	96 x 10 mg	96 x 30 mg	96 x 60 mg	
	CHROMABOND® MULTI 96 HLB (60 µm))			
				738920.060M	1
	CHROMAFIX® MULTI 96 HLB (30 µm)				
		738921.010M	738921.030M		1

CHROMABOND® HR-X

30 mg HR-X polypropylene co	60 mg		200 mg	500 mg	1 g	Pack of			
	olumns (85 um)	100 mg	200 mg	ooo mg	' 9				
730934	(00 p)	730935				30			
	730936		730931	730937		30			
			730938	730939		30			
				730940	730941	20			
CHROMABOND® HR-X polypropylene columns (85 μm) · BIGpacks									
			730931.250			250			
			730938.250	730939.250		250			
CHROMABOND® HR-X polypropylene columns (45 μm)									
730934P45		730935P45				30			
	730936P45		730931P45			30			
LV-HR-X (85 μm)									
			732132			30			
_	LV-HR-X (85 μm)	LV-HR-X (85 μm)	LV-HR-X (85 μm)	V 1 7	V I 7	V 1 7			

	Adsorbent weight	96 x 100 mg	
èn	CHROMABOND® MULTI 96 HR-X (85 µm)		
		738530.100M	1

Modern polymeric CHROMABOND® SPE phases

Ordering information (cont.)

CHROMABOND® HR-XC

	Volume	Adsorbent weight						Pack of
		30 mg	60 mg	100 mg	150 mg	200 mg	500 mg	
TT	CHROMABOND® HR-XC polyp	ropylene columns (85 μ	um)					
	1 mL	730969		730049				30
	3 mL		730956			730952	730953	30
	6 mL				730957		730955	30
ш	CHROMABOND® HR-XC polyp	ropylene columns (45 μ	um)					
	1 mL	730969P45		730049P45				30
	3 mL		730956P45			730952P45		30
	Size	S		М		L		Pack of
	Minimum adsorbent weight	50 mg		140 mg		400 mg		
44	CHROMAFIX® HR-XC cartridge	es (85 µm)						
		731755		731756		731757		50

CHROMABOND® HR-XA

	Volume A	dsorbent weight						Pack of		
	30) mg	60 mg	100 mg	150 mg	200 mg	500 mg			
T	CHROMABOND® HR-X polypropylene columns (85 μm)									
	1 mL 73	80968		730727				30		
	3 mL		730950			730951	730954	30		
₹ ·	6 mL				730958		730966	30		
П	CHROMABOND® HR-XA polypropylene columns (45 µm)									
	1 mL 73	0968P45		730727P45				30		
	3 mL		730950P45			730951P45	730954	30		
	Size	S		М		L		Pack of		
	Minimum adsorbent weight	70 mg		215 mg		510 mg				
I _p	CHROMAFIX® HR-XA cartridges (85 μm)									
		731768		731769		731770		50		

CHROMABOND® HR-XCW

	Volume	Adsorbent weight						Pack of
		30 mg	60 mg	100 mg	150 mg	200 mg	500 mg	
T	CHROMABOND® HR-XCW	polypropylene colu	mns (85 μm)					
	1 mL	730731		730733				30
	3 mL		730735			730739	730741	30
	6 mL				730737		730743	30
	CHROMABOND® HR-XCW	polypropylene colu	mns (45 μm)					
	1 mL	730731P45		730733P45				30
	3 mL		730735P45			730739P45		30
	Size	S		M		L		Pack of
	Minimum adsorbent weigl	nt 60 mg		160 mg		450 mg		
	CHROMAFIX® HR-XCW ca	rtridges (85 µm)						
		731774		731775		731776		50

Ordering information (cont.)

CHROMABOND® HR-XAW

	Volume A	dsorbent weight						Pack of	
	30) mg	60 mg	100 mg	150 mg	200 mg	500 mg		
	CHROMABOND® HR-XAW polypropylene columns (85 μm)								
	1 mL 73	30728		730729				30	
	3 mL		730747			730748	730744	30	
	6 mL				730749		730745	30	
	CHROMABOND® HR-XAW polypropylene columns (45 μm)								
	1 mL 73	30728P45		730729P45				30	
	3 mL		730747P45			730748P45		30	
	Size	S		M		L		Pack of	
	Minimum adsorbent weight	50 mg		120 mg		360 mg			
	CHROMAFIX® HR-XAW cart	ridges (85 µm)							
		731771		731772		731773		50	

Registered trademarks

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