



Phase overview for special separations



Overview

Separation / mechanism	Recommended column	Specification of the phase	Page
Environmental analysis			
Anion exchange chromatography of inorganic anions	NUCLEOGEL® Anion I NUCLEOSIL® Anion II	Strongly basic polymer-based anion exchanger Strongly basic silica-based anion exchanger	230
RP chromatography of PAHs	NUCLEODUR® C ₁₈ PAH NUCLEOSIL® 100-5 C ₁₈ PAH	NUCLEODUR® polymer-coated with C ₁₈ groups USP L1 NUCLEOSIL® 100 polymer-coated with C ₁₈ groups USP L1	227 229
Enantiomer separation			
Polar and π-π interactions	NUCLEOCEL DELTA	Silica-based modified cellulose phases USP L40	233
Formation of inclusion complexes	NUCLEODEX α-PM, β-PM, γ-PM and β-OH	Silica-based permethylated and underivatized cyclodextrin phases USP L45	231
Enantioselective binding to chiral protein surface structures	RESOLVOSIL BSA-7	Silica-based protein phase (BSA)	234
Ligand exchange	NUCLEOSIL® CHIRAL-1	Covalently bonded amino acid – Cu(II) complexes USP L32	235
Charge-transfer, dipole-dipole interactions and others	NUCLEOSIL® CHIRAL-2 NUCLEOSIL® CHIRAL-3	Silica-based brush type phases USP L36	236
Separation of biological macromolecules			
Anion exchange chromatography of oligonucleotides and nucleic acids	NUCLEOGEN® DEAE	Silica-based DEAE anion exchanger	237
Anion exchange chromatography of peptides, large proteins and oligonucleotides	NUCLEOGEL® SAX	Polymer-based strongly basic anion exchanger USP L23	240
Cation exchange chromatography of proteins, peptides and carbohydrates	NUCLEOGEL® SCX	Polymer-based strong cation exchanger USP L22	240
Reversed phase chromatography of proteins, peptides and oligonucleotides	NUCLEOSIL® MPN NUCLEOSIL® PPN NUCLEOGEL® RP 300	Monomerically bonded alkyl chains on silica USP L1 / USP L26 Polymerically bonded alkyl chains on silica USP L1 Polystyrene – divinylbenzene polymer USP L21	243 244 245
Reversed phase chromatography of small molecules	NUCLEOGEL® RP 100	Small pore macroporous PS-DVB polymer USP L21	245
Food analysis · sugars			
RP chromatography of mono- and oligosaccharides	NUCLEOSIL® Carbohydrate	Silica-based special amino phase USP L8	246
Separation of sugars, alcohols, org. acids based on ion exclusion, ion exchange, size exclusion, ligand exchange, NP and RP effects	NUCLEOGEL® SUGAR 810 H, Ca	Resins with sulfonic acid modification in different ionic forms H form USP L17 / Ca form L19 / Pb form L34 / Na form L58	247
Separation of sugars, alcohols, org. acids based on steric exclusion, ligand exchange and partition effects	NUCLEOGEL® SUGAR Ca, Na, Pb NUCLEOGEL® ION 300 OA		248
Gel permeation chromatography (GPC)			
Water-insoluble compounds	NUCLEOGEL® GPC	Polystyrene – divinylbenzene polymer	249



HPLC columns for environmental analyses



NUCLEODUR® C₁₈ PAH special octadecyl phase for PAH analysis · USP L1

Technical data

- Base material NUCLEODUR® silica, particle sizes 1.8 and 3 µm, pore size 110 Å; polymeric coating

Recommended application

- Allows efficient gradient separation of the 16 PAHs according to EPA

Analysis of 16 EPA PAHs with or without acetonitrile

MN Appl. Nos. 123820/123830

Separation with acetonitrile

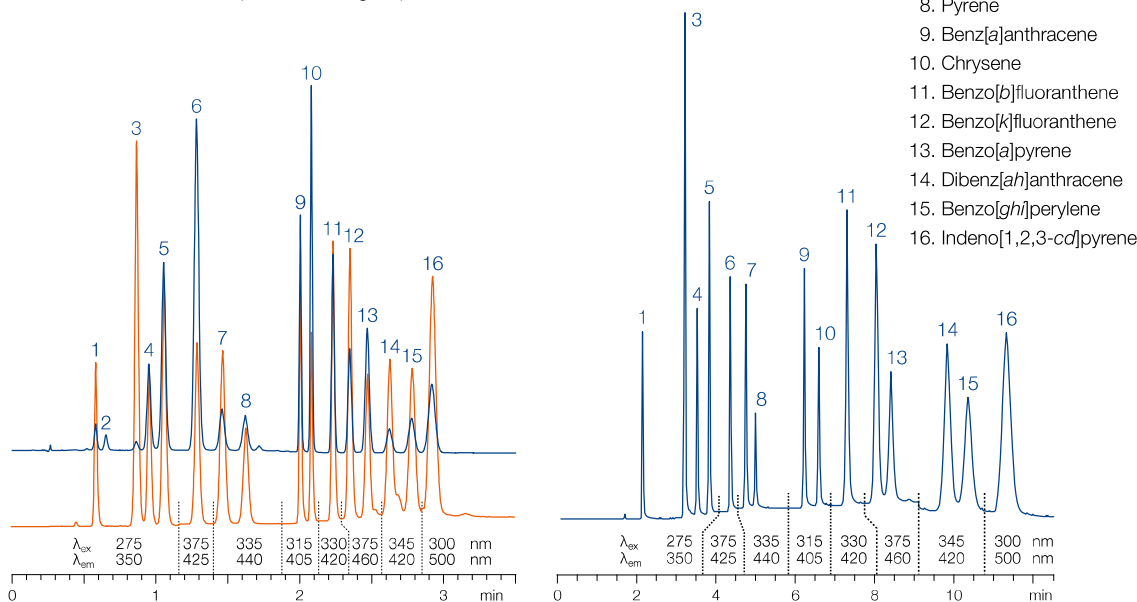
Column: 100 x 4 mm
NUCLEODUR® C18 PAH, 3 µm
Eluent: A) methanol – water (80:20, v/v)
B) acetonitrile 2–20 % B in 1.2 min,
20–100 % B in 0.5 min, 100 % B
for 2.5 min, 100–2 % B in 0.4 min
Flow rate: 2.5 mL/min, temperature 35 °C
Detection: UV, 254 nm
fluorescence (see chromatogram)

Separation without acetonitrile

Column: 125 x 4 mm
NUCLEODUR® C18 PAH, 3 µm
Eluent: A) water
B) methanol 65–97 % B in 6 min,
97 % B for 5 min, 97–65 % B in
0.5 min
Flow rate: 2 mL/min, temperature 35 °C
Detection: fluorescence (see chromatogram)

Peaks:

1. Naphthalene
2. Acenaphthylene (not detectable by fluorescence)
3. Acenaphthene
4. Fluorene
5. Phenanthrene
6. Anthracene
7. Fluoranthene
8. Pyrene
9. Benz[a]anthracene
10. Chrysene
11. Benzo[b]fluoranthene
12. Benzo[k]fluoranthene
13. Benzo[a]pyrene
14. Dibenz[ah]anthracene
15. Benzo[ghi]perylene
16. Indeno[1,2,3-cd]pyrene



Detection of separated PAHs with UV (250–280 nm), diode array or fluorescence detection at different wavelengths for excitation and emission (acenaphthylene cannot be analyzed with fluorescence detection).

Ordering information

Eluent in column acetonitrile – water (70:30, v/v)

ID	Length →	100 mm	125 mm	150 mm	250 mm	EC guard columns*
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NUCLEODUR® C₁₈ PAH, 1.8 µm particle size 1.8 µm · UHPLC

Analytical EC columns

	2 mm	760773.20				761970.20
	3 mm	760773.30				761970.30
	4 mm	760773.40				761970.30

NUCLEODUR® C₁₈ PAH, 3 µm particle size 3 µm

Analytical EC columns

	3 mm	760783.30	760784.30	760785.30	760786.30	761971.30
	4 mm	760783.40	760784.40	760785.40	760786.40	761971.30

Guard column system

Guard columns for EC columns with ID	2 mm	3 mm	4 mm	4.6 mm	Guard column holder
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* Column Protection System (pack of)

EC 4/2 (3)

4/3 (3)

4/3 (3)

4/3 (3)

718966



HPLC columns for environmental analyses



Separation of 18 PAHs on NUCLEODUR® C₁₈ PAH

MN Appl. No. 123840

Column: 125 x 4 mm
NUCLEODUR® C₁₈ PAH, 3 µm

Eluent: A) methanol – water (70:30, v/v); B) acetonitrile
0–20 % B in 1.5 min,
20–50 % B in 1.5 min,
50–100 % B in 1.0 min,
100 % B for 3 min,
100–0 % B in 0.5 min

Flow rate: 1.5 mL/min

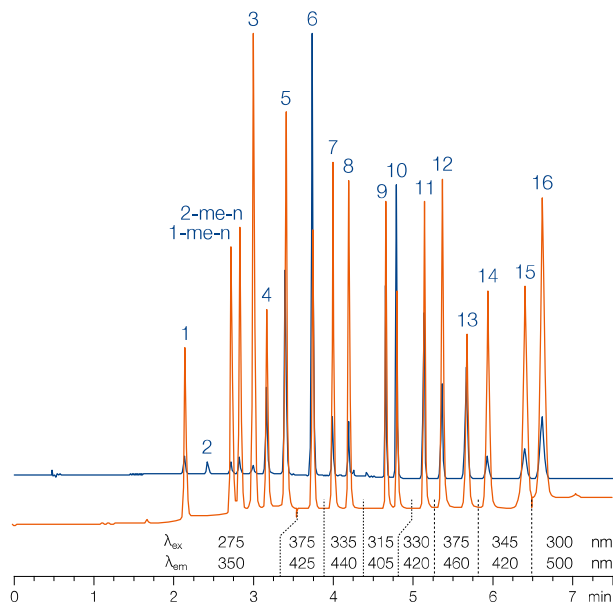
Temperature: 35 °C

Injection: UV: 1 µL,

Fluorescence: 0.5 µL

Detection: UV, 254 nm
fluorescence
(see chromatogram)

Peaks:
(concentrations 10 ng/µL per compound)
1.–16. see page 227
1-me-n: 1-methylnaphthalene
2-me-n: 2-methylnaphthalene

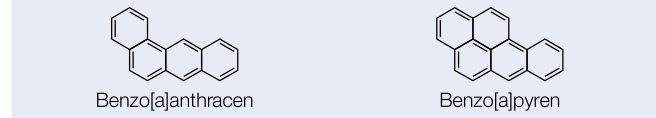


Analysis of polycyclic aromatic hydrocarbons (PAHs) by HPLC

Polycyclic aromatic hydrocarbons (PAHs) are chemical compounds that consist of fused aromatic rings and do not contain heteroatoms or carry substituents. As a pollutant, they are of concern because some compounds have been identified as carcinogenic, mutagenic, and teratogenic. PAHs are natural components of coal or gas. They are delivered to our environment by pyrolysis (incomplete burning) of organic materials like coal, oil, fuel, wood, tobacco, ... and hence can be found globally. Today most PAHs accrue from anthropogenic processes – but also natural origins (forest fire) are possible. Regarding to past pollutions an important impact had production of coke and gas from black coal. Waste products (e.g., tar) from coking or gas plants are often origin of serious ground water pollutions.

Since a number of PAHs (e.g., benzo[a]pyrene, 3-methylcholanthrene and benzanthracene) have been proven to be carcinogenic, control of the PAH content of food, water and soil is an important task for routine analysis. For choice and limiting values of the polycyclics we refer to the governmental regulations, which exist in many countries (e.g., EPA method 610 of the United States Environmental Protection Agency).

PAHs can be determined by different chromatographic techniques (TLC, GC, HPLC). Thus the 6 PAHs according to German drinking water specification (TVO) can, e.g., be analyzed by TLC (see German Standard DIN 38 409), while a much larger number of polycyclic aromatics can be determined by GC or HPLC.



HPLC columns for PAH analysis

For PAH analyses we have developed specially modified C₁₈ phases based on NUCLEODUR® and NUCLEOSIL® which allow efficient gradient separation of 16 PAHs according to EPA. Detection of the separated PAHs can be achieved by UV (250–280 nm), with diode array or with fluorescence detection at different wavelengths for excitation and emission. Acenaphthylene cannot be analyzed with fluorescence detection. For cost-effective routine PAH analysis we recommend applications using methanol instead of acetonitrile as eluent. For rapid analysis NUCLEODUR® C₁₈ PAH (3 µm) in short columns (100 mm) provides excellent results at high flow rates. Hereby separation of 16 PAHs according to EPA can be achieved in less than 3 min.

Tightened regulations require determination of 2 additional PAHs (1- and 2-methylnaphthalene) – so we developed highly efficient methods for 18 PAHs on the NUCLEODUR® C₁₈ PAH.



HPLC columns for environmental analyses



NUCLEOSIL® 100-5 C₁₈ PAH special octadecyl phase for PAH analysis · USP L1

Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å; polymeric coating
- Detection of the separated PAH with UV (250–280 nm), diode array or fluorescence detection at different wavelengths for excitation and emission (acenaphthylene cannot be analyzed with fluorescence detection)

Recommended application

- Efficient gradient separation of the 16 PAHs according to EPA

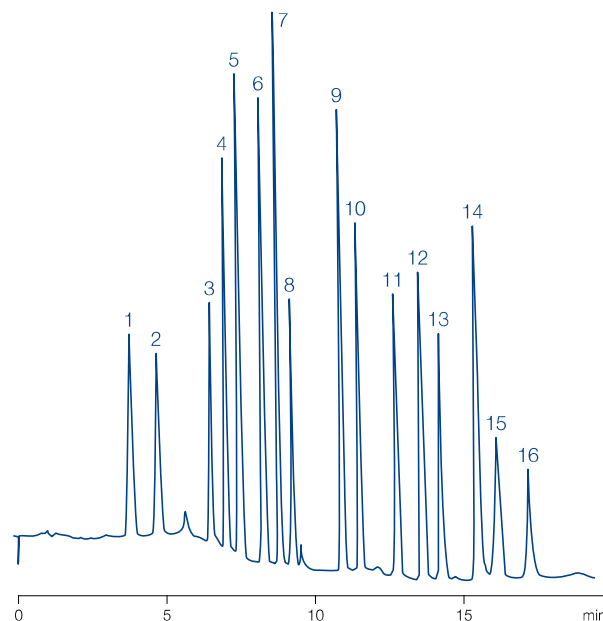
Separation of the PAH standard according to EPA (REF 722393)

MN Appl. No. 115040

Column: 150 x 4 mm NUCLEOSIL® 100-5 C₁₈ PAH
 Eluent: A) methanol – water (80:20)
 B) acetonitrile – tetrahydrofuran (93:7)
 0–100 % B in 10 min, 5 min 100 % B
 Flow rate: 1 mL/min
 Pressure: 140 bar
 Temperature: 20 °C
 Detection: UV, 260 nm

Peaks: (10 µg/mL each in acetonitrile)

1. Naphthalene	10. Chrysene
2. Acenaphthylene	11. Benzo[b]fluoranthene
3. Acenaphthene	12. Benzo[k]fluoranthene
4. Fluorene	13. Benzo[a]pyrene
5. Phenanthrene	14. Dibenz[ah]anthracene
6. Anthracene	15. Benzo[ghi]perylene
7. Fluoranthene	16. Indeno[1,2,3-cd]pyrene
8. Pyrene	
9. Benz[a]anthracene	



Ordering information

Eluent in column acetonitrile – water 70:30

ID	Length →	150 mm	250 mm	EC guard columns*
NUCLEOSIL® 100-5 C₁₈ PAH particle size 5 µm, pore size 100 Å				
Analytical EC columns				
	2 mm	720117.20	721168.20	
	3 mm	720923.30	720117.30	721168.30
	4 mm	720923.40	720117.40	721168.30
	4.6 mm	720117.46	721168.30	

PAH standard according to EPA for HPLC

Analytical EC columns

PAH standard for HPLC	16 PAH according to EPA method 610 in acetonitrile (1 mL) for composition see chromatogram above	722393
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Guard column system

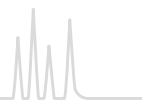
Guard columns for EC columns with ID	2 mm	3 mm	4 mm	4.6 mm	Guard column holder
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	718966

EC columns in packs of 1, guard columns in packs of 3. For details of our column systems see page 250.

This product contains harmful substances which must be specially labeled as hazardous. For detailed information please see SDS.



HPLC columns for environmental analyses



Anion columns for analysis of inorganic anions

NUCLEOGEL® Anion I

Technical data

- Strongly basic polymer-based anion exchanger, particle size 10 µm; pH stability 1–14
- Eluent in column 4 mmol/L salicylate buffer pH 7.8

- Contrary to the silica-based phase also suited for fluoride analysis

NUCLEOSIL® Anion II

Technical data

- Base material NUCLEOSIL® silica, particle size 10 µm, pore size 300 Å strongly basic anion exchanger, exchange capacity 50 µeq/g, pH stability 2–7.5
- Eluent in column 0.15 mol/L $(\text{NH}_4)_2\text{HPO}_4$ buffer pH 5.2 recommended buffer concentration for separation of inorganic anions: 2 mmol/L phthalate

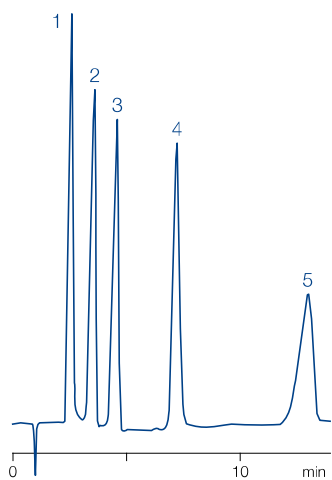
- Preferred method of detection: conductivity or negative UV detection

Separation of an anion standard

MN Appl. No. 106440

Column: 250 x 4 mm NUCLEOSIL® Anion II
 Eluent: 2 mmol/L potassium hydrogen phthalate, pH 5.7
 Flow rate: 2 mL/min
 Detection: UV, 280 nm

Peaks:
 1. H_2PO_4^-
 2. Cl^-
 3. NO_2^-
 4. NO_3^-
 5. SO_4^{2-}

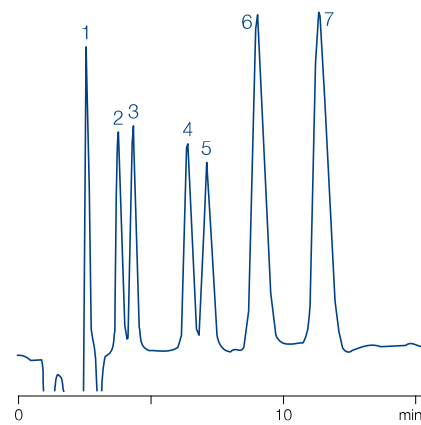


Separation of inorganic anions

MN Appl. No. 115050

Column: 120 x 4.6 mm NUCLEOGEL® Anion I
 Eluent: 4 mmol/L salicylic acid – Tris pH 7.8
 Flow rate: 1 mL/min
 Detection: UV, 254 nm

Peaks:
 1. F^-
 2. Cl^-
 3. NO_2^-
 4. Br^-
 5. NO_3^-
 6. PO_4^{3-}
 7. SO_4^{2-}



Ordering information

ID

Length →
120 mm

250 mm

Guard columns*

NUCLEOGEL® Anion I eluent 4 mmol/L salicylate buffer pH 7.8

Analytical Valco type columns



4.6 mm

719533

719543

NUCLEOSIL® Anion II eluent 0.15 mol/L $(\text{NH}_4)_2\text{HPO}_4$ buffer pH 5.2

Analytical EC columns



4 mm

720094.40

721169.30

* NUCLEOGEL® Anion I Valco type guard columns cartridges are 21 x 4 mm, require guard column holder C, REF 719538, see page 250 (columns in packs of 1, guard columns in packs of 2)

NUCLEOSIL® Anion II guard columns are used with the Column Protection System (REF 718966, see page 251).



HPLC columns for enantiomer separations



NUCLEODEX columns enantiomer separation based on cyclodextrins

NUCLEODEX β -OH β -cyclodextrin ($R = H; n = 2$) · USP L45

Technical data

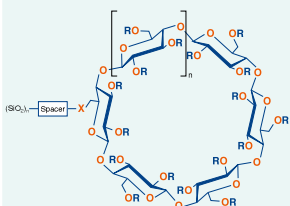
- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Separation based on hydrogen bonds and dipole interactions between functional groups of the analyte and hydroxyl groups of the cyclodextrin

- Examples for successful enantiomer separations: chlorthalidone and other compounds, which require free hydroxyl groups for enantioselective interactions
- Eluent in column CH₃OH – 0.1 % TEAA pH 4 (55:45)

NUCLEODEX α -PM permethylated α -cyclodextrin ($R = CH_3; n = 1$)

Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Examples for successful enantiomer separations: mecoprop and dichlorprop as free carboxylic acids, trans-stilbene oxide, styrene oxide



NUCLEODEX β -PM permethylated β -cyclodextrin ($R = CH_3; n = 2$) · USP L45

Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Examples for successful enantiomer separations: mephobarbital (prominal), pesticide derivatives mecoprop methyl and dichlorprop methyl

- Eluent in column CH₃OH – 0.1 % TEAA pH 4 (65:35)

NUCLEODEX γ -PM permethylated γ -cyclodextrin ($R = CH_3; n = 3$)

Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Examples for successful enantiomer separations: steroids or other larger molecules

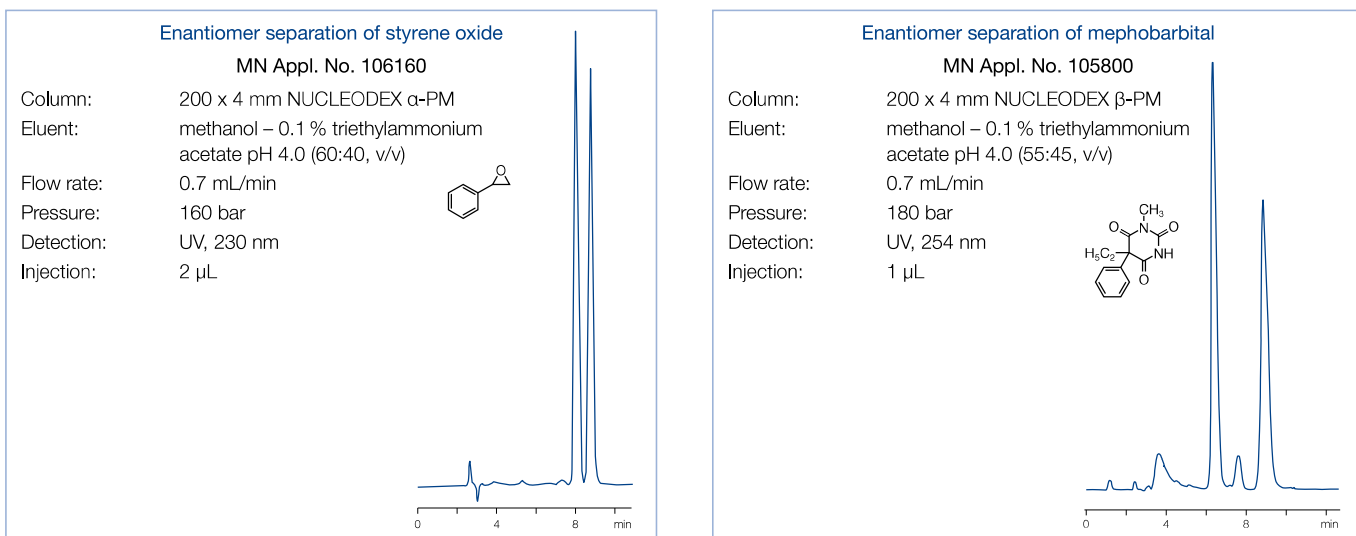
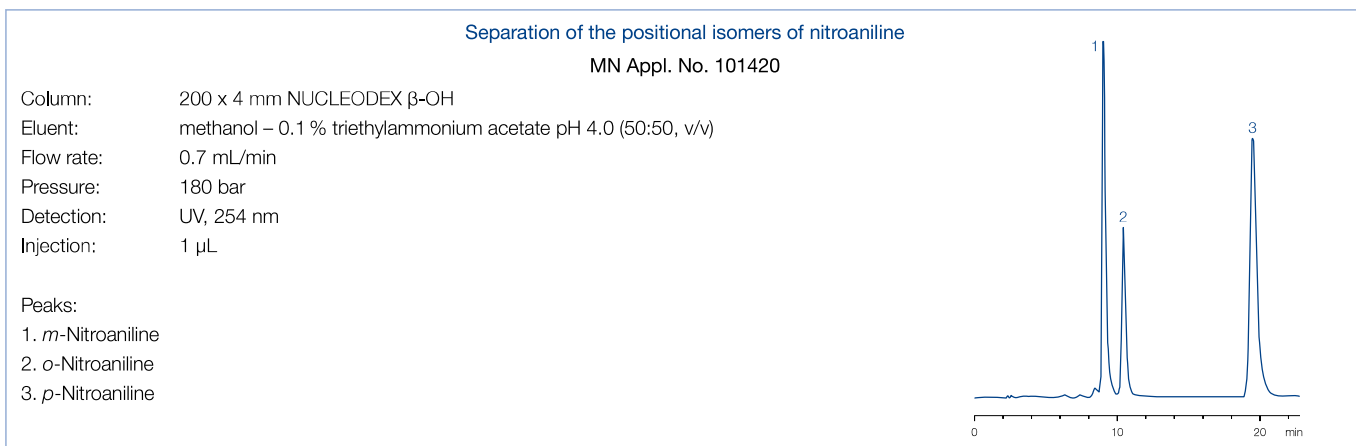
- Eluent in column CH₃OH – 0.1 % TEAA pH 4 (55:45)

✓ Recommended application

- NUCLEODEX phases are especially suited for the control of optical purity, but also for semipreparative separations and for the analysis of positional and cis-trans isomers.
- For numerous separations on NUCLEODEX phases please visit our website: www.mn-net.com/apps



HPLC columns for enantiomer separations



Ordering information			
ID	Length → 200 mm	EC guard columns*	
NUCLEODEX β-OH eluent methanol – 0.1 % TEAA pH 4 (55:45)			
Analytical EC columns			
4 mm	720124.40	721171.30	
NUCLEODEX α-PM eluent methanol – 50 mmol/L phosphate pH 3 (70:30)			
Analytical EC columns			
4 mm	720127.40	721469.30	
NUCLEODEX β-PM eluent methanol – 0.1 % TEAA pH 4 (65:35)			
Analytical EC columns			
4 mm	720125.40	721176.30	
NUCLEODEX γ-PM eluent methanol – 0.1 % TEAA pH 4 (55:45)			
Analytical EC columns			
4 mm	720752.40	721178.30	
NUCLEODEX CC screening kit			
contains one CC 30/4 each with NUCLEODEX β -OH, α -PM, β -PM and γ -PM as well as one CC column holder 30 mm		721920	

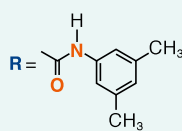
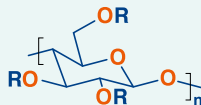
* EC 4/3 guard columns for EC columns with 4 mm ID require the Column Protection System guard column holder (REF 718966, see page 251). Columns and guard columns in packs of 1.



HPLC columns for enantiomer separations



NUCLEOCEL DELTA enantiomer separation based on a cellulose derivative · USP L40



Technical data

- Base material silica, chiral selector cellulose tris-(3,5-dimethylphenylcarbamate)
- High resolution type (S) with 5 µm particle size, allows use of shorter columns (150 mm) for faster separations, pressure stability up to ~150 bar (2000 psi), pH stability 1–9
- NUCLEOCEL DELTA for normal phase applications: eluent in column n-heptane – 2-propanol (90:10, v/v) typical eluents are heptane – propanol mixtures
- NUCLEOCEL DELTA-RP for reversed phase applications: eluent in column acetonitrile – water (40:60, v/v) designed for use either in polar organic mode or with eluents containing high concentrations of chaotropic salts such as perchlorate

Recommended application

- Pharmaceutically active compounds, chiral pollutants (e.g., herbicides, PCB), chiral compounds in food (dyes, preservatives), chiral catalysts and bioorganic compounds

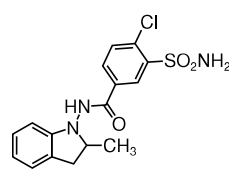
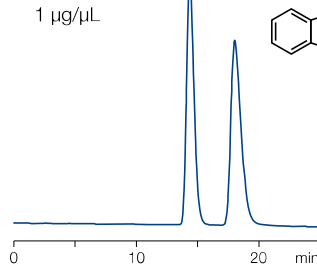
Similar phases: Chiralcel® OD, Kromasil® CelluCoat™, Eurocel® 01, Lux™ Cellulose-1

Enantiomer separation of indapamide

MN Appl. No. 121230

Column: 250 x 4,6 mm NUCLEOCEL DELTA-RP S
Eluent: acetonitrile – water (40:60, v/v)
Flow rate: 0.5 mL/min
Temperature: 40 °C
Detection: UV, 254 nm
Injection: 5 µL
Concentration: 1 µg/µL

$\alpha = 1.3$
 $R_s = 2.6$

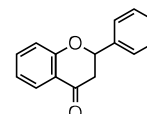
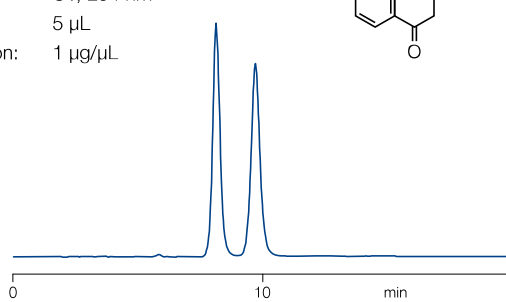


Enantiomer separation of flavanone

MN Appl. No. 121260

Column: 250 x 4,6 mm NUCLEOCEL DELTA S
Eluent: n-heptane – 2-propanol (90:10, v/v)
Flow rate: 1 mL/min
Temperature: 25 °C
Detection: UV, 254 nm
Injection: 5 µL
Concentration: 1 µg/µL

$\alpha = 1.29$
 $R_s = 2.6$



Ordering information

ID

Length →
150 mm

250 mm

EC guard columns*

NUCLEOCEL DELTA S, 5 µm eluent n-heptane – 2-propanol (90:10, v/v)

Analytical EC columns



4.6 mm

720445.46

721185.30

NUCLEOCEL DELTA-RP S, 5 µm eluent acetonitrile – water (40:60, v/v)

Analytical EC columns



4.6 mm

720451.46

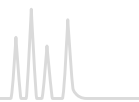
720450.46

721186.30

* EC 4/3 guard column cartridges are used for EC columns of 4.6 mm ID with the Column Protection System guard column holder (REF 718966, see page 251). Columns and guard columns in packs of 1.



HPLC columns for enantiomer separations



RESOLVOSIL BSA-7 protein phase for enantiomer separation · USP L75

Technical data

- Base material NUCLEOSIL® silica, particle size 7 µm, pore size 300 Å chiral selector bovine serum albumin (BSA)
- Separation based on selective interaction of proteins with low molecular compounds, i.e. principles of bioaffinity, including hydrophobic interactions (similar to a true reversed phase), interactions of polar groups and steric effects

Recommended application

- Amino acid derivatives, aromatic amino acids, aromatic sulfoxides, barbiturates, benzodiazepinones, benzoin and benzoin derivatives, β-blockers, coumarin derivatives, and for monitoring stereoselective microbial and enzymatic conversions

Enantiomer separation of *N*-benzoyl-D,L-amino acids

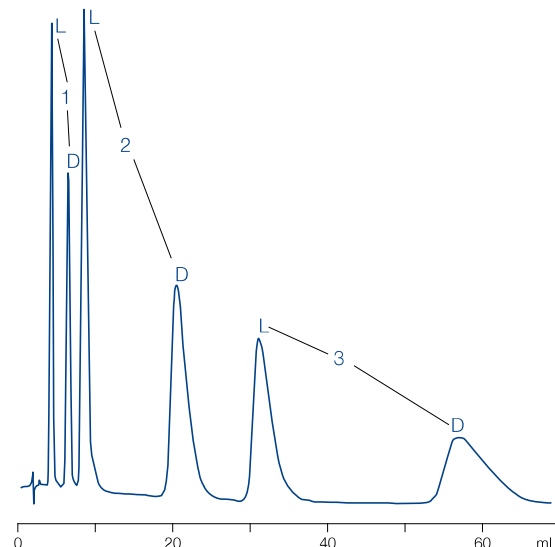
MN Appl. No. 105450

S. Allenmark et al. in "Affinity chromatography and biological recognition" (I. Chaiken, M. Wilchek, and I. Parikh, Eds.), Academic Press, New York, 1983, 259–260

Column: 150 x 4 mm RESOLVOSIL BSA-7
Eluent: 50 mmol/L phosphate buffer pH 6.5
+ 1 % 1-propanol
Flow rate: 0.70 mL/min
Detection: UV, 225 nm

Peaks:

1. Serine
2. Alanine
3. Phenylalanine



Ordering information

Eluent in column 0.1 mol/L phosphate buffer pH 7.5, 2 % 1-propanol

ID

Length →
150 mm

EC guard columns*

RESOLVOSIL BSA-7

Analytical EC columns



4 mm

720046.40

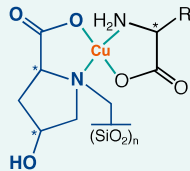
721402.30

* EC 4/3 guard columns for EC columns with 4 mm ID require the Column Protection System guard column holder (REF 718966, see page 251). Columns and guard columns in packs of 1.



HPLC columns for enantiomer separations

NUCLEOSIL® CHIRAL-1 enantiomer separation based on ligand exchange · USP L32



Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 120 Å chiral selector L-hydroxyproline – Cu²⁺ complexes
- Principal interaction mode:
- formation of ternary mixed-ligand complexes with Cu(II) ions; differences in the stability of the diastereomeric complexes cause chromatographic separation

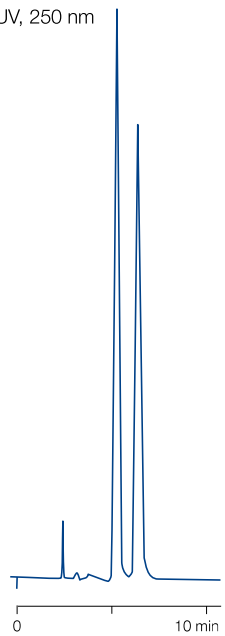
Recommended application

- Enantiomers with two polar functional groups with the correct spacing such as α-amino acids, α-hydroxycarboxylic acids (e.g., lactic acid), N-alkyl-α-amino acids etc.

D,L-alanine enantiomers

MN Appl. No. 105410

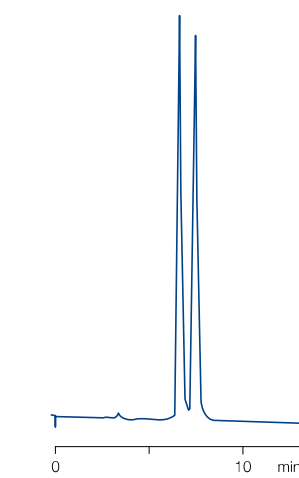
Column: 250 x 4 mm NUCLEOSIL® CHIRAL-1
Eluent: 0.5 mmol/L CuSO₄
Flow rate: 1 mL/min
Pressure: 60 bar
Temperature: 60 °C
Detection: UV, 250 nm



D,L-threonine enantiomers

MN Appl. No. 105410

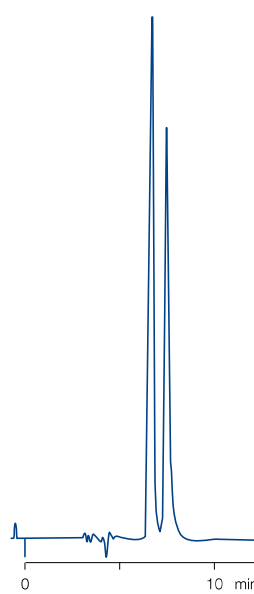
Column: 250 x 4 mm NUCLEOSIL® CHIRAL-1
Eluent: 0.25 mmol/L CuSO₄
Flow rate: 0.8 mL/min
Pressure: 65 bar
Temperature: 60 °C
Detection: UV, 240 nm



Lactic acid enantiomers

MN Appl. No. 105560

Column: 250 x 4 mm NUCLEOSIL® CHIRAL-1
Eluent: 0.5 mmol/L CuSO₄
Flow rate: 0.8 mL/min
Temperature: 60 °C
Detection: UV, 240 nm
Injection: 1 µL



Ordering information

Eluent in column 0.5 mmol/L copper sulfate solution

ID

Length →
250 mm

EC guard columns*

NUCLEOSIL® CHIRAL-1

Analytical EC columns



4 mm

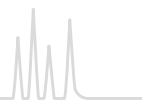
720081.40

721188.30

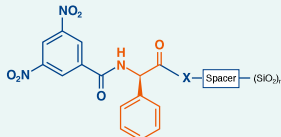
* EC 4/3 guard columns for EC columns with 4 mm ID require the Column Protection System guard column holder (REF 718966, see page 251). Columns and guard columns in packs of 1.



HPLC columns for enantiomer separations



NUCLEOSIL® CHIRAL-2 · CHIRAL-3 enantiomer separation in organic eluent systems · USP L36



Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å chiral selector for NUCLEOSIL® CHIRAL-2 is *N*-(3,5-dinitrobenzoyl)-D-phenylglycine, for CHIRAL-3 the optical antipode is used, "brush type" phases
- Principle interaction modes: charge-transfer interactions, hydrogen bonds, dipole-dipole interactions and steric effects

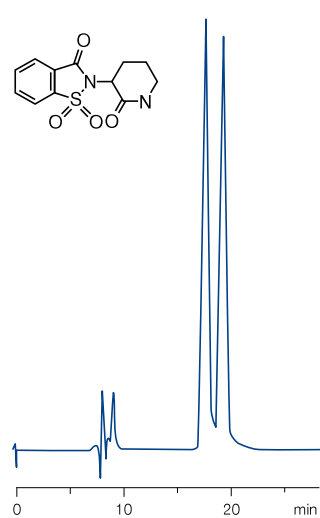
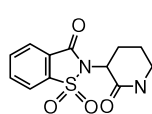
Recommended application

- analysis of stereoisomers such as separation of enantiomers and diastereomers, control of optical purity of plant protectives (pesticides, e.g., propionic acid derived herbicides) pharmaceuticals etc. and for product control in chiral organic syntheses
- For control of optical purity of a substance, the columns NUCLEOSIL® CHIRAL-2 and NUCLEOSIL® CHIRAL-3 allow to select conditions such that the minor enantiomer, present as an impurity, is eluted before the main peak. Overlapping peaks are avoided. This makes an exact quantification of the impurity much easier.

Enantiomer separation of *D,L*-supidimide

MN Appl. No. 105690

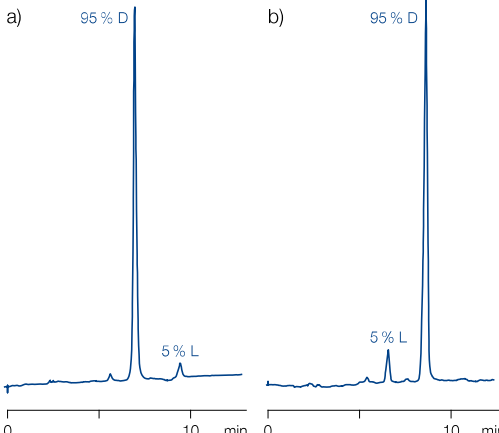
Column: 250 x 4 mm NUCLEOSIL® CHIRAL-2
Eluent: tetrahydrofuran – *n*-heptane (10:3, v/v)
Flow rate: 1.0 mL/min
Detection: UV, 220 nm



Control of optical purity of mecoprop methyl

MN Appl. No. 111360

Columns: a) 250 x 4 mm NUCLEOSIL® CHIRAL-2
b) 250 x 4 mm NUCLEOSIL® CHIRAL-3
Eluent: *n*-heptane – 2-propanol – TFA (100:0.05:0.05, v/v/v)
Flow rate: 1 mL/min, ambient temperature
Detection: UV, 230 nm, Injection 1 µL (sample with 90 % ee)



Ordering information

Eluent in column *n*-heptane – 2-propanol – TFAA (100:0.05:0.05, v/v/v)

ID

Length →
250 mm

EC guard columns*

NUCLEOSIL® CHIRAL-2

Analytical EC columns



4 mm

720088.40

721190.30

NUCLEOSIL® CHIRAL-3

Analytical EC columns



4 mm

720350.40

721190.30

Guard columns for NUCLEOSIL® CHIRAL-2 and CHIRAL-3 are identical.

* EC 4/3 guard columns for EC columns with 4 mm ID require the Column Protection System guard column holder (REF 718966, see page 251). EC columns and EC guard columns in packs of 1.



HPLC columns for biochemical separations

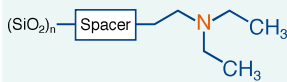
NUCLEOGEN® columns anion exchange chromatography of nucleic acids

NUCLEOGEN® 60-7 DEAE pore size 60 Å

Technical data

- Base material silica, particle size 7 µm; DEAE anion exchanger
- For the separation of oligonucleotides up to chain lengths of 40 bases with recoveries > 95 % capacity 200 A₂₆₀/mL (~ 300 A₂₆₀ for a 125 x 4 mm ID column, 1875 A₂₆₀ for a 125 x 10 mm ID column)

- Preparative separations possible when using higher flow rates and longer gradient times



NUCLEOGEN® 500-7 DEAE pore size 500 Å

Technical data

- Base material silica, particle size 7 µm; DEAE anion exchanger
- For the separation of tRNA, 5S RNA, viroids and messenger RNA in the intermediate molecular weight range (25–1 000 kDa) with recoveries > 95 %

- Capacity 730 A₂₆₀ for a 125 x 6 mm ID column, 1940 A₂₆₀ for a 125 x 10 mm ID column

NUCLEOGEN® 4000-7 DEAE pore size 4000 Å

Technical data

- Base material silica, particle size 7 µm; DEAE anion exchanger
- For the separation of plasmids, DNA restriction fragments, ribosomal RNA, messenger RNA and viral RNA, i.e. very high molecular weight nucleic acids (e.g., 1–50 MDa)

- Capacity 120 A₂₆₀ for a 125 x 6 mm ID column, 350 A₂₆₀ for a 125 x 10 mm ID column

For more separations of deoxyoligonucleotides, plasmids and DNA restriction fragments visit our website
www.mn-net.com/apps



HPLC columns for biochemical separations



Separation of plasmid pBR 322

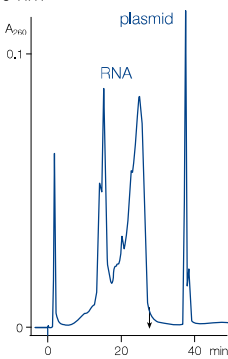
MN Appl. No. 107480

M. Colpan, D. Riesner, private communication

A) isolation of plasmid DNA from a crude cell lysate

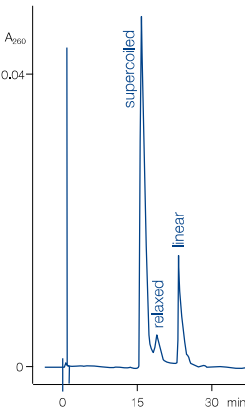
Sample: 5 µg plasmid pBR 322 containing cleared lysate from *E. coli*
Column: 125 x 6 mm NUCLEOGEN® 4000-7 DEAE
Eluent: A) 20 mmol/L K phosphate buffer pH 6.9; 5 mol/L urea
B) eluent A + 1.5 mol/L KCl
20–100 % B in 50 min;
arrow = ionic strength of 850 mmol/L

Flow rate: 1.0 mL/min, 70 bar, ambient temperature
Detection: UV, 260 nm



B) separation of supercoiled plasmid from relaxed and linear forms

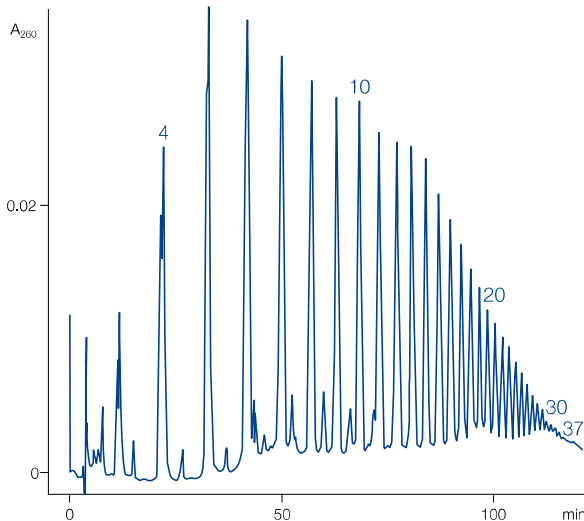
Sample: plasmid pBR 322, supercoiled, relaxed and linear
Column: 125 x 6 mm NUCLEOGEN® 4000-7 DEAE
Eluent: A) 20 mmol/L K phosphate buffer pH 6.8; 6 mol/L urea
B) eluent A + 2 mol/L KCl
42–100 % B in 230 min
Flow rate: 1.5 mL/min, 45 bar, ambient temperature



Separation of oligo(rA)_n

MN Appl. No. 115180

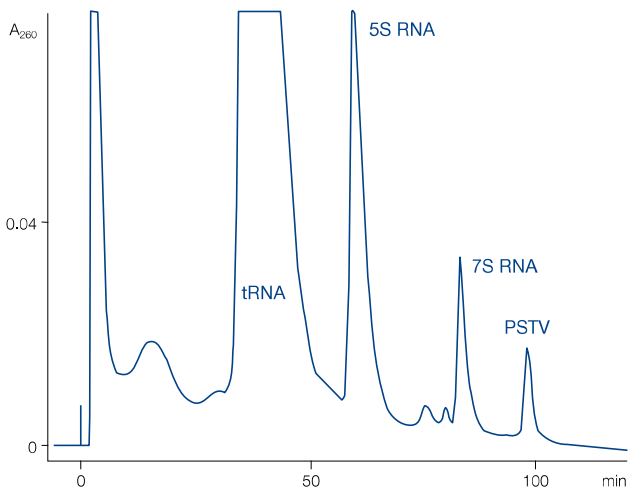
Column: 125 x 4 mm NUCLEOGEN® 60-7 DEAE
Eluent: A) 20 mmol/L phosphate buffer, pH 5.5,
5 mol/L urea
B) buffer A + 1 mol/L KCl
0–100 % B in 200 min
Flow rate: 2 mL/min
Pressure: 110 bar
Temperature: ambient
Detection: UV, 260 nm



Preparative separation of a crude RNA extract of viroid (PSTV) infected tomato plants

MN Appl. No. 107490

D. Riesner, BioEngineering 1 (1988) 42–48
Column: 125 x 6 mm NUCLEOGEN® 500-7 DEAE
Eluent: A) 250 mmol/L KCl, 20 mmol/L phosphate buffer,
pH 6.6, 5 mol/L urea
B) 1 mol/L KCl, 20 mmol/L phosphate buffer, pH 6.6,
5 mol/L urea
0–50 % B in 120 min, 50–100 % B in 250 min
Flow rate: 3 mL/min
Pressure: 40 bar, ambient temperature
Detection: 260 nm





HPLC columns for biochemical separations

Ordering information

Eluent in column methanol

ID	Length → 125 mm	Guard columns*
NUCLEOGEN® 60-7 DEAE particle size 7 µm, pore size 60 Å		
Analytical EC columns		
 4 mm	736596.40	736400.40
Preparative VarioPrep columns		
 10 mm	736597.100	736400.40
NUCLEOGEN® 500-7 DEAE particle size 7 µm, pore size 500 Å		
Analytical Valco type columns		
 6 mm	736598	736400.40
Preparative VarioPrep columns		
 10 mm	736599.100	736400.40
NUCLEOGEN® 4000-7 DEAE particle size 7 µm, pore size 4000 Å		
Analytical Valco type columns		
 6 mm	736601	736400.40
Preparative VarioPrep columns		
 10 mm	736602.100	736400.40

* NUCLEOGEN® guard columns are 30 mm long and require the CC column holder 30 mm (REF 721823).

Columns in packs of 1, guard columns in packs of 2.



HPLC columns for biochemical separations



NUCLEOGEL® SAX anion exchange of biological macromolecules · USP L23

Technical data

- Polymer-based strongly basic anion exchanger -N⁺(CH₃)₃, gel matrix quaternized PEI; particle size 8 µm, pore size 1000 Å
- pH working range 1–13, max. working pressure 200 bar

NUCLEOGEL® SCX cation exchange of biological macromolecules · USP L22

Technical data

- Polymer-based strongly acidic cation exchanger -SO₃⁻, hydrophilic gel matrix; particle size 8 µm, pore size 1000 Å
- pH working range 1–13, max. working pressure 200 bar

Recommended application

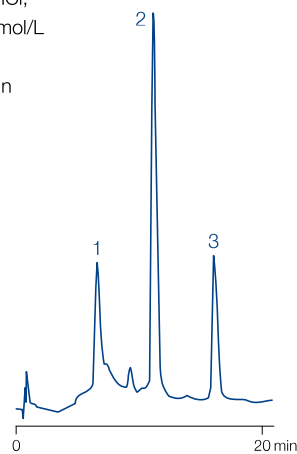
- Purification of peptides, large proteins and oligonucleotides, high capacity for proteins even at pH 10

Separation of hen's egg white

MN Appl. No. 115200

Sample: frozen egg white was thawed, filtered and diluted 1:8 with eluent A
 Column: 50 x 4.6 mm NUCLEOGEL® SAX 1000-8
 Eluent: A) 0.01 mol/L Tris-HCl, pH 7.5; B) A + 0.5 mol/L NaAc, pH 7.5;
 0–100 % B in 20 min
 Flow rate: 1 mL/min
 Inj. volumen: 50 µL
 Detection: UV, 280 nm

Peaks:
 1. Conalbumin
 2. Ovalbumin
 3. not identified

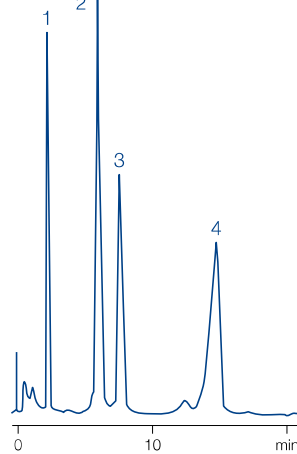


Separation of protein standards

MN Appl. No. 108261

Column: 50 x 4.6 mm NUCLEOGEL® SCX 1000-8
 Eluent: A) 0.02 mol/L KH₂PO₄, pH 6.0
 B) A + 0.5 mol/L NaCl, pH 6.0
 0–100 % B in 20 min
 Flow rate: 1 mL/min
 Detection: UV, 280 nm

Peaks:
 1. Myoglobin
 2. α-Chymotrypsinogen A
 3. Cytochrome C
 4. Lysozyme



Ordering information

Eluent in column 0.1 mol/L Na₂SO₄ + 0.2 % NaN₃

ID

Length →
50 mm

Guard columns*

NUCLEOGEL® SAX pore size 1000 Å

Analytical Valco type columns



4.6 mm

719469

719600

NUCLEOGEL® SCX pore size 1000 Å

Analytical Valco type columns



4.6 mm

719475

719540

* NUCLEOGEL® SAX and SCX Valco type guard columns measure 5 x 3 mm and require the guard column holder B, REF 719539 (see page 250)
 Columns in packs of 1, guard columns in packs of 2.



HPLC columns for biochemical separations

NUCLEODUR® 300 C₁₈ ec · C₄ ec wide pore silica for biochromatography · USP L1 (C₁₈) · USP L26 (C₄)

Key feature

- Reliable wide pore RP phases for daily routine analysis
- Medium density octadecyl or butyl modification with exhaustive endcapping
- Ideal phases for separation of biomolecules

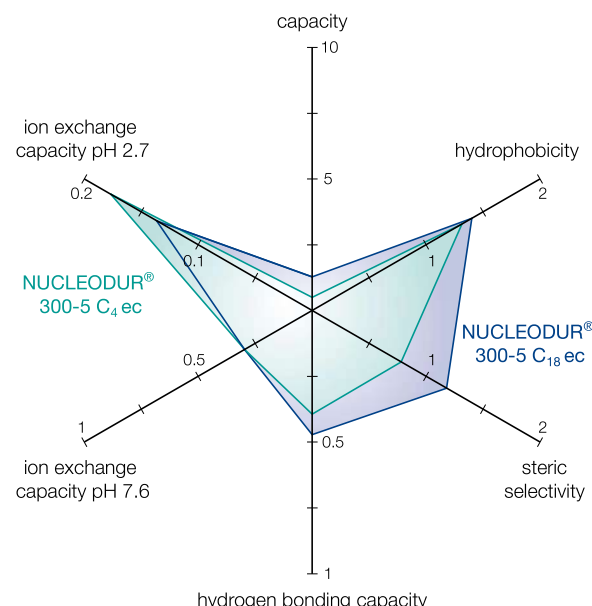
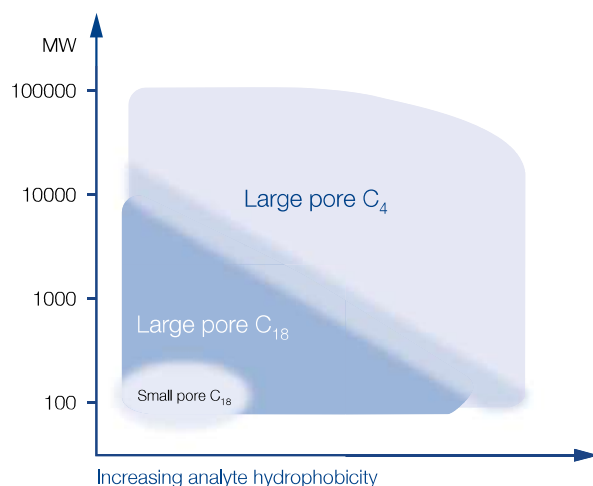
Technical data

- Pore size 300 Å; particle size 5 µm, carbon content 4 % for C₁₈, 2.5 % for C₄; pH stability 1–9; high reproducibility from lot to lot

Recommended application

- Biological macromolecules like proteins or peptides

Column selection by analyte characteristics



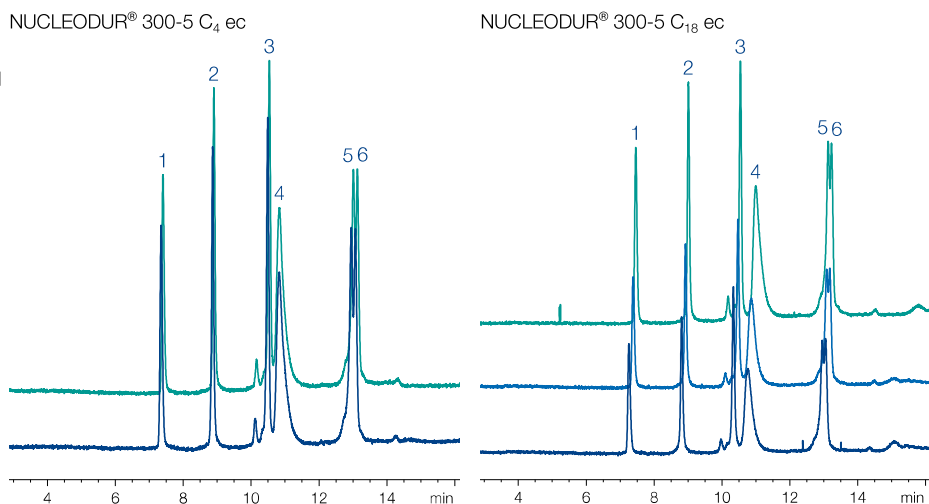
Tanaka plots of NUCLEODUR® wide pore phases

Batch-to-batch reproducibility of NUCLEODUR® 300-5 C₄ ec and NUCLEODUR® 300-5 C₁₈ ec

MN Appl. Nos. 126551 / 126552

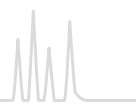
Columns: 250 x 4 mm
 Eluent: A) 0.1 % TFA in water
 B) 0.08 % TFA in acetonitrile
 20–60 % B in 15 min
 Flow rate: 1 mL/min
 Temperature: 25 °C
 Detection: UV, 280 nm

Peaks:
 1. Ribonuclease A
 2. Cytochrome C
 3. Lysozyme
 4. BSA
 5. β-Lactoglobulin
 6. β-Lactoglobulin 2





HPLC columns for biochemical separations



Comparison of narrow and wide pore NUCLEODUR® for the separation of proteins

MN Appl. No. 126590

Columns: 250 x 4.6 mm NUCLEODUR® 300-5 C₁₈ ec
250 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 5 µm

Eluent: A) 0.1 % TFA in water
B) 0.08 % TFA in acetonitrile
20–65 % B in 15 min
(3 min 65 % B)

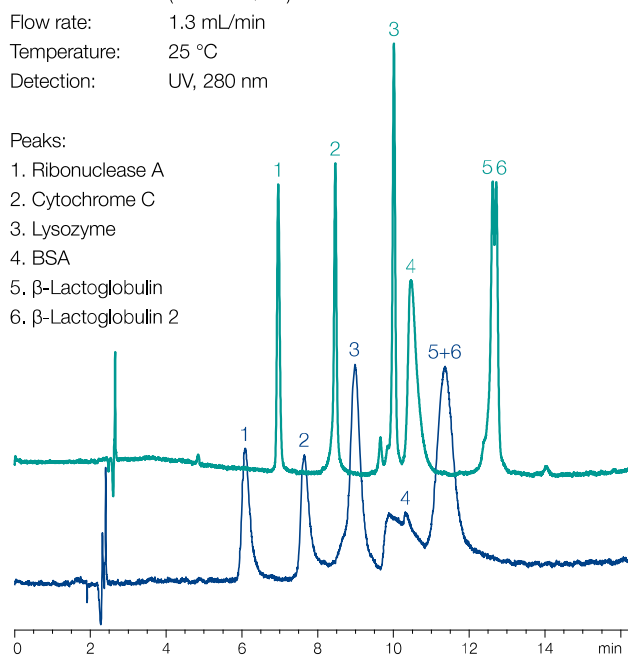
Flow rate: 1.3 mL/min

Temperature: 25 °C

Detection: UV, 280 nm

Peaks:

1. Ribonuclease A
2. Cytochrome C
3. Lysozyme
4. BSA
5. β-Lactoglobulin
6. β-Lactoglobulin 2



Sharper peaks of larger molecules on wide pore material

Tryptic digest of cytochrome C

MN Appl. No. 126600

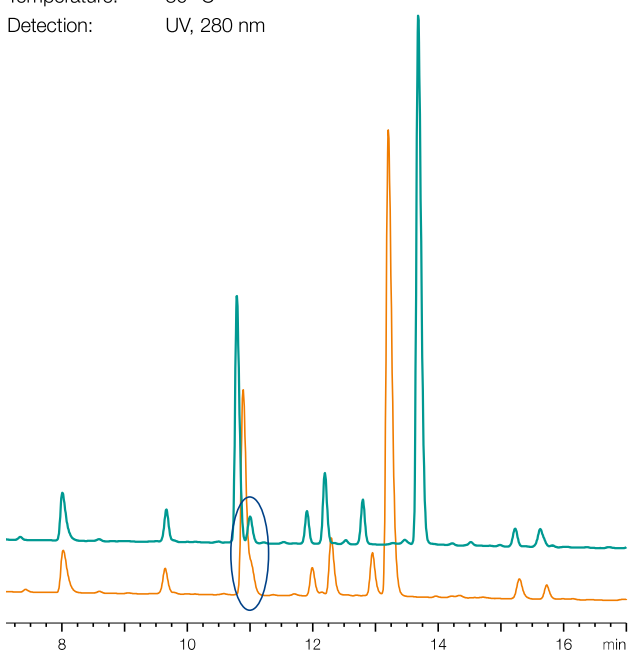
Columns: 250 x 4.6 mm NUCLEODUR® 300-5 C₁₈ ec
250 x 4.6 mm Jupiter® C₁₈, 5 µm

Eluent: A) 0.1 % TFA in water
B) 0.08 % TFA in acetonitrile
5–40 % B in 15 min (1 min 40 % B)

Flow rate: 1.3 mL/min

Temperature: 30 °C

Detection: UV, 280 nm



Less tailing and better separation on NUCLEODUR® 300 C₁₈ ec

Ordering information

Eluent in column acetonitrile – water

ID	Length →	100 mm	125 mm	150 mm	250 mm	EC guard columns*
NUCLEODUR® 300-5 C₁₈ ec octadecyl phase, particle size 5 µm, pore size 300 Å, endcapped, 4 % C						
Analytical EC columns						
	2 mm	760183.20	760184.20	760185.20	760186.20	761988.20
	3 mm	760183.30	760184.30	760185.30	760186.30	761988.30
	4 mm	760183.40	760184.40	760185.40	760186.40	761988.30
	4.6 mm	760183.46	760184.46	760185.46	760186.46	761988.30
NUCLEODUR® 300-5 C₄ ec butyl phase, particle size 5 µm, pore size 300 Å, endcapped, 2.5 % C						
Analytical EC columns						
	2 mm	760193.20	760194.20	760195.20	760196.20	761989.20
	3 mm	760193.30	760194.30	760195.30	760196.30	761989.30
	4 mm	760193.40	760194.40	760195.40	760196.40	761989.30
	4.6 mm	760193.46	760194.46	760195.46	760196.46	761989.30

* EC guard columns require the Column Protection System guard column holder (REF 718966, see page 251).

EC columns in packs of 1, guard columns in packs of 3.



HPLC columns for biochemical separations

NUCLEOSIL® MPN RP chromatography of biological macromolecules

NUCLEOSIL® 100-5 C₁₈ MPN · USP L1

Key feature

- Octadecyl phase, particle size 5 µm; pore size 100 Å
- Dynamic protein binding capacity per g packing: 6 mg BSA, 110 mg cytochrome C
- pH working range 2–8, max. working pressure 250 bar

Technical data

- Silica-based reversed phase materials with monomerically bonded alkyl chains, brush type structure predominantly hydrophobic forces with a small portion of hydrophilic interactions
- Maximum separation efficiency can be achieved when the injected protein mass does not exceed 1–2 % of the maximum protein loading capacity.

NUCLEOSIL® 300-5 C₄ MPN · USP L26

Key feature

- Butyl phase, particle size 5 µm, pore size 300 Å
- Dynamic protein binding capacity per g packing: 14 mg BSA, 27 mg cytochrome C especially suited for the purification of larger, hydrophobic peptides and very different proteins
- pH working range 2–8, max. working pressure 250 bar

Technical data

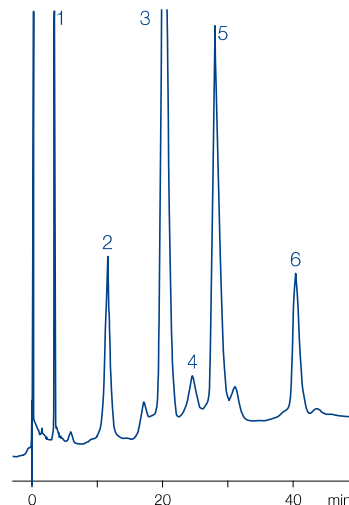
- Silica-based reversed phase materials with monomerically bonded alkyl chains, brush type structure predominantly hydrophobic forces with a small portion of hydrophilic interactions
- Maximum separation efficiency can be achieved when the injected protein mass does not exceed 1–2 % of the maximum protein loading capacity.

Separation of haemoglobin chains

MN Appl. No. 108240

Column: 250 x 4 mm NUCLEOSIL® 300-5 C₄ MPN
 Eluent: A) 20 % acetonitrile, 80 % water, 0.1 % TFA
 B) 60 % acetonitrile, 40 % water, 0.1 % TFA
 40–60 % B in 60 min
 Flow rate: 1 mL/min
 Detection: UV, 220 nm

Peaks:
 1. Hem
 2. β-globin
 3. α-globin
 4. α^T-globin
 5. δ^C-globin
 6. α^L-globin



Ordering information

Eluent in column methanol

ID

Length →
250 mm

EC guard columns*

NUCLEOSIL® 100-5 C₁₈ MPN

Analytical EC columns



4 mm

720231.40

NUCLEOSIL® 300-5 C₄ MPN

Analytical EC columns



4 mm

720245.40

721119.30

* EC guard columns require the Column Protection System guard column holder (REF 718966, see page 251). Columns in packs of 1, guard columns in packs of 2.



HPLC columns for biochemical separations



NUCLEOSIL® PPN RP chromatography of biological macromolecules

NUCLEOSIL® 100-5 C₁₈ PPN · USP L1

Key feature

- Octadecyl phase, particle size 5 µm, pore size 100 Å, dynamic protein binding capacity per g packing: 8 mg BSA, 64 mg cytochrome C; suited for the separation of peptides and proteins up to about 40 kD, also suited for basic peptides

NUCLEOSIL® 500-5 C₁₈ PPN · USP L1

Key feature

- Octadecyl phase, particle size 5 µm, pore size 500 Å, dynamic protein binding capacity per g packing: 22 mg BSA, 40 mg cytochrome C; especially suited for large peptides and medium-size hydrophilic proteins

Separation of a protein standard

MN Appl. No. 108220

Column: 125 x 4 mm NUCLEOSIL® 100-5 C₁₈ PPN

Eluent: A) 0.1 % TFA in H₂O
B) 0.08 % TFA in CH₃CN

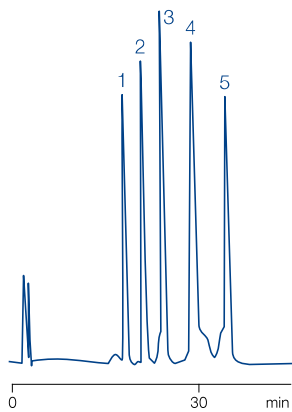
20–60 % B in 10 min

Flow rate: 1.0 mL/min

Detection: UV, 280 nm

Peaks:

1. Ribonuclease
2. Cytochrome C
3. Lysozyme
4. β-Lactoglobulin
5. Ovalbumin



Technical data

- Silica-based reversed phase materials with polymerically bonded alkyl chains; exclusively hydrophobic interactions
- pH working range 1–9, max. working pressure 250 bar

Technical data

- Silica-based reversed phase materials with polymerically bonded alkyl chains; exclusively hydrophobic interactions
- pH working range 1–9, max. working pressure 250 bar

Separation of pancreatic secretion of piglets

MN Appl. No. 108280

Column: 125 x 4 mm NUCLEOSIL® 500-5 C₁₈ PPN

Eluent: A) 0.1 % TFA in H₂O
B) 0.08 % TFA in CH₃CN

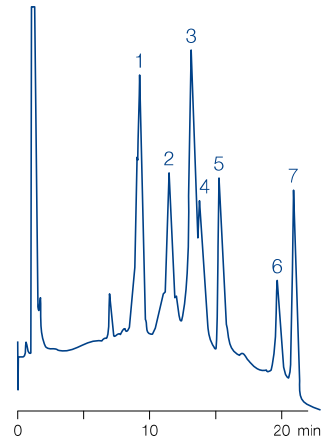
30–50 % B in 14 min, then 50–65 % B in 6 min

Flow rate: 1 mL/min

Detection: UV, 215 nm

Peaks:

1. Trypsin + trypsinogen
2. Proelastase
3. Lipase + α-Chymotrypsin
4. Chymotrypsinogen
5. α-Amylase
- 6., 7. Procarboxypeptidase



Ordering information

Eluent in column methanol

ID

Length →
250 mm

EC guard columns*

NUCLEOSIL® 100-5 C₁₈ PPN particle size 5 µm, pore size 100 Å

Analytical EC columns



4 mm

720252.40

721567.30

NUCLEOSIL® 500-5 C₁₈ PPN particle size 5 µm, pore size 500 Å

Analytical EC columns



4 mm

720258.40

721924.30

* EC guard columns require the Column Protection System guard column holder (REF 718966, see page 251).

Columns in packs of 1, guard columns in packs of 2.



HPLC columns for biochemical separations

NUCLEOGEL® RP columns RP columns for biochemical applications · USP L21

Technical data

- Polystyrene resin cross-linked with divinylbenzene, available particle sizes 5 µm and 8 µm, available pore sizes 100 Å and 300 Å
- pH working range 1–13, max. working pressure 180 bar
- Small pore columns for reversed phase separation of small molecules such as pharmaceuticals with basic properties, e.g., organic heterocycles; also suited for separation of nucleosides and nucleotides up to 5000 Da; allow gradient as well as isocratic elution

- Wide pore columns are especially recommended for large biomolecules higher background hydrophobicity compared to silica phases

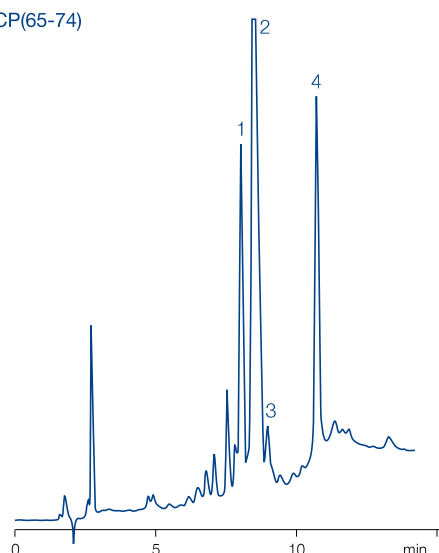
Analysis of the synthetic acyl carrier protein ACP(65-74)

MN Appl. No. 108500

Column: 150 x 4.6 mm NUCLEOGEL® RP 100-8
 Eluant: A) 0.1 % TFA in acetonitrile – water (1:99, v/v)
 B) 0.1 % TFA in acetonitrile – water (99:1, v/v)
 10–60 % B in 20 min
 Flow rate: 1 mL/min
 Detection: UV, 220 nm

Peaks:

1. ACP(66-74)(H-Gln)
2. ACP(65-74)
3. ACP(66-74)(Glp)
4. Thioanisole



Ordering information

Eluent in column acetonitrile – water

ID	Length →	50 mm	150 mm	250 mm	Guard columns*
NUCLEOGEL® RP 100-5 particle size 5 µm, pore size 100 Å					
Analytical Valco type columns					
	4.6 mm		719454	719455	719542
NUCLEOGEL® RP 100-8 particle size 8 µm, pore size 100 Å					
Analytical Valco type columns					
	4.6 mm		719456	719520	719542
NUCLEOGEL® RP 300-5 particle size 5 µm, pore size 300 Å					
Analytical Valco type columns					
	4.6 mm	719459			719542
NUCLEOGEL® RP 300-8 particle size 8 µm, pore size 300 Å					
Analytical Valco type columns					
	4.6 mm	719460			719542

* Valco type guard columns measure 5 x 3 mm and require Guard column holder B, REF 719539, see page 250.
 Columns in packs of 1, guard columns in packs of 2.



HPLC columns for sugar analyses



NUCLEOSIL® Carbohydrate separation of mono- and disaccharides · USP L8

Technical data

- Matrix: NUCLEOSIL® silica with amino modification, particle size 10 µm

Recommended application

- RP separation of mono- and disaccharides

Separation of sugars

MN Appl. No. 102480

Column: 250 x 4 mm NUCLEOSIL® Carbohydrate

Eluent: acetonitrile – water (79:21, v/v)

Flow rate: 2 mL/min

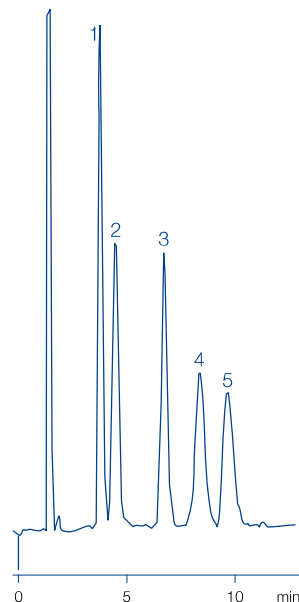
Temperature: 25 °C

Detection: RI

Injection: 10 µL

Peaks:

1. Fructose
2. Glucose
3. Saccharose
4. Maltose
5. Lactose



Ordering information

Eluent in column acetonitrile – water (79:21, v/v)

ID

Length →
250 mm

EC guard columns*

NUCLEOSIL® Carbohydrate

Analytical EC columns



4 mm

720905.40

721170.30

* EC 4/3 guard columns for EC columns with 4 mm ID require the Column Protection System guard column holder (REF 718966, see page 251). Columns and guard columns in packs of 1.



HPLC columns for sugar analyses

NUCLEOGEL® SUGAR 810 separation of sugars · USP L17 (H-Form) · USP L19 (Ca form)

Technical data

- Sulfonated polystyrene - divinylbenzene resins in different ionic forms; due to a different selectivity pattern compared to NUCLEOGEL® SUGAR columns, the range of application is considerably enlarged
- Separation mechanism: ion exclusion, ion exchange, size exclusion, ligand exchange, NP and RP chromatography

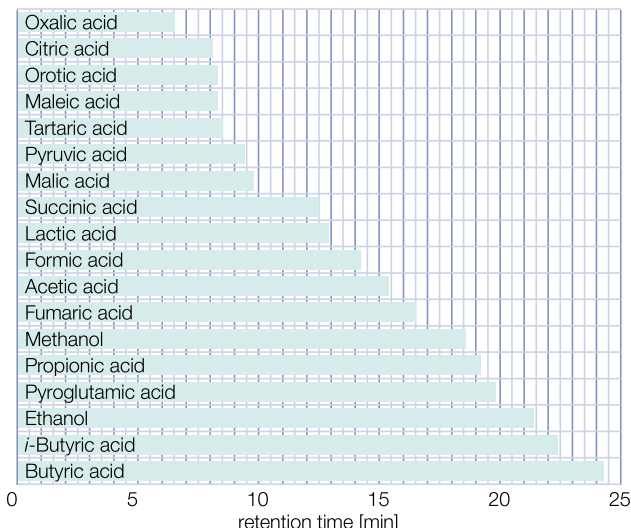
Recommended application

- H^+ form:
Separation of sugars, sugar alcohols and organic acids; eluent in column 5 mmol/L H_2SO_4
- Ca^{2+} form:
Separation of mono-, di- and oligosaccharides; eluent in column water

Organic acids and alcohols

MN Appl. No. 113870

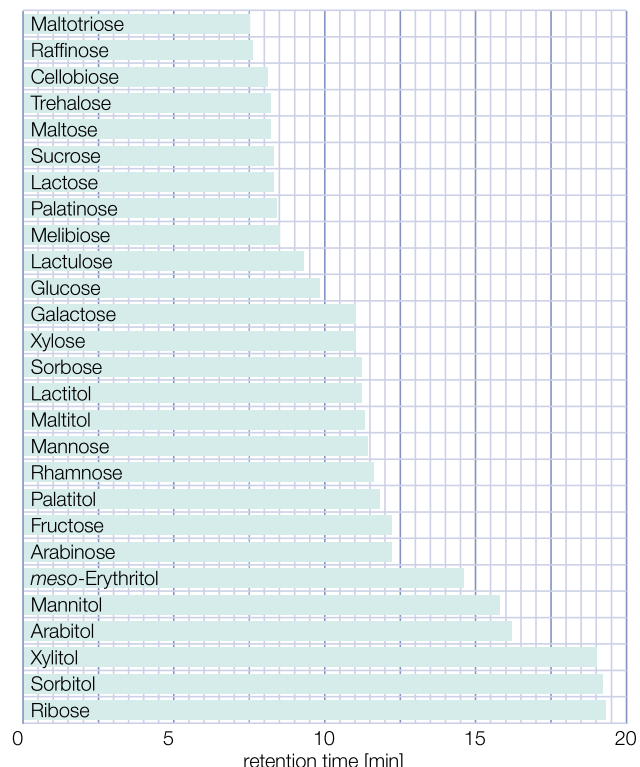
Column: 300 x 7.8 mm NUCLEOGEL® SUGAR 810 H
 Eluent: 5 mmol/L H_2SO_4
 Flow rate: 0.6 mL/min
 Temperature: 35 °C
 Detection: RI
 Injection: 5 μL



Sugars and sugar alcohols

MN Appl. No. 114160

Column: 300 x 7.8 mm NUCLEOGEL® SUGAR 810 Ca
 Eluent: water
 Flow rate: 0.6 mL/min
 Temperature: 85 °C
 Detection: RI



Ordering information

ID

Length →
300 mm

Guard columns*

NUCLEOGEL® SUGAR 810 H eluent in column 5 mmol/L H_2SO_4

Analytical Valco type columns



7.8 mm

719574

719575

NUCLEOGEL® SUGAR 810 Ca eluent in column water

Analytical Valco type columns



7.8 mm

719570

719571

* NUCLEOGEL® SUGAR 810 guard columns measure 30 x 4 mm and require the CC column holder 30 mm (REF 721823). Columns in packs of 1, guard columns in packs of 2.



HPLC columns for sugar analyses



NUCLEOGEL® ION 300 OA / SUGAR

separation of sugars · USP L17 (H form) · USP L19 (Ca form) · USP L34 (Pb form) · USP L58 (Na form)

Technical data

- Sulfonated spherical PS/DVB resins in different ionic forms; mean particle size 10 µm, pore size 100 Å
- Separation mechanism includes steric exclusion, ligand exchange and partition effects, ligand exchange being the predominant force, since the hydrated metal ions form strong interactions with the hydroxyl groups of the sample molecules. The intensity of these interactions decreases in the sequence Pb > Ca > Na
- Recommended operating temperatures: 60–95 °C; maximum pressure 70 bar

Recommended application

- NUCLEOGEL® ION 300 OA:
H⁺ form for separation of sugars, alcohols and organic acids
- NUCLEOGEL® SUGAR:
Ca²⁺ form: separation of mono- and oligosaccharides, sugar alcohols
- Pb²⁺ form: separation of mono- and disaccharides from food and biological samples
- Na⁺ form: separation of oligosaccharides from starch hydrolysates and food

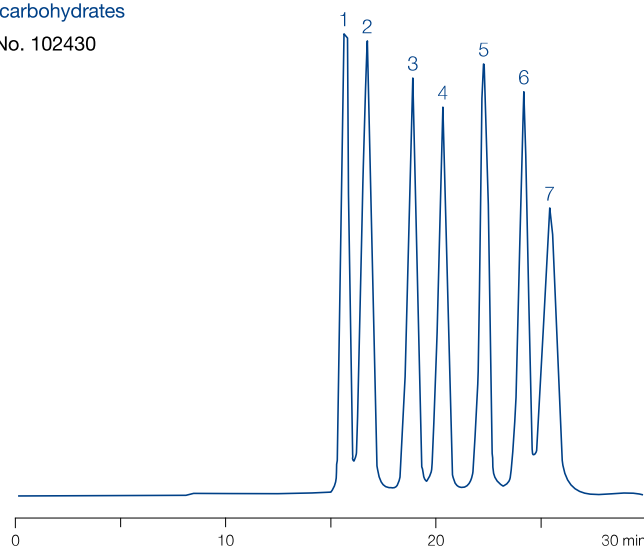
Separation of carbohydrates

MN Appl. No. 102430

Column: 300 x 7.8 mm NUCLEOGEL® SUGAR Pb
Eluent: deionized water
Flow rate: 0.4 mL/min
Temperature: 80 °C
Detection: RI

Peaks:

1. Sucrose
2. Maltose
3. Glucose
4. Xylose
5. Galactose
6. Arabinose
7. Mannose



Ordering information

ID	Length → 300 mm	Guard columns*
NUCLEOGEL® ION 300 OA eluent in column 5 mmol/L H ₂ SO ₄ 5 mmol/L H ₂ SO ₄		
Analytical Valco type columns		
7.8 mm	719501	719537
NUCLEOGEL® SUGAR Ca eluent in column water + 0.02 % azide		
Analytical Valco type columns		
6.5 mm	719531	719535
NUCLEOGEL® SUGAR Pb eluent in column water + 0.02 % azide		
Analytical Valco type columns		
7.8 mm	719530	719534
NUCLEOGEL® SUGAR Na eluent in column water + 0.02 % azide		
Analytical Valco type columns		
7.8 mm	719532	719536

* Valco Type guard columns measure 21 x 4 mm and require the guard column holder C, REF 719538, see page 250.
Columns in packs of 1, guard columns in packs of 2.



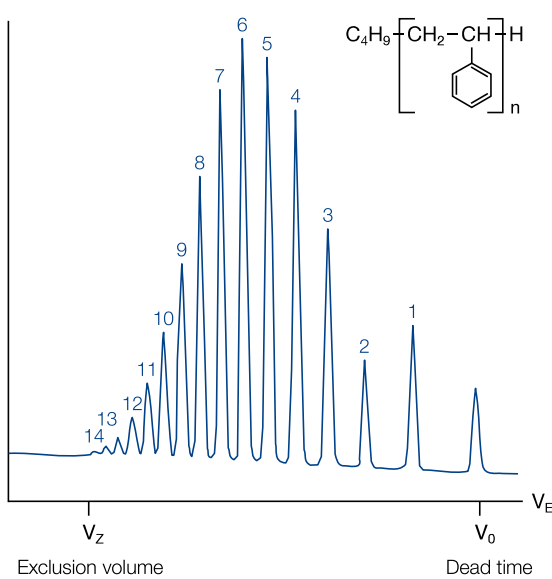
Columns for gel permeation chromatography

NUCLEOGEL® GPC for GPC of water-insoluble substances

Technical data

- Highly crosslinked macroporous, spherical polystyrene – divinylbenzene polymer matrix with good mechanical stability

Chromatogram of styrene oligomers



Working ranges for polystyrene



Ordering information

Eluent in column toluene

Phase	Exclusion limit [kDalton]	Application	Column 300 x 7.7 mm
5 µm particle size			
Analytical Valco type columns			
NUCLEOGEL GPC 50	2	low molecular weight organics	719402
NUCLEOGEL GPC 100	4	oligomers, oils	719403
NUCLEOGEL GPC 500	25	low molecular weight polymers	719404
NUCLEOGEL GPC 103	60	low molecular weight polymers	719405
NUCLEOGEL GPC 104	500	polymers up to 500 kDa	719406
		guard columns 50 x 7.7 mm	719409
10 µm particle size			
Analytical Valco type columns			
NUCLEOGEL GPC 50	2	low molecular weight organics	719410
NUCLEOGEL GPC 100	4	oligomers, oils	719411
NUCLEOGEL GPC 500	25	low molecular weight polymers	719412
NUCLEOGEL GPC 103	60	low molecular weight polymers	719413
NUCLEOGEL GPC 104	500	polymers up to 500 kDa	719414
		guard columns 50 x 7.7 mm	719418

Columns and guard columns in packs of 1.