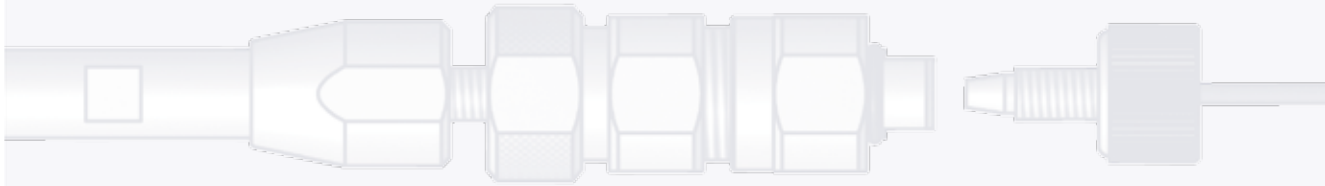


chromatography

NUCLEODUR®

Professional solutions for HPLC



An optimized phase for every field of application



MACHEREY-NAGEL

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MN
Since 1911

Contents / Customer service

NUCLEODUR® high purity silica	1
Overview of NUCLEODUR® phases	2
Summary of available modifications and properties	2
NUCLEODUR® RP phases with 1.8 µm particle size	6
C ₁₈ Gravity / C ₈ Gravity · nonpolar high density phases	8
C ₁₈ Isis · high steric selectivity	12
C ₁₈ Pyramid · for highly aqueous eluents	14
PolarTec · RP phase with embedded polar group	NEW!
PFP · hydrophobic pentafluorophenyl phase	NEW!
Sphinx RP · bifunctional RP phase	20
C ₁₈ HTec · base-deactivated preparative phase	NEW!
C ₁₈ ec / C ₈ ec · nonpolar phases for routine analyses	24
HILIC · zwitterionic ammonium sulfonic acid phase for HILIC	26
CN / CN-RP · cyano-modified high purity silica	28
NH ₂ / NH ₂ -RP · amino-modified high purity silica	30
unmodified high purity NUCLEODUR® silica	31
Applications	32
Pharmaceuticals and drugs	32
Food analysis	42
Biological and natural compounds	46
Pollutants and miscellaneous organics	54
Alphabetical index of analytes	64
Index of application numbers / Trademarks	70
Ordering information of packed columns	71

Meeting your needs

If you have any questions concerning our NUCLEODUR® program or other chromatography products, please feel free to contact us:

Technical support and customer service phone +49 24 21 969-175
e-mail tech-chroma@mn-net.com

The MACHEREY-NAGEL internet catalog with integrated webshop is full of useful information about our wide product range. In addition our online database offers more than 3000 applications which might actually already solve your analytical questions.

www.mn-net.com

	Germany and international	Phone	+49 24 21 969-0
		Toll-free	0800 2616 000
		Fax	+49 24 21 969-199 or -198
		E-mail	info@mn-net.com
	USA	Phone	+1 484 821 0984
		Toll-free	888-321-6224 (MACH)
		Fax	+1 484 821 1272
		E-mail	sales-us@mn-net.com
	France	Phone	+33 388 68 22 68
		Fax	+33 388 51 76 88
		E-mail	sales-fr@mn-net.com
	Switzerland	Phone	+41 62 388 55 00
		Fax	+41 62 388 55 05
		E-mail	sales-ch@mn-net.com



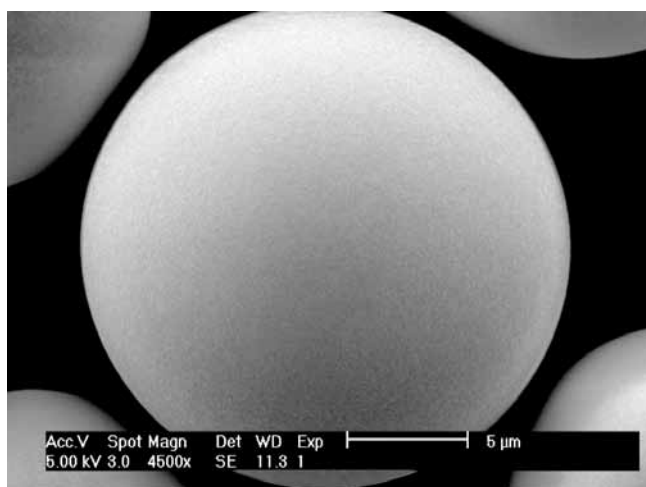
NUCLEODUR®

NUCLEODUR® is a fully synthetical type B silica (silica of 3rd generation) offering highly advanced physical properties like **totally spherical** particle shape, outstanding **surface smoothness**, high pressure stability and **low metal content**.

NUCLEODUR® as a state-of-the-art silica is the ideal base material for modern HPLC phases. It is the result of MACHEREY-NAGEL's pioneering research in chromatography for more than 40 years and worthy successor of MN's world famous NUCLEOSIL® silica.

In RP liquid chromatography the efficiency of the packing is strongly affected by the quality of the base silica itself. Shortcomings in the surface geometry of the particles or metal contaminants are the main reasons for inadequate coverage with the covalently bonded alkylsilanes in the subsequent derivatization steps. It is well known, that poor surface coverage and, in consequence, high activity of residual free silanols often results in peak tailing or adsorption, particularly with basic compounds.

Particle shape and surface symmetry



NUCLEODUR® silicas are synthesized in a unique and carefully controlled manufacturing process which provides silica particles, which are totally spherical. The picture shows the outstanding smoothness of the NUCLEODUR® surface.

Purity

As already mentioned above, a highly pure silica is required for achieving symmetric peak shapes and maximum resolution.

Inclusions of, e.g., iron or alkaline earth metal ions on the silica surface are largely responsible for the unwanted interactions with ionizable analytes, e.g., amines or phenolic compounds (see appl. 118630 on page 63).

NUCLEODUR® is virtually free of metal impurities and low acidic surface silanols. Elemental analysis data of NUCLEODUR® 5 µm measured by AAS are listed below.

Elementary analysis (metal ions) of NUCLEODUR® 100-5

Aluminium	< 5	ppm
Iron	< 5	ppm
Sodium	< 5	ppm
Calcium	< 10	ppm
Titanium	< 1	ppm
Zirconium	< 1	ppm
Arsenic	< 0.5	ppm
Mercury	< 0.05	ppm

Pressure stability

The totally spherical and 100% synthetic silica gel exhibits an outstanding mechanical stability, even at high pressures and elevated eluent flow rates. In addition, after several cycles of repeated packing, no significant drop in pressure can be observed. The latter is of prime importance for preparative and process-scale applications.

Physical data of NUCLEODUR®

Surface area (BET)	340 m ² /g
Pore size	110 Å
Pore volume	0.9 mL/g

NUCLEODUR® modifications

Several different surface modifications based on NUCLEODUR® silica have been developed over the last years providing a full range of specified HPLC phases and an ideal tool for every separation:

- NUCLEODUR® C₁₈ Gravity and C₈ Gravity
- NUCLEODUR® C₁₈ Isis
- NUCLEODUR® C₁₈ Pyramid
- NUCLEODUR® PolarTec
- NUCLEODUR® PFP
- NUCLEODUR® Sphinx RP
- NUCLEODUR® C₁₈ HTec
- NUCLEODUR® C₁₈ ec and C₈ ec
- NUCLEODUR® HILIC
- NUCLEODUR® CN and CN-RP
- NUCLEODUR® NH₂ and NH₂-RP
- unmodified NUCLEODUR®

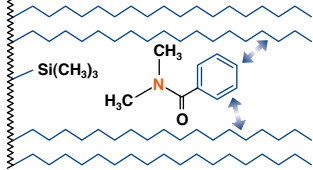
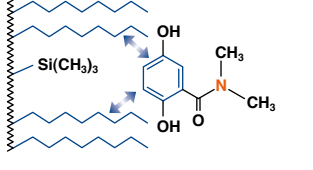
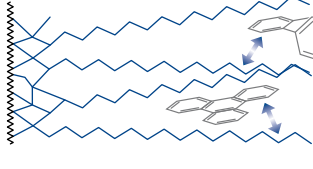
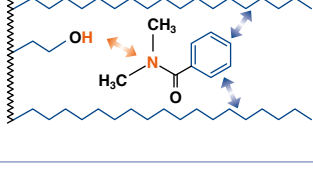
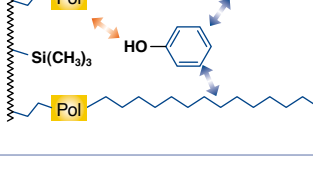
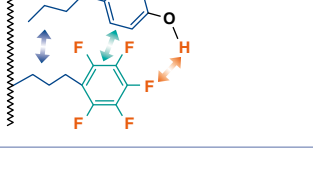
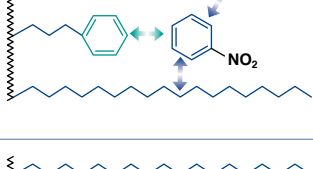
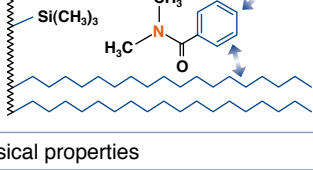
All phases are described in detail on the following pages.

Overview of NUCLEODUR® HPLC phases

Phase	Specification	Characteristics*	Stability	Structure
C ₁₈ Gravity	octadecyl phase, high density coating, multi-endcapping 18% C · USP L1	A	pH stability 1–11, suitable for LC/MS	NUCLEODUR® (Si-O ₂) _n
C ₈ Gravity		B		
C ₁₈ Isis		C		
C ₁₈ Pyramid	C ₁₈ modification with polar endcapping 14% C · USP L1	A	stable against 100% aqueous eluents, pH stability 1–9, suitable for LC/MS	NUCLEODUR® (Si-O ₂) _n
PolarTec		B		
C ₁₈ HTec		C		
PFP	pentafluorophenyl-propyl modification with multi- endcapping 8% C · USP L43	A	pH stability 1–9, suitable for LC/MS	NUCLEODUR® (Si-O ₂) _n
Sphinx RP		B		
C ₁₈ HTec		C		

* A = hydrophobic selectivity, B = polar/ionic selectivity, C = steric selectivity

An optimized phase for every separation

Application	Similar phases**	Separation principle · Retention mechanism
in general compounds with ionizable functional groups such as basic pharmaceuticals and pesticides	NUCLEOSIL® C₁₈ HD XTerra® RP18 / MS C ₁₈ ; Luna® C18(2), Gemini®, Synergi® Max RP; Zorbax® Extend-C18; Inertsil® ODS III; Purospher® STAR RP-18; Hypersil™ BDS	only hydrophobic interactions (van der Waals interactions) 
like C ₁₈ Gravity, however generally shorter retention times for nonpolar compounds	NUCLEOSIL® C₈ HD XTerra® RP8 / MS C ₈ ; Luna® C8; Zorbax® Eclipse XDB-C8	only hydrophobic interactions (van der Waals interactions) 
high steric selectivity, thus suited for separation of positional and structural isomers, planar/nonplanar molecules	NUCLEOSIL® C₁₈ AB Inertsil® ODS-P; Pro C18 RS; Zorbax® SB	steric interactions and hydrophobic interactions 
basic pharmaceutical ingredients, very polar compounds, organic acids	Aqua, Synergi® Hydro-RP; AQ; Atlantis® dC ₁₈	hydrophobic interactions and polar interactions (H bonds) 
basic pharmaceuticals, organic acids, pesticides, amino acids, water-soluble vitamins	NUCLEOSIL® C₁₈ Nautilus ProntoSIL® C18; Zorbax® Bonus-RP, Polaris® Amide-C18; Ascentis® RP Amide; SymmetryShield™ RP18; SUPELCOSIL™ LC-ABZ+; HyPURITY™ ADVANCE	hydrophobic interactions and polar interactions (H bonds) 
aromatic and unsaturated compounds, halogen compounds, phenols, isomers, polar pharmaceuticals, antibiotics	ACQUITY® CSH Fluoro-Phenyl; Hypersil™ GOLD PFP; Luna® PFP(2); Discovery® HS F5; Allure® PFP Propyl, Ultra II PFP Propyl	polar interactions (H bonds), dipole-dipole interactions π-π interactions and hydrophobic interactions 
compounds with aromatic and multiple bond systems	no similar phases	π-π interactions and hydrophobic interactions 
robust and well base deactivated C ₁₈ phase; all separation tasks with preparative potential	XTerra® RP18 / MS C ₁₈ / SunFire™ C ₁₈ ; Luna® C18(2), Gemini®, Synergi® Max RP; Zorbax® Extend-C18; Inertsil® ODS III; Purospher® STAR RP-18; Hypersil™ BDS	only hydrophobic interactions (van der Waals interactions) 

** phases which provide a similar selectivity based on chemical and physical properties



Overview of NUCLEODUR® HPLC phases

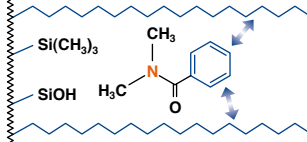
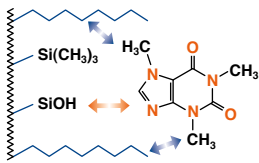
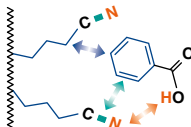
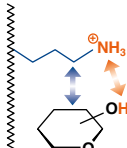
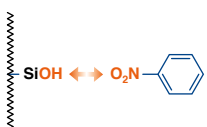
NUCLEODUR®

Phase	Specification	Characteristics*	Stability	Structure
C ₁₈ ec	octadecyl phase, medium density coating endcapping 17.5 % C · USP L1	A B C	pH stability 1–9	NUCLEODUR® (Si-O ₂) _n
C ₈ ec	octyl phase, medium density coating endcapping 10.5 % C · USP L7	A B C		
HILIC	zwitterionic ammonium sulfonic acid modification 7 % C	A B C -		
CN / CN-RP	cyano (nitrile) phase for NP and RP separations 7 % C · USP L10	A B C -	pH stability 1–8, stable towards highly aqueous mobile phases	NUCLEODUR® (Si-O ₂) _n
NH ₂ / NH ₂ -RP	amino phase for NP and RP separations 2.5 % C · USP L8	A B C -	pH stability 2–8, stable towards highly aqueous mobile phases	NUCLEODUR® (Si-O ₂) _n
SiOH	unmodified high purity silica USP L3	A - B n.a. C -	pH stability 2–8	NUCLEODUR® (Si-O ₂) _n

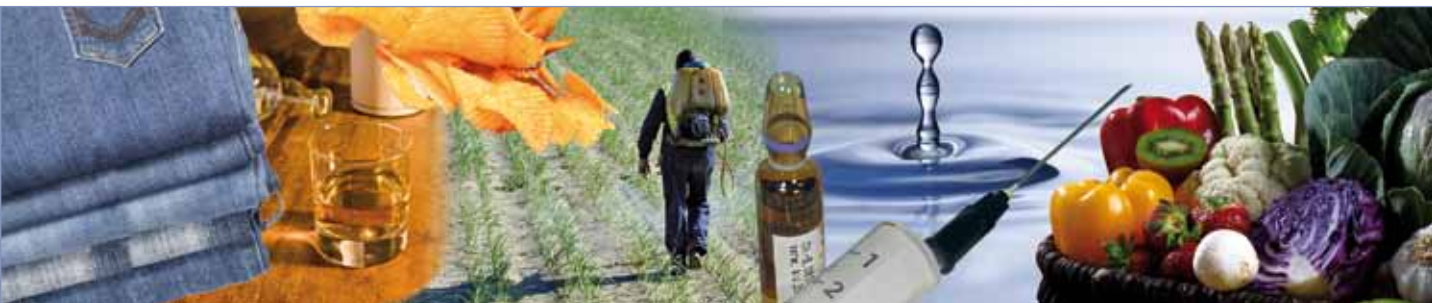
* A = hydrophobic selectivity, B = polar/ionic selectivity, C = steric selectivity



An optimized phase for every separation

Application	Similar phases**	Separation principle · Retention mechanism
robust C ₁₈ phase for routine analyses	NUCLEOSIL® C₁₈ Spherisorb® ODS II; Symmetry® C ₁₈ ; Hypersil™ ODS; Inertsil® ODS II; Kromasil C ₁₈ ; LiChrospher® RP-18	only hydrophobic interactions (van der Waals interactions) some residual silanol interactions 
robust C ₈ phase for routine analyses	NUCLEOSIL® C₈ ec / C₈ Spherisorb® C ₈ ; Symmetry® C ₈ ; Hypersil™ MOS; Kromasil C ₈ ; LiChrospher® RP-8	only hydrophobic interactions (van der Waals interactions) some residual silanol interactions 
hydrophilic compounds such as organic polar acids and bases, polar natural compounds	SeQuant™ ZIC®-HILIC; Obelisc™	ionic / hydrophilic interactions, electrostatic interactions 
polar organic compounds (basic drugs), molecules containing π electron systems	NUCLEOSIL® CN / CN-RP	π-π interactions, polar interactions (H bonds), hydrophobic interactions 
sugars, sugar alcohols and other hydroxy compounds, DNA bases, polar compounds in general	NUCLEOSIL® NH₂ / NH₂-RP	polar / ionic interactions, hydrophobic interactions 
polar compounds in general	unmodified NUCLEOSIL®	polar / ionic interactions 

** phases which provide a similar selectivity based on chemical and physical properties



1.8 µm particle size

Key features

- Decrease of analysis time (ultra fast HPLC)
 - Shorter columns with high separation efficiency
 - Significant improvement of resolution and detection sensitivity
 - Suitable for LC/MS due to low bleeding characteristics
- NUCLEODUR® 1.8 µm particles are fractionated to limit the increase in back pressure

NUCLEODUR® phases available in 1.8 µm:

C₁₈ Gravity
C₈ Gravity
C₁₈ Isis
C₁₈ Pyramid
Sphinx RP
C₁₈ HTec
HILIC

Advantages of 1.8 µm particle size

Miniaturization in HPLC has a long history. It started in the early stage of HPLC development with the reduction of particle size from 10 µm via 7 µm to standard 5 µm – which is still the most widely used particle diameter in analytical HPLC – to 3 µm spherical particles which so far was the smallest particle size available for gaining higher theoretical plates and efficiencies. With the introduction of 1.8 µm NUCLEODUR® particles researchers have turned over a new leaf in HPLC column technology. Columns packed with these microspherical particles show extraordinary improvements in terms of plate numbers, column efficiencies and resolution compared with their 3 µm counterparts.

Features of 1.8 µm NUCLEODUR® silica particles

Increase of separation efficiency by higher number of theoretical plates (N):

50 x 4.6 mm NUCLEODUR® C₁₈ Gravity

3 µm: N ≥ 100 000 plates/m (h value ≤ 10)

1.8 µm: N ≥ 166 667 plates/m (h value ≤ 6)

Increase of the plate number by app. 67% offers the possibility of using shorter columns with equal plate numbers resulting in a decrease of analysis time.

Significant improvement in resolution

Use of 1.8 µm instead of 3 µm particles leads to an increase of resolution by a factor 1.29 (29%) since the resolution is inversely proportional to the square root of the particle size:

$$R_s = \frac{\sqrt{N}}{4} \left(\frac{\alpha - 1}{\alpha} \right) \left(\frac{k_i'}{k_i' + 1} \right)$$

R_s = resolution

α = selectivity (separation factor)

k_{i'} = retention

N = plate number with N ∝ 1/d_p

d_p = particle size

Resolution as a function of particle size

Column: 50 x 4 mm NUCLEODUR® C₁₈ Gravity
A) 3 µm, B) 1.8 µm

Eluent: acetonitrile – water (80:20, v/v)

Flow rate: 2 mL/min, pressure: A) 80 bar, B) 160 bar

Detection: UV, 254 nm

Peaks:

1. Naphthalene

2. Ethylbenzene

A) 3 µm

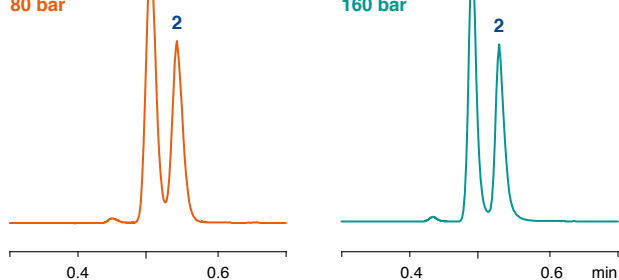
R_s = 1.11

80 bar

B) 1.8 µm

R_s = 1.42

160 bar



Increase in separation efficiency

Column back pressure

Due to the smaller particle size the back pressure will increase according to

$$\Delta_P = \frac{\Phi \cdot L_C \cdot \eta \cdot u}{d_P^2}$$

Δ_P = pressure drop
 Φ = flow resistance (nondimensional)
 L_C = column length
 η = viscosity
 u = linear velocity
 d_P = particle diameter

Because of the high sphericity of the NUCLEODUR® particles and the very narrow particle size distribution we were able to keep the back pressure on a moderate level. Nevertheless the use of columns packed with sub 2 µm particles generally makes special demands on the HPLC equipment. Pumps should be designed for pressures of 250–1000 bars and the entire system should feature the lowest possible dead volume.

Comparison of back pressures:

Eluent: 100% methanol
Flow rate: 1.5 mL/min
Temperature: 22 °C
Column dimension: 50 x 4.6 mm

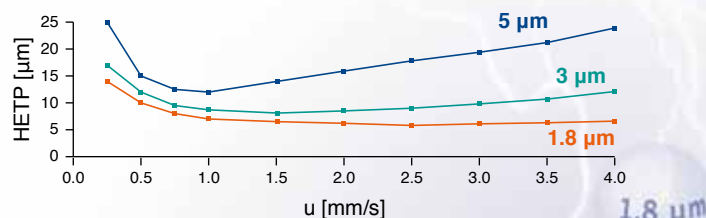
	NUCLEODUR® C ₁₈ Gravity	Competitor A
3 µm	70 bar	–
1.8 µm	130 bar	170 bar

Higher flow rates and shorter run times

optimal flow rate for 1.8 µm particles is higher than for 3 and 5 µm particles (see figures – the flow rate should be at the van-Deemter minimum)

Van-Deemter plot

column 50 x 4.6 mm, acetonitrile – water (50:50, v/v), analyte toluene



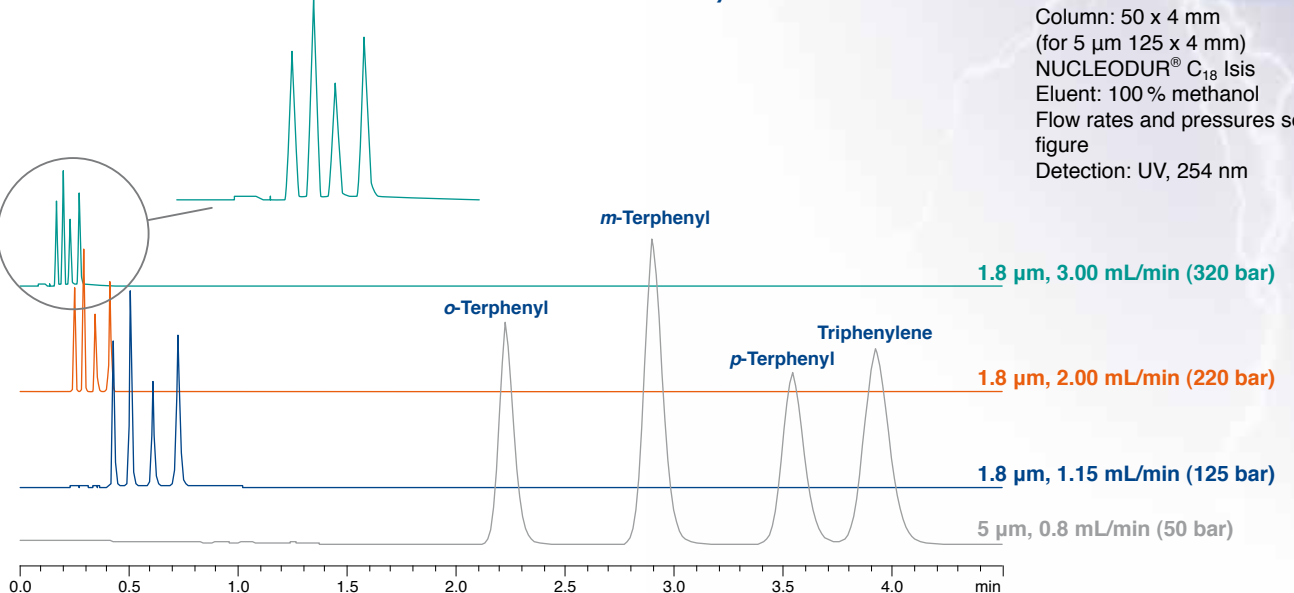
Technical requirements

To gain the best result in ultra fast HPLC based on 1.8 µm particles certain technical demands on the instrument are made. Pumps for pressures of 250–1000 bar realizing a flow rate of 2–3 mL are required. The dead volume of the LC system has to be reduced to a minimum. In addition, fast data recording is necessary for an optimum chromatographic result.

Currently the following NUCLEODUR® premium phases (C₁₈ Gravity, C₈ Gravity, C₁₈ Isis, C₁₈ Pyramid, Sphinx RP, C₁₈ HTec, HILIC) are available in 1.8 µm. The description of each phase and its selectivity can be found in the individual chapters.

More applications on NUCLEODUR® 1.8 µm can be found in the "Applications" section from page 32.

Reduction of analysis time



C₁₈ Gravity / C₈ Gravity

Key features:

- Suitable for LC/MS and HPLC at pH extremes (pH 1–11)
- Superior base deactivation
- Ideal for method development

Technical characteristics:

Available as octadecyl (C₁₈) and octyl (C₈), multi-endcapped; pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm for C₁₈, 1.8 and 5 µm for C₈; 7, 10, 12 and 16 µm particles for preparative purposes on request; carbon content 18% for C₁₈, 11% for C₈

Recommended application:

Overall sophisticated analytical separations

Compound classes separated include: pharmaceuticals, e.g., analgesics, anti-inflammatory drugs, antidepressants; herbicides; phytopharmaceuticals; immunosuppressants

USP L1 (C₁₈) / USP L7 (C₈)

Base deactivation

NUCLEODUR® C₁₈ Gravity and NUCLEODUR® C₈ Gravity are based on the ultrapure NUCLEODUR® silica.

A unique derivatization process generates a homogeneous surface with a high density of bonded silanes (carbon content ~18% for C₁₈, ~11% for C₈). The following thorough endcapping suppresses any unwanted polar interactions between the silica surface and the sample, which makes "Gravity" particularly suitable for the separation of basic and other ionizable analytes. The figure on the right shows a comparison study, where the strongly basic amitriptyline is eluted on various highly base deactivated C₁₈ phases under isocratic conditions. For a discussion of the different retention behavior of octadecyl phases compared to octyl phases see page 25.

Tanaka diagrams

Several NUCLEODUR® phases have been examined in accordance with Tanaka et al. [J. Chromatogr. Sci. 27 (1989) 721] and Johnson et al. [Chromatographia 44 (1997) 151] with respect to the following parameters:

Capacity = k' (pentylbenzene)

Hydrophobicity = α (pentylbenzene, butylbenzene)

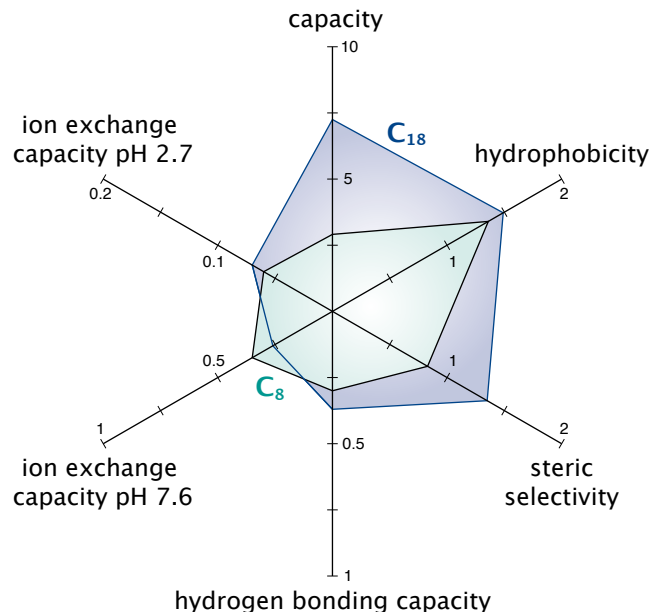
Steric selectivity = α (triphenylene, *o*-terphenyl)

Hydrogen bonding capacity (silanol capacity) = α (caffeine, phenol)

Ion exchange capacity at 2 different pH values (2.7 and 7.6) = α (benzylamine, phenol)

The resulting Tanaka plots are shown with the respective phases.

Tanaka plots of NUCLEODUR® C₈ and C₁₈ Gravity



Nonpolar high density phases

Comparison of different base deactivated phases

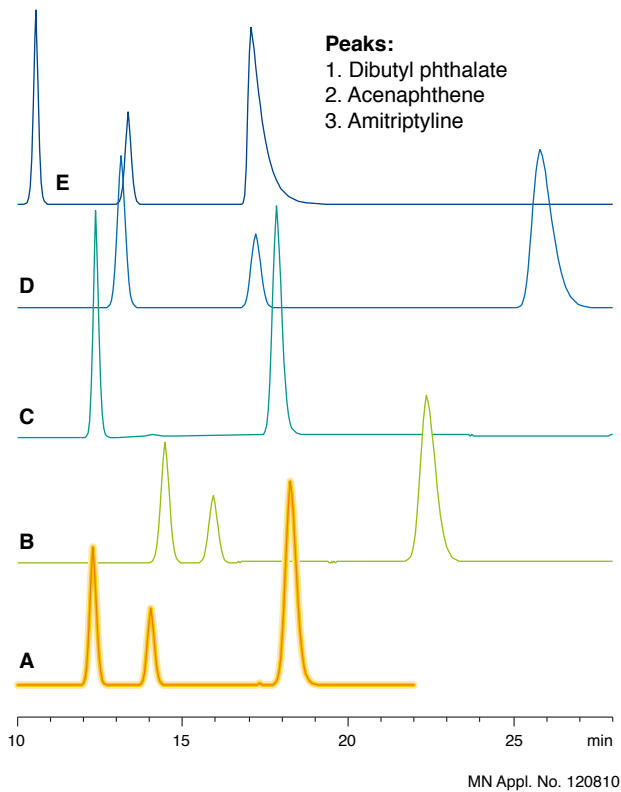
Columns: 250 x 4 mm, all phases C₁₈, 5 µm
 A) NUCLEODUR Gravity
 B) phase I
 C) phase L (1 and 2 overlap)
 D) phase P
 E) phase S

Eluent: methanol – 20 mM KH₂PO₄, pH 7.0 (75:25, v/v)

Flow rate: 1.0 mL/min

Temperature: 30 °C

Detection: UV, 254 nm



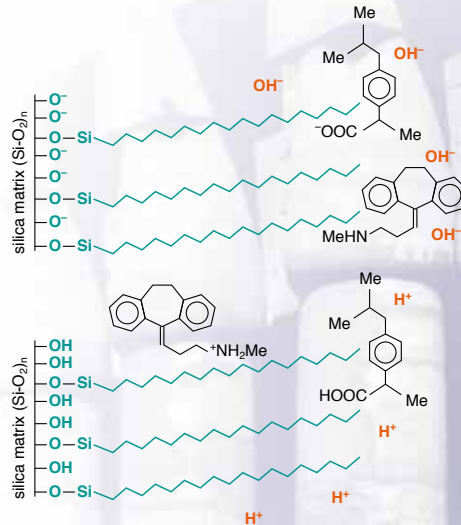
Enhanced pH stability

One major disadvantage of using silica stationary phases is the limited stability at strongly acidic or basic pH ranges. Cleavage of the siloxane bonding by hydrolysis, or dissolution of the silica will rapidly lead to a considerable loss in column performance. Therefore conventional RP phases are usually not recommended to be run with mobile phases at pH > 8 or pH < 2 for extended periods of time. The special surface bonding technology and the low concentration of trace elements of NUCLEODUR® C₈ / C₁₈ Gravity allow for use at an expanded pH range from pH 1 to 11.

When is enhanced pH stability beneficial?

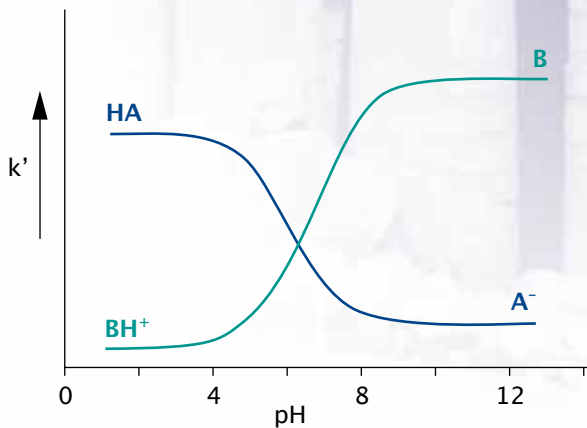
The option to work at an expanded pH range is often required in method development. Many nitrogen containing compounds like basic drugs are protonated at acidic or neutral pH and exhibit poor retention on a standard C₁₈ phase. The retention behavior can be improved by working at a higher pH, where the analyte is no longer protonated, but formally neutrally charged, as a rule between pH 9-10. For acidic analytes it is exactly in inverse proportion, maximum retention can be attained at low pH.

Surface silanols at different pH values



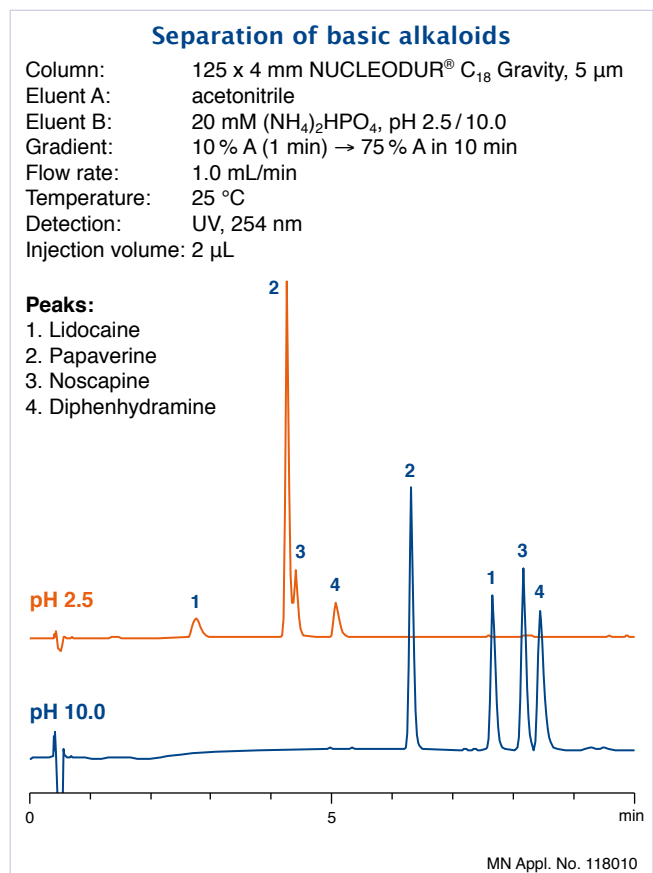
The figure above shows the extent of protonation of surface silanols and of two exemplary analytes at acidic and alkaline pH. The following graph explains the general correlation between retention and pH.

Correlation between retention and pH for basic and acidic compounds

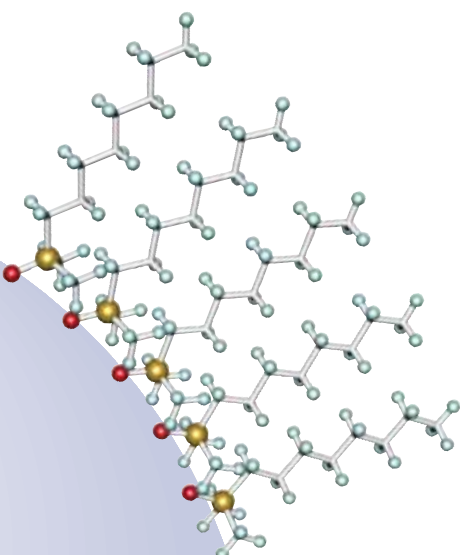


C₁₈ Gravity / C₈ Gravity

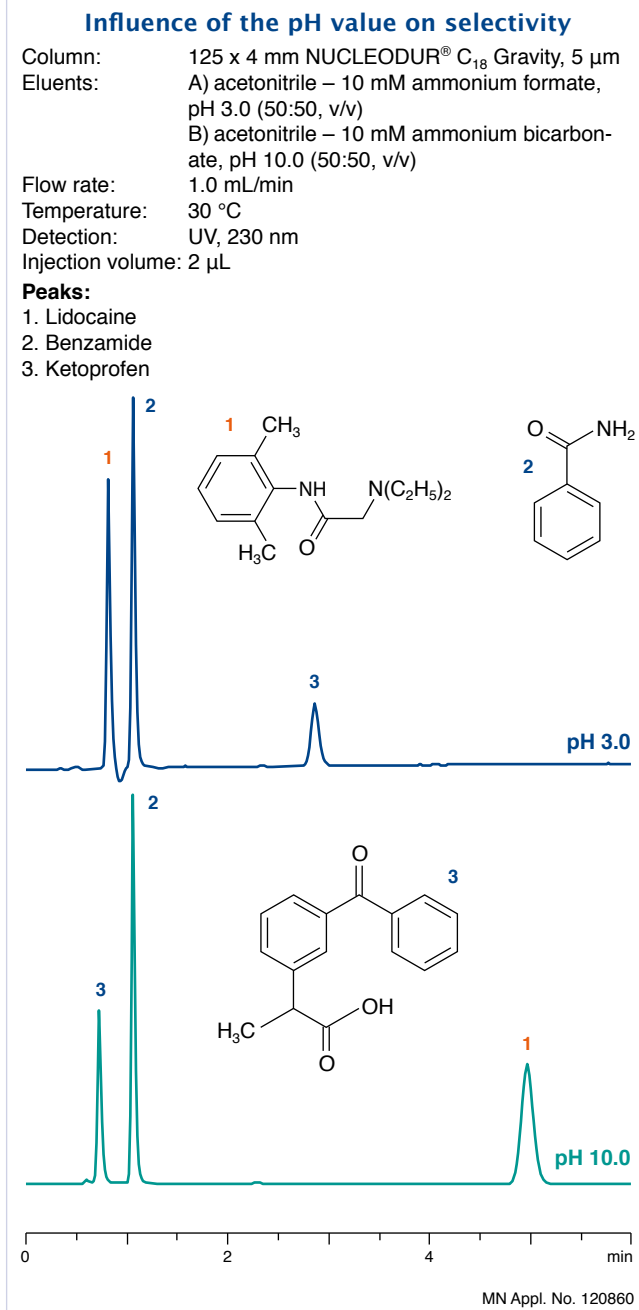
As it was previously mentioned, pH stability of the stationary phase can be helpful for improving selectivity in method development. The figure below shows the separation of 4 basic drugs under acidic and basic conditions.



At pH 2.5 the protonated analytes exhibit poor retention (early elution) and in addition an inadequate resolution for papaverine and noscapine, whilst the formally non ionized molecules can be baseline separated due to the better retention pattern at alkaline pH.



A further example how selectivity can be controlled by the pH value is demonstrated below. The sample mixture consists of an acid (ketoprofen), a base (lidocaine) and benzamide. Under acidic conditions the protonated lidocaine is eluted very fast due to lack of sufficiently strong hydrophobic interactions between analyte and C₁₈ chains, in contrary to the formally neutral ketoprofen, which is eluted after about 3 minutes. However at pH 10 a reversal of the elution order, with a visibly longer retention time for the basic lidocaine, can be achieved.



Nonpolar high density phases

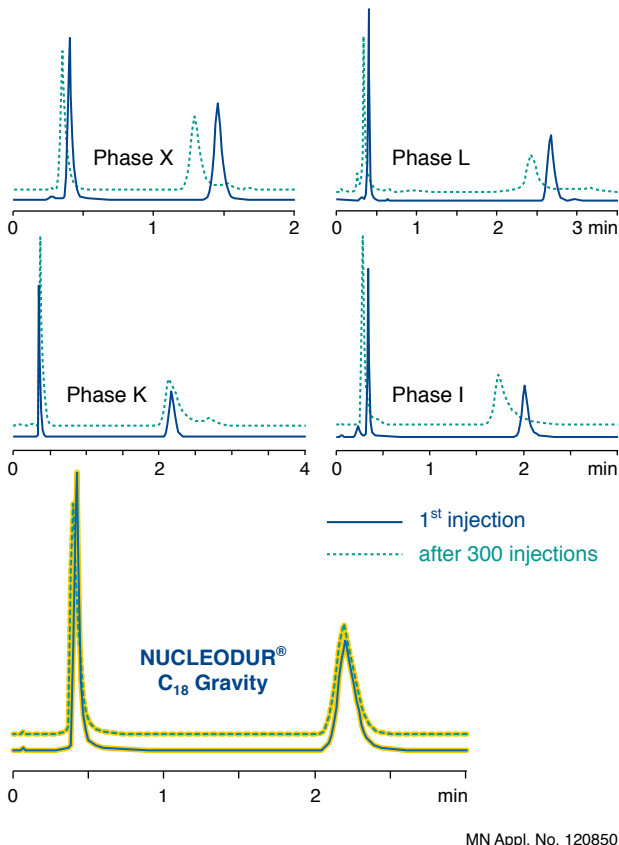
The following chromatograms demonstrate the stability of NUCLEODUR® C₁₈ Gravity under alkaline conditions in comparison with 4 commercially available modern RP18 phases. Again, the ultrapure Gravity with its unique high density surface bonding technology withstands strong alkaline mobile phase conditions. Even after 300 injections no loss of column efficiency, identified, e.g., by peak broadening or decrease in retention times, could be observed.

Stability of NUCLEODUR® C₁₈ Gravity under alkaline conditions compared with different C₁₈ phases

Columns: 50 x 4.6 mm
Eluent: methanol – water – ammonia (20:80:0.5, v/v/v), pH 11
Flow rate: 1.3 mL/min
Temperature: 30 °C
Detection: UV, 254 nm
Injection volume: 2.0 µL

Peaks:

1. Theophylline
2. Caffeine



MN Appl. No. 120850

The pH stability of silica under alkaline conditions is mainly a kinetic effect and based on the velocity of the dissolution of the silica support. It is worth mentioning, that this phenomenon also depends on type and concentration of buffers, as well as on the temperature. It is well known, that the use of phosphate buffers, particularly at elevated temperatures, can reduce column lifetime even at moderate pH. If possible, phosphate buffers should be replaced by less harmful alternatives.

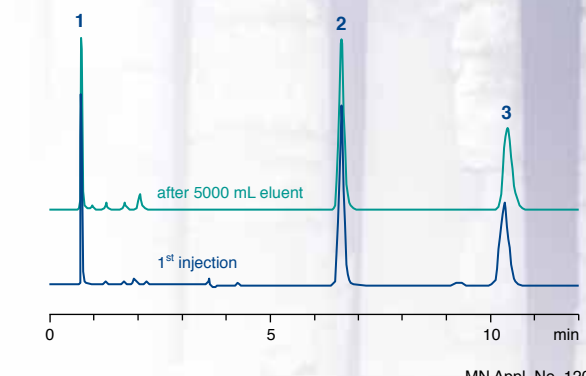
The following chromatograms show the excellent column stability of NUCLEODUR® C₁₈ Gravity in acidic conditions. The retention time of all three compounds in the column performance test remains consistent and virtually unchanged, even after the column is run with 5000 mL eluent. Due to the extremely stable surface modification, no cleavage of the Si-O-Si bonding occurs, column deterioration is therefore successfully prevented.

Stability of NUCLEODUR® C₁₈ Gravity at pH 1.5

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
Eluent: acetonitrile – 1 % TFA in water (50:50, v/v), pH 1.5
Flow rate: 1.0 mL/min
Temperature: 30 °C,
Detection: UV, 230 nm
Injection volume: 5 µL

Peaks:

1. Pyridine
2. Toluene
3. Ethylbenzene



MN Appl. No. 120840

For comparison of the selectivity of NUCLEODUR® C₈ Gravity and C₁₈ Gravity please also see the application "Retention behavior of different NUCLEODUR® phases" on page 15. Some general selection criteria and principles of different retention and selectivity of C₁₈ and C₈ columns can be found on page 25.

C₁₈ Isis

Key features:

- Exceptional steric selectivity
- Outstanding surface deactivation
- Suitable for LC/MS and HPLC at pH 1-10

Technical characteristics:

C₁₈ phase with special polymeric, crosslinked surface modification; pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm; carbon content 20%

Recommended application:

Steroids,
(o, p, m-) substituted aromatics,
fat-soluble vitamins
USP L1

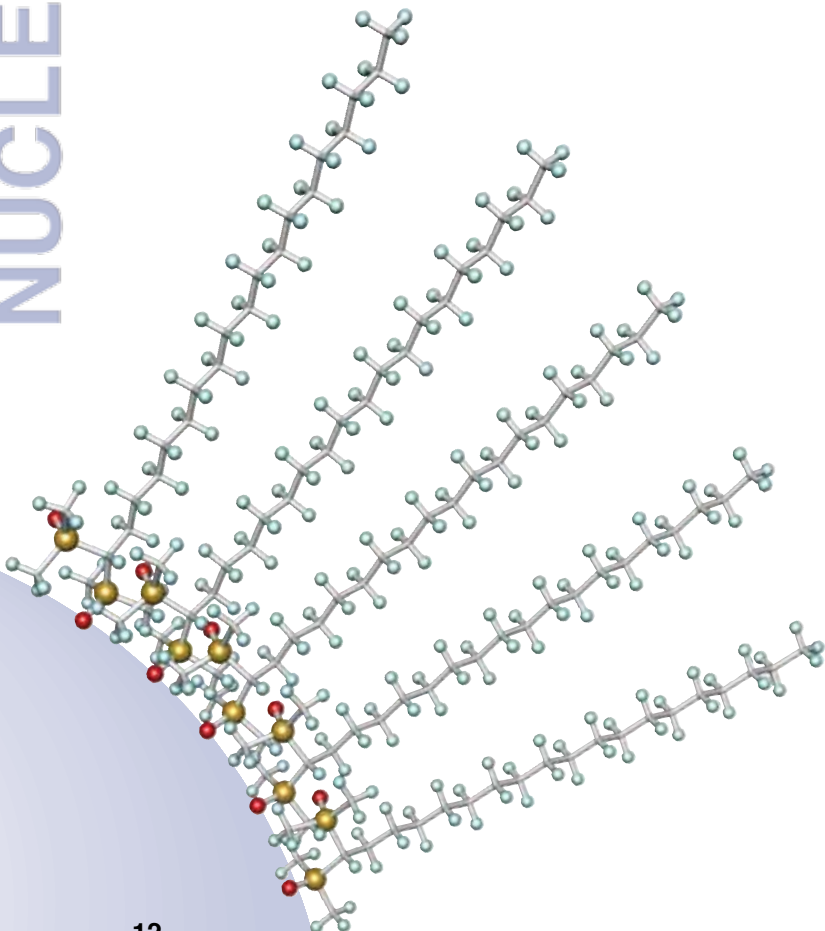
Surface modification

By use of specific C₁₈ silanes and appropriate polymeric bonding technologies a dense shield of alkyl chains protects the subjacent silica matrix. Elemental analysis of NUCLEODUR® C₁₈ Isis shows a carbon load of 20%.

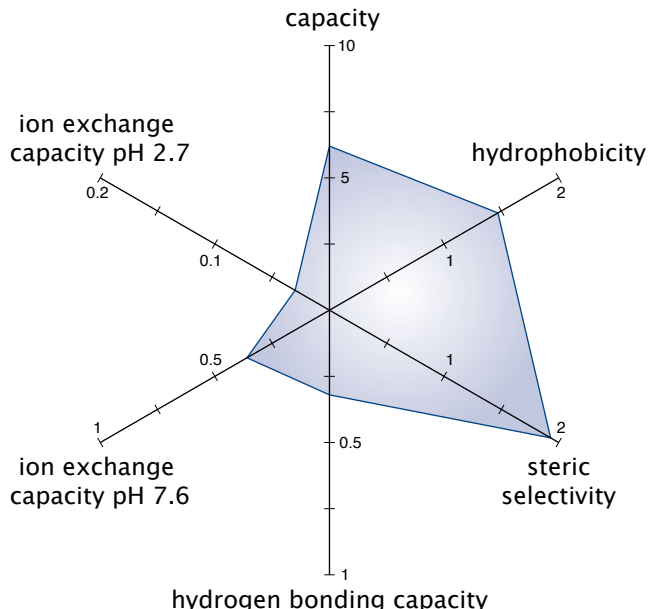
The target crosslinking of the C₁₈ chains on the surface enables the separation of compounds with similar molecular structure but different stereochemical properties. The technical term for this feature is steric selectivity.

The chromatograms on the right reveal the improved resolution for positional isomers in a test mixture of aromatic compounds on NUCLEODUR® C₁₈ Isis (1) in direct comparison with monomerically coated (2) and polar endcapped (3) C₁₈ columns.

NUCLEODUR®



Tanaka plot of NUCLEODUR® C₁₈ Isis



Steric selectivity of NUCLEODUR® C₁₈ Isis

Columns: 125 x 4 mm; NUCLEODUR® C₁₈ Isis, monomerically coated C₁₈ phase, polar endcapped phase

Eluent: methanol – water (90:10, v/v)

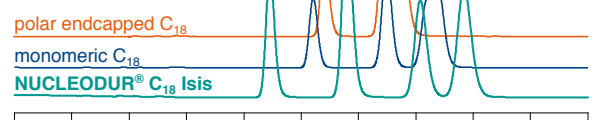
Flow rate: 1 mL/min, temperature: 35 °C

Detection: UV, 254 nm

Injection volume: 5 µL

Peaks:

1. o-Terphenyl
2. m-Terphenyl
3. p-Terphenyl
4. Triphenylene



The separation of o-terphenyl and triphenylene is a concrete example to evaluate the selectivity potential of a reversed phase column in terms of the different shape of two molecules. The phenyl rings of o-terphenyl are twisted out of plane while triphenylene has a planar geometry.

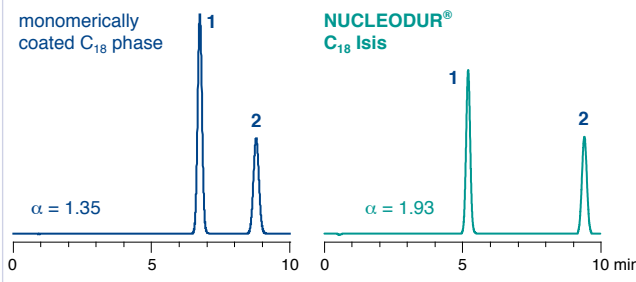
High steric selectivity

The separation factor (α -value) is a measure for the steric selectivity. As is shown in the following chromatograms the α -value is considerably larger on NUCLEODUR® C₁₈ Isis compared to a conventional C₁₈ column.

Steric selectivity of NUCLEODUR® C₁₈ Isis

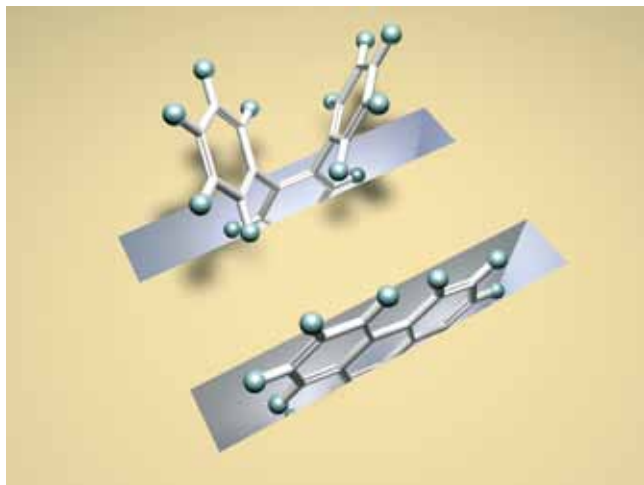
Columns: 125 x 4 mm
Eluent: methanol – water (80:20, v/v)
Flow rate: 1 mL/min, temperature: 40 °C
Detection: UV, 254 nm, injection volume: 1 µL

Peaks: 1. o-Terphenyl, 2. Triphenylene



Sander and Wise [LCGC 8 (1990) 378–390] proposed a model for the retention of aromatic compounds based on molecular shape, which is referred to as "Slot Model". This model pictures the bonded C₁₈ phase on the silica surface with slots which the analytes have to penetrate during retention. Planar molecules are able to penetrate these slots deeper than nonplanar molecules of similar molecular weight and length-to-breadth ratio. Thus triphenylene is longer retained than o-terphenyl.

Slot model



Surface deactivation

The chromatography of basic analytes requires a high density of surface-bonded C₁₈ silanes combined with a thorough endcapping procedure to keep silanol activity at a minimum. This ensures tailing-free elution of even strongly basic amino-containing compounds (see appl. 121210 on page 35).

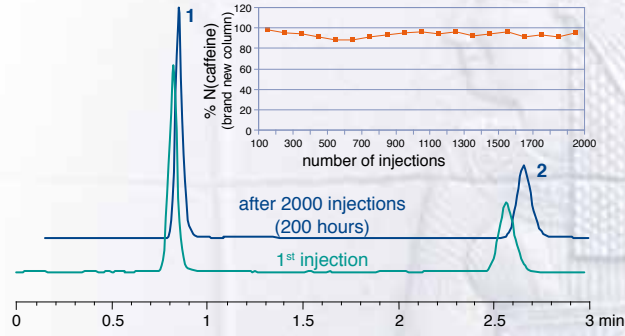
Stability

The applied special surface bonding technology also provides improved stability features for the NUCLEODUR® C₁₈ Isis phase. The proof for this was given in a long-term test in which the decrease of plate counts for caffeine at pH 10 and 50 °C has been observed over a period of 200 hours and 2000 sample injections, respectively. In addition retention and peak shape of caffeine and theophylline were compared with the chromatographic performance of the brand new column.

Stability of NUCLEODUR® C₁₈ Isis at pH 10

Column: 125 x 4 mm NUCLEODUR® C₁₈ Isis, 5 µm
Eluent: methanol – 50 mM triethylamine, pH 10 (25:80, v/v)
Flow rate: 1 mL/min, temperature: 50 °C
Detection: UV, 254 nm, injection volume: 5 µL

Peaks: 1. Theophylline, 2. Caffeine

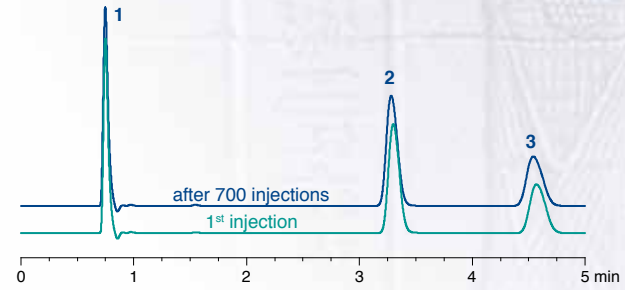


The following chromatograms exhibit the excellent stability of NUCLEODUR® C₁₈ Isis at pH 1 and 80 °C. After 700 column runs retention time and peak shape of the three test compounds remain almost unchanged.

Stability of NUCLEODUR® C₁₈ Isis at pH 1

Column: 125 x 4 mm NUCLEODUR® C₁₈ Isis, 5 µm
Eluent: acetonitrile – 1 % TFA, pH 1 (50:50, v/v)
Flow rate: 1 mL/min, temperature: 80 °C
Detection: UV, 254 nm, injection volume: 5 µL

Peaks: 1. Pyridine, 2. Toluene, 3. Ethylbenzene



- NUCLEODUR® C₁₈ Isis does not show any degradation under the applied mobile phase conditions. An enhanced pH stability in the range from pH 1 to 10 can be certified for this phase.

C₁₈ Pyramid

Key features:

- Stable in 100% aqueous mobile phase systems
- Interesting polar selectivity features
- Excellent base deactivation; suitable for LC/MS due to low bleeding characteristics

Technical characteristics:

Special phase with polar endcapping; pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm (7 and 10 µm particles for preparative purposes on request); carbon content 14%; pH stability 1-9

Recommended application:

Analgesics, penicillin antibiotics, nucleic acid bases, water-soluble vitamins, complexing agents, organic acids

USP L1

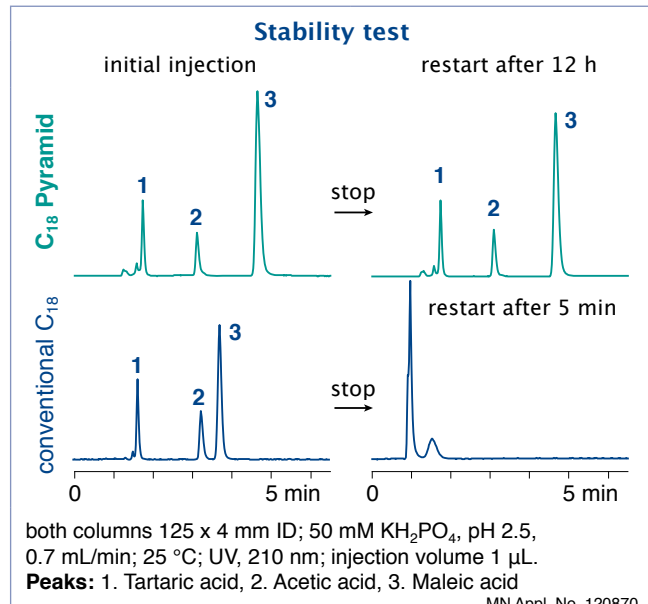
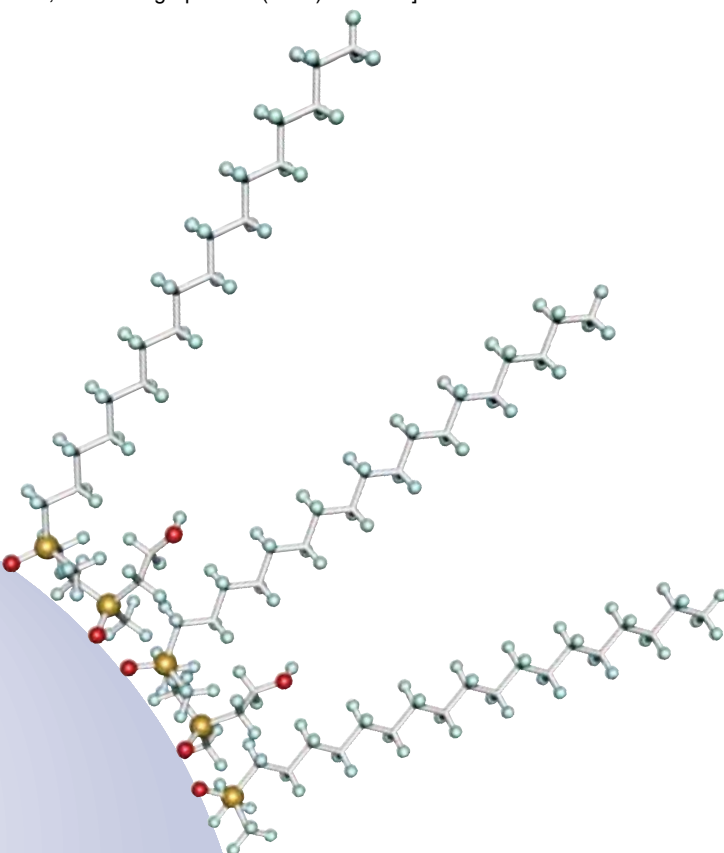
RP-HPLC with highly aqueous mobile phases

The efforts to neutralize unwanted activity of unreacted surface silanols often results in well base-deactivated phases with high carbon load, but a limited scope of selectivity beyond non-polar interactions. In particular polar compounds like carboxylic acids, drug metabolites, etc. show only weak retention on densely bonded reversed phase columns due to distinct hydrophobic properties but low polar interactions. Very polar analytes require highly aqueous mobile phases for solubility and retention. Conventional reversed phase columns often display stability problems in eluent systems with high percentage of water (> 95%) as evidenced by a sudden decrease of retention time and overall poor reproducibility. This phenomenon is described as phase collapse caused by the mobile phase expelled from the pores due to the fact, that hydrophobic RP phases are incompletely wetted with the mobile phase [U. D. Neue et al., Chromatographia 54 (2001) 169-177].

Different approaches can be used to increase column stability with highly aqueous mobile phase systems. The most promising concepts are incorporating a polar group in the hydrophobic alkyl chain, or using hydrophilic endcapping procedures to improve the wettability of the reversed phase modification. NUCLEODUR® PolarTec may be taken as an example for the embedded polar group strategy, in which a C₁₈ silane with a polar function is successfully linked to the silica surface.

Stability features

NUCLEODUR® C₁₈ Pyramid is a silica phase with hydrophilic endcapping, designed especially for use in eluent systems of up to 100% water. The figure below shows the retention behavior of tartaric, acetic and maleic acid under purely aqueous conditions on NUCLEODUR® C₁₈ Pyramid in comparison with a conventionally bonded RP phase.



It can be shown that the retention times for NUCLEODUR® C₁₈ Pyramid remain nearly unchanged between initial injection and restart after the flow has been stopped for 12 hours, whilst the performance of the conventional RP column already collapsed totally after 5 min.

for highly aqueous eluents

Retention characteristics

Based on the ultrapure NUCLEODUR® silica the polar surface derivatization exhibits retention characteristics, which differentiate the "Pyramid" from conventional C₁₈ stationary phases. The chromatogram below shows the improved retention behavior of very polar compounds such as short chain organic acids, which are insufficiently retained on RP columns with predominantly hydrophobic surface properties. For more separations on NUCLEODUR® C₁₈ Pyramid see the "Applications" section from page 32.

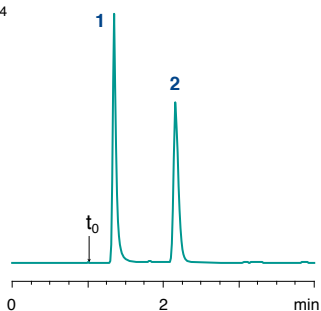
Separation of very polar compounds

Column: 125 x 4 mm NUCLEODUR® C₁₈ Pyramid,
5 µm
Eluent: 0.2% H₃PO₄
Flow rate: 1.0 mL/min
Temperature: 22 °C
Detection: UV, 202 nm
Injection volume: 2 µL

Peaks:

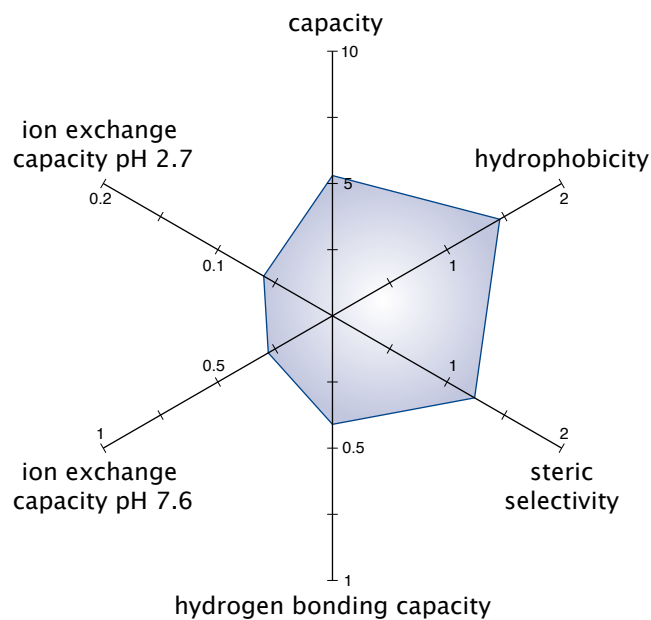
- 1. Formic acid
- 2. Acetic acid

MN Appl. No. 119170



In addition to the exceptional polar selectivity NUCLEODUR® C₁₈ Pyramid also provides adequate hydrophobic retention (see application 119190 at www.mn-net.com). The capacity factors of the non-polar, alkyl-substituted benzenes toluene and ethylbenzene do not go too far in comparison with standard C₁₈ phases.

Tanaka plot of NUCLEODUR® C₁₈ Pyramid



Base deactivation

The perceptible increase in polarity has no impact on the retention behavior of ionizable analytes. Even with the strongly basic compounds of the tricyclic antidepressant drug test mixture, no unwanted interactions or a so-called lack in base deactivation are observed (see application 119200 on page 35).

Retention behavior of polar and non-polar compounds on different NUCLEODUR® RP columns

Columns: 250 x 4 mm, 5 µm particles

Eluent: methanol – 25 mM NH₄H₂PO₄, pH 7 (65:35, v/v)

Flow rate: 0.8 mL/min

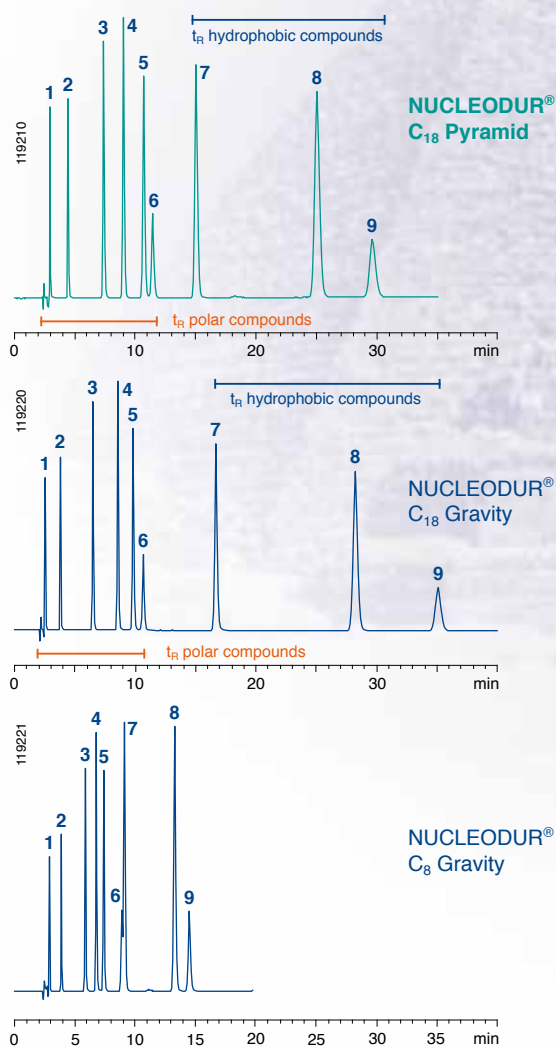
Temperature: 40 °C

Detection: UV, 254 nm

Injection volume: 5 µL

Peaks:

- | | |
|-----------------------|-----------------|
| 1. Chlorpheniramine | 6. Lidocaine |
| 2. Dimethyl phthalate | 7. Naphthalene |
| 3. Benzamide | 8. Biphenyl |
| 4. Ethyl benzoate | 9. Acenaphthene |
| 5. Benzophenone | |



Key features:

- Excellent base deactivation
- Suitable for LC/MS and stable in 100% aqueous mobile phases
- Pronounced steric selectivity

Technical characteristics:

Phase with embedded polar group; pore size 110 Å; particle sizes 3 µm and 5 µm; carbon content 17%; pH stability 1–9

Recommended application:

Exceptional selectivity for carbocyclic acids, phenols and nitrogen containing compounds, polar compounds like basic pharmaceuticals, organic acids, pesticides, amino acids, water soluble vitamins, etc.

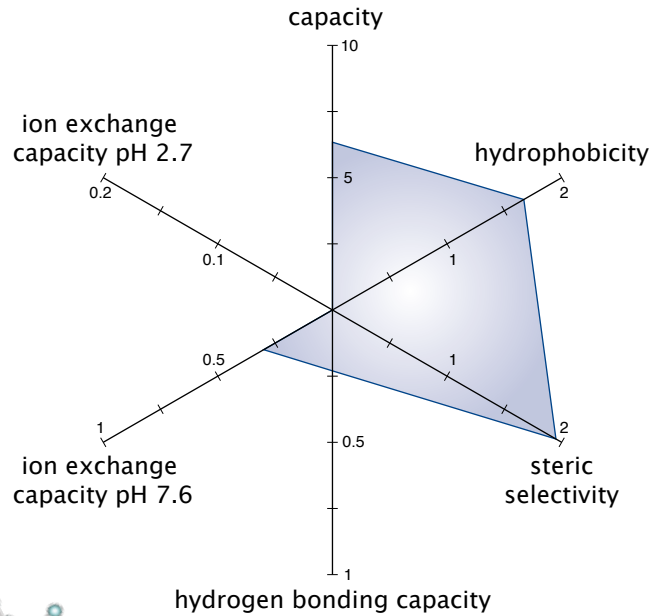
USP L1 and L60

RP-HPLC under 100% aqueous conditions

The dominant form of interactions of conventional C₁₈ phases are non-polar London dispersion forces. Besides non-polar interactions phases with embedded polar groups possess the ability to show polar interactions (dipole-dipole, hydrogen-bondings, π-π, etc.) These interactions enhance retention and selectivity for polar compounds like carbocyclic acids, phenols and nitrogen containing compounds (see applications).

In order to increase retention for polar compounds it is often necessary to decrease the organic ratio of the mobile phase to zero. Under these conditions many conventional C₁₈ phases display the so-called dewetting effect which means that the mobile phase is expelled from the pores. This phenomenon leads to a dramatic loss in retention. NUCLEODUR® PolarTec is stable in 100% aqueous mobile phases and therefore especially suited for the separation of polar compounds like organic acids (see appl. 124562 on page 55).

Tanaka plot of NUCLEODUR® PolarTec



Due to the shielding effect of the embedded group NUCLEODUR PolarTec shows an excellent base deactivation, which is at the top-notch of embedded polar group phases on the market. The pronounced steric selectivity (see Tanaka plot) is an additional tool for the separation of complex mixtures.

RP phase with embedded polar group

Bleeding of NUCLEODUR® PolarTec

Columns: 150 x 3 mm NUCLEODUR® PolarTec, 5 μ m
 150 x 3 mm Waters SymmetryShield™ RP18, 5 μ m

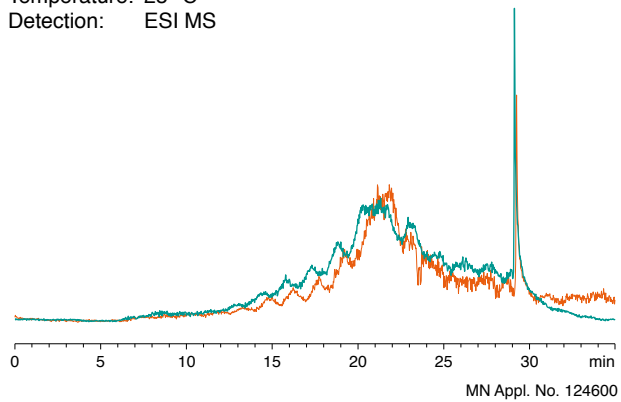
Eluent: A) acetonitrile, B) water

Gradient: 10 % B \rightarrow 90 % B in 10 min

Flow rate: 0.2 mL/min

Temperature: 25 °C

Detection: ESI MS



Due to low bleeding characteristics NUCLEODUR® PolarTec is also suitable for LC/MS.

Separation of histidine

Column: 150 x 3 mm NUCLEODUR® PolarTec, 3 μ m

Eluent: 1.0 mM perfluoropentanoic acid in water –
 0.5 mM perfluoropentanoic acid in acetonitrile
 (99.5:0.5, v/v)

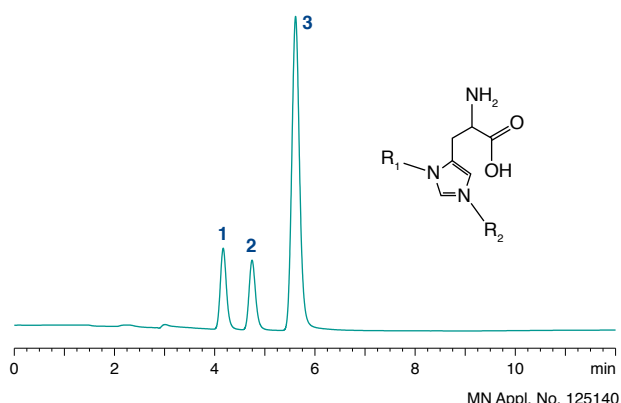
Flow rate: 0.4 mL/min

Temperature: 20 °C

Detection: UV, 230 nm

Peaks:

1. 3-Methylhistidine R₁ = H, R₂ = CH₃
 2. Histidine R₁ = R₂ = H
 3. 1-Methylhistidine R₁ = CH₃, R₂ = H



Even after days or weeks of operation in purely aqueous eluents the C₁₈ chains of NUCLEODUR® PolarTec are neither folded nor show any collapsing. A significant reduction of retention time cannot be observed.

Stability of NUCLEODUR® PolarTec

Column: 150 x 3 mm NUCLEODUR® PolarTec, 3 μ m

Eluent: 30 mM KH₂PO₄, pH 3.0

Flow rate: 0.5 mL/min

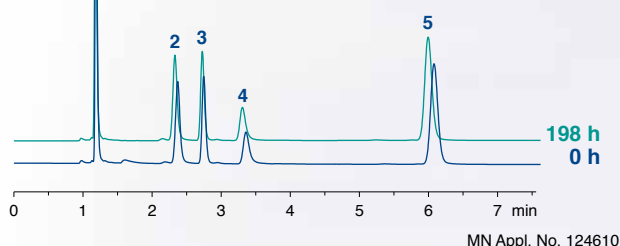
Temperature: 30 °C

Detection: UV, 220 nm

Peaks:

1. Cytosine
 2. Uracil
 3. Adenine
 4. Guanine
 5. Thymine

measurement every 14 h;
 in between flow was stopped



MN Appl. No. 124610

In spite of the polar character of the embedded functional group NUCLEODUR® exhibits sufficient hydrophobic properties and is very well suited for analyzing basic compounds. Good base deactivation of NUCLEODUR® PolarTec is illustrated by the separation of pyridine and phenol.

Phase comparison of NUCLEODUR® PolarTec

Columns: 150 x 3 mm NUCLEODUR® PolarTec, 5 μ m
 150 x 3 mm Waters SymmetryShield™ RP18, 5 μ m

Eluent: acetonitrile – water (50:50, v/v)

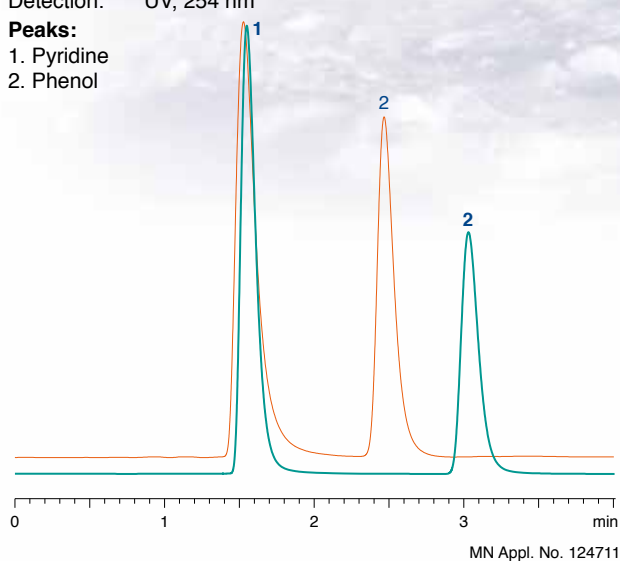
Flow rate: 0.56 mL/min

Temperature: 25 °C

Detection: UV, 254 nm

Peaks:

1. Pyridine
 2. Phenol



MN Appl. No. 124711

Key features:

- Hydrophobic phase with alternative selectivity in comparison to classical C₁₈ modifications
- Separation principle based on 4 retention mechanisms:
 - polar interactions (H bonds)
 - dipole-dipole interactions
 - π-π interactions
 - hydrophobic interactions
- Suitable for LC/MS due to low bleeding characteristics

Technical characteristics:

Phase with pentafluorophenyl-propyl modification and multi-endcapping; pore size 110 Å; particle sizes 3 µm and 5 µm; carbon content 8%; pH stability 1–9

Recommended application:

Aromatic and unsaturated compounds, phenols, halogenated compounds, isomers, polar compounds like pharmaceuticals, antibiotics; high retention of basic compounds

USP L43

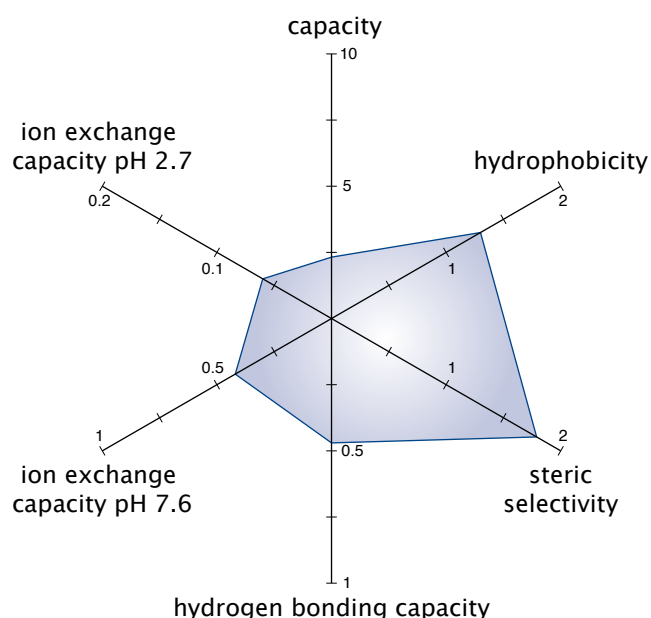
Orthogonality in selectivity

Fluorinated stationary phases in HPLC have gained increasing interest over the last years. Most common representative of fluorinated silica phases is the pentafluorophenyl modification (PFP or F5). Especially the orthogonal selectivity compared to traditional alkyl phases widens the scope in analytical HPLC. Thus NUCLEODUR® PFP offers an excellent selectivity especially for highly polar analytes like aromatic and unsaturated compounds, phenols or halogenated hydrocarbons.

Halogen substitutes in molecules result often in an increase of their polarity accompanied by a decrease of typical retention characteristics in RP-HPLC.

While a typical C₁₈ phase just provides hydrophobic interactions between stationary phase and analyte NUCLEODUR® PFP offers four different retention mechanisms: polar interactions (H bonds), dipole-dipole interactions, π-π interactions and hydrophobic interactions. Especially the pronounced ion exchange capacity and distinct steric selectivity are typical for the character of fluorinated phases.

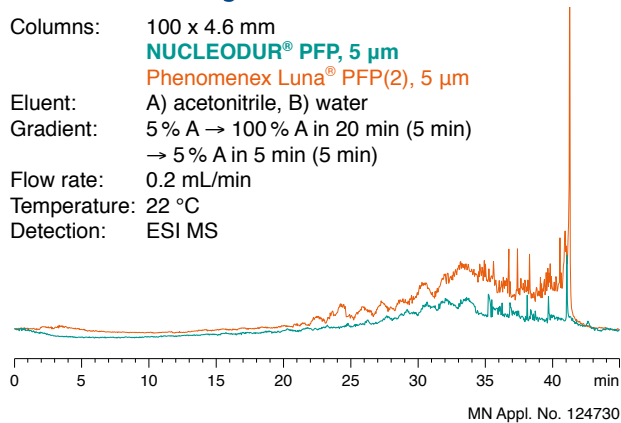
Tanaka plot of NUCLEODUR® PFP



Due to low bleeding characteristics NUCLEODUR® PFP is also suitable for LC/MS.

Bleeding of NUCLEODUR® PFP

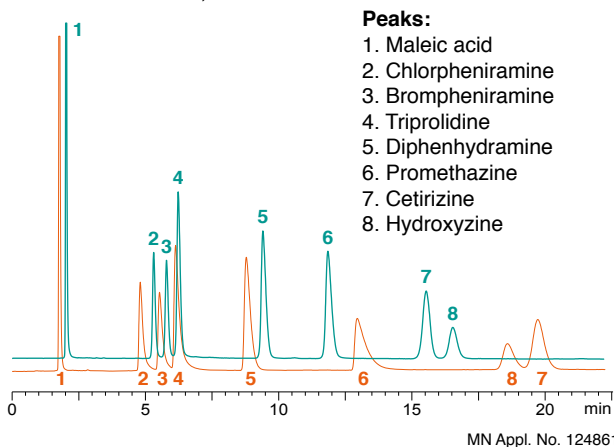
Columns: 100 x 4.6 mm
NUCLEODUR® PFP, 5 µm
Phenomenex Luna® PFP(2), 5 µm
 Eluent: A) acetonitrile, B) water
 Gradient: 5 % A → 100 % A in 20 min (5 min)
 → 5 % A in 5 min (5 min)
 Flow rate: 0.2 mL/min
 Temperature: 22 °C
 Detection: ESI MS



Hydrophobic pentafluorophenyl phase

Separation of antihistamines

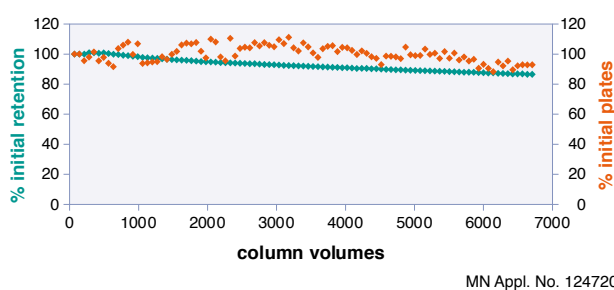
Columns: **250 x 3 mm NUCLEODUR® PFP, 5 µm**
250 x 3 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: acetonitrile – 20 mM KH₂PO₄ (30:70, v/v)
 Flow rate: 0.563 mL/min
 Temperature: 30 °C
 Detection: UV, 210 nm



Based on a special surface modification procedure NUCLEODUR® PFP offers highest stability also at low pH values.

Stability of NUCLEODUR® PFP

Column: **125 x 4 mm NUCLEODUR® PFP, 5 µm**
 Eluent: acetonitrile – water, 0.1 % TFA, pH 1 (50:50, v/v)
 Flow rate: 1 mL/min
 Temperature: 80 °C
 Detection: UV, 254 nm
 Injection volume: 2 µL
 Sample: ethylbenzene



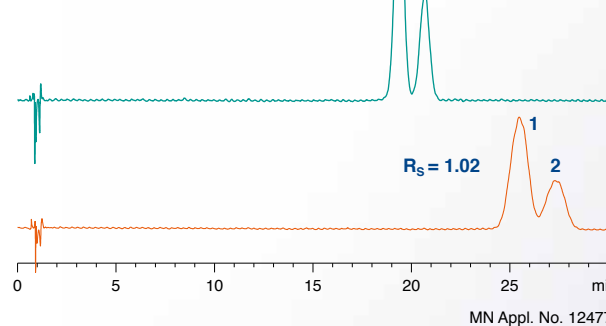
Compared to fluorinated HPLC columns of other manufacturers NUCLEODUR® PFP shows excellent separation capabilities and shorter retention times e.g., for critical isomers like beta- and dexamethasone.

Separation of beta- and dexamethasone

Columns: **100 x 4.6 mm NUCLEODUR® PFP, 5 µm**
100 x 4.6 mm Phenomenex Luna® PFP(2), 5 µm
 Eluent: acetonitrile – water (20:80, v/v)
 Flow rate: 1.3 mL/min
 Temperature: 30 °C
 Detection: UV, 260 nm

Peaks:

1. Betamethasone
2. Dexamethasone



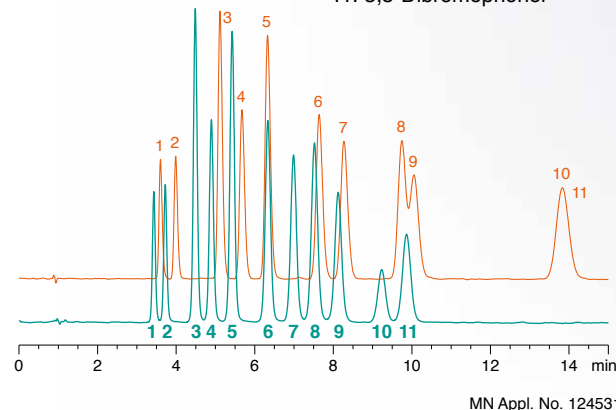
NUCLEODUR® PFP offers a completely different retention behavior compared to alkyl modified silica and is often used successfully for separations which provide just insufficient results on traditional C₁₈ phases. More and more applications in the areas of (bio-) pharma, natural compounds and environment are published and show the broad application field for fluorinated phases.

Separation of phenol isomers

Columns: **125 x 4 mm NUCLEODUR® PFP, 5 µm**
125 x 4 mm NUCLEODUR® C₁₈ HTec, 5 µm
 Eluent: acetonitrile, 0.1 % formic acid – water, 0.1 % formic acid (35:65, v/v)
 Flow rate: 1 mL/min
 Temperature: 35 °C
 Detection: UV, 280 nm

Peaks:

- | | |
|-----------------------|-----------------------|
| 1. o-Cresol | 6. 2,6-Dichlorophenol |
| 2. m-Cresol | 7. 2,3-Dichlorophenol |
| 3. 3,4-Dimethylphenol | 8. 2,4-Dichlorophenol |
| 4. 3,5-Dimethylphenol | 9. 3,4-Dichlorophenol |
| 5. 2,5-Dimethylphenol | 10. 2,4-Dibromophenol |
| | 11. 3,5-Dibromophenol |



Sphinx RP

Key features:

- Distinct selectivity based on well-balanced bifunctional surface coverage
- Widens the scope for method development based on additional π - π interactions
- Suitable for LC/MS due to low bleeding characteristics

Technical characteristics:

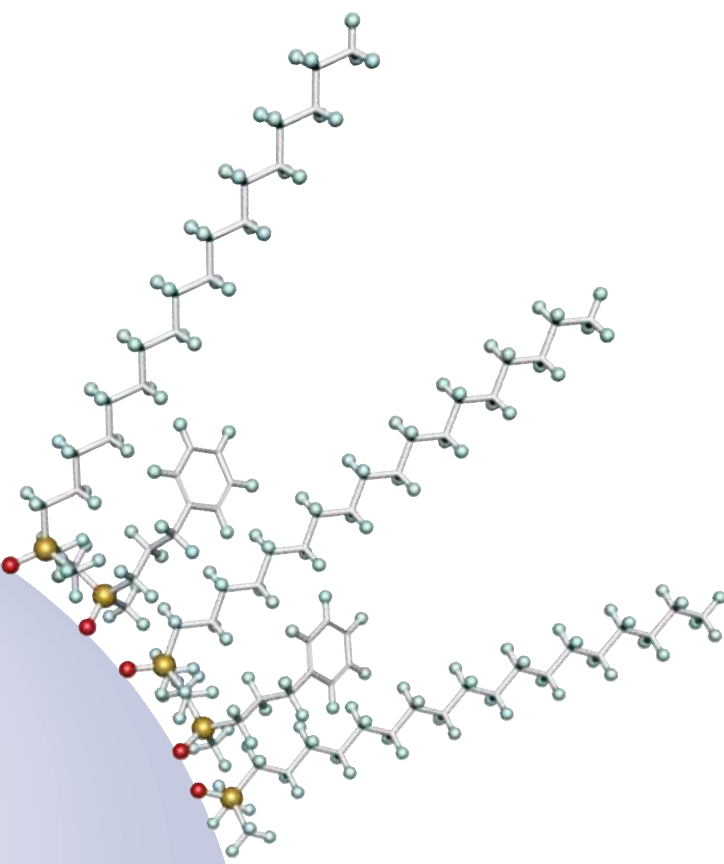
Octadecyl and propylphenyl modified silica; pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm; carbon content 15%; pH stability 1-10; high reproducibility and consistent quality

Recommended application:

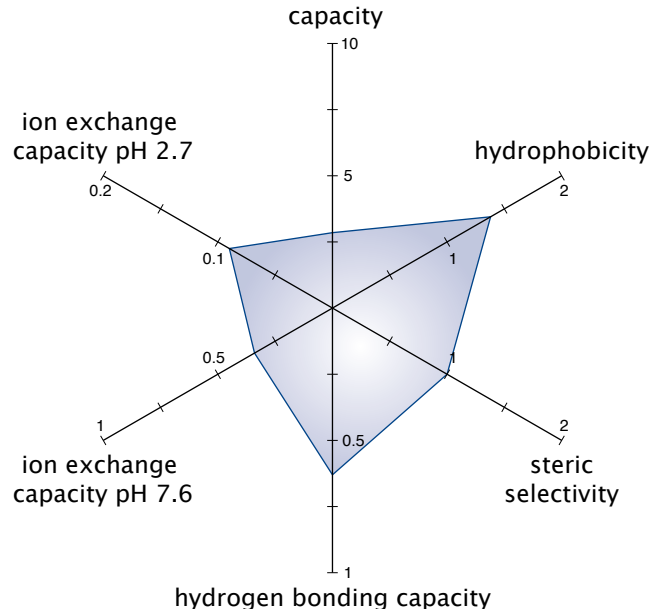
Quinolone antibiotics, sulfonamides, xanthines, substituted aromatics
USP L1 and L11

Alternative RP selectivity

NUCLEODUR® Sphinx RP is characterized by exceptional selectivity features generated by a **well-balanced ratio of covalently bonded octadecyl and phenyl groups**. The combination of classical hydrophobic with π - π interactions (aromatic ring system) expands the scope of selectivity in comparison with conventional reversed phase packings. NUCLEODUR® Sphinx RP is particularly suited for the separation of molecules containing aromatic and multiple bonds. For the separation of polar compounds NUCLEODUR® Sphinx RP can be especially recommended and can also outperform many customary C₁₈ phases. In addition, exhaustive endcapping steps minimize unwanted surface silanol activity and guarantee excellent peak shapes even for strong basic analytes.



Tanaka plot of NUCLEODUR® Sphinx RP

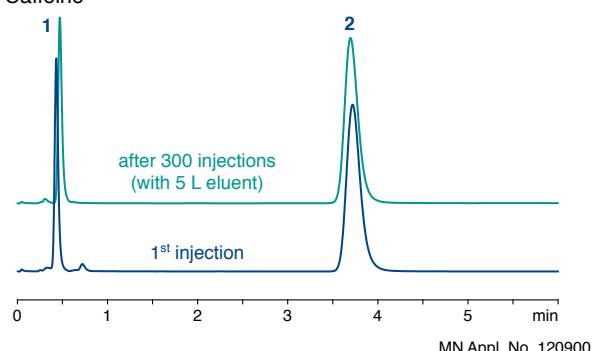


Stability of NUCLEODUR® Sphinx RP at pH 10

Column: 50 x 4.6 mm NUCLEODUR® Sphinx RP, 5 µm
Eluent: methanol – dil. NH₃, pH 10 (20:80, v/v)
Flow rate: 1.0 mL/min
Temperature: 30 °C
Detection: UV, 275 nm
Injection volume: 3 µL

Peaks:

1. Theophylline
2. Caffeine



MN Appl. No. 120900

Bifunctional RP phase

Different from standard phenyl phases, NUCLEODUR® Sphinx RP is far more stable towards hydrolysis and is also suggested for LC/MS applications.

Due to the additional intermolecular interactions NUCLEODUR® Sphinx RP is an interesting replenishment to the high density bonded phases NUCLEODUR® C₈ / C₁₈ Gravity and the polar endcapped NUCLEODUR® C₁₈ Pyramid.

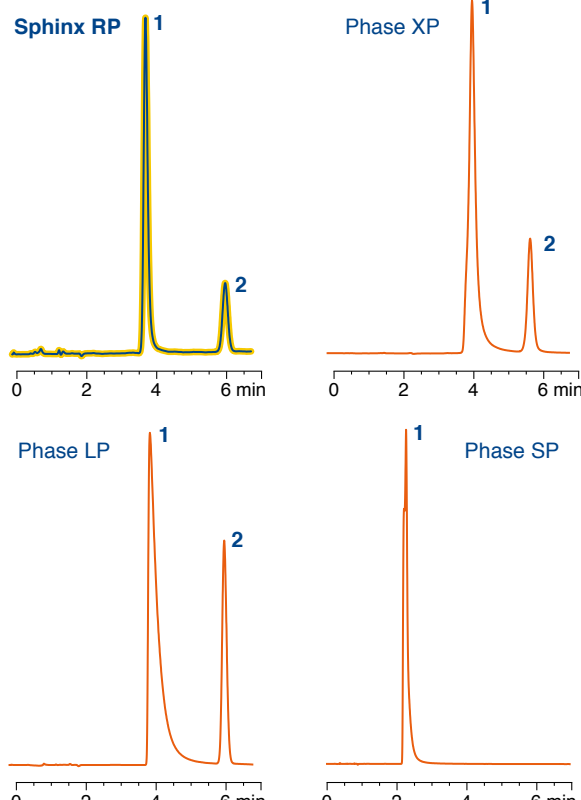
Comparison of surface deactivation of different phenyl modified RP phases

Columns: 150 x 4.6 mm
 NUCLEODUR® Sphinx RP, 5 µm
 Competitor 1 (column XP)
 Competitor 2 (column LP)
 Competitor 3 (column SP)

Eluent: methanol – water (30:70, v/v)
 Flow rate: 1 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 2 µL

Peaks:

1. Pyridine
2. Phenol



The selectivity advantage of NUCLEODUR® Sphinx RP is impressively shown in the flavonoid application below.

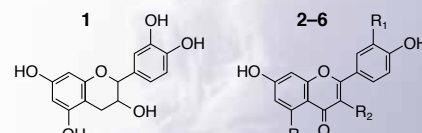
While a baseline separation of kaempferol and isorhamnetin can be achieved on NUCLEODUR® Sphinx RP, the two compounds are not or just poorly separated on NUCLEODUR® C₈ Gravity or C₁₈ Gravity. The additional π-π interactions of the aromatic ring systems provide the necessary difference in retention to outperform the classical C₁₈ and C₈ phases.

Separation of flavonoids on 3 different NUCLEODUR® phases

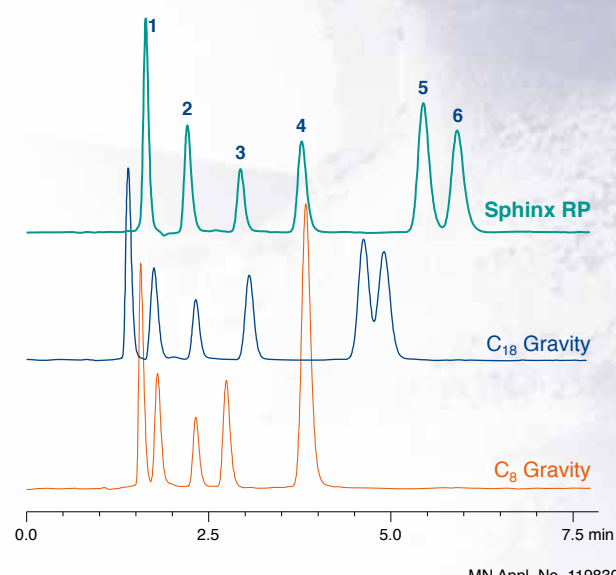
Columns: 150 x 4.6 mm
NUCLEODUR® Sphinx RP, 5 µm
 NUCLEODUR® C₁₈ Gravity, 5 µm
NUCLEODUR® C₈ Gravity, 5 µm

Eluent: water – methanol (40:60, v/v)
 Flow rate: 1 mL/min
 Temperature: 30 °C
 Detection: UV, 270 nm
 Injection volume: 3 µL

Peaks:



1. Catechin
2. Rutin R₁ = R₃ = OH, R₂ = O-rutinose
3. Fisetin R₁ = R₂ = OH, R₃ = H
4. Quercetin R₁ = R₂ = R₃ = OH
5. Kaempferol R₁ = H, R₂ = R₃ = OH
6. Isorhamnetin R₁ = OCH₃, R₂ = R₃ = OH



Key features:

- Reliable and durable standard RP phase for up-scaling to preparative scale, suited for LC/MS
- High loadability and excellent stability
- Outstanding base deactivation

Technical characteristics:

High density octadecyl modification (C₁₈); pore size 110 Å; particle sizes 1.8 µm, 3 µm, 5 µm, 7 µm and 10 µm for analytical and preparative separations; carbon content 18%; pH stability 1-11

Recommended application:

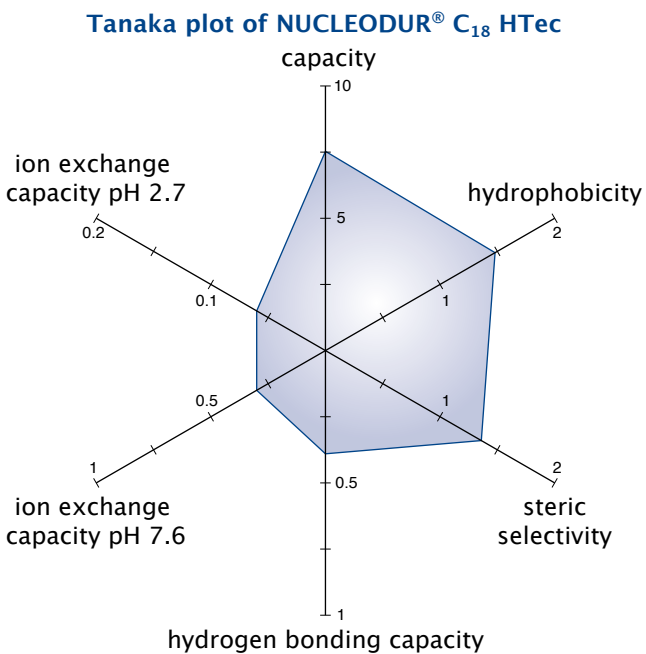
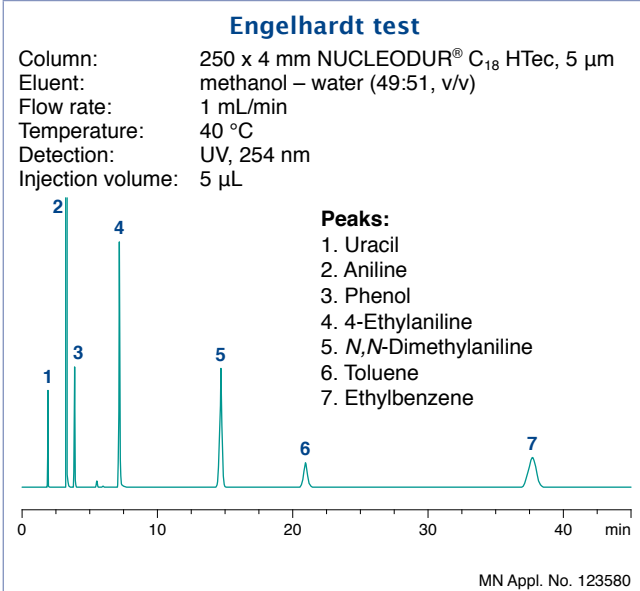
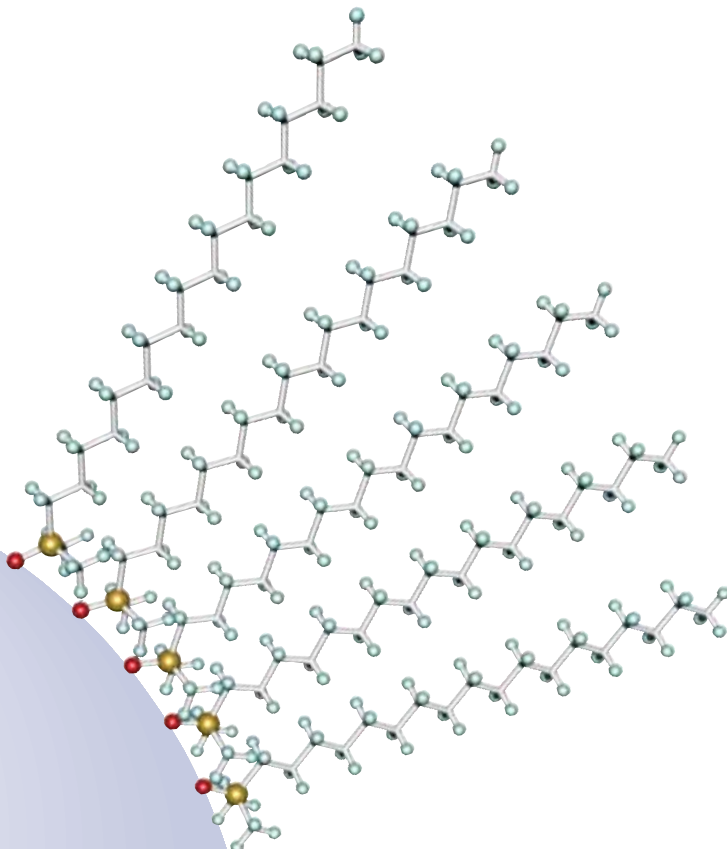
Sophisticated analytical and preparative separations of basic, neutral and acidic pharmaceuticals, derivatized amino acids, pesticides, fat-soluble vitamins, aldehydes, ketones and phenolic compounds

USP L1

Preparative separations place high demands on silica based HPLC materials. Apart from excellent selectivity and base deactivation, robustness (pH, pressure stability, ...) and capacity are vital criteria for optimal and efficient separation at the preparative scale.

Selectivity and base deactivation

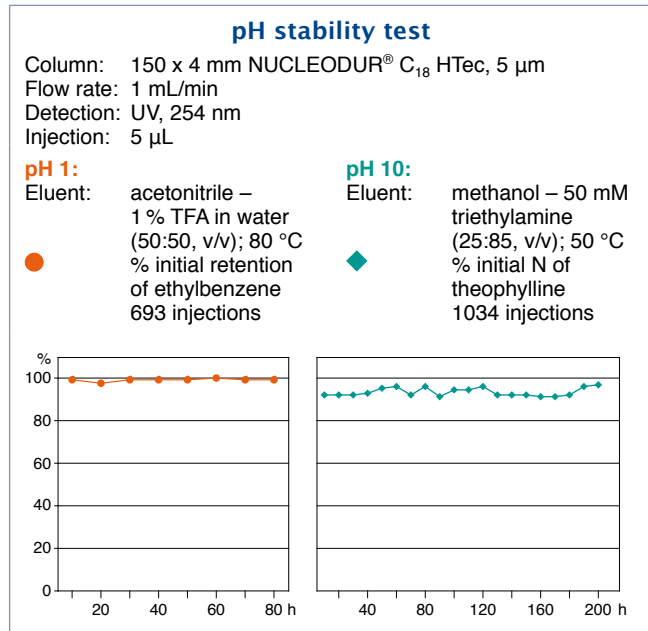
The innovative and special endcapping procedure leads to exceptionally good base deactivation – the Engelhardt test demonstrates superb selectivity, peak symmetry and peak shape over the entire polarity range. In addition NUCLEODUR® C₁₈ HTec scores in low bleed characteristics and is therefore highly suitable for LC/MS.



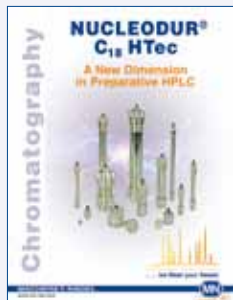
Base-deactivated preparative phase

Stability and lifetime

Based on fully synthetic and extremely robust totally spherical NUCLEODUR® silica, NUCLEODUR® C₁₈ HTec offers outstanding mechanical rigidity and is thus the perfect choice also for self-packing of prep-columns. The special surface modification and endcapping procedure result in high chemical stability even at extreme chromatographic conditions like high flow rates, temperature or critical solvents (DMSO). Furthermore, NUCLEODUR® C₁₈ HTec columns show a remarkably long lifetime in acidic (pH 1) as well as basic (pH 10) mobile phases.



Due to innovative surface coating procedures NUCLEODUR® C₁₈ HTec offers excellent analytical separation properties and is the first choice for up-scaling to preparative column dimensions.



Please ask for our special NUCLEODUR® C₁₈ HTec brochure for preparative HPLC separation.

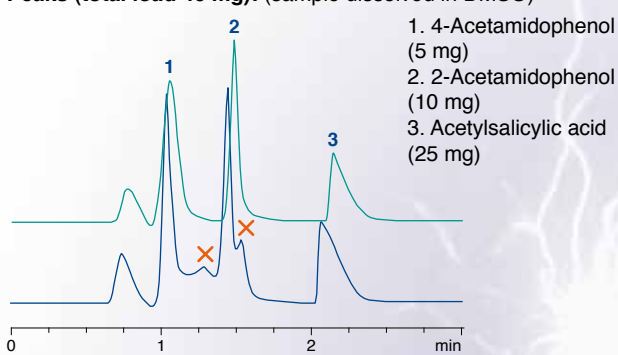
Capacity

A vital criterion for efficiency in preparative HPLC is the capacity of the separation medium. NUCLEODUR® C₁₈ HTec is characterized by a notably high loadability under both basic and acidic conditions, while competitor columns show overload effects even at lower loads (x).

Loadability under acidic conditions

Columns: VP 100 x 21 mm NUCLEODUR® C₁₈ HTec, 5 µm
100 x 21.2 mm AXIA™ Gemini® 5 µm C₁₈ 110 Å
Eluent: acetonitrile – formic acid in H₂O pH 3.0 (30:70, v/v)
Flow rate: 28 mL/min; temperature 22 °C; pressure 124 bar
Detection: UV, 254 nm

Peaks (total load 40 mg): (sample dissolved in DMSO)



Up-scaling

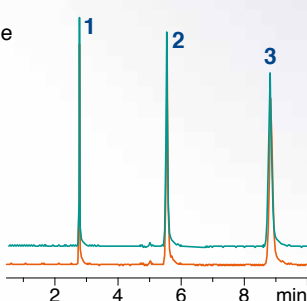
Due to highest quality standards in our silica production and phase chemistry combined with optimized packing technology, NUCLEODUR® C₁₈ HTec delivers exceptional transferability from analytical to preparative scale. This doesn't just apply to the use of different particle sizes (e.g., 5, 7 or 10 µm) but also for diverse column dimensions (e.g., ID 4.6 to 21 mm).

Up-scaling with NUCLEODUR® C₁₈ HTec

Columns: EC 250 x 4.6 mm NUCLEODUR® C₁₈ HTec, 5 µm
VP 250 x 21 mm NUCLEODUR® C₁₈ HTec, 5 µm
Eluent: acetonitrile – water (80:20, v/v)
Flow rates: 1.3 mL/min, 27 mL/min
Temperature: 22 °C
Pressure: 84 bar, 109 bar
Detection: UV, 254 nm
Inj. volume: 3 µL, 60 µL

Peaks: (1 mg/mL of each compound)

1. Phenol
2. Naphthalene
3. Anthracene



MN Appl. No. 123780

C₁₈ ec / C₈ ec

Key features:

- Ideal and reliable standard RP phase for daily routine analysis and up-scaling for preparative HPLC
- Medium density octadecyl (C₁₈) and octyl (C₈) modification with exhaustive endcapping
- Wide range of application areas

Technical characteristics:

Pore size 110 Å; particle sizes 3 µm and 5 µm; 7 µm, 10 µm, 12 µm, 16 µm, 20 µm, 30 µm and 50 µm for preparative separations; carbon content 17.5 % for C₁₈, 10.5 % for C₈; pH stability 1–9, high reproducibility from lot to lot

Recommended application:

Basic, neutral or acidic drugs
derivatized amino acids
pesticides
fat-soluble vitamins
aldehydes and ketones
phenolic compounds
USP L1 (C₁₈) / L7 (C₈)

NUCLEODUR® C₁₈ ec for daily routine analysis and up-scaling for preparative HPLC

The efficiency of a separation is controlled by particle size and selectivity of the stationary phase. The exceptional surface coverage of monomeric bonded alkylsilanes, combined with an exhaustive endcapping, results in a surface with lowest silanol activity. This allows the tailing-free elution of polar compounds such as basic drugs. NUCLEODUR® C₁₈ ec is available in 9 different particle sizes (3, 5, 7, 10, 12, 16, 20, 30 and 50 µm) which cover the whole range from high speed analytical HPLC up to medium and low pressure prep LC. NUCLEODUR® C₁₈ ec is also an ideal tool for scale-up purposes.

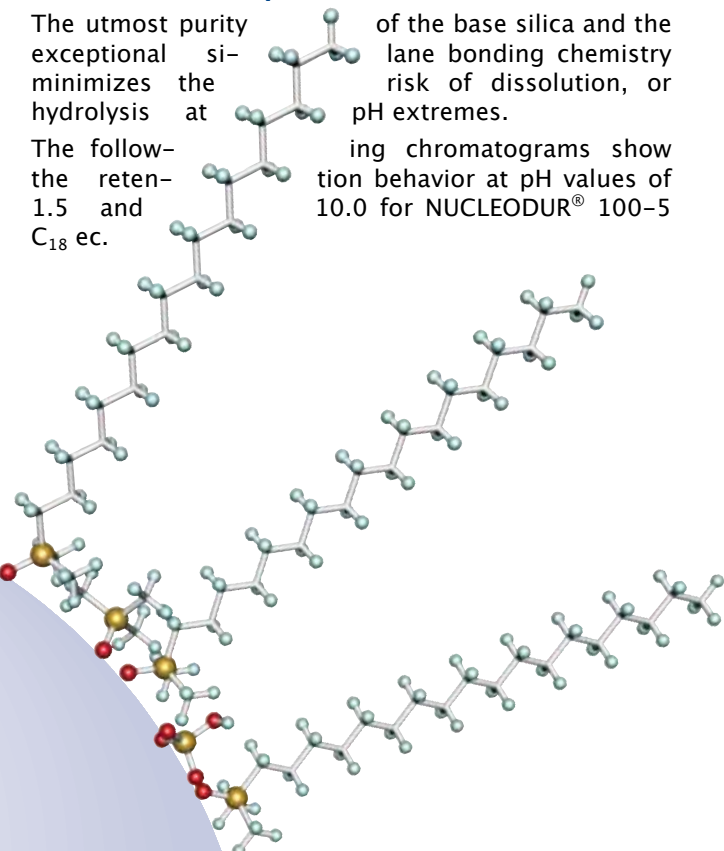
Chemical stability

The utmost purity and exceptional silanol passivation minimizes the risk of hydrolysis at

of the base silica and the lane bonding chemistry risk of dissolution, or pH extremes.

ing chromatograms show tion behavior at pH values of 10.0 for NUCLEODUR® 100-5

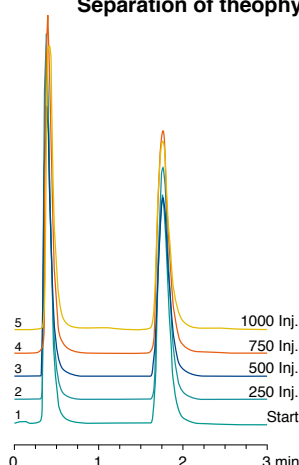
The follow-
the reten-
1.5 and
C₁₈ ec.



pH stability of NUCLEODUR® C₁₈ ec

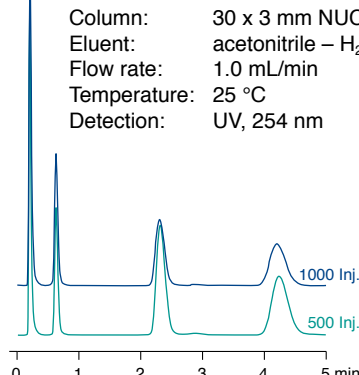
Separation of theophylline and caffeine at pH 10

Column: 30 x 3 mm
NUCLEODUR® 100-5 C₁₈ ec
Eluent:
methanol – aq. NH₃
(20:80, v/v), pH 10
Flow rate: 0.5 mL/min
Temperature: 25 °C
Detection: UV, 254 nm



Separation of uracil, veratrol, toluene and ethylbenzene at pH 1.5

Column: 30 x 3 mm NUCLEODUR® 100-5 C₁₈ ec
Eluent: acetonitrile – H₂O (65:35, v/v), TFA, pH 1.5
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm

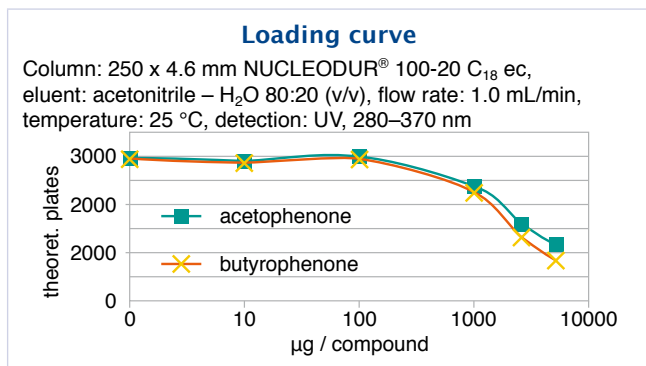


Loadability

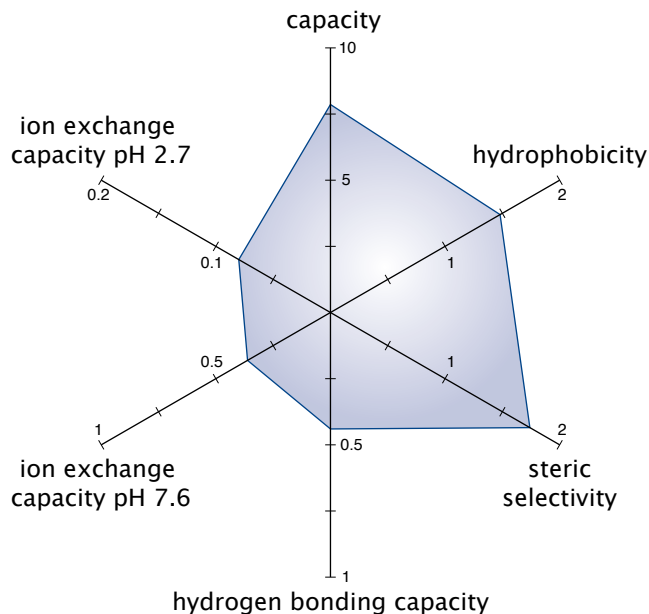
Loadability, probably the most important feature for preparative LC applications, is determined by pore size, pore volume and surface area of the packing. However, it can also be influenced by the molecular weight of the analytes. In the figure below the mass loading curve for acetophenone and butyrophenone on a NUCLEODUR®

Nonpolar phases for routine analyses

100–20 C₁₈ ec column describes the correlation between the increase of column loading and the decrease of separation efficiency.



Tanaka plot of NUCLEODUR® C₁₈ ec



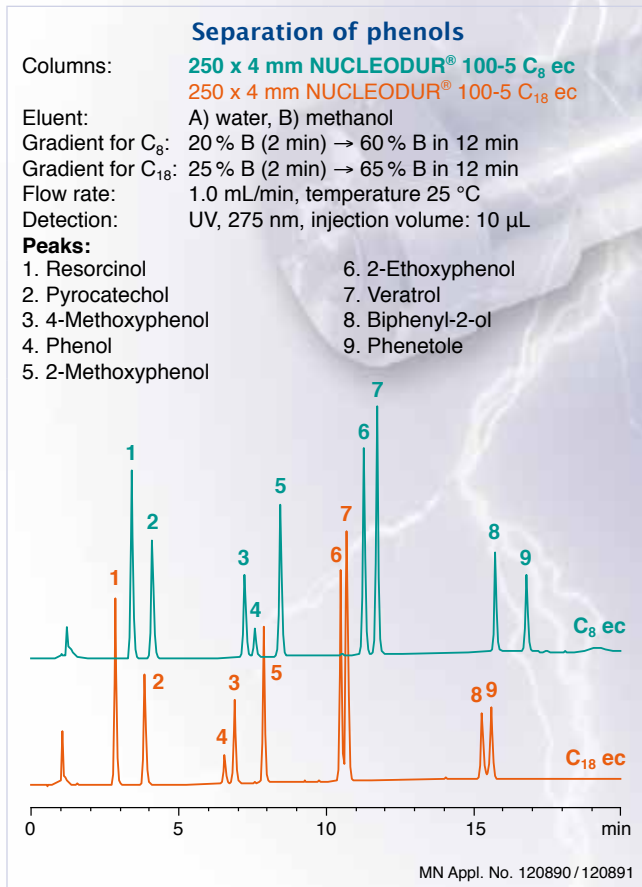
NUCLEODUR® octyl phases

In addition to the program of NUCLEODUR® C₁₈ phases MACHEREY-NAGEL offers the corresponding octyl modified NUCLEODUR® C₈ Gravity and NUCLEODUR® C₈ ec columns to expand the reversed phase tool box effectively. Based on the same totally spherical and highly pure silica the C₈ phases exhibit the same excellent chemical and mechanical stability features as the C₁₈ counterparts. Indeed NUCLEODUR® C₈ Gravity can also be run at pH extremes (pH 1–11) by choosing appropriate elution parameters. Due to the shorter chain and less hydrophobic properties of the stationary phase the retention of non-polar compounds is decreased, and in consequence a reduction in time of analysis can be achieved. Moreover a stronger polar selectivity, particularly with the separation of ionizable analytes is frequently observed (as distinct from the C₁₈ phases).

NUCLEODUR® C₈ ec and NUCLEODUR® C₈ Gravity are most suitable for the development of new methods but also for robust routine analysis.

C₁₈ or C₈ – the best of both worlds

Chromatographers now might wonder about the differences between C₈ and C₁₈ phases and the preferred range of application. Indeed there are no general guidelines which could make the choice easier but it will always be beneficial to add both phases to the existing pool of reversed phase columns in the laboratory. However, comparative studies reveal some different selectivity patterns of NUCLEODUR® C₈ ec and NUCLEODUR® C₁₈ ec. The separation of phenols below shows baseline separation for 2-ethoxyphenol and dimethoxybenzene (veratrol) and in addition a reversal of the elution order of phenol and 4-methoxyphenol can be shown on the octyl phase.



Some general principles are:

- High density C₈ and C₁₈ phases allow tailing-free elution, also for very polar compounds
- Octyl phases (C₈) show superior polar selectivity
- Octadecyl phases (C₁₈) show superior hydrophobic selectivity
- Hydrophobic compounds show shorter retention times on C₈ phases

Key features:

- Ideal for reproducible and stable chromatography of highly polar analytes
- Suitable for analytical and preparative applications as well as LC/MS
- Very short column conditioning period

Technical characteristics:

Ammonium – sulfonic acid modified silica; pore size 110 Å; particle sizes 1.8, 3 and 5 µm; carbon content 7%; pH stability 2–8.5

Recommended application:

Hydrophilic compounds such as organic polar acids and bases, polar natural compounds, nucleosides, oligonucleotides, amino acids, peptides, water soluble vitamins

NUCLEODUR® HILIC

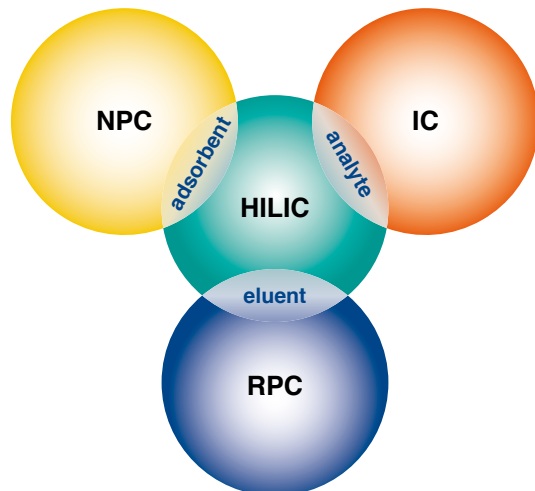
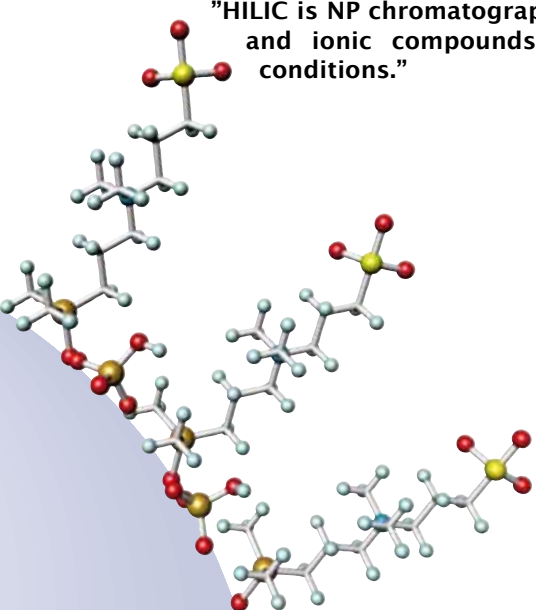
Separation science is always looking for new and effective strategies to accomplish the tasks of modern analytics. Especially for polar compounds reversed phase HPLC – the most common analytical method – is often limited. Here, hydrophilic stationary phases provide an additional tool for the separation of polar analytes in HPLC.

The expression **HILIC** (Hydrophilic Interaction Liquid Chromatography) was firstly published by Andrew Alpert in 1990 – since then it took quite some efforts to develop robust and reproducible hydrophilic HPLC phases for HILIC chromatography [A. Alpert, J. Chromatography 499 (1990), 177–196].

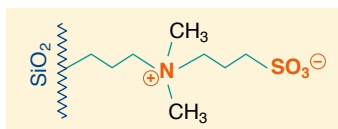
HILIC combines the characteristics of the 3 major methods in liquid chromatography – reversed phase (RPC), normal phase (NPC) and ion chromatography (IC):

- stationary phases (adsorbents) are mostly polar modifications of silica or polymers (SiOH, NH₂, Diol, (zwitter) ions, ...) – like in NPC
- mobile phases (eluents) are mixtures of aqueous buffer systems and organic modifier like acetonitrile or methanol – like in RPC
- fields of application include quite polar compounds as well as organic and inorganic ions – like in IC

"HILIC is NP chromatography of polar and ionic compounds under RP conditions."



NUCLEODUR® HILIC is a special zwitterionic modified stationary phase based on ultra spherical NUCLEODUR® particles. The betaine character of the ammonium-sulfonic acid ligands results in total charge equalization and in an overall neutrally charged but highly polar surface.



Retention characteristic

Commonly HILIC is described as partition chromatography or liquid/liquid extraction system between the mobile and stationary phase. Versus a water-poor mobile phase a water-rich layer on the surface of the polar stationary phase is formed. Thus, a distribution of the analytes between these two layers will occur.

Furthermore HILIC includes weak electrostatic mechanisms as well as hydrogen donor interactions between neutral polar molecules under high organic elution conditions. This distinguishes HILIC from ion exchange chromatography – main principle for HILIC separation is based on compound's polarity and degree of solvation. More polar compounds will have stronger interaction with the stationary aqueous layer than less polar compounds – resulting in a stronger retention. Nonpolar

Zwitterionic phase

compounds exhibit faster elution profiles due to minor hydrophobic interactions. Thus, as shown for the separation of uracil and naphthalene the elution order is quite often inverse on HILIC columns compared to RP columns.

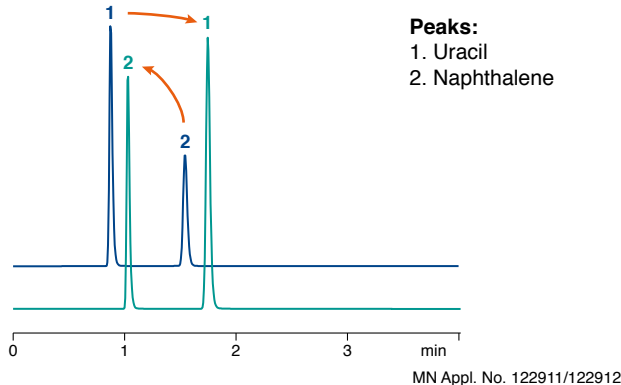
Separation of uracil and naphthalene

Columns: **125 x 4 mm NUCLEODUR® C₁₈ Pyramid, 3 µm**
125 x 4 mm NUCLEODUR® HILIC, 3 µm

Eluent: acetonitrile – water (90:10, v/v)

Flow rate: 1.0 mL/min, temperature 25 °C

Detection: UV, 254 nm



In comparison with medium polar aminopropyl phases or modification with less balanced charge equalization NUCLEODUR® HILIC shows a superb separation and peak shape for critical compounds like adenosine phosphates.

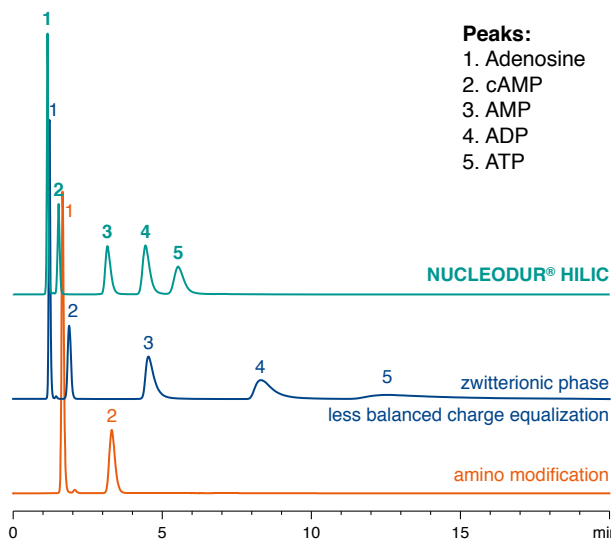
Separation of adenosine and phosphates

Columns: **125 x 4 mm NUCLEODUR® HILIC, 5 µm**
125 x 4 mm zwitterionic phase with quat. ammonium – sulfonic acid ratio 1:0.9
125 x 4 mm amino-modified silica

Eluent: acetonitrile – 100 mM ammonium acetate, pH 5.3 (70:30, v/v)

Flow rate: 1.3 mL/min, temperature 25 °C

Detection: UV, 259 nm



Stability features

Due to an advanced and unique surface modification procedure (pat. pend.) NUCLEODUR® HILIC columns provide short equilibration times – after just 20 min equilibration already the 2nd injection shows stable and reproducible results. Beyond this, NUCLEODUR® HILIC columns are characterized by an outstanding column life time – even after nearly 800 runs the columns show no loss of pristine performance – peak shape and retention are still immaculate.

Stability and equilibration

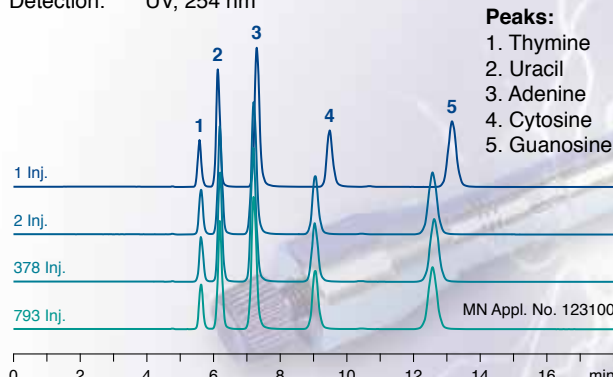
Column: **250 x 4 mm NUCLEODUR® HILIC, 5 µm**

Eluent: acetonitrile – 5 mM ammonium acetate (80:20, v/v)

Flow rate: 0.6 mL/min

Temperature: 25 °C

Detection: UV, 254 nm



Separation of growth regulators

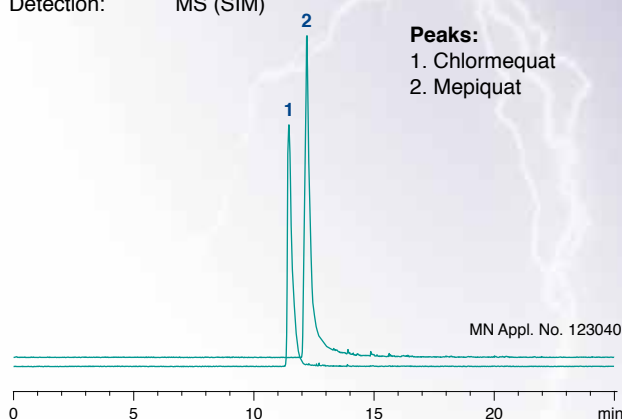
Column: **125 x 2 mm NUCLEODUR® HILIC, 3 µm**

Eluent: acetonitrile – 25 mM ammonium acetate, pH 6.82 (80:20, v/v)

Flow rate: 0.3 mL/min, temperature 25 °C

Detection: MS (SIM)

Peaks:
 1. Chlormequat
 2. Mepiquat



Due to its high loadability NUCLEODUR® HILIC is absolutely suitable for preparative and semi-preparative applications.

Overall NUCLEODUR® HILIC provides excellent chromatographic features and is hereby the perfect choice for separation of polar or charged compounds.

Key features:

- High retention capacity especially for very polar and unsaturated compounds
- Multi-mode column (RP and NP) widens scope of selectivity
- Stable against hydrolysis at low pH (working range pH 1–8)

Technical characteristics:

Cyanopropyl-modified high purity silica; pore size 110 Å; particle sizes 3 µm and 5 µm; carbon content 7%; special endcapping; high reproducibility from lot to lot; different retention characteristics in comparison to C₈ and C₁₈

Recommended application:

Tricyclic antidepressants,
steroids,
organic acids
USP L10

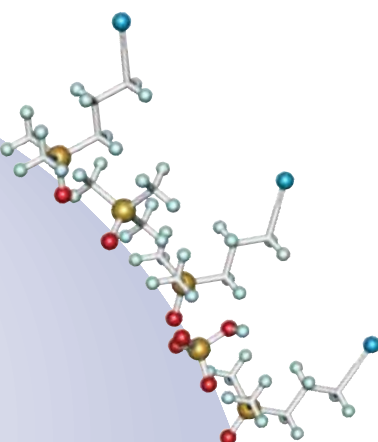
Alternative bonded-phase functionality

In reversed phase HPLC it is fairly common to start with C₁₈ or C₈ columns, if new methods have to be developed. However, superior polarity and selectivity properties often required for more sophisticated separations, are not always sufficiently provided by classical RP phases, which are usually characterized by a hydrophobic layer of monomeric or polymeric bonded alkylsilanes.

One approach to improve the resolution of compounds poorly separated on nonpolar stationary phases, is to change bonded-phase functionality.

The fully endcapped and highly reproducible (see figure top right) NUCLEODUR® CN-RP phase has cyanopropyl groups on the surface able to generate a clearly recognizable different retention behavior compared to purely alkyl-functionalized surface modifications (see figure down right).

The polarity of the NUCLEODUR® CN-RP phase can be classified as intermediate based on multiple retention mechanisms such as dipole-dipole, π-π, and also hydrophobic interactions [C. S. Young and R. J. Weigand, LCGC 20 (2002) 464–473]. Therefore, this phase shows a distinct selectivity for polar organic compounds as well as for molecules containing π-electron systems (e.g., analytes with double bonds, tricyclic antidepressants) [V. R. Meyer, Practical High Performance Liquid Chromatography (John Wiley & Sons, New York, 3rd. ed., 1999)].

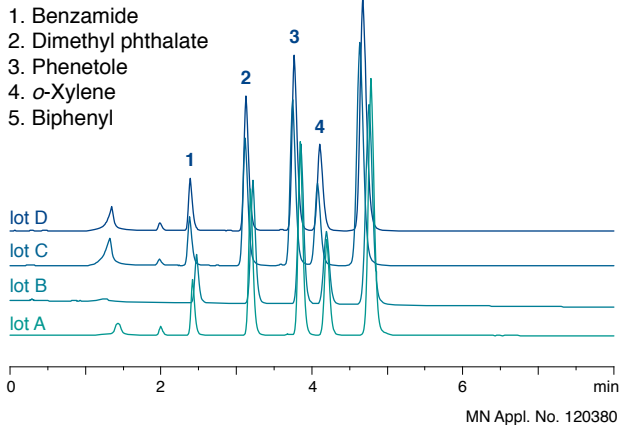


Reproducibility of NUCLEODUR® CN-RP

Column: 250 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: acetonitrile – water (60:40, v/v)
Flow rate: 1.0 mL/min, temperature 20 °C
Detection: UV, 254 nm, injection volume: 5 µL

Peaks:

1. Benzamide
2. Dimethyl phthalate
3. Phenetole
4. o-Xylene
5. Biphenyl

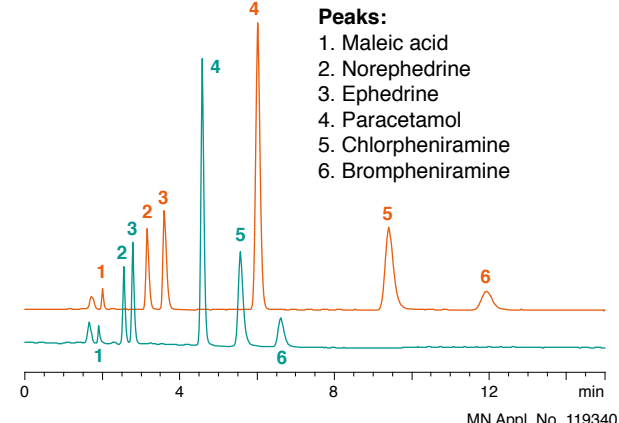


Separation of cold medicine ingredients on two different NUCLEODUR® phases

Columns: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
250 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: acetonitrile – 100 mM sodium citrate, pH 2.5 (15:85, v/v)
Flow rate: 1.0 mL/min, temperature 25 °C
Detection: UV, 270 nm, injection volume: 10 µL

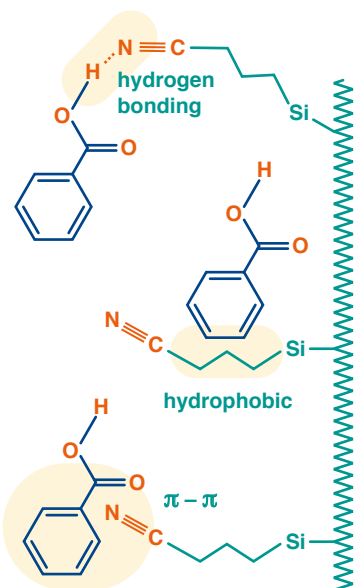
Peaks:

1. Maleic acid
2. Norephedrine
3. Ephedrine
4. Paracetamol
5. Chlorpheniramine
6. Brompheniramine



Cyano-modified high purity silica

Interactions on cyano-modified silica



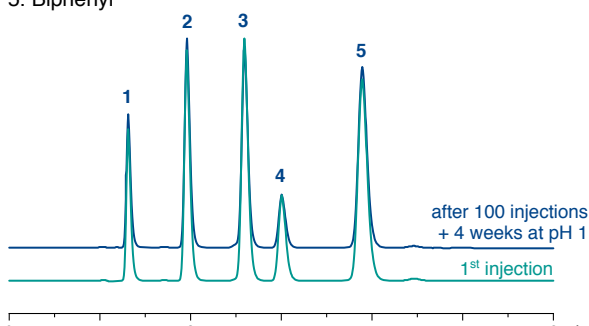
Short-chain bonded phases are sometimes suspected of revealing shortcomings in stability towards hydrolysis at low pH [J. J. Kirkland, LCGC 14 (1996) 486–500]. The following chromatograms show that even after 100 sample injections and four weeks storage at pH 1 (blue curve), neither a considerable shift in retention, nor a visible change in peak symmetry could be noticed (green curve = new column).

Stability of NUCLEODUR® CN-RP at pH 1

Column: 125 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: acetonitrile – water, 2% TFA, pH 1 (50:50, v/v)
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm
Injection volume: 5 µL

Peaks:

1. Benzamide
2. Dimethyl phthalate
3. Phenetole
4. *o*-Xylene
5. Biphenyl



MN Appl. No. 119350

Due to the exceptional polarity features the cyano phase can also be run in the normal phase mode. NUCLEODUR® CN columns for normal phase applications are shipped in *n*-heptane. The drastic change in selectivity and order of elution for a mixture of various steroids in normal and reversed phase mode is displayed in following figure. Moreover the high coverage combined with a thorough endcapping makes NUCLEODUR® 100-5 CN-RP suitable for the separation of ionizable compounds such as basic drugs.

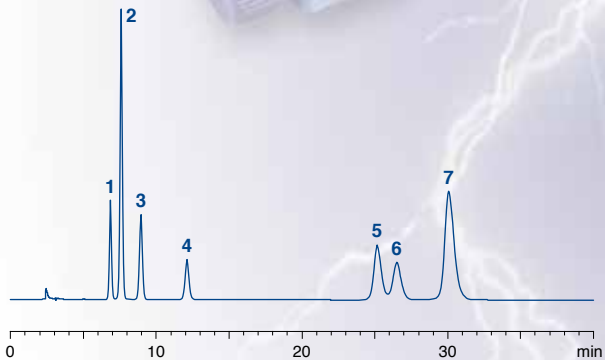
Separation of steroids in normal phase and reversed phase mode

Normal phase mode

Column: 250 x 4 mm NUCLEODUR® 100-5 CN
Eluent: *n*-heptane – 2-propanol (90:10, v/v)
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm
Injection volume: 10 µL

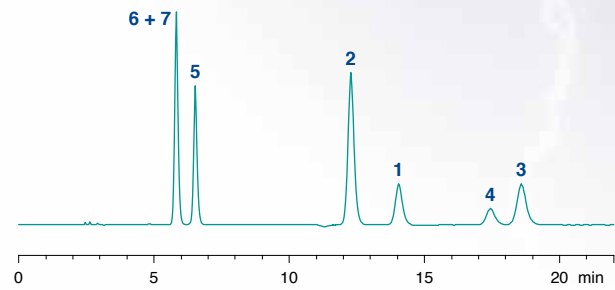
Peaks:

1. Methyltestosterone
2. Testosterone
3. Norgestrel
4. Medrysone
5. Cortisone
6. Hydrocortisone
7. Prednisolone



Reversed phase mode

Column: 250 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: acetonitrile – water (25:75, v/v)
other conditions as for normal phase mode



MN Appl. Nos. 119271 / 119272

● **Key features:**

- Multi-mode columns (for RP, NP and IC)
- Stable against hydrolysis at low pH (working range pH 2–8), 100% stable in water; suitable for LC/MS
- Widens scope of analytical HPLC into the polar range

● **Technical characteristics:**

Aminopropyl-modified high purity silica; pore size 110 Å; particle sizes 3, 5 and 7 µm; carbon content 2.5%; not endcapped

● **Recommended application:**

Polar compounds under RP conditions (sugars, DNA bases), hydrocarbons under NP conditions
USP L8

- **Normal phase chromatography (NP)** with hexane, dichloromethane or 2-propanol as mobile phase for polar compounds such as substituted anilines, esters, chlorinated pesticides
- **Reversed phase chromatography (RP)** of polar compounds in aqueous-organic eluent systems
- **Ion exchange chromatography** of anions and organic acids using conventional buffers and organic modifiers

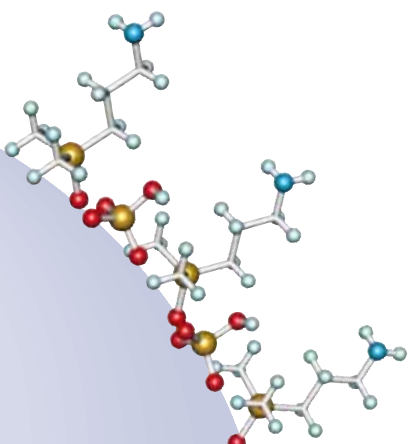
Some compounds, especially polar substances, cannot be sufficiently resolved on C₁₈ phases. Polar-modified silica phases offer alternative selectivities such expanding the spectrum of analytical HPLC into the polar range.

Multi-mode columns

Besides cyano modifications, amino modifications belong to the most frequently used polar silica phases – both feature the important advantage, that they can be run in the RP mode using aqueous-organic eluent mixtures as well as in the NP mode e.g., with hexane as mobile phase. NUCLEODUR® NH₂, too, belongs to the so-called multi-mode columns.

It can be used for reversed phase chromatography (RP) of polar compounds such as sugars in aqueous-organic eluent systems, for normal phase chromatography (NP) of substituted aromatics or chlorinated pesticides with organic mobile phases such as hexane, dichloromethane or 2-propanol, but also for ion exchange chromatography of anions and organic acids using conventional buffers and organic modifiers.

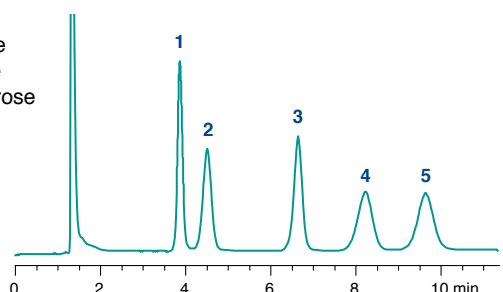
Main field of application of NUCLEODUR® NH₂ is the separation of simple and complex sugars, sugar alcohols and other hydroxy compounds under RP conditions as well as hydrocarbons under NP conditions.



Reversed phase separation of sugars

Column: 250 x 4 mm NUCLEODUR® 100-5 NH₂-RP
Eluent: acetonitrile – water (79:21, v/v)
Flow rate: 2 mL/min
Detection: RI

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

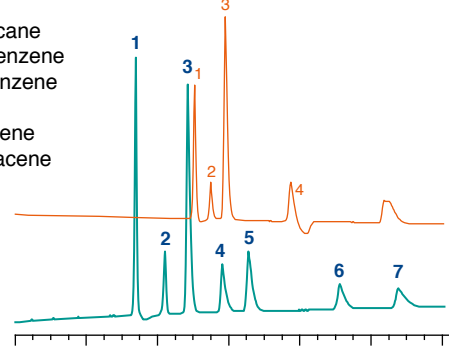


MN Appl. No. 122160

Normal phase separation of middle distillates in accordance with DIN EN 12916

Columns: 250 x 4 mm NUCLEODUR® 100-5 NH₂
conventional aminopropyl phase
Eluent: heptane
Flow rate: 1 mL/min
Detection: RI

Peaks:
1. Cyclohexane
2. 1-Phenyldodecane
3. 1,2-Dimethylbenzene
4. Hexamethylbenzene
5. Naphthalene
6. Dibenzothiophene
7. 9-Methylanthracene



MN Appl. No. 122180

SiOH — Unmodified high purity silica

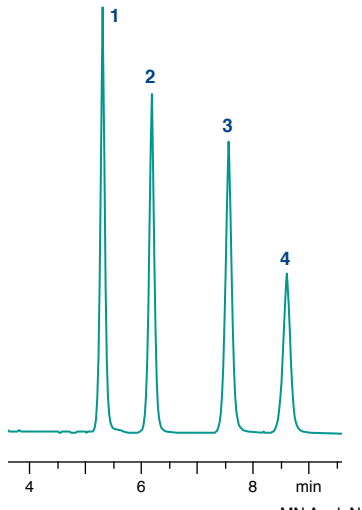
Even at lower flow rates than for C₁₈ phases, NUCLEODUR® NH₂ achieves good separations of polar compounds such as DNA bases – this reduces the back pressure as well as the solvent consumption. Even very polar compounds like streptomycin are retained sufficiently for quantitative and qualitative analysis.

Separation of DNA bases

Column: 250 x 4 mm NUCLEODUR® 100-5 NH₂-RP
Eluent: acetonitrile – water (80:20, v/v)
Flow rate: 0.6 mL/min
Temperature: 35 °C
Pressure: 30 bar
Detection: UV, 254 nm

Peaks:

1. Thymine
2. Uracil
3. Cytosine
4. Adenine



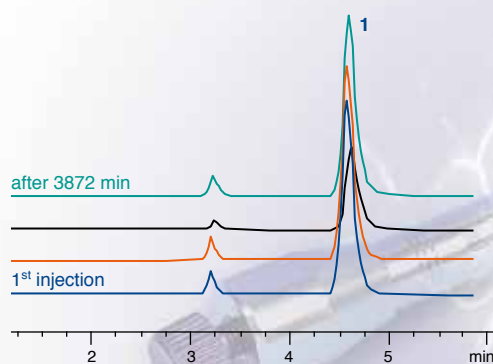
The example below proves the enhanced pH stability of the NUCLEODUR® amino phase and also the outstanding suitability of this column for the separation of total herbicides (AMPA, glyphosate, glufonilate, ...) – you may find the complete application 122190 in the "Applications" section on page 60.

Resistance towards hydrolysis for NUCLEODUR® NH₂-RP

Column: 250 x 4 mm NUCLEODUR® 100-5 NH₂-RP
Eluent: acetonitrile – 50 mmol KH₂PO₄, pH 1.75 (50:50, v/v)
Flow rate: 0.6 mL/min
Detection: UV, 254 nm

Peaks:

1. Aminomethyl-phosphonic acid (AMPA)



One of the main problems with conventional amino phases is insufficient resistance towards hydrolysis. Due to a special method of surface modification NUCLEODUR® NH₂ features a pronounced stability at higher as well as at lower pH values. The figure at right shows, that even after several days of exposure of the column material at pH 1.75 good separation efficiency and peak symmetry are maintained. The resulting high column life allows cost reduction due to lower column consumption.

Based on the superspherical silica NUCLEODUR® this amino phase – like all other members of the NUCLEODUR® family – features a very good pressure stability, which makes it the perfect choice for preparative separations as well as for LC-MS applications. Additionally, the high batch-to-batch reproducibility of NUCLEODUR® NH₂ offers the advantage of reliable analyses especially for routine work.

SiOH

Key features:

- Totally spherical high purity silica
- Pressure stable up to 600 bar
- Suitable for analytical and preparative separation of polar and midpolar compounds

Technical characteristics:

Unmodified high purity silica; pore size 110 Å; particle sizes 3 to 50 µm; pore volume 0.9 mL/g, surface area (BET) 340 m²/g; pH stability 2–8; metal content < 10 ppm (see table on page 1)

Recommended application:
Polar and mid-polar compounds under normal phase conditions
USP L3

Applications

NUCLEODUR®

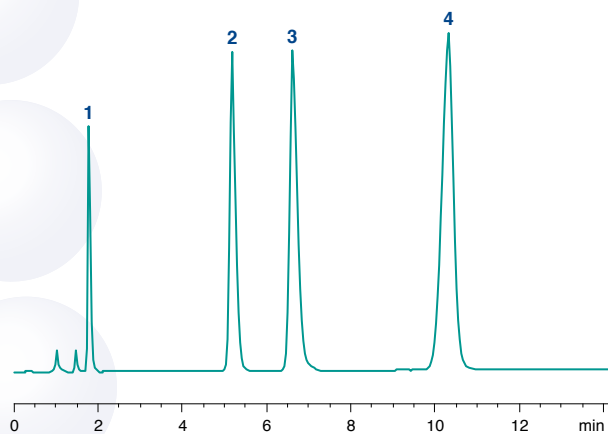
Anesthetics

MN Appl. No. 119410

Column: 125 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: methanol – 20 mM KH₂PO₄, pH 6.95 (65:35, v/v)
 Flow rate: 1 mL/min
 Temperature: 30 °C
 Detection: UV, 254 nm
 Injection volume: 13 µL

Peaks:

1. Benzocaine
2. Lidocaine
3. Tetracaine
4. Butacaine



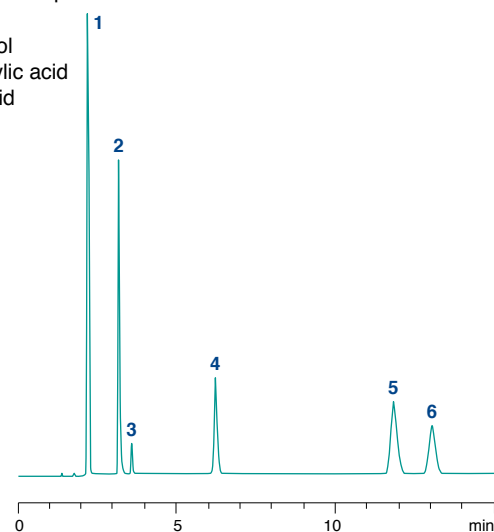
Analgesics

MN Appl. No. 118600

Column: 125 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
 Eluent: methanol – 0.1 % phosphoric acid (40:60, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 240 nm
 Injection volume: 13 µL

Peaks:

1. Paracetamol
2. Caffeine
3. 2-Acetamidophen
4. Acetanilide
5. Acetylsalicylic acid
6. Phenacetin



For a fast separation (< 1 min) of acetylsalicylic acid and salicylic acid on NUCLEODUR® 100-5 C₁₈ ec see application 117780 at www.mn-net.com.

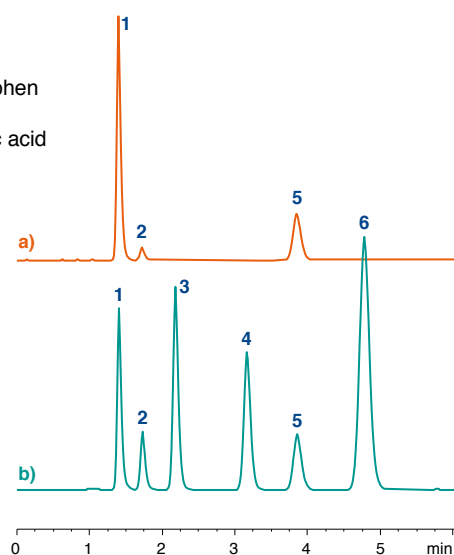
Analgesics

MN Appl. No. 117770

Column: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile – 20 mM KH₂PO₄, pH 2.5 (50:50, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 230 nm
 Injection volume: 5 µL

Peaks:

1. Paracetamol
2. Acetylsalicylic acid
3. Salicylic acid
4. Ketoprofen
5. Diclofenac
6. Ibuprofen



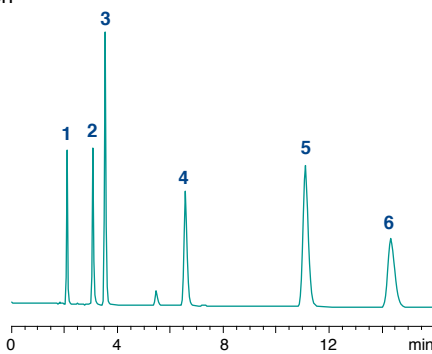
Analgesics

MN Appl. No. 119160

Column: 250 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: acetonitrile – 0.1 % TFA (50:50, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

1. Paracetamol
2. Acetylsalicylic acid
3. Methyl-4-hydroxybenzoate
4. Ketoprofen
5. Flurbiprofen
6. Ibuprofen



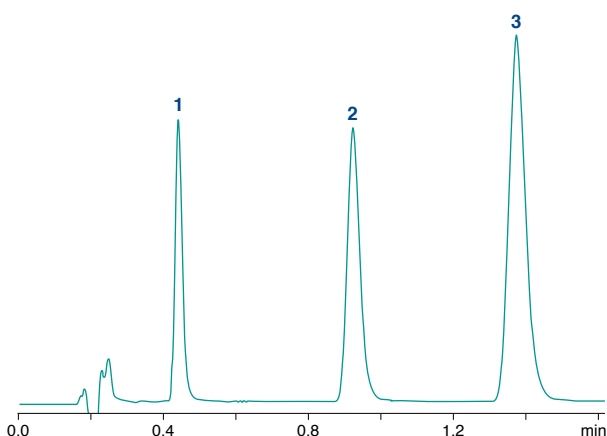
Anti-inflammatory drugs

MN Appl. No. 122130

Column: 50 x 3 mm NUCLEODUR® C₁₈ Pyramid, 1.8 µm
 Eluent: phosphate buffer, pH 2.5 – acetonitrile – methanol (425:475:100, v/v/v)
 Flow rate: 1.0 mL/min
 Temperature: 50 °C
 Detection: UV, 240 nm
 Injection volume: 2 µL

Peaks:

1. Chlorocresol
2. Clobetasol 17-propionate
3. Beclometasone dipropionate



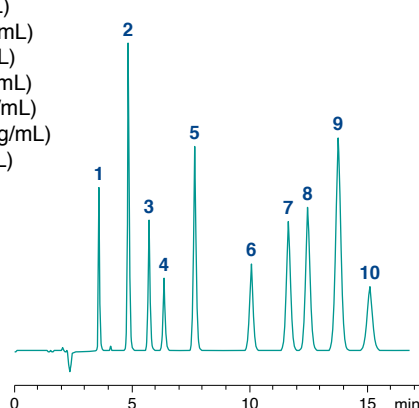
Analgesic and anti-inflammatory drugs

MN Appl. No. 118590

Column: 250 x 4 mm NUCLEODUR® 100-5 C₈ ec
 Eluent: acetonitrile – 1 % acetic acid (48:52, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 230 nm
 Injection volume: 10 µL

Peaks:

1. Acetylsalicylic acid (1.6 µg/mL)
2. Tolmetin (26 µg/mL)
3. Piroxicam (26 µg/mL)
4. Suprofen (26 µg/mL)
5. Naproxen (0.64 µg/mL)
6. Diflunisal (1.6 µg/mL)
7. Fenoprofen (26 µg/mL)
8. Flurbiprofen (26 µg/mL)
9. Indomethacin (52 µg/mL)
10. Ibuprofen (52 µg/mL)



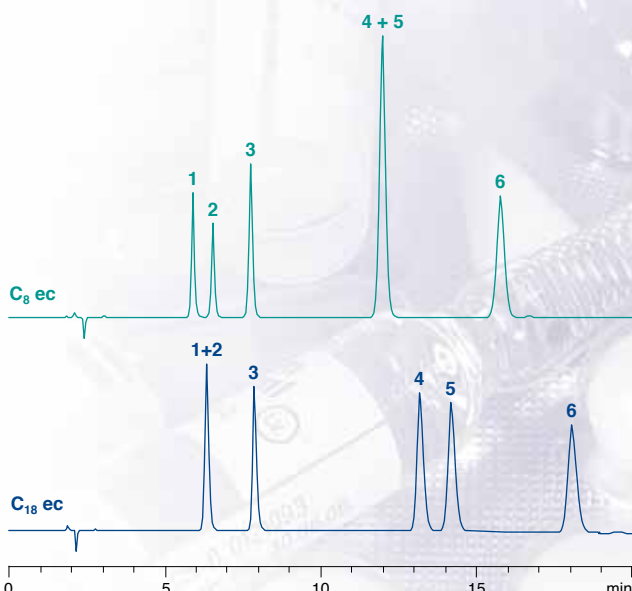
Anti-inflammatory drugs

MN Appl. No. 120880/120881

Columns: 50 x 4 mm NUCLEODUR® 100-5 C₈ ec
 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile – water, 1 % acetic acid (48:52, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 230 nm
 Injection volume: 10 µL

Peaks:

1. Piroxicam
2. Suprofen
3. Ketoprofen
4. Carprofen
5. Fenoprofen
6. Diclofenac



This separation of various non-steroidal anti-inflammatory drugs illustrates the differences in polarity between C₈ and C₁₈ and the resulting impact on efficiency. NUCLEODUR® C₈ ec exhibits enhanced selectivity and excellent resolution for the polar compounds piroxicam and suprofen which co-elute on the C₁₈ column. However due to the longer alkyl chain NUCLEODUR® C₁₈ ec shows a distinct hydrophobic selectivity that leads to baseline separation of the more non-polar analytes carprofen and fenoprofen with superior peak shapes.

Applications

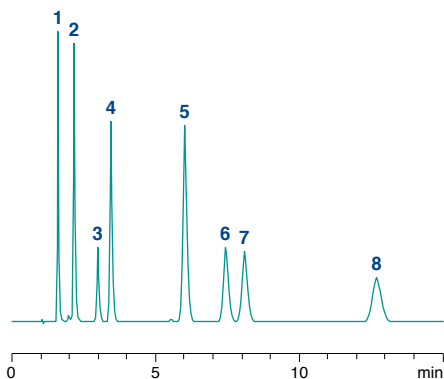
Anti-inflammatory drugs

MN Appl. No. 117830

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile – 20 mM KH₂PO₄, pH 2.5 (45:55, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 22 °C
 Detection: UV, 230 nm
 Injection volume: 5 µL

Peaks:

1. Acetylsalicylic acid
2. Sulindac
3. Tolmetin
4. Ketoprofen
5. Flurbiprofen
6. Diclofenac
7. Ibuprofen
8. Meclofenamic acid

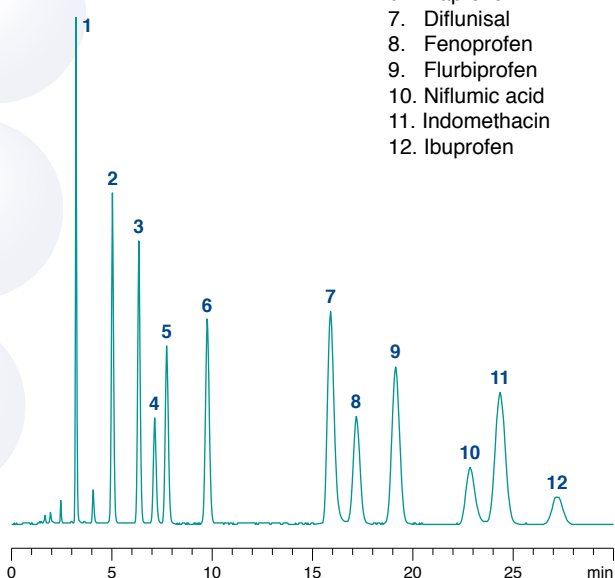


MN App. No. 122550

Column: 250 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Flow rate: 1.3 mL/min
 other conditions as above

Peaks:

1. Acetylsalicylic acid
2. Sulindac
3. Piroxicam
4. Suprofen
5. Tolmetin
6. Naproxen
7. Diflunisal
8. Fenoprofen
9. Flurbiprofen
10. Niflumic acid
11. Indometacin
12. Ibuprofen



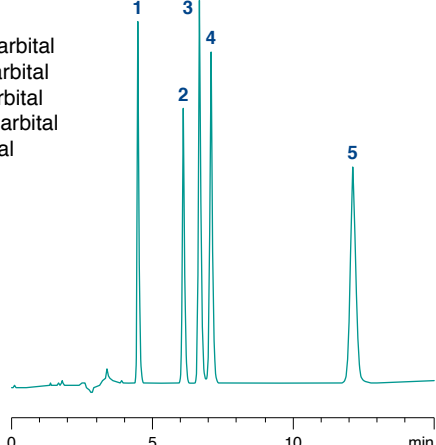
Barbiturates

MN Appl. No. 117820

Column: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile – water (50:50, v/v)
 Flow rate: 0.7 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

1. Phenobarbital
2. Pentobarbital
3. Hexobarbital
4. Mephobarbital
5. Thiameyal



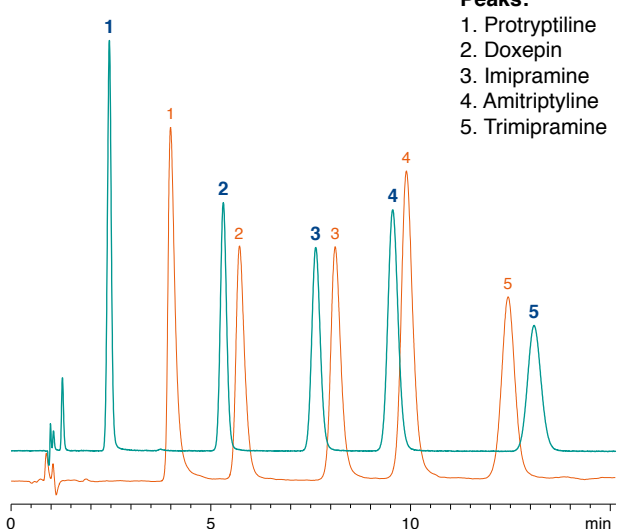
Tricyclic antidepressants

MN Appl. No. 124622

Columns: 150 x 3 mm NUCLEODUR® PolarTec, 5 µm
 150 x 3 mm Waters SymmetryShield™ RP18, 5 µm
 Eluent: methanol – 25 mM KH₂PO₄, pH 7 (70:30, v/v)
 Flow rate: 0.66 mL/min
 Temperature: 30 °C
 Detection: UV, 254 nm
 Injection volume: 1 µL

Peaks:

1. Protryptiline
2. Doxepin
3. Imipramine
4. Amitriptyline
5. Trimipramine



Excellent endcapping of NUCLEODUR® PolarTec displays significantly better peak shapes and less tailing for strong basic components compared to other phases with embedded polar group.

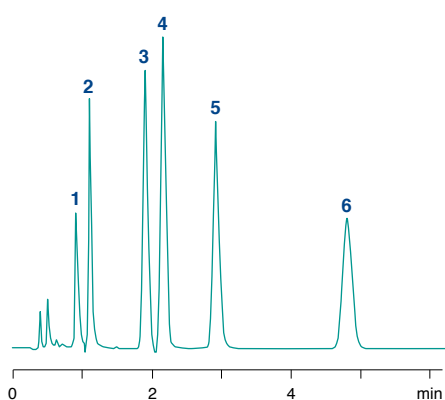
Tricyclic antidepressants

MN Appl. No. 117800

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: acetonitrile – 20 mM KH₂PO₄, pH 7.0 (65:35, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 2 µL

Peaks:

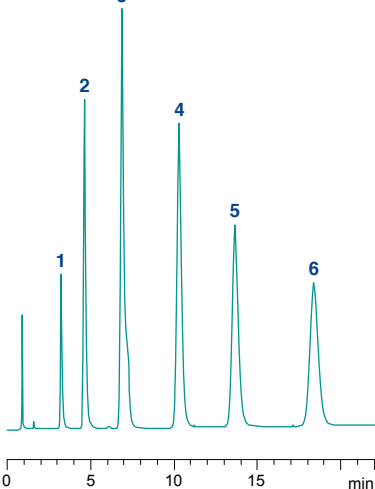
1. Protriptyline
2. Nortriptyline
3. Doxepin
4. Imipramine
5. Amitriptyline
6. Trimipramine



MN Appl. No. 118520

Column: 125 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
 Eluent: methanol – 20 mM KH₂PO₄, pH 7.0 (65:35, v/v)
 Temperature: 25 °C

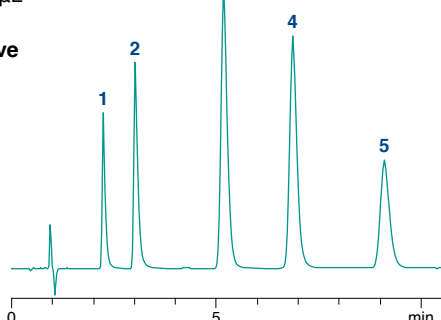
Peaks and other conditions as above



MN Appl. No. 119200

Column: 125 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: methanol – 20 mM NH₄H₂PO₄, pH 6.95 (70:30, v/v)
 Temperature: 40 °C
 Injection volume: 5 µL

Peaks and other conditions as above



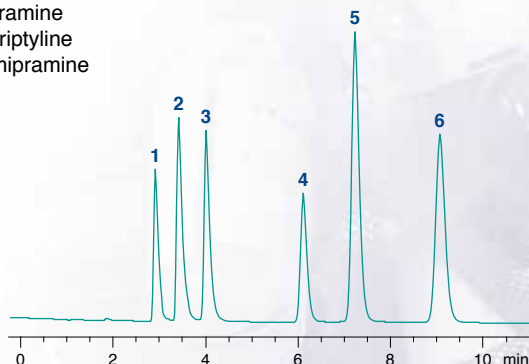
Tricyclic antidepressants

MN Appl. No. 121210

Column: 150 x 4 mm NUCLEODUR® C₁₈ Isis, 5 µm
 Eluent: methanol – 20 mM KH₂PO₄, pH 7 (75:25, v/v)
 Flow rate: 1 mL/min
 Temperature: 40 °C
 Detection: UV, 230 nm
 Injection volume: 8 µL

Peaks:

1. Protriptyline
2. Maprotiline
3. Nortriptyline
4. Imipramine
5. Amitriptyline
6. Clomipramine



Peak symmetry at 10 % of peak height:

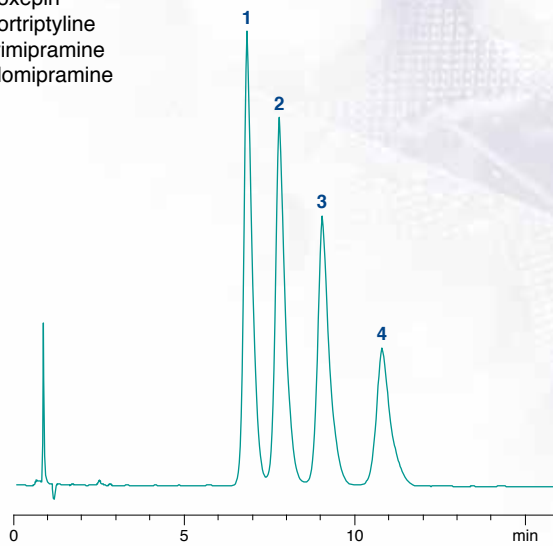
A_s (imipramine): 1.29
 A_s (amitriptyline): 1.26
 A_s (clomipramine): 1.16

MN Appl. No. 119280

Column: 250 x 4 mm NUCLEODUR® 100-5 CN-RP
 Eluent: acetonitrile – 20 mM KH₂PO₄, pH 6.5 (55:45, v/v)
 Flow rate: 1 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 2.5 µL (25 µg/mL)

Peaks:

1. Doxepin
2. Nortriptyline
3. Trimipramine
4. Clomipramine



Applications

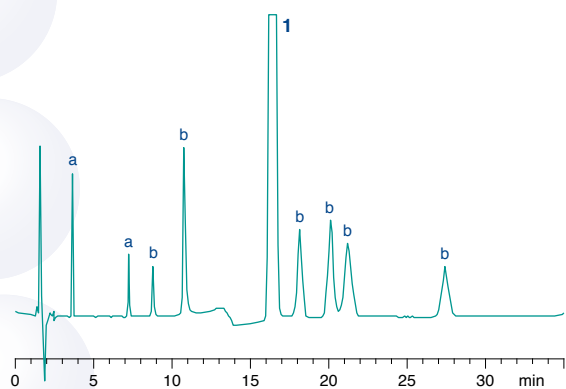
Neuroleptics

MN Appl. No. 121612

Column: 250 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
 Eluent: acetonitrile – 6.0 g/L KH₂PO₄, 2.9 g/L sodium dodecylsulfate, 9.0 g/L tetra-n-butylammonium bromide pH 8 (40:60, v/v)
 Flow rate: 1.5 mL/min
 Temperature: 40 °C
 Detection: 237 nm
 Injection volume: 5 µL

Peaks:

1. Chloroprothixene hydrochloride
- a. Additives
- b. Impurities



For separation on NUCLEODUR® C₁₈ Gravity see application 121611 at www.mn-net.com.

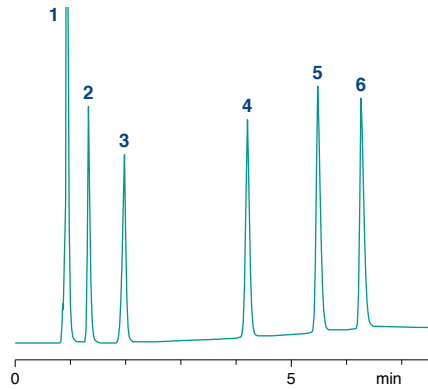
Cold medicine

MN Appl. No. 117810

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluents: A) 50 mM KH₂PO₄ + 5 mM pentanesulfonate (Na salt), pH 2.5; B) methanol
 Flow rate: 1.0 mL/min
 Temperature: 40 °C
 Detection: UV, 230 nm
 Injection volume: 5 µL

Peaks:

1. Maleic acid
2. Paracetamol
3. Pseudoephedrine
4. Benzoic acid
5. Chlorpheniramine
6. Dextromethorphan



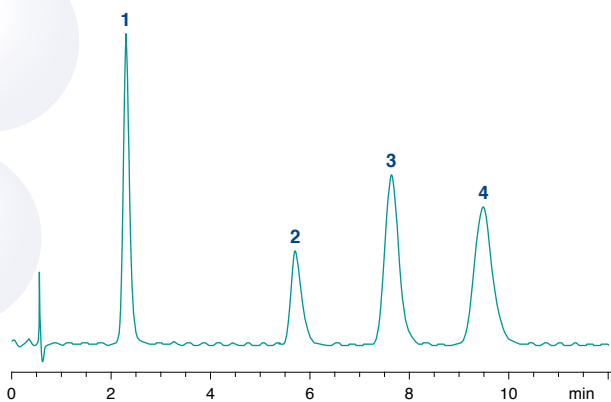
Gastric acid inhibitors

MN Appl. No. 122520

Column: 75 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 3 µm
 Eluent: methanol – 20 mM KH₂PO₄, pH 7 with TEA (20:80, v/v)
 Flow rate: 1.3 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 10 µL

Peaks:

1. Famotidine
2. Cimetidine
3. Nizatidine
4. Pirenzepine hydrochloride



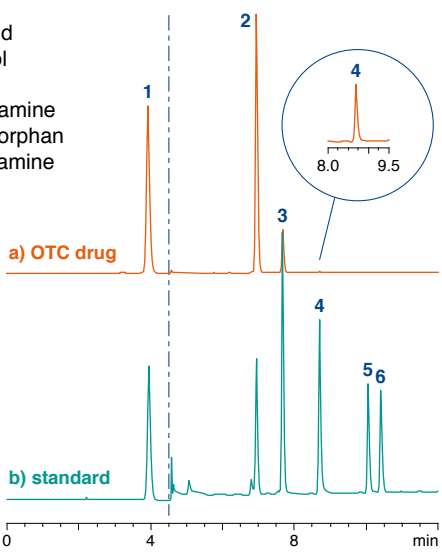
Cold medicine ingredients

MN Appl. No. 119110/119120

Column: 250 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: A) 50 mM NH₄H₂PO₄, pH 2.5; B) acetonitrile 0 % B → 60 % B in 13 min
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 230 nm for 4.5 min, then 261 nm
 Injection volume: a) 2 µL, b) 4 µL

Peaks:

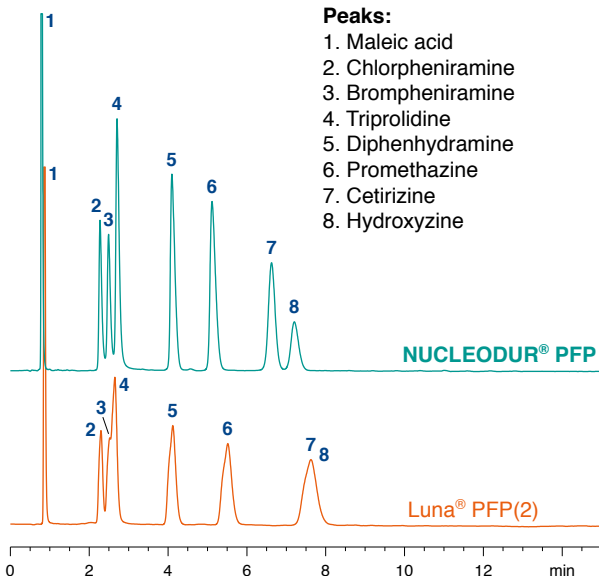
1. Ascorbic acid
2. Paracetamol
3. Caffeine
4. Chlorpheniramine
5. Dextromethorphan
6. Diphenhydramine



Antihistamines

MN Appl. No. 124851

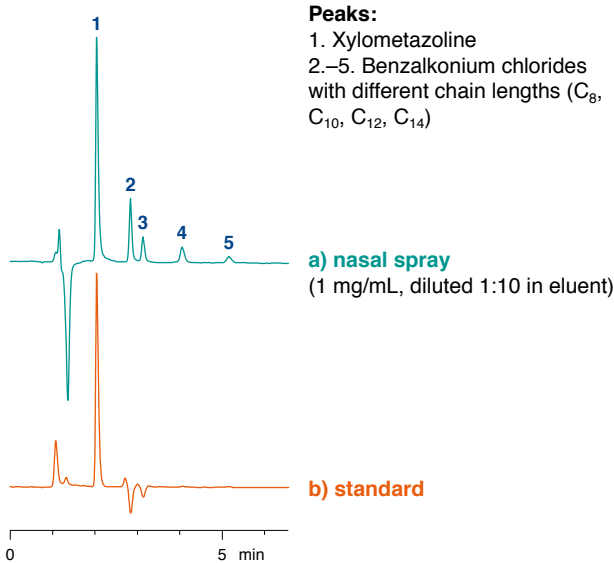
Columns: 100 x 4.6 mm NUCLEODUR® PFP, 5 µm
 Eluent: 100 x 4.6 mm Phenomenex Luna® PFP(2), 5 µm
 acetonitrile – 20 mM KH₂PO₄, pH 3.0 (30:70, v/v)
 Flow rate: 1.3 mL/min
 Temperature: 30 °C
 Detection: UV, 210 nm
 Injection volume: 1 µL



Xylometazoline in nasal spray

MN Appl. No. 120390

Column: 125 x 4 mm NUCLEODUR® 100-5 CN-RP
 Eluent: acetonitrile – 50 mM Na citrate, pH 3.0 (50:50, v/v)
 Flow rate: 0.8 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 100 µL



β₂-Agonists in human urine by LC-MS/MS

MN Appl. No. 119760

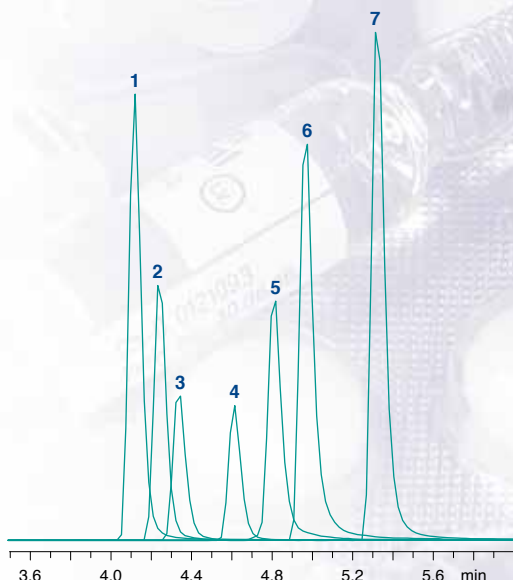
Column: 75 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Sample prep.: please refer to Thevis et al., J. Mass Spectrom 38 (2003) 1197–1206
 Eluents: A) 5 mM ammonium acetate with 0.1 % acetic acid, pH 3.5; B) acetonitrile; 0 % B → 100 % B in 6 min, reequilibration at 100 % A for 3.5 min
 Flow rate: 0.8 mL/min
 Temperature: 25 °C
 Detection: electrospray ionization / multiple reaction monitoring (MRM) on an Applied Biosystems API 2000
 Injection volume: 20 µL

LC-MS/MS chromatogram

2 mL urine aliquot fortified with 200 ng each

Peaks:

1. Reproterol (4.12 min)
2. Fenoterol (4.24 min)
3. Ritodrine (4.34 min)
4. Ractopamine (4.61 min)
5. Clenbuterol (4.81 min)
6. Bambuterol (4.97 min)
7. Mapenterol (5.32 min)



Courtesy of M. Thevis and W. Schänzer, Institute of Biochemistry, German Sport University, Cologne, Germany.

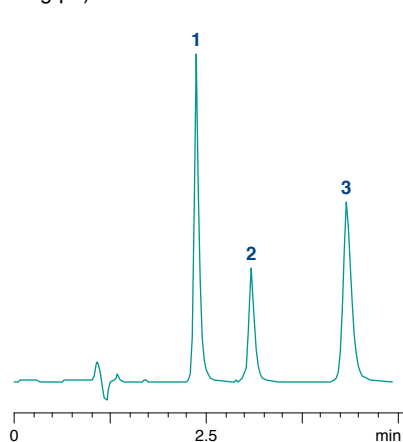
Applications

Basic drugs

MN Appl. No. 119320

Column: 125 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: acetonitrile – 20 mM KH₂PO₄, pH 6.5 (50:50, v/v)
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm
Injection volume: 1.0 µL

Peaks:	Tailing factor
1. Procainamide (5 ng/µL)	1.3
2. Clonidin (10 ng/µL)	1.2
3. Clenbuterol (12 ng/µL)	1.2

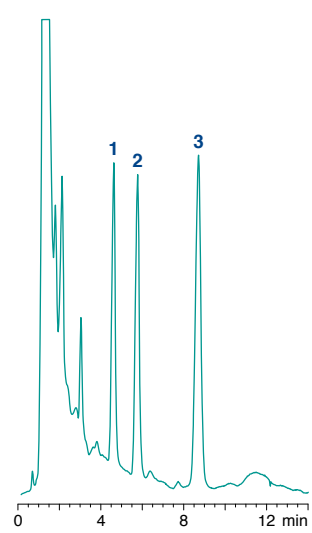


Benzodiazepine midazolam and metabolite from plasma

MN Appl. No. 118470

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 3 µm
Eluent: 127 mL KH₂PO₄ (9.1 g/L H₂O) + 309 mL Na₂HPO₄ (11.9 g/L H₂O) + 852 mL methanol + 0.15 g octanesulfonic acid, pH 5.56
Flow rate: 0.7 mL/min
Temperature: 25 °C
Detection: UV, DAD

Peaks:	
1. α-Hydroxymidazolam (metabolite)	
2. Midazolam (250 ng/mL)	
3. Internal standard	



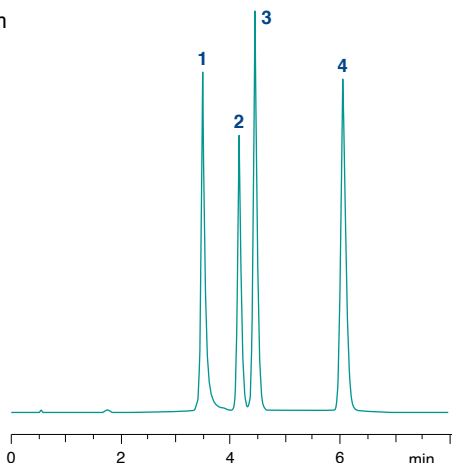
Courtesy of Mrs. Richter, Institute of Anesthetics, Biochemical Laboratory, University of Erlangen, Germany

Benzodiazepines

MN Appl. No. 117850

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
Eluent: acetonitrile – 20 mM KH₂PO₄, pH 6.5 (45:55, v/v)
Flow rate: 1.0 mL/min
Temperature: 22 °C
Detection: UV, 254 nm
Injection volume: 5 µL

Peaks:	
1. Bromazepam	
2. Oxazepam	
3. Lorazepam	
4. Temazepam	

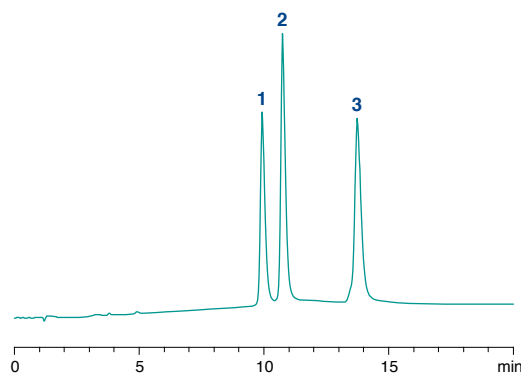


Sedative drugs

MN Appl. No. 119300

Column: 125 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: A) methanol
B) 50 mM ammonium acetate, pH 5.0
70 % B → 50 % B in 10 min (10 min)
Flow rate: 1.5 mL/min
Temperature: 30 °C
Detection: UV, 254 nm
Injection volume: 1 µL (1 + 2: 670 µg/mL, 3: 335 µg/mL)

Peaks:	
1. Promethazine	
2. Promazine	
3. Chlorpromazine	



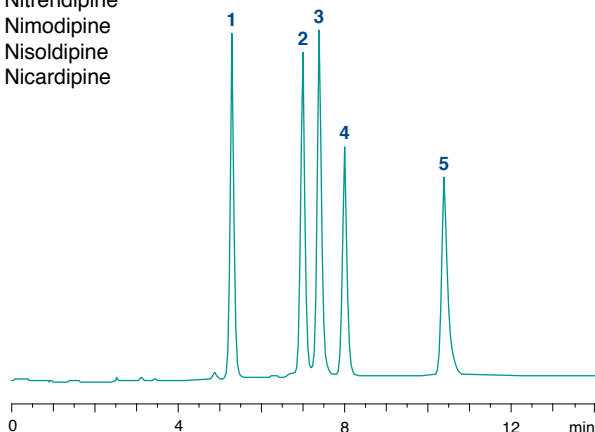
Coronary therapeutic drugs (Ca-antagonists)

MN Appl. No. 119310

Column: 125 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: A) acetonitrile, B) 20 mM KH₂PO₄, pH 6.5
30 % B → 50 % B in 7.5 min (7.5 min)
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm
Injection volume: 2.5 µL (25 µg/mL each)

Peaks:

1. Nifedipine
2. Nitrendipine
3. Nimodipine
4. Nisoldipine
5. Nicardipine



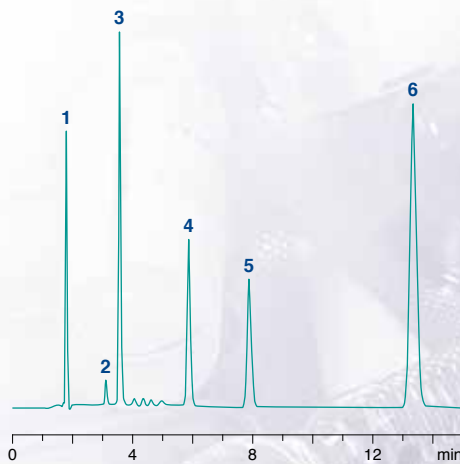
Antibacterial drugs

MN Appl. No. 117870

Column: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
Eluent: acetonitrile – water (40:60, v/v) 0.05 % TFA
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm
Injection volume: 5 µL

Peaks:

1. Ofloxacin
2. Ciprofloxacin
3. Cinoxacin
4. Penicillin G
5. Penicillin V
6. Cloxacillin



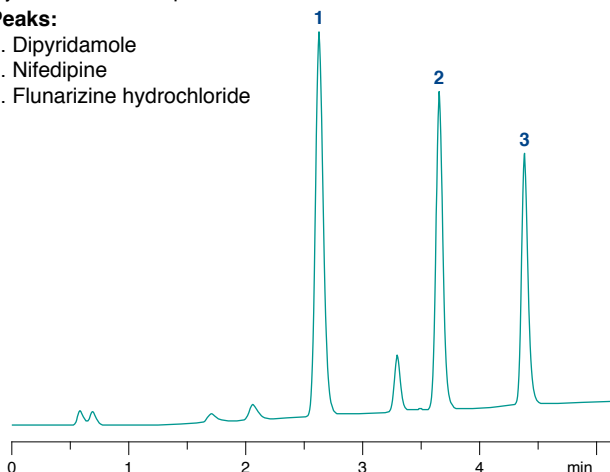
Cardiovascular drugs

MN Appl. No. 122560

Column: 75 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 3 µm
Eluent: A) 50 mM KH₂PO₄ + Na pentanesulfonate pH 2.5
B) methanol
45 % B → 90 % B in 6 min
Flow rate: 1.3 mL/min
Temperature: 35 °C
Detection: UV, 230 nm
Injection volume: 5 µL

Peaks:

1. Dipyridamole
2. Nifedipine
3. Flunarizine hydrochloride



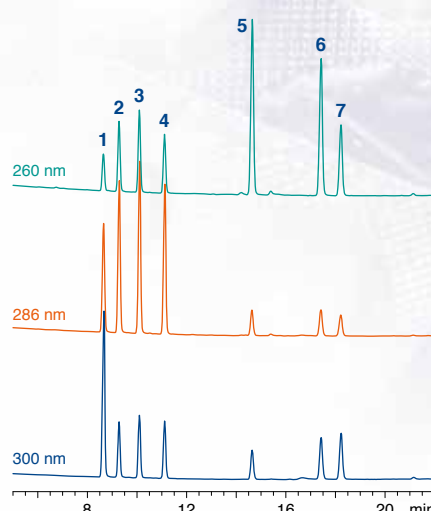
Gyrase inhibitors

MN Appl. No. 120400

Column: 150 x 3 mm NUCLEODUR® Sphinx RP, 5 µm
Eluent: A) 0.05 M H₃PO₄, B) acetonitrile
5 % B → 50 % B in 20 min
Flow rate: 0.5 mL/min
Detection: UV DAD, 260 nm, 286 nm and 300 nm
Injection volume: 20 µL (0.625 ng/µL of each compound)

Peaks:

1. Marbofloxacin
2. Ciprofloxacin
3. Enrofloxacin
4. Sarafloxacin
5. Oxolinic acid
6. Nalidixic acid
7. Flumequine



Courtesy of R. Lippold, Chemical and Veterinary Research Agency, Freiburg, Germany.

Applications

NUCLEODUR®

Quinolone antibiotics

MN Appl. No. 120460/120470

Columns: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 125 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
 125 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm

Eluent: methanol – 0.2% formic acid (40:60, v/v)

Flow rate: 1.0 mL/min

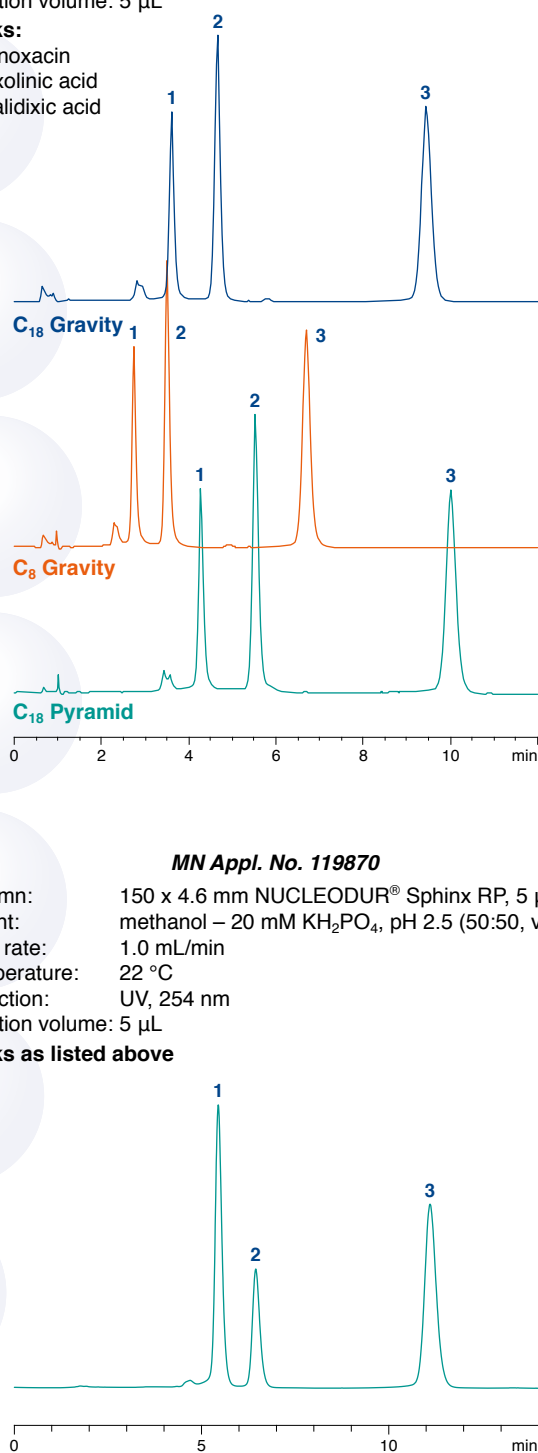
Temperature: 30 °C

Detection: UV, 254 nm

Injection volume: 5 µL

Peaks:

1. Cinoxacin
2. Oxolinic acid
3. Nalidixic acid



Penicillin antibiotics

MN Appl. No. 117860

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec

Eluent: acetonitrile – 20 mM KH₂PO₄, pH 3.0 (40:60, v/v)

Flow rate: 1.0 mL/min

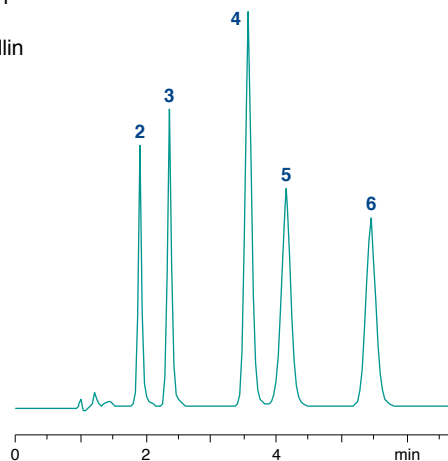
Temperature: 22 °C

Detection: UV, 254 nm

Injection volume: 5 µL

Peaks:

1. Amoxicillin
2. Penicillin G
3. Penicillin V
4. Cloxacillin
5. Nafcillin
6. Dicloxacillin



MN Appl. No. 119150

Column: 250 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm

Eluent: acetonitrile – 0.1 % TFA (50:50, v/v)

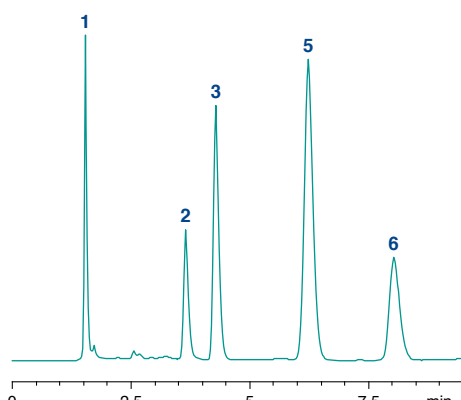
Flow rate: 1.0 mL/min

Temperature: 25 °C

Detection: UV, 254 nm

Injection volume: 1 µL

Peaks as listed above



MN Appl. No. 119870

Column: 150 x 4.6 mm NUCLEODUR® Sphinx RP, 5 µm

Eluent: methanol – 20 mM KH₂PO₄, pH 2.5 (50:50, v/v)

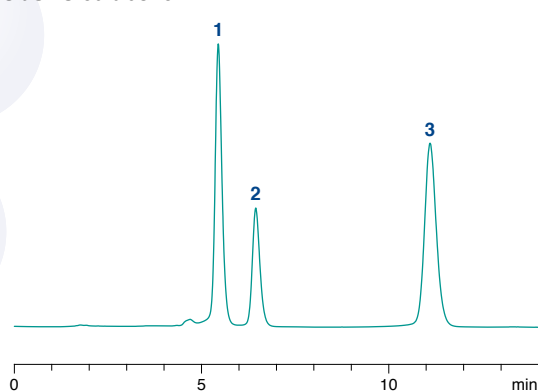
Flow rate: 1.0 mL/min

Temperature: 22 °C

Detection: UV, 254 nm

Injection volume: 5 µL

Peaks as listed above



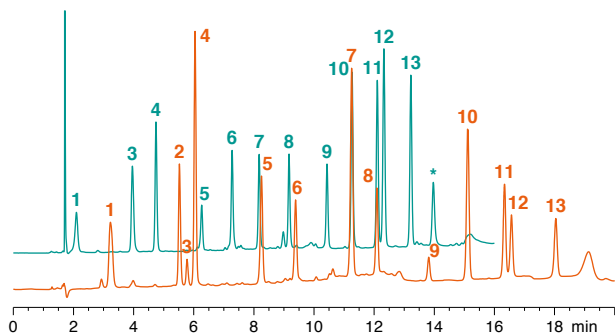
β -lactam antibiotics

MN Appl. No. 124840

Column: 150 x 3 mm NUCLEODUR® PFP, 5 μ m
 Eluents: A) acetonitrile, B) 25 mM KH₂PO₄, pH 2.7
 10% A → 60% A in 15 min
 A) acetonitrile, B) 0.1% TFA
 10% A → 45% A in 15 min (5 min)
 Flow rate: 0.563 mL/min
 Temperature: 40 °C, 50 °C
 Detection: UV, 220 nm, UV, 240 nm
 Injection volume: 1 μ L

Peaks:

- | | | |
|----------------|-----------------|-------------------|
| 1. Amoxicillin | 6. Cefamandole | 11. Cloxacillin |
| 2. Cephalexin | 7. Cefalotin | 12. Nafticillin |
| 3. Ampicillin | 8. Piperacillin | 13. Dicloxacillin |
| 4. Cefotaxime | 9. Penicillin V | * Impurity |
| 5. Cefoxitin | 10. Oxacillin | |

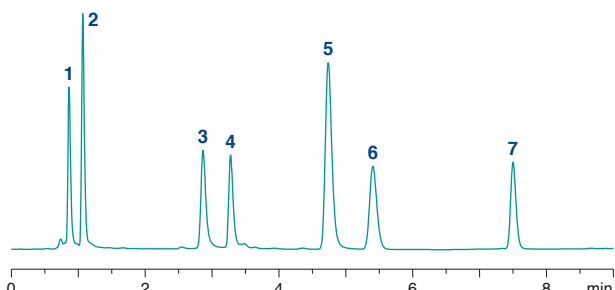


MN Appl. No. 123760

Column: 125 x 4 mm NUCLEODUR® C₁₈ HTec, 5 μ m
 Eluent: A) acetonitrile, B) 0.05% TFA in water
 70% B (1 min) → 60% B in 0.5 min (3.5 min) →
 50% B in 1 min → 37.5% B in 4 min
 Flow rate: 0.9 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 10 μ L
 Concentration: 300 μ g/mL

Peaks:

1. Amoxicillin
2. Enrofloxacin
3. Cinoxacin
4. Oxolinic acid
5. Nalidixic acid
6. Penicillin V
7. Cloxacillin



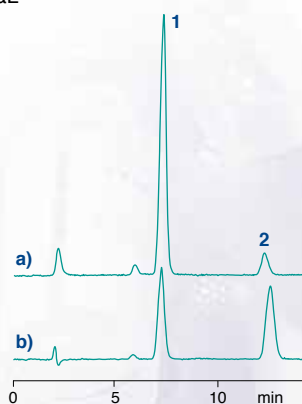
Anticoccidial drugs (polyether antibiotics)

MN Appl. No. 118760

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 3 μ m
 Eluent: methanol – 50 mmol phosphate buffer pH 3.0 – methylheptylamine (900:99:1, v/v/v)
 Flow rate: 0.7 mL/min
 Temperature: 23 °C
 Detection: UV/VIS, 600 nm after post column derivatization with dimethylaminobenzaldehyde (0.4 mL/min)
 Injection volume: 100 μ L

Peaks:

1. Monensin sodium
2. Salinomycin sodium



a): Sample spiked with monensin sodium and content of salinomycin sodium

b): Standard of monensin sodium and salinomycin sodium

Courtesy of J. Schönher, Saxon State Institute for Agriculture, Leipzig, Germany

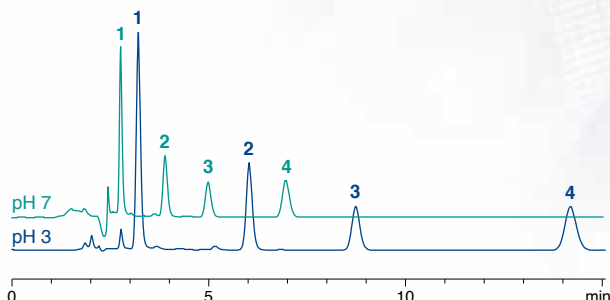
Cephalosporin antibiotics

MN Appl. No. 122580/122590

Column: 150 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 5 μ m
 Eluent: acetonitrile – 25 mM KH₂PO₄ (20:80, v/v)
 pH 3 with H₃PO₄, pH 7
 Flow rate: 0.8 mL/min
 Temperature: 35 °C
 Detection: UV, 254 nm
 Injection volume: 2 μ L

Peaks:

1. Cefotaxime
2. Cefoxitin
3. Cefamandole
4. Cephalothin



Protonation causes a drastic increase in retention time, but an improved peak symmetry.

Applications

NUCLEODUR®

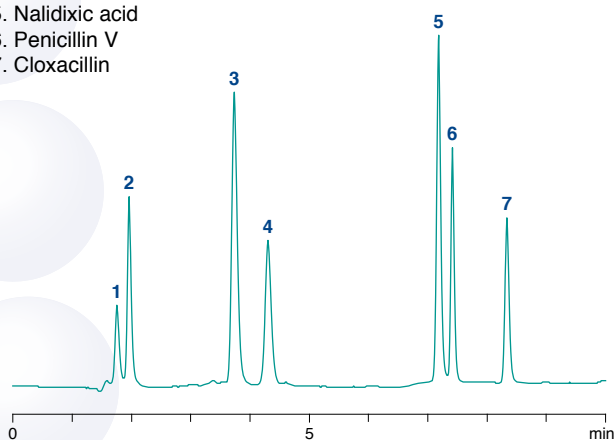
Antibacterial drugs

MN Appl. No. 122470

Column: 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: A) acetonitrile, B) water + 0.05 % TFA
 60 % B (4 min) → 40 % B in 1 min (5 min)
 Flow rate: 0.9 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

1. Amoxicillin
2. Enrofloxacin
3. Cinoxacin
4. Oxolinic acid
5. Nalidixic acid
6. Penicillin V
7. Cloxacillin



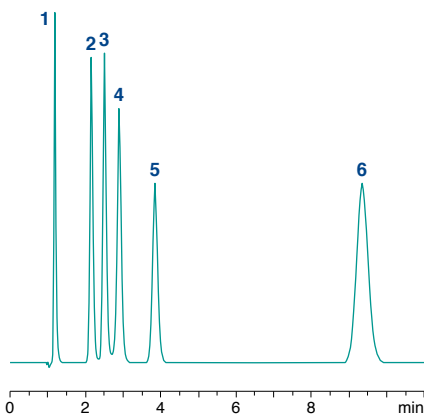
Sulfonamides

MN Appl. No. 117880

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: methanol – 0.1 % TFA (20:80, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 22 °C
 Detection: UV, 230 nm
 Injection volume: 4 µL

Peaks:

1. Sulfanilamide
2. Sulfadiazine
3. Sulfathiazole
4. Sulfamerazine
5. Sulfadimidine
6. Succinylsulfathiazole



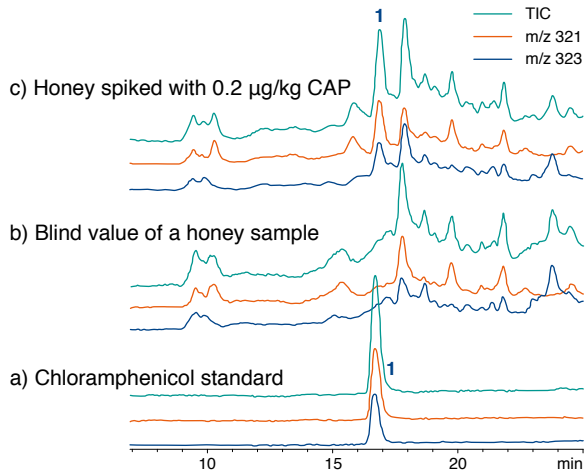
Determination of chloramphenicol residues in honey by microbore HPLC

MN Appl. No. 119810

Column: 100 x 1 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: A) methanol, B) water
 15 % A → 80 % A in 9 min (15 min) → 15 % A in 1 min; injection after 7 min
 Flow rate: 60 µL/min
 Detection: MS
 Injection volume: 1 µL

Peaks:

1. Chloramphenicol (CAP)



S. Oepkemeier, H.D. Winkler, GIT 46 (2002) 982–985.

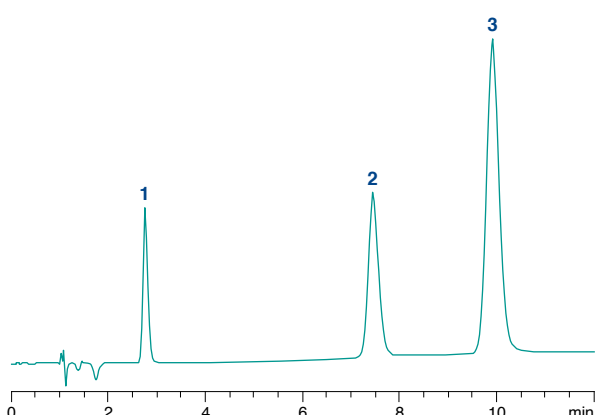
Separation of theobromine, vanillin and caffeine

MN Appl. No. 119920

Column: 125 x 4 mm NUCLEODUR® Sphinx RP, 5 µm
 Eluent: methanol – 1.25 % acetic acid (20:80, v/v)
 Flow rate: 1 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 0.8 µL

Peaks:

1. Theobromine
2. Caffeine
3. Vanillin



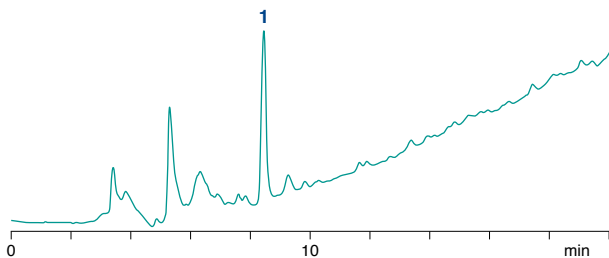
Alkaline tannic acid mixture

MN Appl. No. 120450

Column: 250 x 4.6 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: A) 0.425 % H₃PO₄, pH 1.4, B) acetonitrile
 5 % B → 25 % B in 15 min (5 min) → 5 % B in
 2 min (3 min)
 Flow rate: 0.8 mL/min
 Detection: UV, 275 nm (optimized for gallic acid)
 Injection volume: 10 µL

Peaks:

1. Gallic acid



Soft drink additives

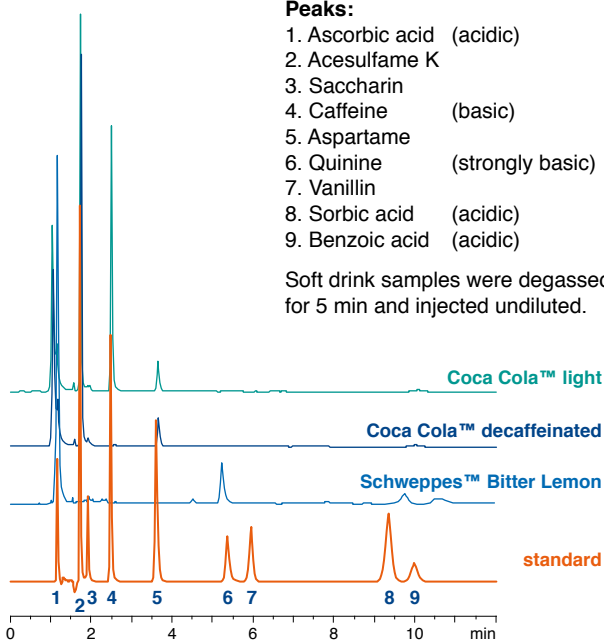
MN Appl. No. 118560

Column: 150 x 4.6 mm NUCLEODUR® 100-5 C₈ ec
 Eluent: 20 mM KH₂PO₄, pH 3 – acetonitrile (5:1, v/v)
 Flow rate: 1.9 mL/min
 Temperature: 25 °C
 Detection: UV, 220 nm
 Injection volume: 10 µL

Peaks:

1. Ascorbic acid (acidic)
2. Acesulfame K
3. Saccharin
4. Caffeine (basic)
5. Aspartame
6. Quinine (strongly basic)
7. Vanillin
8. Sorbic acid (acidic)
9. Benzoic acid (acidic)

Soft drink samples were degassed for 5 min and injected undiluted.



For fast separation of sweeteners on NUCLEODUR® 100-5 C₁₈ ec see appl. 117940 at www.mn-net.com.

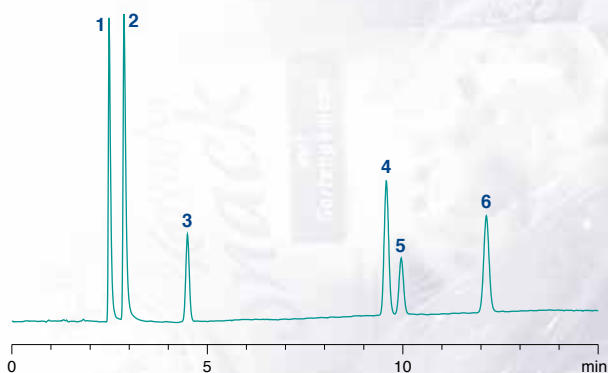
Sweeteners

MN Appl. No. 123750

Column: 150 x 4.6 mm NUCLEODUR® C₁₈ HTec, 5 µm
 Eluent: A) acetonitrile, B) 25 mM KH₂PO₄, pH 3.5
 15 % A (2.5 min) → 25 % A in 9.5 min (3 min)
 Flow rate: 1.3 mL/min
 Temperature: 40 °C
 Detection: UV, 220 nm
 Injection volume: 5 µL
 Concentration: 0.1 mg/mL each

Peaks:

1. Acesulfame K
2. Saccharin
3. Aspartame
4. Benzoic acid
5. Sorbic acid
6. Dehydroacetic acid



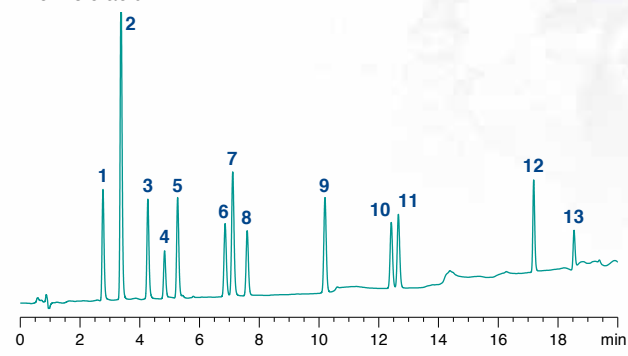
Preservatives

MN Appl. No. 124590

Column: 150 x 3 mm NUCLEODUR® PolarTec, 3 µm
 Eluent: A) acetonitrile, 0.1 % TFA, B) water, 0.1 % TFA
 20 % A → 50 % A in 12 min → 65 % A in 2 min →
 95 % A in 6 min
 Flow rate: 0.9 mL/min
 Temperature: 45 °C
 Detection: UV, 220 nm
 Injection volume: 5 µL

Peaks:

1. Benzyl alcohol
2. Phenoxyethanol
3. Dehydroacetic acid
4. p-Anisic acid
5. Methyl paraben
6. Salicylic acid
7. Benzoic acid
8. Ethyl paraben
9. Propyl paraben
10. Isobutyl paraben
11. Butyl paraben
12. Irgasan
13. 3,3,4-Triclocarbanilid



Applications

NUCLEODUR®

Food dyes

MN Appl. No. 122500/122510

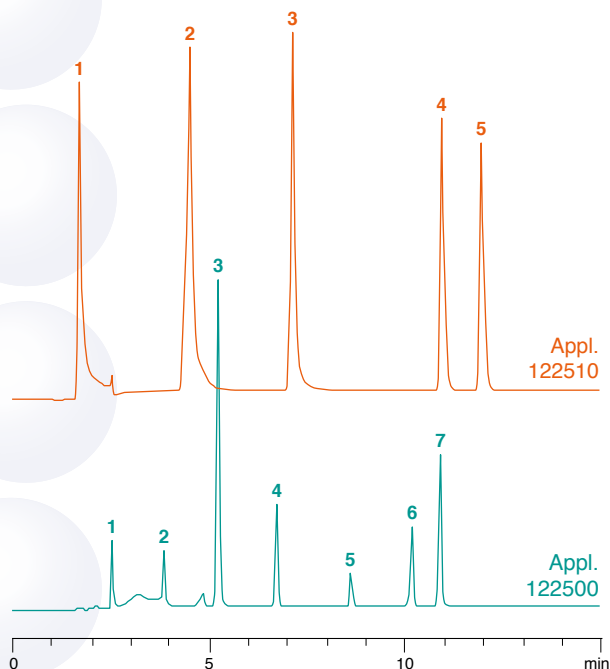
Column: 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: A) acetonitrile, B) 20 mM KH₂PO₄, pH 5
 95 % B → 50 % B in 20 min → 20 % B in 5 min
 → 95 % B in 1 min (4 min)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks application 122510

1. Ponceau 6R (E 126)
2. Ponceau 4R (E 124)
3. Azorubine (E 122)
4. Erythrosine (E 127)
5. Fast Red E

Peaks application 122500

- 1., 2. Tartrazine (E 102)
3. Fast Yellow
- 4.–6. Quinoline yellow (E 104)
7. Yellow orange S (sunset yellow CFC, E 110)



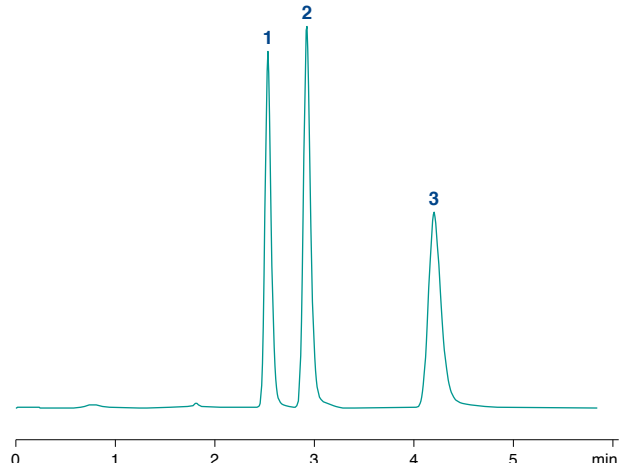
Acrylamide, methacrylamide and methacrylic acid

MN Appl. No. 123010

Column: 125 x 4 mm NUCLEODUR® HILIC, 5 µm
 Eluent: acetonitrile – 0.1 % formic acid (98:2, v/v)
 Flow rate: 0.6 mL/min
 Temperature: 22 °C
 Detection: UV, 210 nm
 Injection volume: 0.5 µL

Peaks:

1. Methacrylamide
2. Acrylamide
3. Methacrylic acid



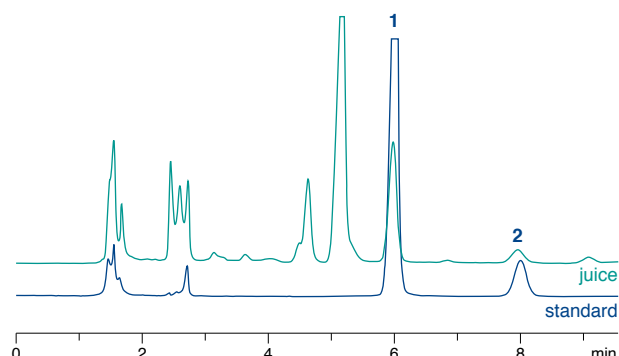
Patulin and hydroxymethylfurfural in apple juice

MN Appl. No. 121800

Column: 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm + 8 x 4 mm guard column
 Sample prep.: see appl. 121800 at www.mn-net.com
 Eluent: water – acetonitrile (95:5, v/v)
 Flow rate: 1.5 mL/min
 Detection: UV, 276 nm
 Injection volume: 10 µL

Peaks:

1. Hydroxymethylfurfural
2. Patulin



Courtesy of A. Gessler, Wesergold Getränkeindustrie GmbH & Co. KG, Rinteln, Germany.

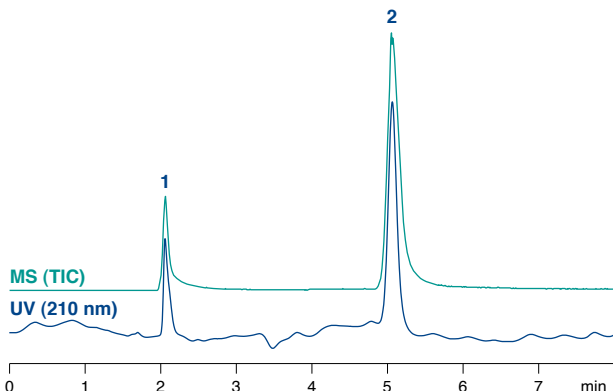
Melamine and cyanuric acid

MN Appl. No. 123070

Column: 125 x 2 mm NUCLEODUR® HILIC, 5 µm
 Eluent: acetonitrile – 10 mM ammonium formate, pH 4 (90:10, v/v)
 Flow rate: 0.2 mL/min
 Temperature: 25 °C
 Detection: UV, 210 nm and MS (TIC)
 Injection volume: 2 µL

Peaks:

1. Melamine (10 µg/mL)
2. Cyanuric acid (490 µg/mL)

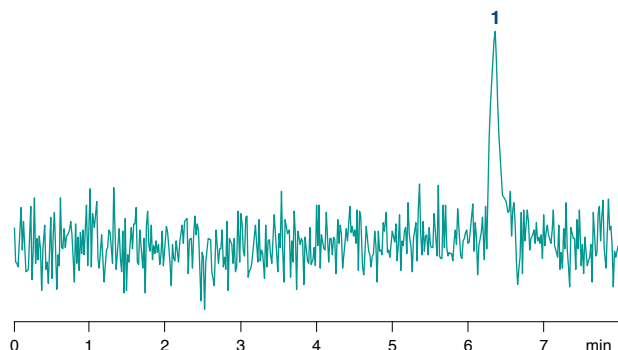


MN Appl. No. 123090

Column: 125 x 4 mm NUCLEODUR® HILIC, 5 µm
 Eluent: acetonitrile – 10 mM ammonium formate, pH 4 (90:10, v/v)
 Flow rate: 0.6 mL/min
 Temperature: 25 °C
 Detection: MS
 Injection volume: 1 µL

Peaks:

1. Melamine in milk (100 pg/injection)



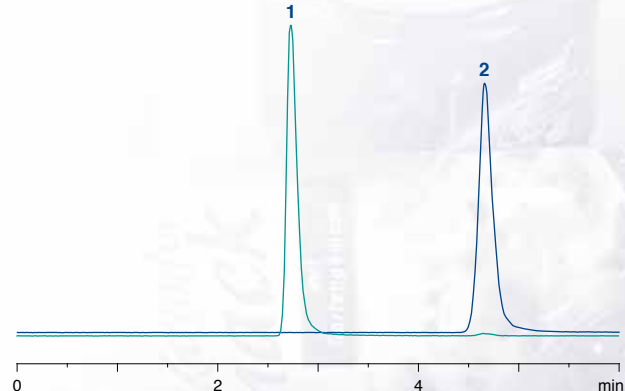
Creatinine and creatine

MN Appl. No. 123000

Column: 125 x 2 mm NUCLEODUR® HILIC, 3 µm
 Eluent: acetonitrile – 10 mM ammonium acetate, pH 4 (70:30, v/v)
 Flow rate: 0.2 mL/min
 Temperature: 25 °C
 Detection: MS
 Injection volume: 5 µL (30 ng/µL)

Peaks:

1. Creatinine
2. Creatine



For UV detection see appl. 122990 at www.mn-net.com.

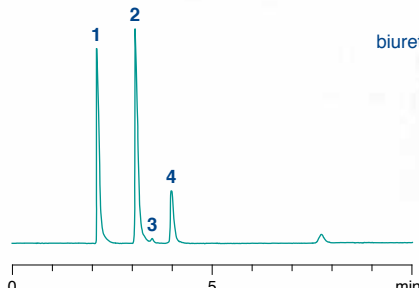
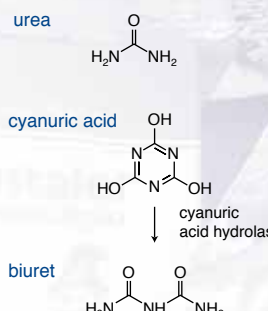
Separation of urea, biuret and cyanuric acid

MN Appl. No. 120440

Column: 250 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: water (100 %)
 Flow rate: 1 mL/min
 Detection: UV, 190 nm
 Injection volume: 10 µL

Peaks:

1. Urea (0.5 mg/mL)
2. Biuret (0.09 mg/mL)
3. impurity in biuret
4. Cyanuric acid (0.05 mg/mL)



Courtesy of C. Greve, Institute of Chemical Engineering, University of Clausthal, Germany.

Applications

Amino acids as OPA derivatives

MN Appl. No. 118450

Column: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec

Eluent: A) methanol – acetonitrile (50:50, v/v), B) Na acetate buffer pH 6.5 + 5 % A)

100 % B (2.9 min) → 95 % B in 3.1 min → 85 % B in 11 min → 83 % B in 3 min → 70 % B in 10 min → 62 % B in 8 min → 35 % B in 7 min → 0 % B in 1 min (2 min) → 80 % B in 0.5 min (2.5 min) → 100 % B in 0.1 min

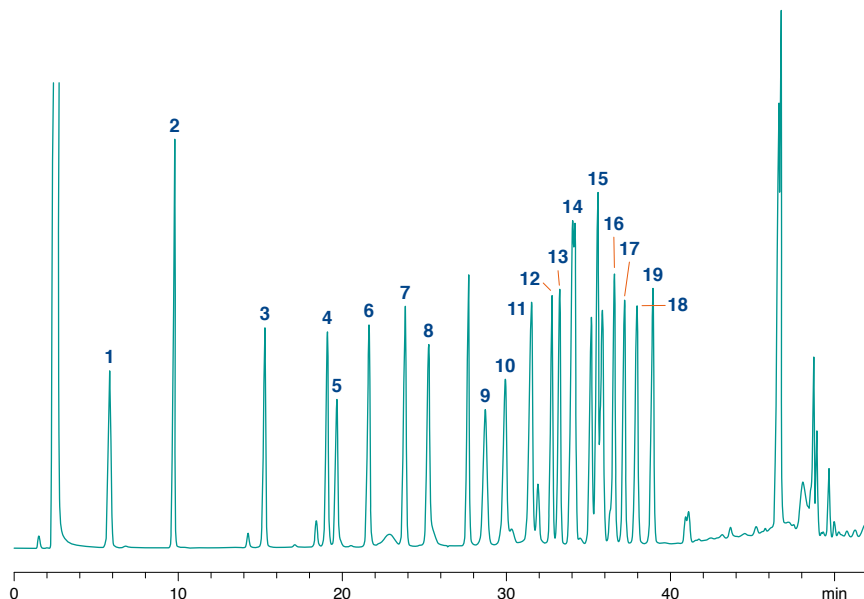
Flow rate: 1 mL/min

Detection: fluorescence,

λ_{ex} 230 nm, λ_{em} 450 nm

Peaks:

1. Aspartic acid
2. Glutamic acid
3. Asparagine
4. Serine
5. Glutamine
6. Histidine
7. Glycine
8. Alanine
9. Arginine
10. γ -Aminobutyric acid
11. Tyrosine
12. Valine
13. Methionine
14. Norvaline (int. std.)
15. Tryptophan
16. Phenylalanine
17. Isoleucine
18. Leucine
19. Lysine



Courtesy of Mr. Zürcher, Technical University of Munich, Chair of Brewing Technology, Freising Weihenstephan, Germany.

For separation of amino acids also see appl. 120510 at www.mn-net.com.

Determination of physiological amino acids from supernatants of cell cultures

MN Appl. No. 118980

Column: 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 3 μ m

Sample preparation: supernatants from cell cultures are deproteinated, derivatized with phenylisothiocyanate and filtered

Eluent: A) 70 mM sodium acetate, pH 6.5, 2.5 % acetonitrile, 1 ppm EDTA; B) acetonitrile – water – methanol (45:40:15, v/v/v)
3 % B (2 min) → 7.5 % B in 16 min (8 min) → 44 % B in 49 min
(washing: 100 % B for 10 min; equilibration: 3 % B for 5 min)

Flow rate: 0.8 mL/min

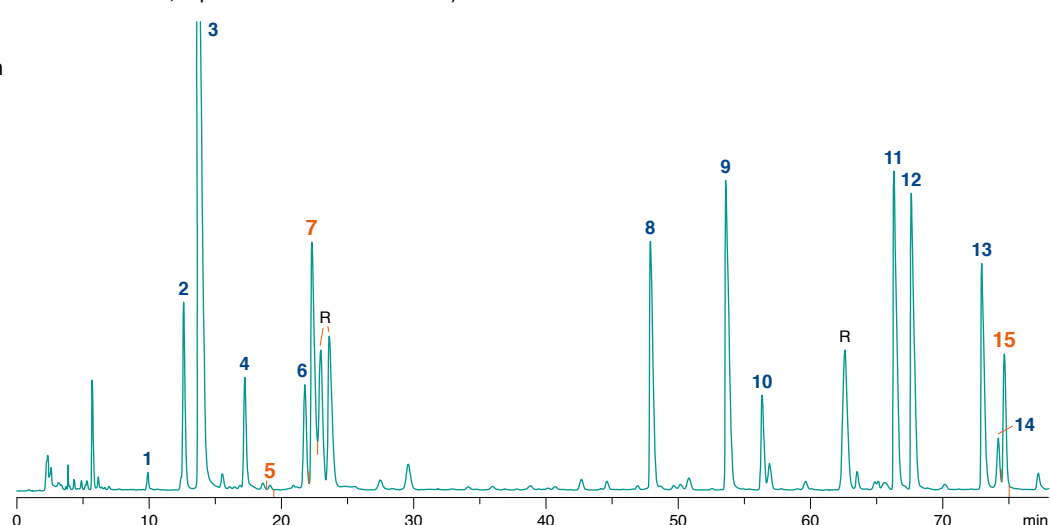
Temperature: 47 ± 1 °C

Detection: UV, 254 nm

Injection volume: 40 μ L

Peaks:

1. 4-Hydroxyproline
2. Serine
3. Glutamine
4. Histidine
5. Citrulline (5.90 μ mol/L)
6. Threonine
7. Arginine (504.33 μ mol/L)
8. Tyrosine
9. Valine
10. Methionine
11. Isoleucine
12. Leucine
13. Phenylalanine
14. Tryptophan
15. Ornithine (143.54 μ mol/L)



Citrulline, arginine and ornithine were determined quantitatively.

Courtesy of Dr. J. Weinreich, Center for Medical Research, Clinic for General Surgery, University Clinical Center, Tübingen, Germany.

Biological and natural compounds

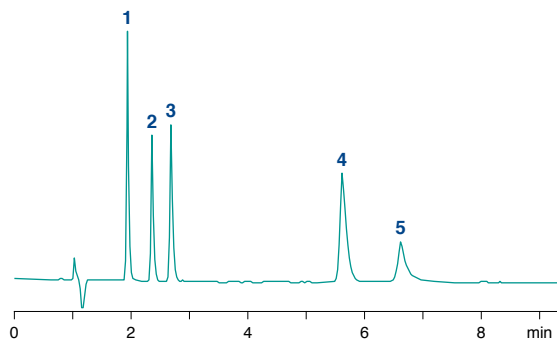
Aromatic amino acids and histamine

MN Appl. No. 122980

Column: 125 x 4 mm NUCLEODUR® HILIC, 3 µm
 Eluent: acetonitrile – 100 mM ammonium acetate, pH 4 (75:25, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 218 nm
 Injection volume: 0.5 µL

Peaks:

1. Phenylalanine
2. Phenylglycine
3. Tyrosine
4. Histamine
5. Histidine



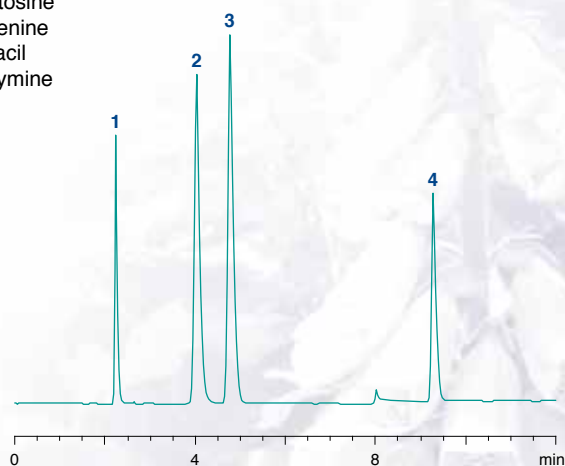
Nucleic acid bases

MN Appl. No. 119140

Column: 250 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: A) 50 mM NH₄H₂PO₄, pH 2.5, B) acetonitrile 100% A (2.5 min) → 90% A in 10 min
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 3 µL

Peaks:

1. Cytosine
2. Adenine
3. Uracil
4. Thymine



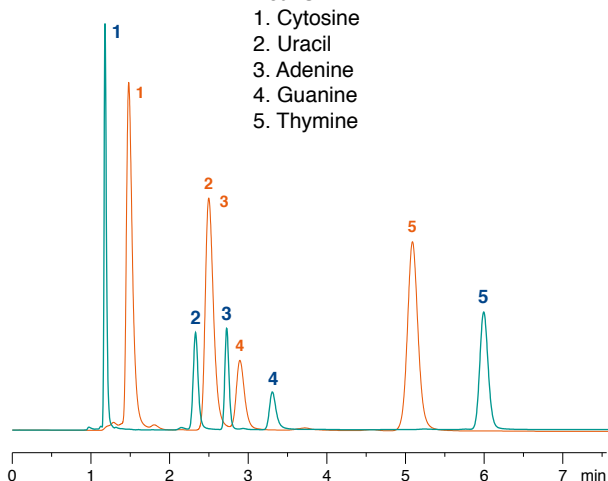
Nucleic acid bases

MN Appl. No. 124672

Columns: 150 x 3 mm NUCLEODUR® PolarTec, 5 µm
 150 x 3 mm Waters SymmetryShield™ RP18, 5 µm
 Eluent: 30 mM KH₂PO₄, pH 3.0
 Flow rate: 0.5 mL/min
 Temperature: 30 °C
 Detection: UV, 220 nm
 Injection volume: 1 µL, 3 µL

Peaks:

1. Cytosine
2. Uracil
3. Adenine
4. Guanine
5. Thymine



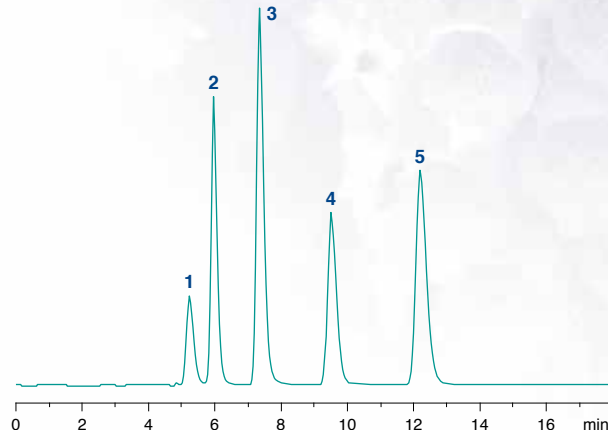
Separation of all five nucleic acid bases just on NUCLEODUR® PolarTec

MN Appl. No. 122950

Column: 125 x 4 mm NUCLEODUR® HILIC, 5 µm
 Eluent: acetonitrile – 5 mM ammonium acetate (80:20, v/v)
 Flow rate: 0.3 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm

Peaks:

1. Thymine
2. Uracil
3. Adenine
4. Cytosine
5. Guanosine



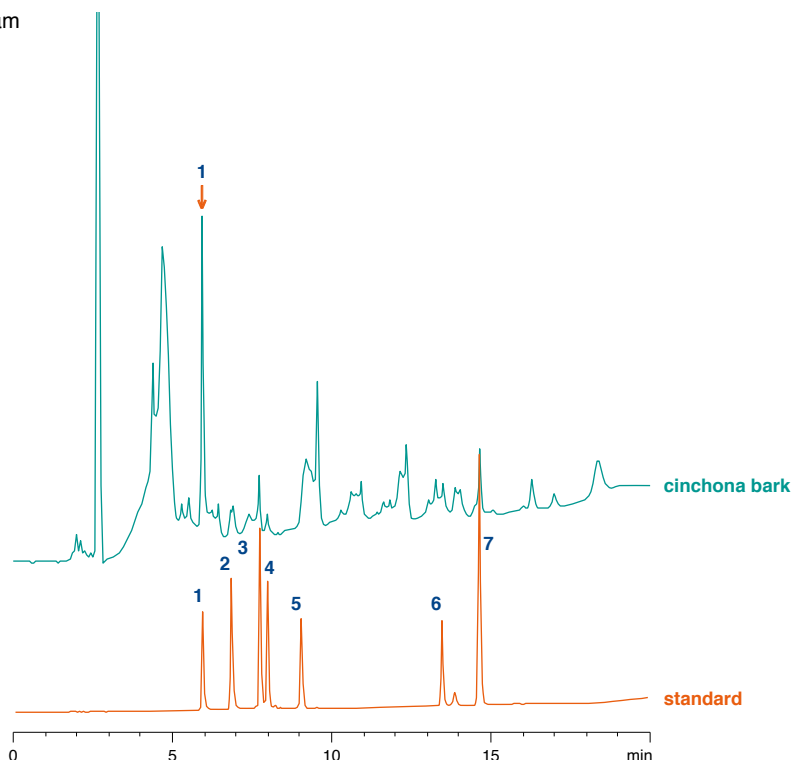
Applications

Determination of quinine in cinchona bark

MN Appl. No. 118580

Column: 125 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
Eluent: A) 20 mM NH₄H₂PO₄, pH 2
B) acetonitrile
Gradient: 10 % B → 30 % B in 15 min
Flow rate: 1 mL/min
Temperature: 30 °C
Detection: UV, 210 nm

Peaks:
1. Quinine
2. Scopolamine
3. Brucine
4. Strychnine
5. Atropine
6. Papaverine
7. Noscapine

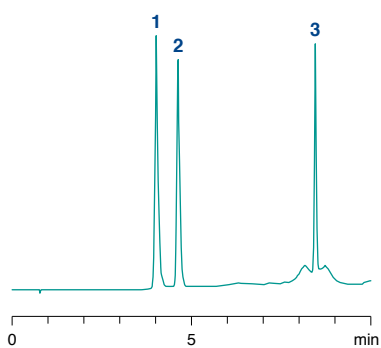


Quinine alkaloids

MN Appl. No. 117960

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
Eluent: A) methanol, B) 20 mM KH₂PO₄, pH 2.5
90 % B → 70 % B in 4 min → 30 % B in 7 min
Flow rate: 1.3 mL/min
Temperature: 25 °C
Detection: UV, 240 nm
Injection volume: 10 µL

Peaks:
1. Chloroquine
2. Quinine
3. Mefloquine

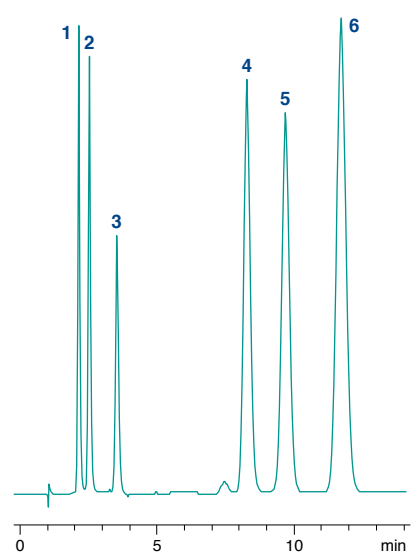


Steroids

MN Appl. No. 118540

Column: 125 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
Eluent: acetonitrile – water (60:40, v/v)
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 240 nm

Peaks:
1. Cortisone
2. Hydrocortisone
3. Hydrocortisone 21-acetate
4. 6α-Methyl-11β-hydroxyprogesterone
5. 6α-Methyl-17α-hydroxyprogesterone
6. 6α-Methyl-17α-hydroxyprogesterone acetate



Biological and natural compounds

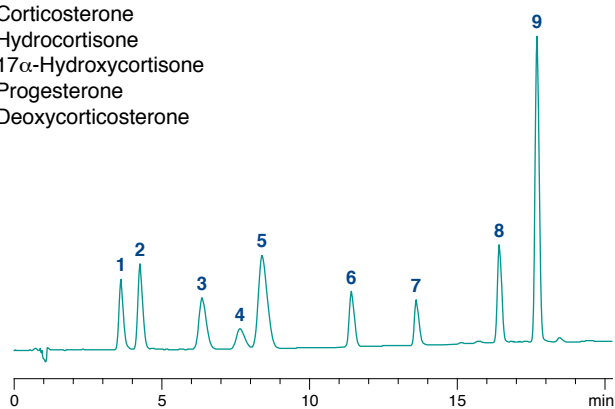
Steroids

MN Appl. No. 122530

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: A) acetonitrile, B) water
 70 % B (7 min) → 20 % B in 16 min → 70 % B in 2 min
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 240 nm
 Injection volume: 3 µL

Peaks:

1. Cortisone
2. Prednisolone
3. 6α-Methylprednisolone
4. Dexamethasone
5. Corticosterone
6. Hydrocortisone
7. 17α-Hydroxycorticosterone
8. Progesterone
9. Deoxycorticosterone



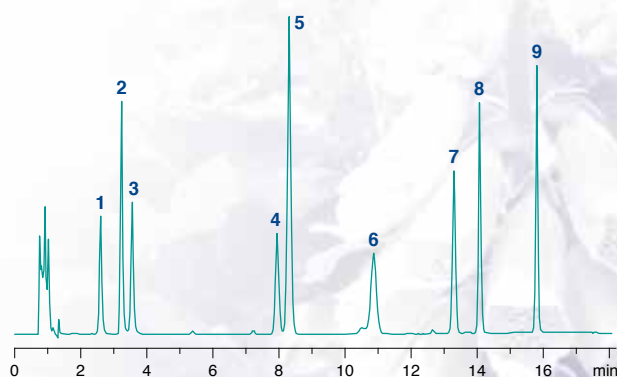
Steroids

MN Appl. No. 123710

Column: 125 x 4 mm NUCLEODUR® C₁₈ HTec, 5 µm
 Eluent: A) acetonitrile, B) water
 70 % B → 60 % B in 5 min (5 min) → 35 % B in 5 min (5 min)
 Flow rate: 1.0 mL/min
 Temperature: 35 °C
 Detection: UV, 254 nm
 Injection volume: 10 µL

Peaks:

- | | |
|-----------------|--------------------------------------|
| 1. Estriol | 6. Estrone |
| 2. Prednisolone | 7. 6α-Methyl-11β-hydroxyprogesterone |
| 3. Cortisone | 8. 6α-Methyl-17α-hydroxyprogesterone |
| 4. Estradiol | 9. Progesterone |
| 5. Testosterone | |



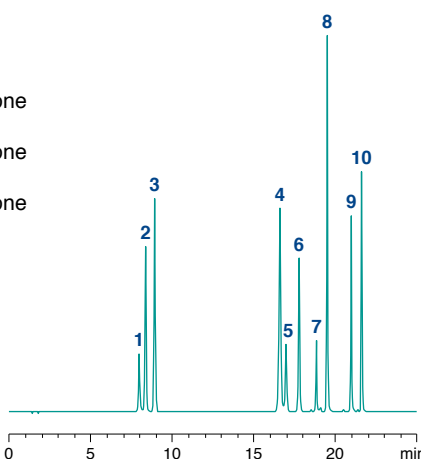
Steroids

MN Appl. No. 118550

Column: 125 x 4 mm NUCLEODUR® 100-5 C₈ ec
 Eluent: A) water, B) methanol
 20 % B (1 min) → 35 % B in 10 min (3 min) → 60 % B in 6 min (5 min)
 Flow rate: 1.0 mL/min
 Temperature: 30 °C
 Detection: UV, 230 nm
 Injection volume: 10 µL (each ~10–50 µg/mL)

Peaks:

1. Estriol
2. Prednisolone
3. Cortisone
4. Testosterone
5. 6α-Methyl-11β-hydroxyprogesterone
6. 6α-Methyl-17α-hydroxyprogesterone acetate
7. 6α-Methyl-17α-hydroxyprogesterone acetate
8. Estradiol
9. Estrone
10. Progesterone



Hydroxytestosterones from cytochrome P450

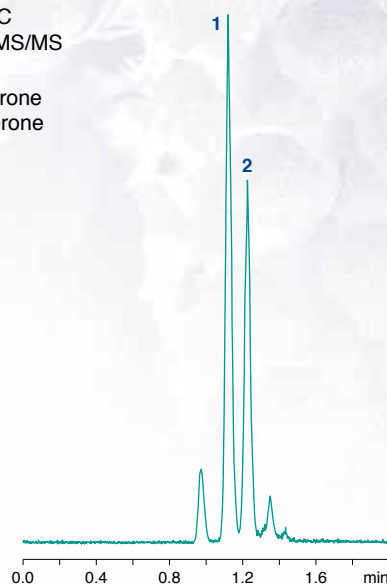
MN Appl. No. 122140

Column: 50 x 2 mm NUCLEODUR® C₁₈ Isis, 1.8 µm
 Eluent: A) water + 0.1 % formic acid; B) acetonitrile – methanol + 0.3 % formic acid
 75 % A → 60 % A in 1.1 min → 0 % A in 0.05 min, → 75 % A in 0.05 min (0.95 min)

Flow rate: 0.9 mL/min
 Temperature: 70 °C
 Detection: LC-MS/MS

Peaks:

1. 6β-Hydroxytestosterone
2. 7α-Hydroxytestosterone



Applications

NUCLEODUR®

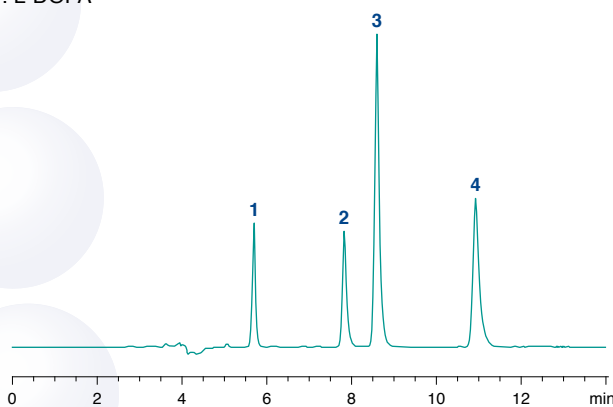
Catecholamines

MN Appl. No. 123030

Column: 250 x 4 mm NUCLEODUR® HILIC, 3 µm
 Eluent: acetonitrile – 25 mM ammonium formate, pH 3 (75:25, v/v)
 Flow rate: 0.8 mL/min
 Temperature: 25 °C
 Detection: UV, 218 nm
 Injection: 5 µL, 30 ng/µL

Peaks:

1. Norephedrine
2. Dopamine
3. Adrenaline
4. L-DOPA



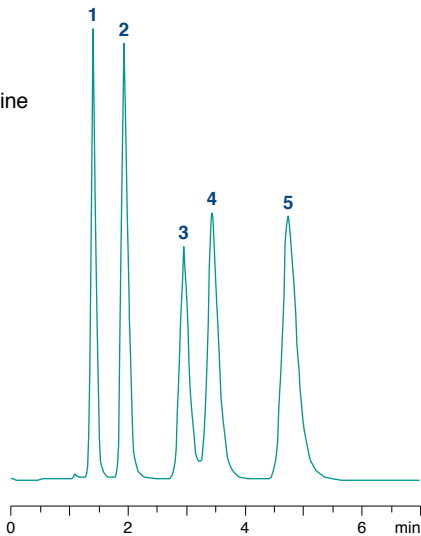
Catecholamines

MN Appl. No. 117930

Column: 125 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: 100 mM NaH₂PO₄, pH 3.0
 Flow rate: 0.8 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

1. Norephedrine
2. Adrenaline
3. Dihydroxyphenylalanine
4. Hydroxytyramine
5. Tyrosine



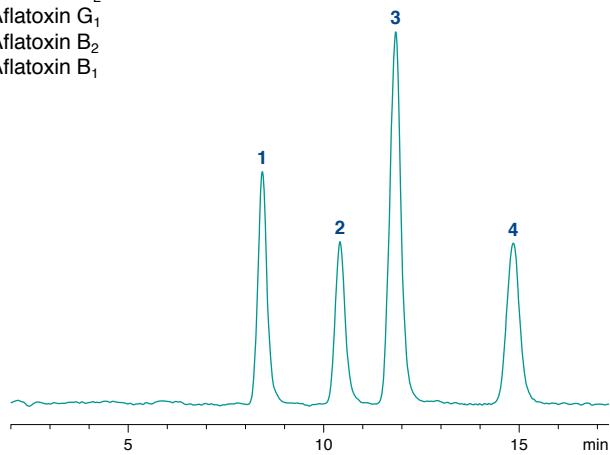
Analysis of aflatoxins from baby food

MN Appl. No. 120780

Column: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: methanol – acetonitrile – water (26:17:57, v/v/v) with 119 mg KBr and 100 µL HNO₃ (65 %) per liter
 Flow rate: 1.0 mL/min
 Detection: fluorescence, λ_{ex} 362 nm, λ_{em} 440 nm, post column derivatization in a CoBrA cell (Dr. Weber Consulting Kft)
 Injection volume: 100 µL

Peaks:

1. Aflatoxin G₂
2. Aflatoxin G₁
3. Aflatoxin B₂
4. Aflatoxin B₁



Analysis of mycotoxins

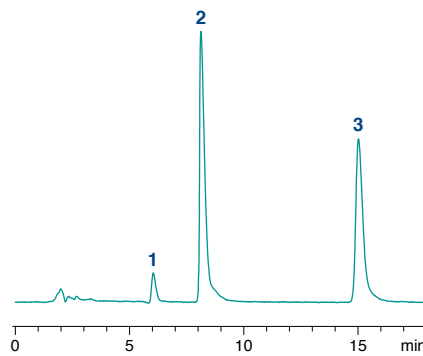
MN Appl. No. 119800

Column: 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Guard column: 8 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: acetonitrile – water (45:55, v/v), 2 mL conc. H₃PO₄/L, adjusted to pH 2.6 with NaOH
 Flow rate: 0.9 mL/min

Detection: fluorescence 273 nm and 455 nm
 Injection volume: 40 µL (7.5 ng of each substance)

Peaks:

1. β-Zearalenol
2. α-Zearalenol
3. Zearalenone



Courtesy of K.H. Ueberschär, Federal Agricultural Research Centre, Institute of Animal Feed, Celle, Germany.

Biological and natural compounds

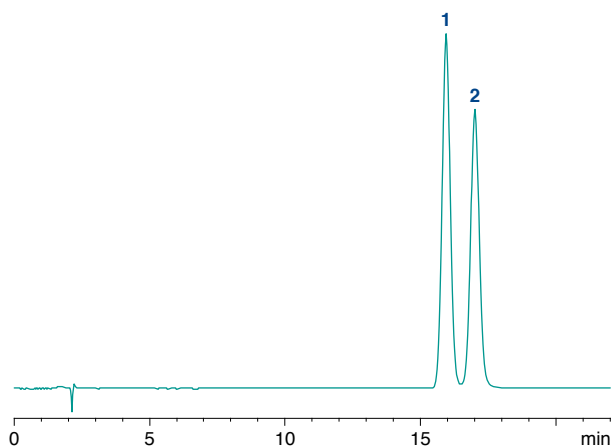
Dexa- and betamethasone

MN Appl. No. 121170

Column: 250 x 4 mm NUCLEODUR® C₁₈ Isis, 5 µm
 Eluent: acetonitrile – water (30:70, v/v)
 Flow rate: 1 mL/min
 Temperature: 25 °C
 Detection: UV, 260 nm
 Injection volume: 5 µL

Peaks:

1. Betamethasone
2. Dexamethasone



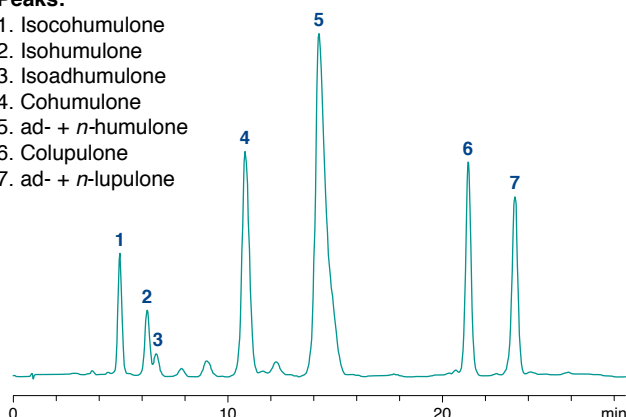
Determination of iso-alpha-acids, alpha- and beta-acids in isomerized hop pellets

MN Appl. No. 121100

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: A) methanol, B) methanol – water – H₃PO₄
 (75:24:1, v/v/v); 100 % B (17 min) → 65 % B in
 8 min → 100 % B in 5 min
 Flow rate: 1.0 mL/min
 Temperature: 35 °C
 Detection: UV, 9 min 270 nm, then 314 nm

Peaks:

1. Isocohumulone
2. Isohumulone
3. Isoadhumulone
4. Cohumulone
5. ad- + n-humulone
6. Colupulone
7. ad- + n-lupulone



M. Biendl et al., European Brewery Convention, J. of the Institute of Brewing **110** (2004) 242–243.

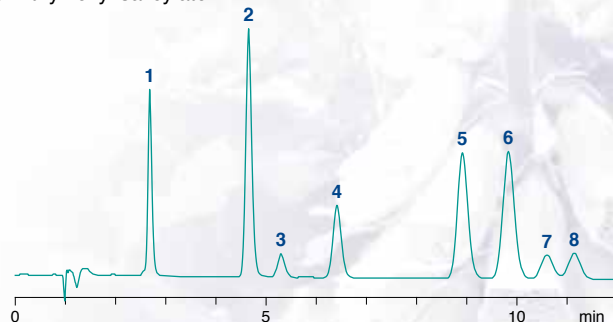
Sunscreen ingredients

MN Appl. No. 121500

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: methanol – 0.5 % H₃PO₄ (82:18, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 42 °C
 Detection: UV, 300 nm
 Injection volume: 10 µL

Peaks:

1. Benzimidazolecarboxylic acid
2. Benzophenone-3
3. 4-Methylbenzylidine camphor
4. Octocrylene
5. Ethylhexyldimethyl PABA
6. Ethylhexyl methoxycinnamate
7. Butyl methoxydibenzoylmethane (BMDBM)
8. Ethylhexyl salicylate



For separation on C₁₈ Gravity see appl. 122660 at www.mn-net.com.

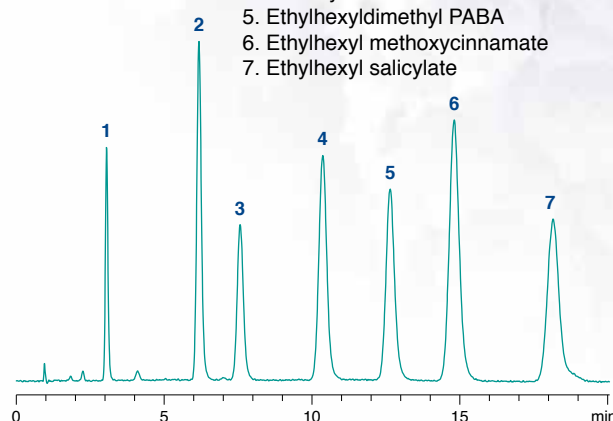
Sunscreen ingredients

MN Appl. No. 123640

Column: 125 x 4 mm NUCLEODUR® C₁₈ HTec, 5 µm
 Eluent: methanol – 100 mM ammonium acetate, pH 4.5
 (80:20, v/v)
 Flow rate: 0.9 mL/min
 Temperature: 35 °C
 Detection: UV, 275 nm
 Injection volume: 12 µL

Peaks:

1. Benzophenone
2. 4-Methylbenzylidine camphor
3. Uvinil Plus
4. Octocrylene
5. Ethylhexyldimethyl PABA
6. Ethylhexyl methoxycinnamate
7. Ethylhexyl salicylate



Applications

NUCLEODUR®

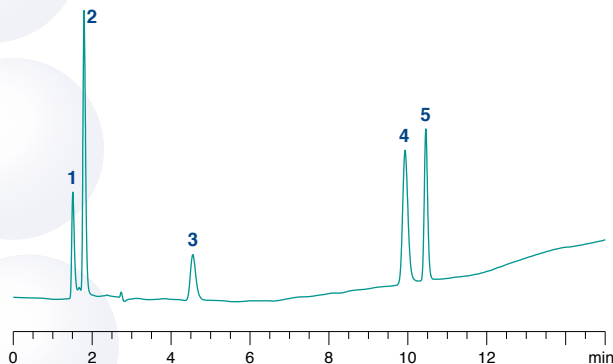
Water-soluble vitamins

MN Appl. No. 124570

Column: 150 x 3 mm NUCLEODUR® PolarTec, 5 µm
 Eluent: A) 25 mM KH₂PO₄, pH 3.0, B) acetonitrile
 10% B (3 min) → 40% B in 12 min
 Flow rate: 0.7 mL/min
 Temperature: 30 °C
 Detection: UV, 220 nm
 Injection volume: 5 µL

Peaks:

1. Vitamin B₁
2. Vitamin B₆
3. Panthothenic acid
4. p-Aminobenzoic acid
5. Vitamin B₂



Water-soluble vitamins

MN Appl. No. 119770

Column: 125 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: A) water, 15 mM heptanesulfonic acid (Na salt),
 25 mM NaH₂PO₄, 0.25% CH₃COOH, 0.005% triethylamine (pH 3.5),
 B) acetonitrile – water (40:60, v/v), 15 mM heptanesulfonic (Na salt), 0.25% CH₃COOH, 0.005% triethylamine (pH ~ 3.5); multistep gradient:
 0% B (5 min) → 10% B in 2.5 min → 25% B in 2.5 min → 50% B in 8 min → 70% B in 7 min → 0% B in 1 min

Flow rate: 1.0 mL/min

Temperature: 25 °C

Detection: UV, 254, 275 and 361 nm

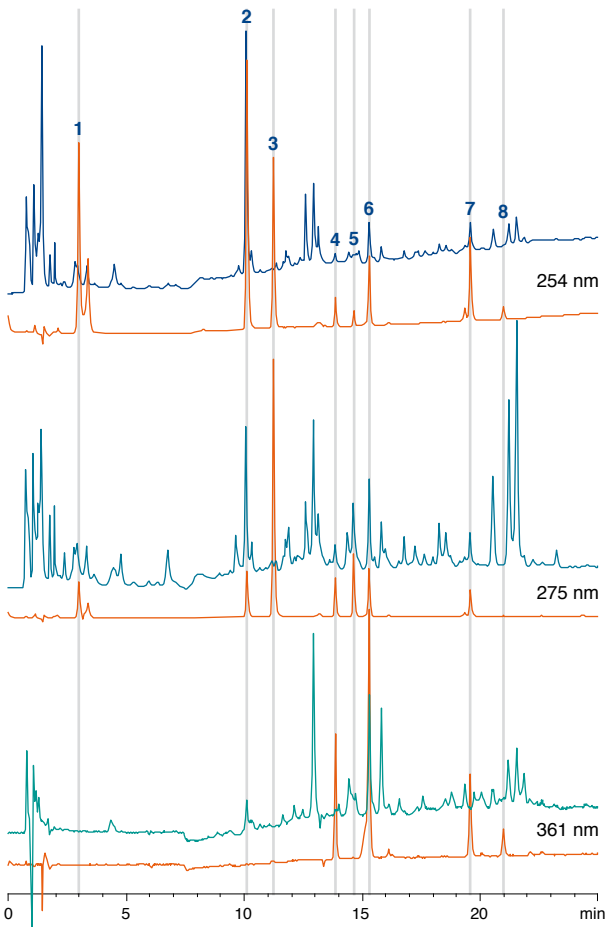
Peaks:

1. Nicotinic acid (0.12 mg/mL)
2. Nicotinamide (0.12 mg/mL)
3. 4-Aminobenzoic acid (0.03 mg/mL)
4. Folic acid (0.24 mg/mL)
5. Vitamin B₆ (pyridoxine hydrochloride, 0.06 mg/mL)
6. Vitamin B₂ (riboflavin, 0.012 mg/mL)
7. Vitamin B₁ (thiamine hydrochloride, 0.06 mg/mL)
8. Rutin (0.012 mg/mL)

orange curves: vitamin test mixture (in eluent A)

blue curves: multivitamin juice (undiluted)

both detected at three different wave lengths



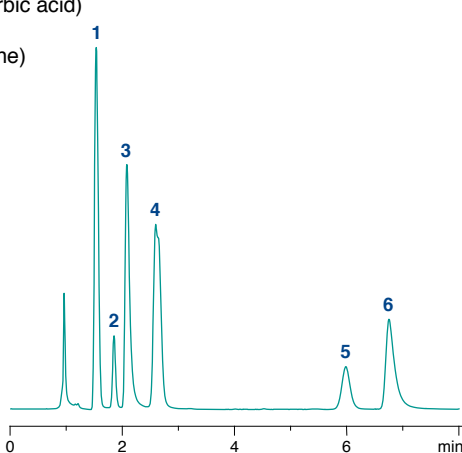
Water-soluble vitamins

MN Appl. No. 122970

Column: 125 x 4 mm NUCLEODUR® HILIC, 3 µm
 Eluent: A) acetonitrile, B) 25 mM ammonium acetate, pH 4
 80% A (1 min) → 70% A in 1 min (11 min)
 Flow rate: 1.0 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 30 µL

Peaks:

1. Nicotinamide
2. Vitamin B₇ (vitamin B₈, vitamin H, biotin)
3. Vitamin B₆ (pyridoxine)
4. Vitamin C (ascorbic acid)
5. Vitamin B₁₂ (cyanocobalamin)
6. Vitamin B₁ (thiamine)



Biological and natural compounds

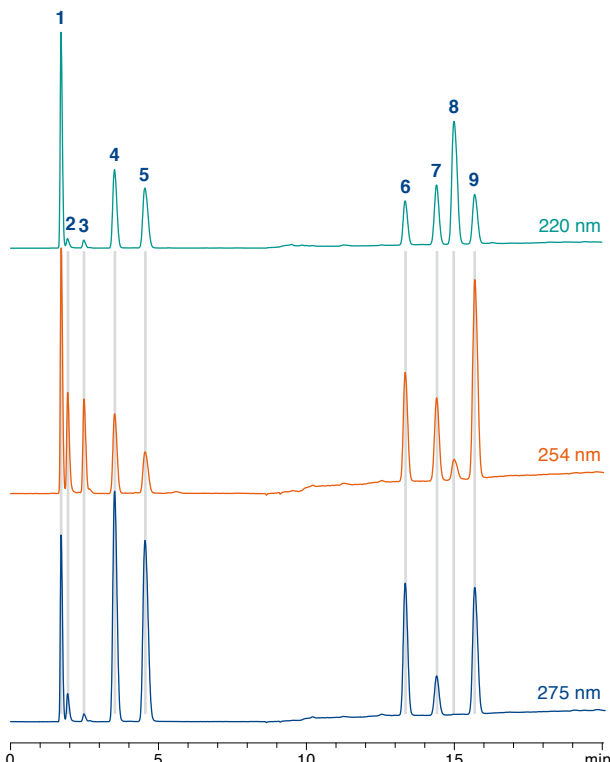
Water-soluble vitamins

MN Appl. No. 122450

Column: 125 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: A) 50 mM KH₂PO₄, pH 3,
 B) methanol – acetonitrile (70:30, v/v)
 0 % B (6 min) → 15 % B in 2 min → 35 % B in
 10 min (5 min)
 Flow rate: 0.6 mL/min
 Temperature: 40 °C
 Detection: UV, 218, 254 and 275 nm
 Injection volume: 10 µL

Peaks:

1. Vitamin B₁ (thiamine)
2. Pyridoxamine
3. Vitamin C (ascorbic acid)
4. Pyridoxal
5. Vitamin B₆ (pyridoxine)
6. Vitamin B₉ (vitamin M, folic acid)
7. Vitamin B₁₂ (cyanocobalamin)
8. Vitamin B₂ (vitamin B₈, vitamin H, (+)-biotin)
9. Vitamin B₂ (riboflavin)



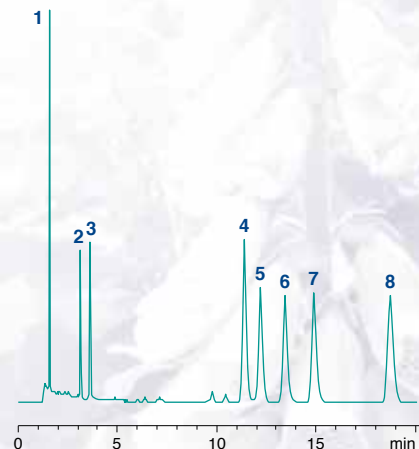
Fat-soluble vitamins and tocopherols

MN Appl. No. 117890

Column: 250 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile
 Flow rate: 1.5 mL/min
 Temperature: 30 °C
 Detection: UV, 280 nm
 Injection volume: 4 µL

Peaks:

1. Vitamin K₃
2. Vitamin A
3. Vitamin A acetate
4. Vitamin D₂
5. Vitamin D₃
6. Vitamin E
 (α-tocopherol)
7. Vitamin E acetate
 (α-tocopherol acetate)
8. Vitamin K₁

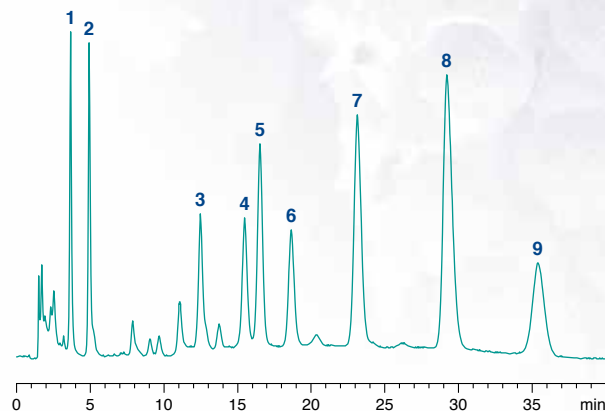


MN Appl. No. 121160

Column: 125 x 2 mm NUCLEODUR® C₁₈ Isis, 5 µm
 Eluent: acetonitrile – water (100:5, v/v)
 Flow rate: 0.2 mL/min
 Temperature: 25 °C
 Detection: UV, 275 nm
 Injection volume: 5 µL

Peaks:

- | | |
|---------------------------|---|
| 1. Vitamin A | 6. γ-Tocopherol |
| 2. Vitamin A acetate | 7. Vitamin E (α-tocopherol) |
| 3. Vitamin K ₂ | 8. Vitamin E acetate (α-tocopherol acetate) |
| 4. Vitamin D ₂ | 9. Vitamin K ₁ |
| 5. Vitamin D ₃ | |



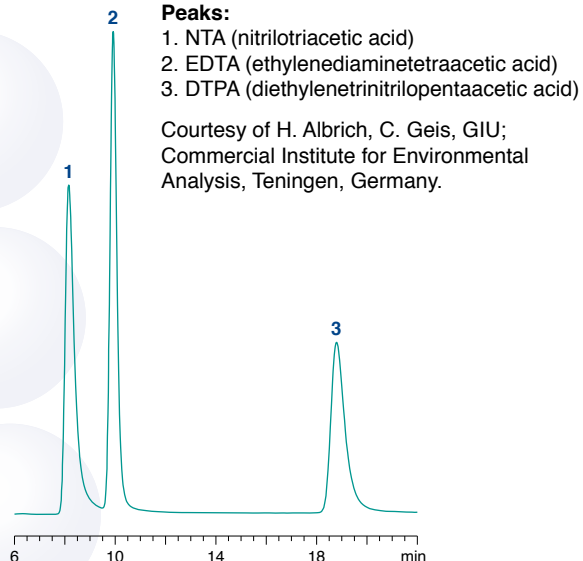
For separation of tocopherols on NUCLEODUR® 100-5 C₁₈ ec see
 appl. 117910 at www.mn-net.com.

Applications

Complexing agents acc. to DIN 38 413-8

MN Appl. No. 119780

Column: 250 x 4 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
Eluent: 0.6 mM HNO₃, 7.53 mM N(C₄H₉)₄HSO₄,
2.6 mM N(C₄H₉)₄OH, 37 µM Fe³⁺
Flow rate: 0.6 mL/min
Temperature: 20 °C
Detection: UV, 260 nm
Injection volume: 50 µL



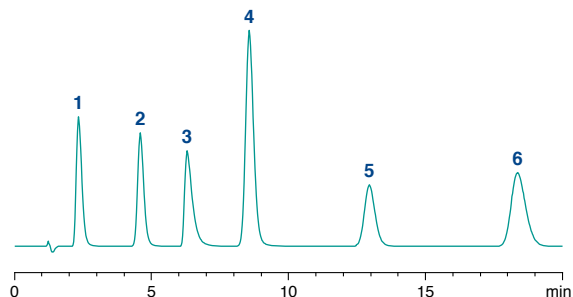
Aromatic acids

MN Appl. No. 121180

Column: 125 x 2 mm NUCLEODUR® C₁₈ Isis, 5 µm
Eluent: methanol – 50 mM KH₂PO₄, pH 3 (10:90, v/v)
Flow rate: 0.25 mL/min
Temperature: 30 °C
Detection: UV, 254 nm
Injection volume: 5 µL

Peaks:

1. Gallic acid
2. 3,4-Dihydroxybenzoic acid
3. 2,5-Dihydroxybenzoic acid
4. 4-Hydroxybenzoic acid
5. Syringic acid
6. Vanillic acid



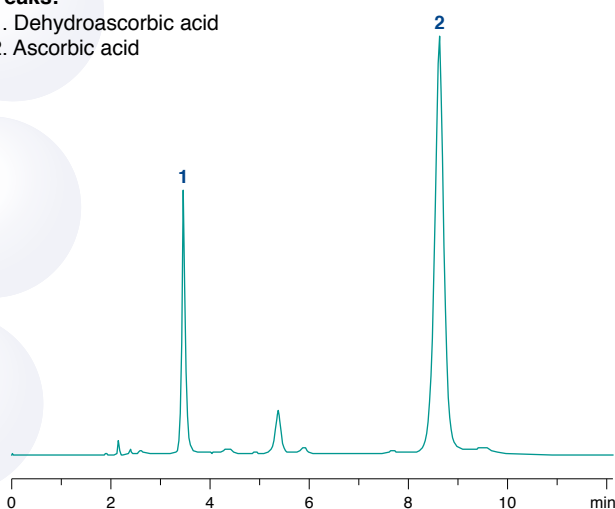
Ascorbic acid and dehydroascorbic acid

MN Appl. No. 122940

Column: 250 x 4 mm NUCLEODUR® HILIC, 5 µm
Eluent: acetonitrile – 100 mM ammonium acetate
(70:30, v/v)
Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 240 nm

Peaks:

1. Dehydroascorbic acid
2. Ascorbic acid



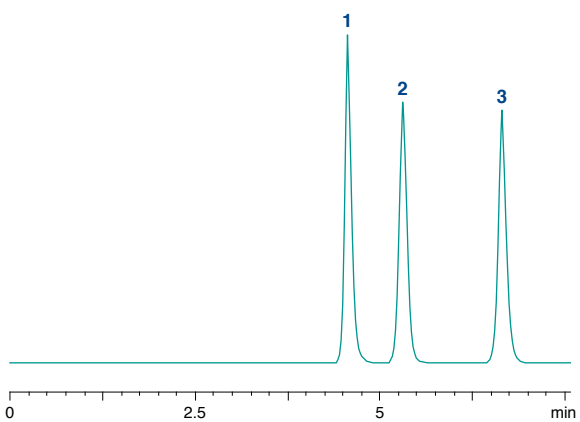
Organic acids

MN Appl. No. 119290

Column: 250 x 4 mm NUCLEODUR® 100-5 CN-RP
Eluent: 25 mM KH₂PO₄, pH 4.0
Flow rate: 0.5 mL/min
Temperature: 30 °C
Detection: UV, 210 nm
Injection volume: 15 µL

Peaks:

1. Aspartic acid
2. Fumaric acid
3. Maleic acid



Pollutants and miscellaneous organics

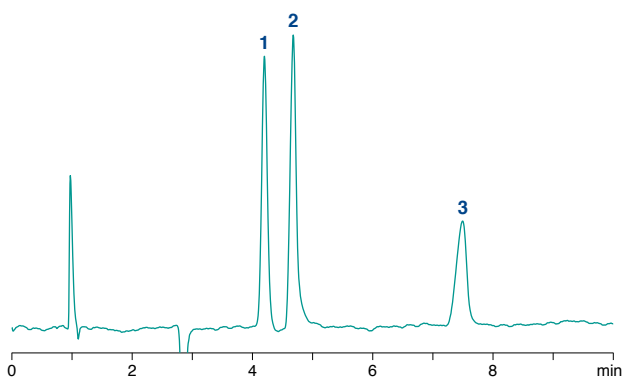
Organic acids

MN Appl. No. 122930

Column: 125 x 4 mm NUCLEODUR® HILIC, 3 µm
 Eluent: acetonitrile – 200 mM ammonium acetate, pH 6.8 (70:30, v/v)
 Flow rate: 1 mL/min
 Temperature: 25 °C
 Detection: UV, 220 nm
 Injection volume: 0.5 µL

Peaks:

1. Fumaric acid
2. Oxalic acid
3. Citric acid

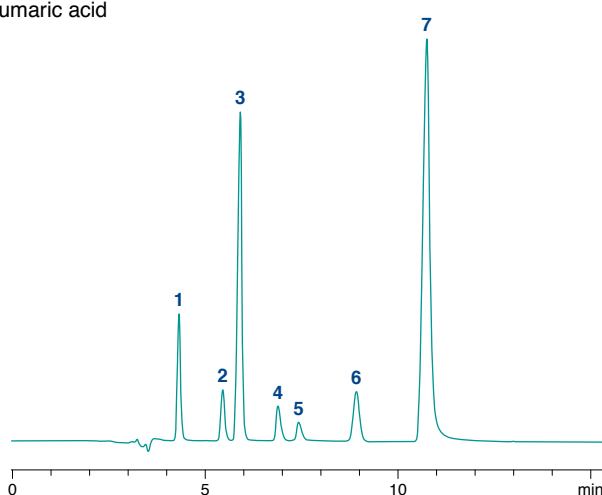


MN Appl. No. 120500

Column: 250 x 4.6 mm NUCLEODUR® C₁₈ Pyramid, 5 µm
 Eluent: 20 mM KH₂PO₄, pH 2.6
 Flow rate: 0.7 mL/min
 Detection: UV, 210 nm
 Injection volume: 20 µL

Peaks:

1. Tartaric acid
2. Malic acid
3. Shikimic acid
4. Lactic acid
5. Acetic acid
6. Citric acid
7. Fumaric acid



Also see appl. 119180 at www.mn-net.com.

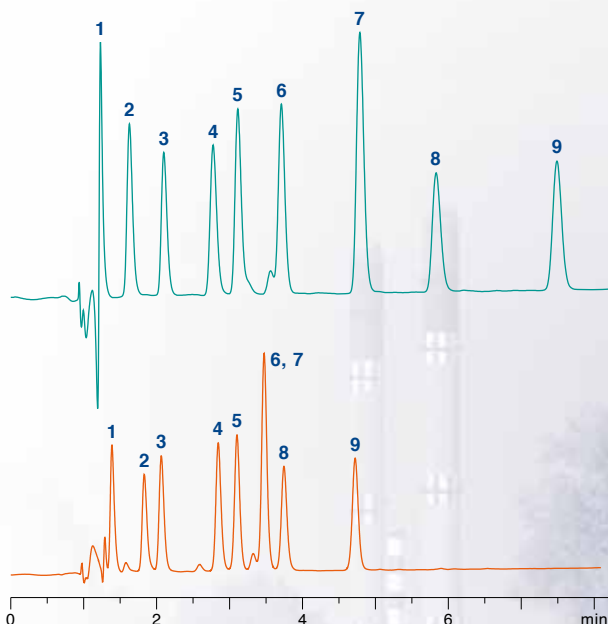
Organic acids

MN Appl. No. 124562

Columns: 150 x 3 mm NUCLEODUR® PolarTec, 5 µm
 Eluent: A) acetonitrile, 0.1% TFA, B) water, 0.1% TFA
 20 % A → 60 % A in 12 min
 Flow rate: 0.73 mL/min
 Temperature: 20 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

- | | |
|-------------------------------|------------------------------|
| 1. Dihydroxymandelic acid | 6. Vanillic acid |
| 2. Gallic acid | 7. 3-Hydroxybenzoic acid |
| 3. Dihydroxyphenylacetic acid | 8. 2,5-Dihydroxybenzoic acid |
| 4. 3,4-Dihydroxybenzoic acid | 9. 2,4-Dihydroxybenzoic acid |
| 5. Syringic acid | |



Applications

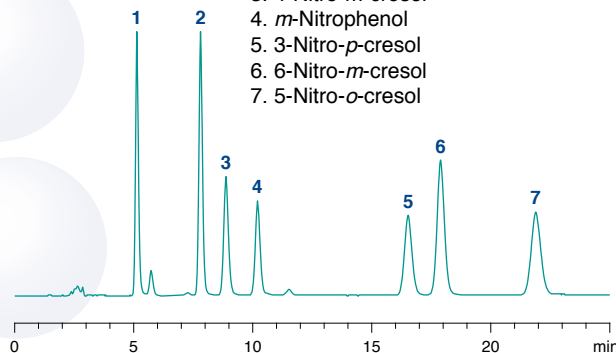
Nitrophenols

MN Appl. No. 122650

Column: 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: methanol – 20 mM NaH₂PO₄, pH 5 (49:51, v/v)
 Flow rate: 0.8 mL/min
 Temperature: 20 °C
 Detection: UV, 235 nm
 Injection volume: 2.0 µL

Peaks:

1. *p*-Nitrophenol
2. *o*-Nitrophenol
3. 4-Nitro-*m*-cresol
4. *m*-Nitrophenol
5. 3-Nitro-*p*-cresol
6. 6-Nitro-*m*-cresol
7. 5-Nitro-*o*-cresol



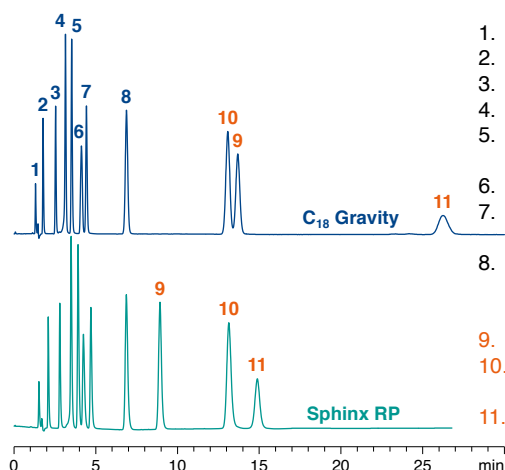
Substituted aromatics

MN Appl. No. 119840/119850

Columns: 150 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 150 x 4.6 mm NUCLEODUR® Sphinx RP, 5 µm
 Eluent: methanol – water (55:45, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 2 µL

Peaks:

1. Uracil
2. Benzamide
3. Phenol
4. Benzaldehyde
5. Aceto-phenone
6. 2-Nitrophenol
7. Nitrobenzene
8. Propyl 4-hydroxy-benzoate
9. Toluene
10. Benzo-phenone
11. Xylene



Nitroaromatics EPA 8330

MN Appl. No. 124490/124500

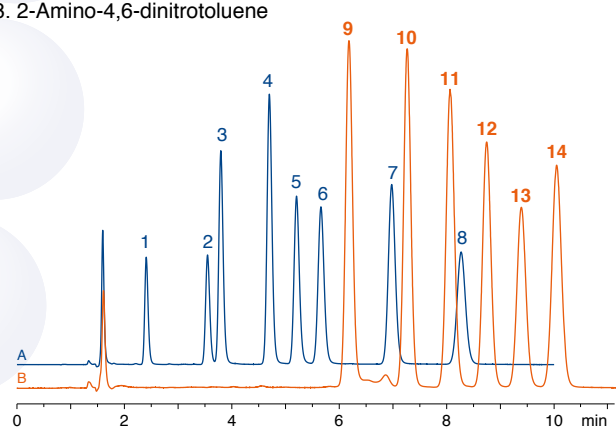
Column: 150 x 3 mm NUCLEODUR® PolarTec, 5 µm
 Eluent:
 Mix A: water – methanol (50:50, v/v)
 Mix B: water, 0.1% formic acid – methanol (45:55, v/v)
 Flow rate: 0.46 mL/min
 Temperature: 50 °C, 60 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks Mix A:

1. Octogen (HMX)
2. Hexogen (RDX)
3. 1,3,5-Trinitrobenzene
4. 1,3-Dinitrobenzene
5. Nitrobenzene
6. 2,4,6-Trinitrotoluene
7. 2,4-Dinitrotoluene
8. 2-Amino-4,6-dinitrotoluene

Peaks Mix B:

9. *N*-Methyl-*N*-2,4,6-tetranitro-aniline (Tetryl)
10. 4-Amino-2,6-dinitrotoluene
11. 2,6-Dinitrotoluene
12. 2-Nitrotoluene
13. 4-Nitrotoluene
14. 3-Nitrotoluene



For separation of Mix A and Mix B in a single run see appl. 124510 at www.mn-net.com.

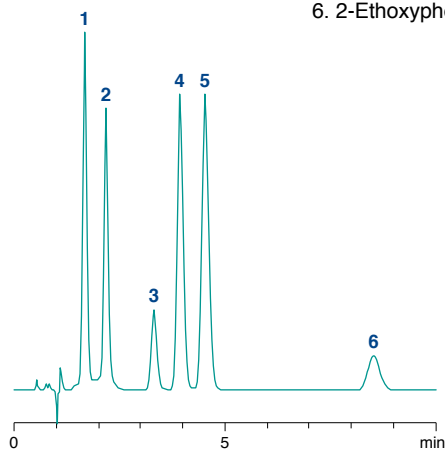
Phenolic compounds

MN Appl. No. 117970

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: methanol – water, 0.1% H₃PO₄ (40:60, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 22 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

1. Resorcinol
2. Pyrocatechol
3. 4-Methoxyphenol
4. Phenol
5. 2-Methoxyphenol
6. 2-Ethoxyphenol



Pollutants and miscellaneous organics

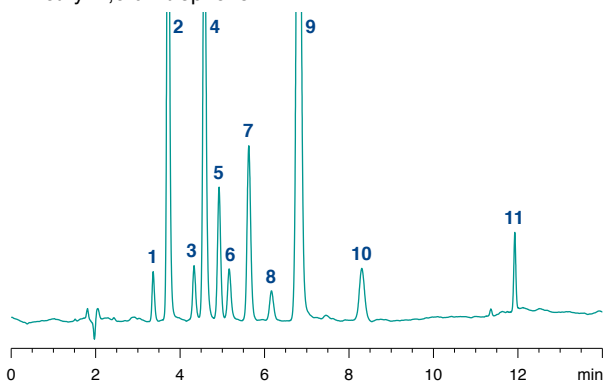
Phenolic compounds

MN Appl. No. 124740

Column: 250 x 3 mm NUCLEODUR® PFP, 5 µm
 Eluent: A) acetonitrile + 1 % acetic acid
 B) water + 1 % acetic acid
 Gradient: 45 % A (7 min) → 70 % A in 2 min (5 min)
 Flow rate: 0.6 mL/min
 Temperature: 45 °C
 Detection: UV, 254 nm
 Injection volume: 1 µL

Peaks:

1. Phenol
2. 4-Nitrophenol
3. 2-Nitrophenol
4. 2,4-Dinitrophenol
5. 2-Chlorophenol
6. 2-Methyl-4,6-dinitrophenol
7. 4-Chloro-3-methylphenol
8. 2,4-Dichlorophenol
9. 2,4-Dimethylphenol
10. 2,4,6-Trichlorophenol
11. Pentachlorophenol



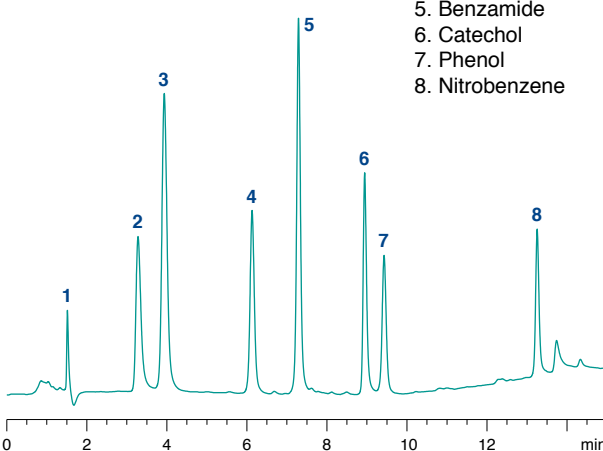
Phenolic compounds

MN Appl. No. 124750

Column: 100 x 2 mm NUCLEODUR® PFP, 3 µm
 Eluent: A) methanol, B) 10 mM ammonium acetate, pH 6.8
 5 % A → 80 % A in 15 min
 Flow rate: 0.25 mL/min
 Temperature: 18 °C
 Detection: UV, 230 nm
 Injection volume: 1 µL

Peaks:

1. Uracil
2. Pyrogallol
3. Phloroglucinol
4. Resorcinol
5. Benzamide
6. Catechol
7. Phenol
8. Nitrobenzene



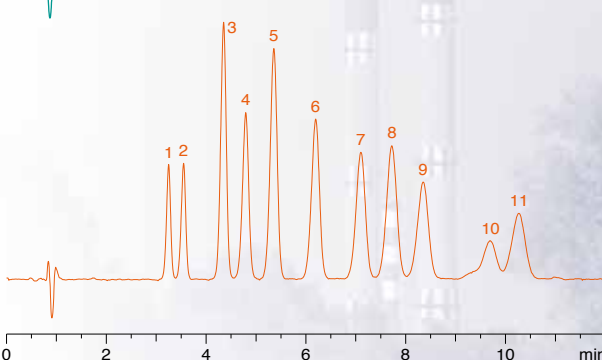
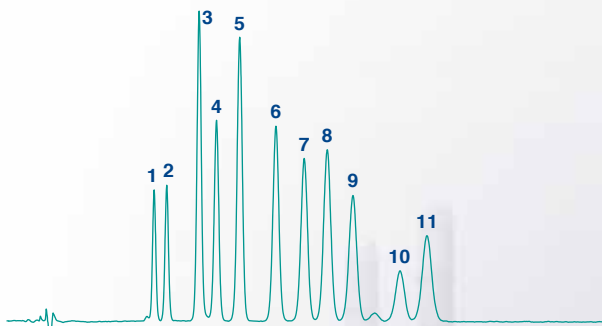
Separation of phenol isomers

MN Appl. No. 124541

Columns: 100 x 4.6 mm NUCLEODUR® PFP, 5 µm
 100 x 4.6 mm Phenomenex Luna® PFP(2), 5 µm
 Eluent: acetonitrile, 0.1 % formic acid – water, 0.1 %
 formic acid (35:65, v/v)
 Flow rate: 1.3 mL/min
 Temperature: 35 °C
 Detection: UV, 280 nm

Peaks:

1. *o*-Cresol
2. *m*-Cresol
3. 3,4-Dimethylphenol
4. 3,5-Dimethylphenol
5. 2,5-Dimethylphenol
6. 2,6-Dichlorophenol
7. 2,3-Dichlorophenol
8. 2,4-Dichlorophenol
9. 3,4-Dichlorophenol
10. 2,4-Dibromophenol
11. 3,5-Dibromophenol



NUCLEODUR® PFP provides under identical conditions a better separation than Luna® PFP(2). While on Luna® PFP(2) for peaks 10 and 11 only a resolution of 1.27 is obtained, on NUCLEODUR® PFP the peaks are baseline separated ($R_S = 1.56$).

Applications

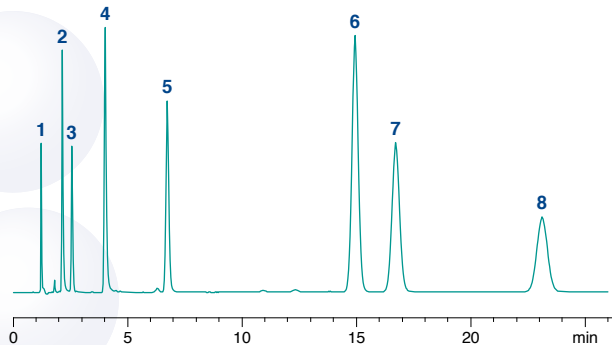
Amines

MN Appl. No. 121200

Column: 150 x 4.6 mm NUCLEODUR® C₁₈ Isis, 5 µm
 Eluent: acetonitrile – 50 mM K₂HPO₄ (40:60, v/v), pH 8
 Flow rate: 1 mL/min
 Temperature: 25 °C
 Detection: UV, 254 nm
 Injection volume: 8 µL

Peaks:

1. Uracil
2. Pyridine
3. Desethylatrazine
4. 4-Acetylpyridine
5. 4-Ethylaniline
6. N,N-Dimethylaniline
7. 4-Aminoanthraquinone
8. 3,5-Dinitro-(1-phenylethylbenzamide)



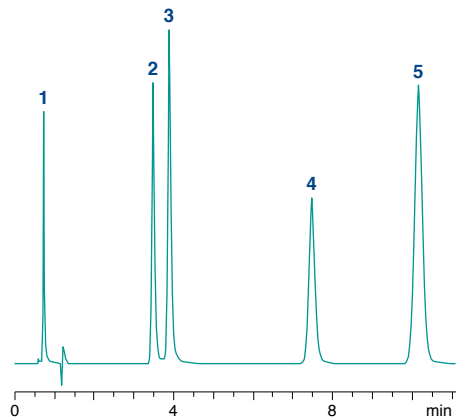
Aromatic aldehydes

MN Appl. No. 117990

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile – water, pH 6.0 (22:78, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 22 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL (~10–50 µg/mL)

Peaks:

1. p-Carboxybenzaldehyde
2. p-Hydroxybenzaldehyde
3. Vanillin
4. 4-Ethoxyvanillin
5. Benzaldehyde



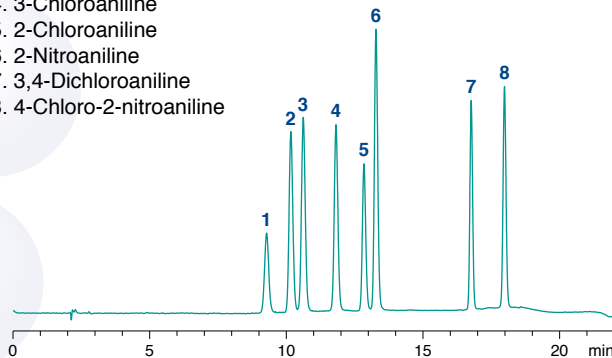
Aromatic amines

MN Appl. No. 124800

Column: 250 x 4 mm NUCLEODUR® PFP, 5 µm
 Eluent: A) methanol, B) 20 mM K₂HPO₄, pH 3
 40 % A (5 min) → 70 % A in 10 min (7 min)
 Flow rate: 1 mL/min
 Temperature: 20 °C
 Detection: UV, 254 nm
 Injection volume: 1 µL

Peaks:

1. 4-Chloroaniline
2. 4-Nitroaniline
3. 3-Nitroaniline
4. 3-Chloroaniline
5. 2-Chloroaniline
6. 2-Nitroaniline
7. 3,4-Dichloroaniline
8. 4-Chloro-2-nitroaniline



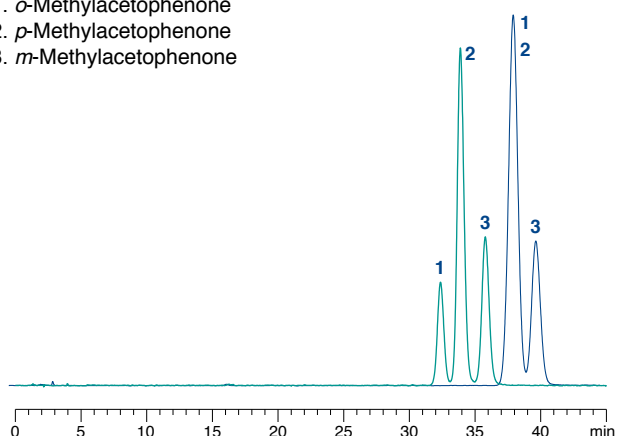
Aromatic ketones

MN Appl. No. 124761

Columns: 250 x 4 mm NUCLEODUR® PFP, 5 µm
 250 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: methanol – water (35:65, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 35 °C
 Detection: UV, 254 nm
 Injection volume: 1 µL

Peaks:

1. o-Methylacetophenone
2. p-Methylacetophenone
3. m-Methylacetophenone



Distinct steric selectivity of NUCLEODUR® PFP provides separation of all three regioisomers, while on NUCLEODUR® C₁₈ Gravity baseline separation can't be achieved.

Pollutants and miscellaneous organics

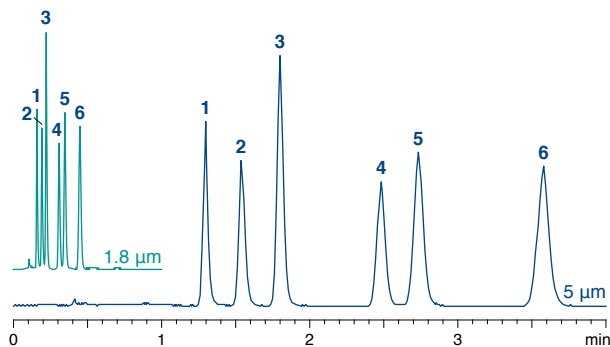
Aromatic ketones

MN Appl. No. 122720/122730

Column: 125 x 2 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Eluent: 50 x 2 mm NUCLEODUR® C₁₈ Gravity, 1.8 µm acetonitrile – water (60:40, v/v)
 Flow rate: 0.33 mL/min, 1.25 mL/min
 Temperature: 25 °C
 Detection: UV, 230 nm

Peaks:

1. Acetophenone
2. Eugenol
3. Propiophenone
4. Butyrophenone
5. Benzophenone
6. Valerophenone

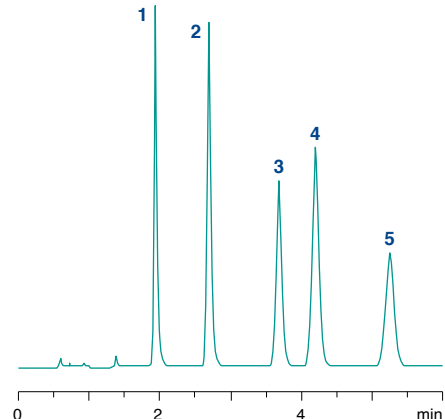


MN Appl. No. 117980

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Eluent: acetonitrile – water (60:40, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 22 °C
 Detection: UV, 230 nm
 Injection volume: 2 µL (~10–50 µg/mL)

Peaks:

1. Acetophenone
2. Propiophenone
3. Butyrophenone
4. Benzophenone
5. Valerophenone



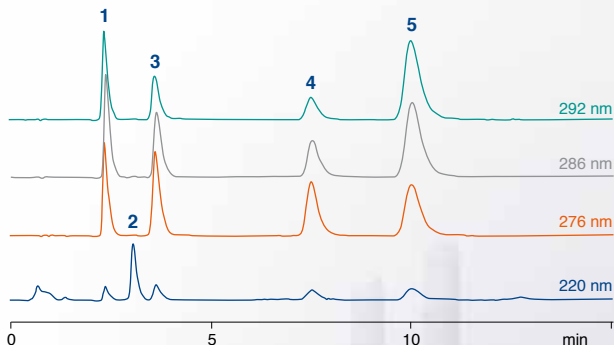
Furfurol and related compounds in transformer oil in accordance with DIN EN 61198-B

MN Appl. No. 121662

Column: 125 x 4 mm NUCLEODUR® 100-5 C₁₈ ec
 Sample prep.: SPE see appl. 304180 at www.mn-net.com
 Eluent: methanol – water (10:90, v/v)
 Flow rate: 2.0 mL/min
 Detection: UV, 220, 276, 286 and 292 nm

Peaks:

1. 5-Hydroxymethyl-2-furfurol
2. 2-Furfuryl alcohol
3. 2-Furfurol
4. 2-Acetyl furan
5. 5-Methyl-2-furfurol



A. Heiseler et al., GIT (2004) 504–505

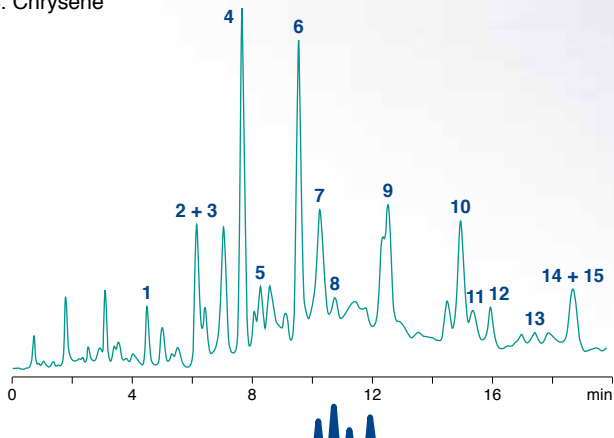
PAHs from tar

MN Appl. No. 120740

Column: 150 x 4.6 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Sample prep.: see Appl. 120740 at www.mn-net.com
 Eluent: A) water, B) acetonitrile
 60 % B (3 min) → 100 % B in 27 min
 Flow rate: 1.0 mL/min
 Detection: UV, 220 nm

Peaks:

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



Applications

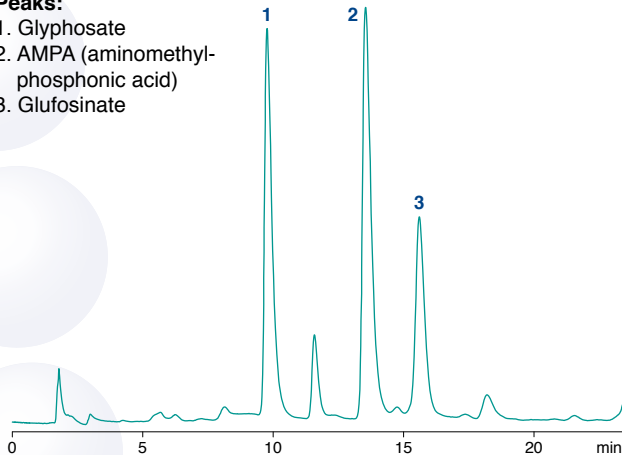
Organophosphorus herbicides

MN Appl. No. 120490

Column: 250 x 3 mm NUCLEODUR® C₁₈ Gravity, 5 µm
 Sample prep.: for SPE see Appl. 303780 at www.mn-net.com; derivatization with FMOC-Cl
 Eluent: A) acetonitrile, B) H₃PO₄, pH 1.2
 30 % A → 35 % A in 27 min → 90 % A in 3 min (6 min) → 30 % A in 2 min (7 min)
 Flow rate: 0.5 mL/min
 Temperature: 30 °C
 Detection: fluorescence, λ_{ex} 263 nm, λ_{em} 317 nm

Peaks:

1. Glyphosate
2. AMPA (aminomethyl-phosphonic acid)
3. Glufosinate



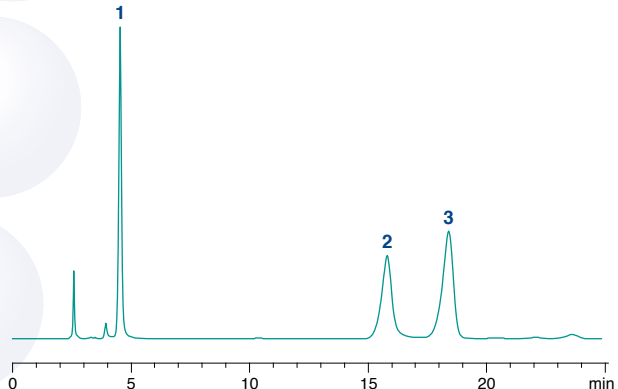
Courtesy of Mr. Schüssler, Mrs. Mikler, Bavarian State Agency for Water Management, Munich.

MN Appl. No. 122190

Column: 250 x 4 mm NUCLEODUR® 100-5 NH₂-RP
 Sample prep.: derivatization with FMOC, concentration of each pesticide 0.3 mg/mL
 Eluent: acetonitrile – 50 mM KH₂PO₄, pH 4.6 (60:40, v/v)
 Flow rate: 0.8 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 5 µL

Peaks:

1. AMPA
2. Glyphosate
3. Glufosinate



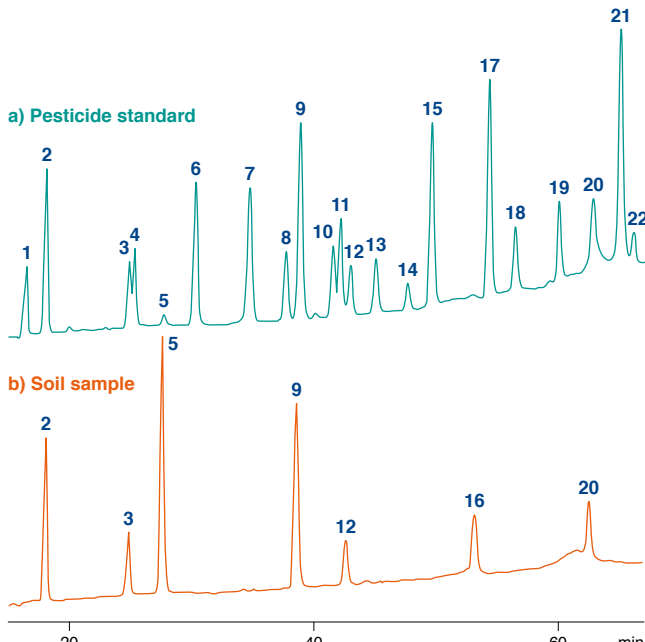
Pesticides from soil

MN Appl. No. 119890

Column: 250 x 4 mm NUCLEODUR® 100-3 C₈ ec
 Eluent: A) water, B) acetonitrile,
 10 % B → 25 % B in 10 min → 30 % B in 10 min (5 min) → 40 % B in 20 min → 50 % B in 20 min (10 min)
 Flow rate: 0.8 mL/min
 Temperature: 35 °C
 Detection: UV, 230 nm

Peaks:

1. Metamitron
2. Desethylatrazine
3. Hexazinone
4. Metoxuron
5. Simazine
6. Cyanazine
7. Methabenzthiazuron (Tribunil®)
8. Chlortoluron
9. Atrazine
10. Monolinuron
11. Isoproturon
12. Diuron
13. Metobromuron
14. Metazachlor
15. Sebutylazine
16. Dichlobenil
17. Terbutylazine
18. Linuron
19. Chloroxuron
20. Propyzamid
21. Terbutryn
22. Metolachlor



Courtesy of E. Marek, LUFA Center for Analyses, Münster, Germany.

Also see appl. 118010 at www.mn-net.com.

Pollutants and miscellaneous organics

Pesticides

MN Appl. No. 120481: triazines

MN Appl. No. 120482: phenylurea derivatives

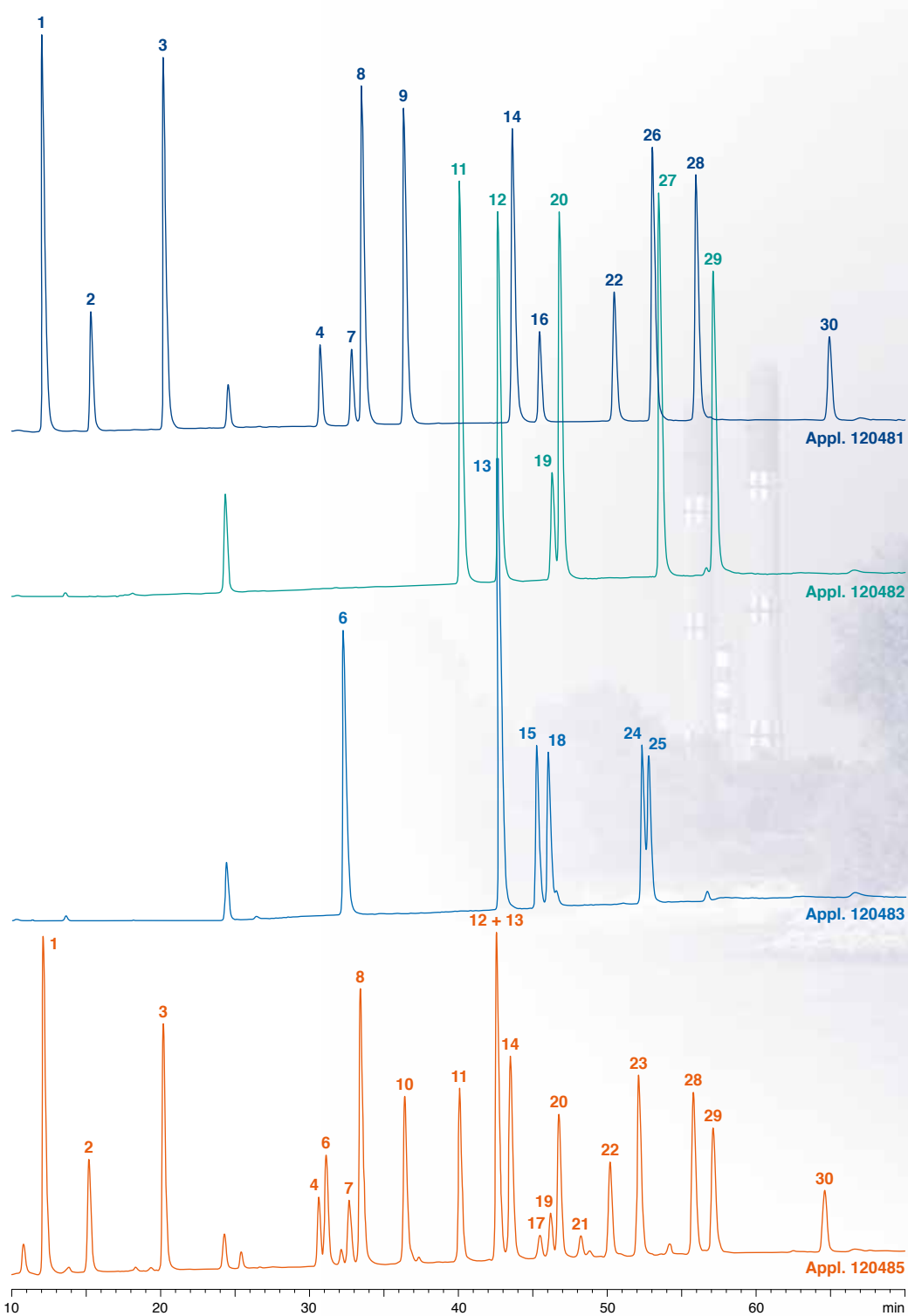
MN Appl. No. 120483: phenoxycarboxylic acids

MN Appl. No. 120485: 21 pesticides

Column: 250 x 4 mm NUCLEODUR® 100-3 C₈ ec with 8 x 4 mm guard column
 Eluent: A) acetonitrile, B) 20 mM KH₂PO₄ + 1 mL conc. H₃PO₄, 85 % B (5 min) → 47 % B in 60 min
 Flow rate: 0.7 mL/min
 Temperature: 35 °C
 Detection: UV, 218 nm
 Injection volume: 50 µL

Peaks:

1. Desisopropylatrazine
2. 2,4-Dichlorobenzamide
3. Desethylatrazine
4. Hexazinone
5. Metoxuron
6. Dicamba
7. Bromacil
8. Simazine
9. Desethylterbutylazine
10. Cyanazine
11. Methabenzthiazuron
12. Chlortoluron
13. Bentazone
14. Atrazine
15. 2,4-D
16. Metalaxyl
17. Monolinuron
18. MCPA
19. Isoproturon
20. Diuron
21. Metobromuron
22. Metazachlor
23. Sebutylazine
24. Dichlorprop
25. Mecoprop
26. Propazine
27. Dimefuron
28. Terbutylazine
29. Linuron
30. Metolachlor



Courtesy of C. Geis, GIU;
 Commercial Institute for
 Environmental Analysis,
 Teningen, Germany.

Applications

NUCLEODUR®

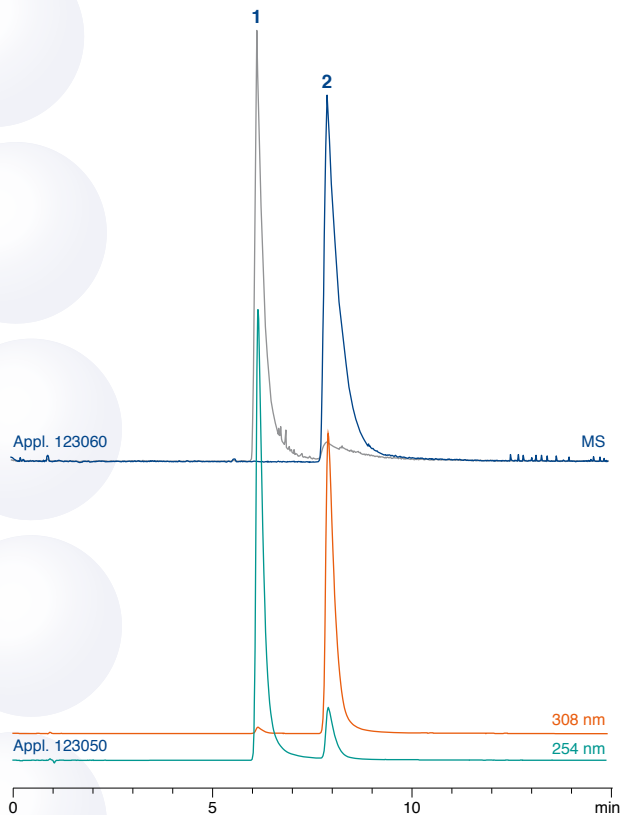
Herbicides

MN Appl. No. 123050/123060

Column: 125 x 2 mm NUCLEODUR® HILIC, 3 µm
 Eluent: acetonitrile – 50 mM ammonium formate, pH 3.2 (80:20, v/v)
 Flow rate: 0.3 mL/min
 Temperature: 45 °C
 Detection: UV, 254 and 308 nm; MS
 Injection volume: 1 µL, 0.5 mg/mL

Peaks:

1. Paraquat
2. Diquat



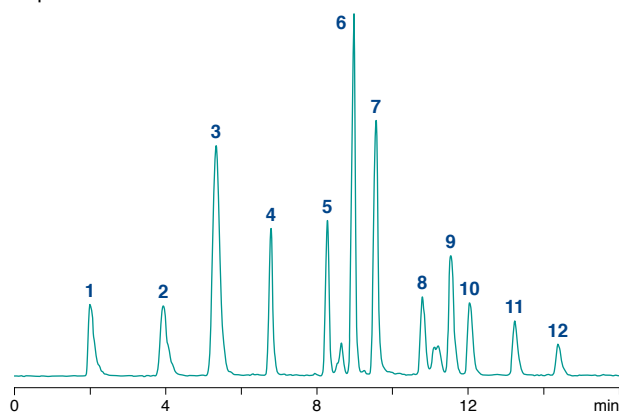
Perfluorinated surfactants in water

MN Appl. No. 121590

Column: 125 x 2 mm NUCLEODUR® Sphinx RP, 3 µm
 Sample prep.: see appl. 121590 at www.mn-net.com
 Eluent: A) 10 mM NH₄ acetate in water – methanol (75:25, v/v); B) 10 mM NH₄ acetate in acetonitrile – methanol (75:25, v/v); 10 % B → 30 % B in 3 min → 55 % B in 8 min → 70 % B in 4 min
 Flow rate: 0.3 mL/min; temperature 50 °C
 Detection: LC-MS-MS; injection volume 50 µL

Peaks:

1. Perfluorobutanoic acid
2. Perfluoropentanoic acid
3. K perfluorobutanesulfonate
4. Perfluorohexanoic acid
5. Perfluoroheptanoic acid
6. K perfluorohexansulfonate
7. Perfluoroctanoic acid
8. Perfluorononanoic acid
9. K perfluooctanesulfonate
10. Perfluorodecanoic acid
11. Perfluoroundecanoic acid
12. Perfluorododecanoic acid



D. Skutlarek et al., Environ Sci Pollut Res 13 (2006) 299–307

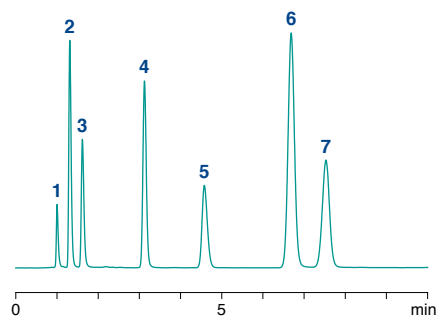
Selectivity test

MN Appl. No. 119880

Column: 125 x 4 mm NUCLEODUR® Sphinx RP, 5 µm
 Eluent: methanol – 25 mM NH₄H₂PO₄, pH 7 (65:35, v/v)
 Flow rate: 1.0 mL/min
 Temperature: 40 °C
 Detection: UV, 254 nm
 Injection volume: 6 µL

Peaks:

1. Uracil
2. 2,7-Dihydroxynaphthalene
3. 2,3-Dihydroxynaphthalene
4. Ethyl benzoate
5. Lidocaine
6. Biphenyl
7. Acenaphthene



Pollutants and miscellaneous organics

Test for metal ions in silica adsorbent

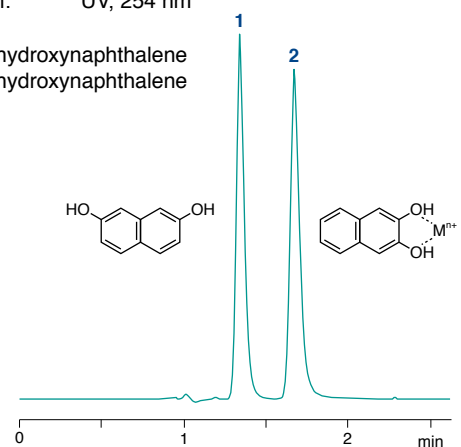
MN Appl. No. 118630

Column: 125 x 4 mm NUCLEODUR® C₈ Gravity, 5 µm
Eluent: methanol – 20 mM KH₂PO₄, pH 7 (65:35, v/v)

Flow rate: 1.0 mL/min
Temperature: 25 °C
Detection: UV, 254 nm

Peaks:

1. 2,7-Dihydroxynaphthalene
2. 2,3-Dihydroxynaphthalene



The ratio of the asymmetry factors of 2,3-dihydroxynaphthalene (2) and 2,7-dihydroxynaphthalene (1) is a measure for the metal ion content of the silica phase, because (2) can form complexes with metal ions, resulting in broad peaks for this compound.

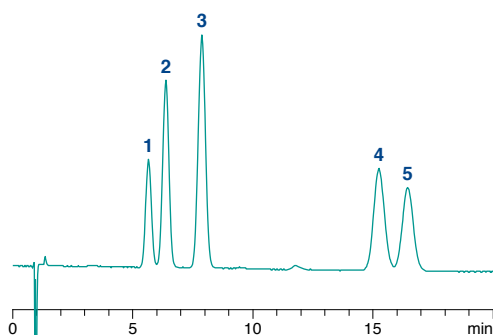
Dihydroxynaphthalenes

MN Appl. No. 121190

Column: 125 x 4 mm NUCLEODUR® C₁₈ Isis, 5 µm
Eluent: methanol – 0.5 % H₃PO₄ (30:70, v/v)
Flow rate: 1 mL/min
Temperature: 30 °C
Detection: UV, 254 nm
Injection volume: 15 µL

Peaks:

1. 1,5-Dihydroxynaphthalene
2. 1,6-Dihydroxynaphthalene
3. 2,7-Dihydroxynaphthalene
4. 1,3-Dihydroxynaphthalene
5. 2,3-Dihydroxynaphthalene



Substance index

A

Acenaphthene	C ₁₈ Gravity, 5 µm C ₁₈ Pyramid, C ₁₈ Gravity, C ₈ Gravity	9 15
	Sphinx RP, 5 µm	62
Acenaphthylene	C ₁₈ Gravity, 5 µm	59
Acesulfame K	100-5 C ₈ ec	43
	C ₁₈ HTec, 5 µm	43
2-Aacetamidophen	C ₈ Gravity, 5 µm	32
Acetamidophenol isomers	C ₁₈ HTec, 5 µm	23
Acetaminophen see Paracetamol		
Acetanilide	C ₈ Gravity, 5 µm	32
Acetic acid	C ₁₈ Pyramid, 5 µm	14, 15, 55
Acetophenone	100-5 C ₁₈ ec	59
	C ₁₈ Gravity, 1.8 vs. 3 µm	59
	C ₁₈ Gravity, Sphinx RP, 5 µm	56
2-Acetyl furan	100-5 C ₁₈ ec	59
4-Acetylpyridine	C ₁₈ Isis, 5 µm	58
Acetylsalicylic acid	100-5 C ₈ ec	33
	100-5 C ₁₈ ec	32, 34
	C ₈ Gravity, 5 µm	32
	C ₁₈ Gravity, 5 µm	34
	C ₁₈ HTec, 5 µm	23
	C ₁₈ Pyramid, 5 µm	32
Acrylamide	HILIC, 5 µm	44
Adenine	100-5 NH ₂ -RP	31
	C ₁₈ Pyramid, 5 µm	47
	HILIC, 5 µm	27, 47
	PolarTec, 3 µm	17
	PolarTec, 5 µm	47
Adhumulone, adlupulone	100-5 C ₁₈ ec	51
Adrenaline	C ₁₈ Gravity, 5 µm	50
	HILIC, 3 µm	50
Aflatoxins	100-5 C ₁₈ ec	50
Alanine	100-5 C ₁₈ ec	46
4-Amino-2,6-dinitrotoluene	PolarTec, 5 µm	56
2-Amino-4,6-dinitrotoluene	PolarTec, 5 µm	56
4-Aminoanthraquinone	C ₁₈ Isis, 5 µm	58
4-Aminobenzoic acid	C ₁₈ Pyramid, 5 µm	52
p-Aminobenzoic acid	PolarTec, 5 µm	52
γ-Aminobutyric acid	100-5 C ₁₈ ec	46
Aminomethylphosphonic acid see AMPA		
Amitriptyline	C ₈ Gravity, C ₁₈ Isis, C ₁₈ Pyramid	35
	C ₁₈ Gravity, 5 µm	9, 35
	PolarTec, 5 µm	34
Amoxicillin	C ₁₈ Gravity, 5 µm	42
	C ₁₈ HTec, 5 µm	41
	C ₁₈ Pyramid, 5 µm	40
	PFP, 5 µm	41
AMPA	100-5 NH ₂ -RP	31, 60
	C ₁₈ Gravity, 5 µm	60
Ampicillin	PFP, 5 µm	41
Aniline	C ₁₈ HTec, 5 µm	22
p-Anisic acid	PolarTec, 3 µm	43
Anthracene	C ₁₈ Gravity, 5 µm	59
	C ₁₈ HTec, 5 µm; EC vs. VP	23
Arginine	100-5 C ₁₈ ec	46
	C ₁₈ Gravity, 3 µm	46
Ascorbic acid	100-5 C ₈ ec	43
	C ₁₈ Pyramid, 5 µm	36, 53
	HILIC, 3 µm	52
	HILIC, 5 µm	54
Asparagine	100-5 C ₁₈ ec	46
Aspartame	100-5 C ₈ ec	43
	C ₁₈ HTec, 5 µm	43
Aspartic acid	100-5 C ₁₈ ec	46
	100-5 CN-RP	54
Atrazine	100-3 C ₈ ec	60, 61

B

Atropine	C ₈ Gravity, 5 µm	48
Azorubine	C ₁₈ Gravity, 5 µm	44
Bambuterol	C ₁₈ Pyramid, 5 µm	37
Beclometasone dipropionate	C ₁₈ Pyramid, 1.8 µm	33
Bentazone	100-3 C ₈ ec	61
Benzaldehyde	100-5 C ₁₈ ec	58
	C ₁₈ Gravity, Sphinx RP, 5 µm	56
Benzalkonium chlorides	100-5 CN-RP	37
Benzamide	100-5 CN-RP	28, 29
	C ₁₈ Gravity, 5 µm	10, 15, 56
	C ₁₈ Pyramid, C ₈ Gravity, 5 µm	15
	PFP, 3 µm	57
	Sphinx RP, 5 µm	56
Benz[a]anthracene	C ₁₈ Gravity, 5 µm	59
Benzocaine	C ₁₈ Pyramid, 5 µm	32
Benzo[b]fluoranthene	C ₁₈ Gravity, 5 µm	59
Benzo[k]fluoranthene	C ₁₈ Gravity, 5 µm	59
Benzoic acid	100-5 C ₈ ec	43
	C ₁₈ Gravity, 5 µm	36
	C ₁₈ HTec, 5 µm	43
	PolarTec, 3 µm	43
Benzo[ghi]perylene	C ₁₈ Gravity, 5 µm	59
Benzophenone	100-5 C ₁₈ ec	59
	C ₁₈ Gravity, 1.8 vs. 3 µm	59
	C ₁₈ Gravity, 5 µm	15, 56
	C ₁₈ HTec, 5 µm	51
	C ₁₈ Pyramid, C ₈ Gravity, 5 µm	15
	Sphinx RP, 5 µm	56
Benzophenone-3	100-5 C ₁₈ ec	51
Benzo[a]pyrene	C ₁₈ Gravity, 5 µm	59
Benzyl alcohol	PolarTec, 3 µm	43
Betamethasone	C ₁₈ Isis, 5 µm	51
Biotin	PFP, 5 µm	19
	C ₁₈ Pyramid, 5 µm	53
Biphenyl	HILIC, 3 µm	52
	100-5 CN-RP	28, 29
	C ₁₈ Pyramid, C ₁₈ Gravity, C ₈ Gravity	15
	Sphinx RP, 5 µm	62
Biphenyl-2-ol	100-5 C ₈ ec / C ₁₈ ec	25
Biuret	C ₁₈ Pyramid, 5 µm	45
Bromacil	100-3 C ₈ ec	61
Bromazepam	100-5 C ₁₈ ec	38
Brompheniramine	100-5 C ₁₈ ec / 100-5 CN-RP	28
	PFP, 5 µm	37
	PFP, C ₁₈ Gravity, 5 µm	19
Brucine	C ₈ Gravity, 5 µm	48
Butacaine	C ₁₈ Pyramid, 5 µm	32
Butyl methoxydibenzoylmethane	100-5 C ₁₈ ec	51
	PolarTec, 3 µm	43
Butyl paraben	100-5 C ₁₈ ec	59
Butyrophenone	C ₁₈ Gravity, 1.8 vs. 3 µm	59
C		
Caffeine	100-5 C ₈ ec	43
	100-5 C ₁₈ ec	24
	C ₈ Gravity, 5 µm	32
	C ₁₈ Gravity, 5 µm	11
	C ₁₈ Isis, 5 µm	13
	C ₁₈ Pyramid, 5 µm	36
	Sphinx RP, 5 µm	20, 42
4-Carboxybenzaldehyde	100-5 C ₁₈ ec	58
Carprofen	100-5 C ₈ ec / C ₁₈ ec	33
Catechin	Sphinx RP, C ₁₈ Gravity, C ₈ Gravity	21
Catechol	PFP, 3 µm	57

Alphabetical index of analytes

Cefalotin	PFP, 5 µm	41	Cytosine	100-5 NH ₂ -RP	31
Cefamandole	C ₁₈ Gravity, PFP, 5 µm	41		C ₁₈ Pyramid, 5 µm	47
Cefotaxime	C ₁₈ Gravity, PFP, 5 µm	41		HILIC, 5 µm	27, 47
Cefoxitin	C ₁₈ Gravity, PFP, 5 µm	41		PolarTec, 3 µm	17
Cephalexin	PFP, 5 µm	41		PolarTec, 5 µm	47
Cephalothin	C ₁₈ Gravity, 5 µm	41			
Cetirizine	PFP, 5 µm	37			
	PFP, C ₁₈ Gravity, 5 µm	19	D		
Chloramphenicol	C ₁₈ Gravity, 5 µm	42	2,4-D	100-3 C ₈ ec	61
Chlormequat	HILIC, 3 µm	27	Dehydroacetic acid	C ₁₈ HTec, 5 µm	43
4-Chloro-2-nitroaniline	PFP, 5 µm	58	Dehydroascorbic acid	HILIC, 5 µm	54
4-Chloro-3-methylphenol	PFP, 5 µm	57	Deoxycorticosterone	C ₁₈ Gravity, 5 µm	49
Chloroaniline isomers	PFP, 5 µm	58	Desethylatrazine	100-3 C ₈ ec	60, 61
Chlorocresol	C ₁₈ Pyramid, 1.8 µm	33		C ₁₈ Isis, 5 µm	58
2-Chlorophenol	PFP, 5 µm	57	Desethylterbutylazine	100-3 C ₈ ec	61
Chloroprothixene hydrochloride			Desisopropylatrazine	100-3 C ₈ ec	61
	C ₈ Gravity, 5 µm	36	Dexamethasone	C ₁₈ Gravity, 5 µm	49
Chloroquine	C ₁₈ Gravity, 5 µm	48		C ₁₈ Isis, 5 µm	51
Chloroxuron	100-3 C ₈ ec	60		PFP, 5 µm	19
Chlorpheniramine	100-5 C ₁₈ ec / 100-5 CN-RP	28	Dextromethorphan	C ₁₈ Gravity, 5 µm	36
	C ₈ Gravity, 5 µm	15		C ₁₈ Pyramid, 5 µm	36
	C ₁₈ Gravity, 5 µm	15, 36	Dibenz[ah]anthracene	C ₁₈ Gravity, 5 µm	59
	C ₁₈ Pyramid, 5 µm	15, 36	Dibenzothiophene	100-5 NH ₂	30
	PFP, 5 µm	37	Dibromophenol isomers	PFP, 5 µm	57
	PFP, C ₁₈ Gravity, 5 µm	19		PFP, C ₁₈ HTec, 5 µm	19
Chlorpromazine	100-5 CN-RP	38	Dibutyl phthalate	C ₁₈ Gravity, 5 µm	9
Chlortoluron	100-3 C ₈ ec	60, 61	Dicamba	100-3 C ₈ ec	61
Chrysene	C ₁₈ Gravity, 5 µm	59	Dichlobenil	100-3 C ₈ ec	60
Cimetidine	C ₁₈ Gravity, 3 µm	36	3,4-Dichloroaniline	PFP, 5 µm	58
Cinoxacin	100-5 C ₁₈ ec	39	2,4-Dichlorobenzamide	100-3 C ₈ ec	61
	C ₁₈ Gravity, 5 µm	42	2,4-Dichlorophenol	PFP, 5 µm	57
	C ₁₈ Gravity, C ₈ Gravity, C ₁₈ Pyramid	40	Dichlorophenol isomers	PFP, 5 µm	57
	C ₁₈ HTec, 5 µm	41		PFP, C ₁₈ HTec, 5 µm	19
	Sphinx RP, 5 µm	40	Dichlorprop	100-3 C ₈ ec	61
Ciprofloxacin	100-5 C ₁₈ ec	39	Diclofenac	100-5 C ₈ ec	33
	Sphinx RP, 5 µm	39	Dicloxacillin	100-5 C ₁₈ ec, C ₁₈ Pyramid, 5 µm	32, 33, 34
Citric acid	C ₁₈ Pyramid, 5 µm	55		100-5 C ₁₈ ec	40
	HILIC, 3 µm	55		PFP, 5 µm	41
Citrulline	C ₁₈ Gravity, 3 µm	46	Diethylenetrinitriopentaacetic acid		
Clenbuterol	100-5 CN-RP	38		C ₁₈ Pyramid, 5 µm	54
	C ₁₈ Pyramid, 5 µm	37	Diflunisal	100-5 C ₈ ec	33
Clobetasol 17-propionate	C ₁₈ Pyramid, 1.8 µm	33		C ₁₈ Gravity, 5 µm	34
Clomipramine	100-5 CN-RP; C ₁₈ Isis, 5 µm	35	Dihydroxybenzoic acid isomers		
Clonidin	100-5 CN-RP	38		C ₁₈ Isis, 5 µm	54
Cloxacillin	100-5 C ₁₈ ec	39, 40		PolarTec, 5 µm	55
	C ₁₈ Gravity, 5 µm	42	Dihydroxymandelic acid	PolarTec, 5 µm	55
	C ₁₈ HTec, 5 µm	41	Dihydroxynaphthalene isomers		
	PFP, 5 µm	41		C ₈ Gravity, 5 µm	63
Cohumulone, colupulone	100-5 C ₁₈ ec	51		C ₁₈ Isis, 5 µm	63
Corticosterone	C ₁₈ Gravity, 5 µm	49	Dihydroxyphenylacetic acid	Sphinx RP, 5 µm	62
Cortisone	100-5 C ₈ ec	49			
	100-5 CN / CN-RP	29	Dihydroxyphenylalanine	PolarTec, 5 µm	55
	C ₈ Gravity, 5 µm	48		C ₁₈ Gravity, 5 µm	50
	C ₁₈ Gravity, 5 µm	49	Dimefuron	100-3 C ₈ ec	61
	C ₁₈ HTec, 5 µm	49	N,N-Dimethylaniline	C ₁₈ HTec, 5 µm	22
Creatine	HILIC, 3 µm	45		C ₁₈ Isis, 5 µm	58
Creatinine	HILIC, 3 µm	45	1,2-Dimethylbenzene	100-5 NH ₂	30
o-, m-Cresol	PFP, 5 µm	57	2,4-Dimethylphenol	PFP, 5 µm	57
	PFP, C ₁₈ HTec, 5 µm	19	Dimethylphenol isomers	PFP, 5 µm	57
Cyanazine	100-3 C ₈ ec	60, 61		PFP, C ₁₈ HTec, 5 µm	19
Cyanocobalamine	C ₁₈ Pyramid, 5 µm	53	Dimethyl phthalate	100-5 CN-RP	28, 29
	HILIC, 3 µm	52		C ₁₈ Pyramid, C ₁₈ Gravity, C ₈ Gravity	15
Cyanuric acid	C ₁₈ Pyramid, 5 µm	45	1,3-Dinitrobenzene	PolarTec, 5 µm	56
	HILIC, 5 µm	45	2,4-Dinitrophenol	PFP, 5 µm	57
Cyclohexane	100-5 NH ₂	30	3,5-Dinitro-(1-phenylethylbenzamide)	C ₁₈ Isis, 5 µm	58
				PolarTec, 5 µm	56



Substance index

Diphenhydramine	C ₁₈ Gravity, 5 µm C ₁₈ Pyramid, 5 µm PFP, 5 µm	10 36 37	Fumaric acid	100-5 CN-RP C ₁₈ Pyramid, 5 µm HILIC, 3 µm	54 55 55
Dipyridamole	PFP, C ₁₈ Gravity, 5 µm	19	2-Furfurol	100-5 C ₁₈ ec	59
Diquat	C ₁₈ Gravity, 3 µm	39	2-Furfuryl alcohol	100-5 C ₁₈ ec	59
Diuron	HILIC, 3 µm	62			
L-DOPA	100-3 C ₈ ec	60, 61	G		
Dopamine	HILIC, 3 µm	50	Gallic acid	C ₁₈ Isis, 5 µm C ₁₈ Pyramid, 5 µm PolarTec, 5 µm	54 43 55
Doxepin	100-5 CN-RP; C ₈ Gravity, C ₁₈ Gravity, C ₁₈ Pyramid, 5 µm PolarTec, 5 µm	35 34	Glucose	100-5 NH ₂ -RP 100-5 NH ₂ -RP	30 60
DTPA	C ₁₈ Pyramid, 5 µm	54	Glufosinate	C ₁₈ Gravity, 5 µm 100-5 C ₁₈ ec 100-5 C ₁₈ ec	60 46 46
E			Glutamic acid	C ₁₈ Gravity, 3 µm 100-5 C ₁₈ ec	46 46
E 102/104/110/122/124/126/127	C ₁₈ Gravity, 5 µm C ₁₈ Pyramid, 5 µm	44 54	Glutamine	100-5 C ₁₈ ec C ₁₈ Gravity, 5 µm PolarTec, 3 µm	46 60 47
EDTA	C ₁₈ Gravity, 5 µm	42	Glycine	100-5 NH ₂ -RP 100-5 NH ₂ -RP	30 60
Enrofloxacin	C ₁₈ HTec, 5 µm	41	Glyphosate	C ₁₈ Gravity, 5 µm PolarTec, 3 µm	60 17
Ephedrine	Sphinx RP, 5 µm	39	Guanine	PolarTec, 5 µm HILIC, 5 µm	47
Erythrosine	100-5 C ₁₈ ec / 100-5 CN-RP	28	Guanosine	HILIC, 5 µm	27, 47
Estradiol	C ₁₈ Gravity, 5 µm	44			
Estriol	100-5 C ₈ ec; C ₁₈ HTec, 5 µm	49	H		
Estrone	100-5 C ₈ ec; C ₁₈ HTec, 5 µm	49	Hexamethylbenzene	100-5 NH ₂	30
2-Ethoxyphenol	100-5 C ₈ ec	25	Hexazinone	100-3 C ₈ ec	60, 61
4-Ethoxyvanillin	100-5 C ₁₈ ec	25, 56	Hexobarbital	100-5 C ₁₈ ec	34
4-Ethylaniline	100-5 C ₁₈ ec	58	Hexogen (RDX)	PolarTec, 5 µm	56
Ethylbenzene	C ₁₈ HTec, 5 µm C ₁₈ Isis, 5 µm 100-5 C ₁₈ ec	22 58 24	Histamine	HILIC, 3 µm	47
	C ₁₈ Gravity, 1.8 vs. 3 µm	6	Histidine	100-5 C ₁₈ ec C ₁₈ Gravity, 3 µm	46 46
	C ₁₈ Gravity, 5 µm	11	n-Humulone	HILIC, 3 µm PolarTec, 3 µm	47 17
	C ₁₈ HTec, 5 µm	22	Hydrocortisone	100-5 C ₁₈ ec 100-5 CN/CN-RP	51 29
	C ₁₈ Isis, 5 µm	13		C ₈ Gravity, 5 µm	48
Ethyl benzoate	C ₁₈ Pyramid, C ₁₈ Gravity, C ₈ Gravity Sphinx RP, 5 µm	15 62		C ₁₈ Gravity, 5 µm	49
Ethylenediaminetetraacetic acid	C ₁₈ Pyramid, 5 µm	54	Hydrocortisone acetate	C ₈ Gravity, 5 µm	48
Ethylhexyldimethyl PABA	100-5 C ₁₈ ec; C ₁₈ HTec, 5 µm	51	4-Hydroxybenzaldehyde	100-5 C ₁₈ ec	58
Ethylhexyl methoxycinnamate	100-5 C ₁₈ ec; C ₁₈ HTec, 5 µm	51	4-Hydroxybenzoic acid	C ₁₈ Isis, 5 µm	54
Ethylhexyl salicylate	100-5 C ₁₈ ec; C ₁₈ HTec, 5 µm	51	3-Hydroxybenzoic acid	PolarTec, 5 µm	55
Ethyl paraben	PolarTec, 3 µm	43	17 α -Hydroxycortisone	C ₁₈ Gravity, 5 µm	49
Eugenol	C ₁₈ Gravity, 1.8 vs. 3 µm	59	Hydroxymethylfurfural	C ₁₈ Gravity, 5 µm	44
F			5-Hydroxymethyl-2-furfural	100-5 C ₁₈ ec	44
Famotidine	C ₁₈ Gravity, 3 µm	36	α -Hydroxymidazolam	C ₁₈ Gravity, 3 µm	59
Fast Red E	C ₁₈ Gravity, 5 µm	44	4-Hydroxyproline	C ₁₈ Gravity, 3 µm	38
Fast Yellow	C ₁₈ Gravity, 5 µm	44	Hydroxytestosterone isomers	C ₁₈ Gravity, 3 µm	46
Fenoprofen	100-5 C ₈ ec; 100-5 C ₁₈ ec	33 33		C ₁₈ Isis, 1.8 µm	49
	C ₁₈ Gravity, 5 µm	33	Hydroxytyramine	C ₁₈ Gravity, 5 µm	50
Fenoterol	C ₁₈ Pyramid, 5 µm	34	Hydroxyzine	PFP, 5 µm	37
Fisetin	Sphinx RP, C ₁₈ Gravity, C ₈ Gravity	21		PFP, C ₁₈ Gravity, 5 µm	19
Flumequine	Sphinx RP, 5 µm	39	I		
Flunarizine hydrochloride	C ₁₈ Gravity, 3 µm	39	Ibuprofen	100-5 C ₈ ec	33
Fluoranthene	C ₁₈ Gravity, 5 µm	59		100-5 C ₁₈ ec	32, 34
Fluorene	C ₁₈ Gravity, 5 µm	59		C ₁₈ Gravity, 5 µm	34
Flurbiprofen	100-5 C ₈ ec	33	Imipramine	C ₁₈ Pyramid, 5 µm	32
	100-5 C ₁₈ ec	34		C ₈ Gravity, C ₁₈ Gravity, C ₁₈ Isis,	
	C ₁₈ Gravity, 5 µm	34		C ₁₈ Pyramid, 5 µm	
	C ₁₈ Pyramid, 5 µm	32	Indeno[1,2,3-cd]pyrene	C ₁₈ Gravity, 5 µm	35
Folic acid	C ₁₈ Pyramid, 5 µm	52, 53	Indomethacin	PolarTec, 5 µm	34
Formic acid	C ₁₈ Pyramid, 5 µm	15		C ₁₈ Gravity, 5 µm	59
Fructose	100-5 NH ₂ -RP	30	Irgasan	100-5 C ₈ ec	33
				C ₁₈ Gravity, 5 µm	34
			Isoadhumulone	PolarTec, 3 µm	43
			Isobutyl paraben	100-5 C ₁₈ ec	51
			Isocohumulone	PolarTec, 3 µm	43
			Isohumulone	100-5 C ₁₈ ec	51
				100-5 C ₁₈ ec	51

Alphabetical index of analytes

Isoleucine	100-5 C ₁₈ ec C ₁₈ Gravity, 3 µm	46 46	6α-Methyl-11β-hydroxyprogesterone 100-5 C ₈ ec; C ₁₈ HTec, 5 µm C ₈ Gravity, 5 µm	49 48	
Isoproturon	100-3 C ₈ ec	60, 61	6α-Methyl-17α-hydroxyprogesterone /+ acetate 100-5 C ₈ ec; C ₁₈ HTec, 5 µm C ₈ Gravity, 5 µm	49 48	
Isorhamnetin	Sphinx RP, C ₁₈ Gravity, C ₈ Gravity	21			
K					
Kaempferol	Sphinx RP, C ₁₈ Gravity, C ₈ Gravity	21	Methyl paraben PolarTec, 3 µm	43	
Ketoprofen	100-5 C ₈ ec 100-5 C ₁₈ ec C ₁₈ Gravity, 5 µm C ₁₈ Pyramid, 5 µm	33 32, 33, 34 10 32	6α-Methylprednisolone Methyltestosterone Metabromuron Metolachlor Metoxuron Midazolam Monensin sodium Monolinuron	49 49 29 60, 61 60, 61 60, 61 38 41 60, 61	
L					
Lactic acid	C ₁₈ Pyramid, 5 µm	55			
Lactose	100-5 NH ₂ -RP	30			
Leucine	100-5 C ₁₈ ec C ₁₈ Gravity, 3 µm	46 46			
Lidocaine	C ₈ Gravity, 5 µm C ₁₈ Gravity, 5 µm C ₁₈ Pyramid, 5 µm Sphinx RP, 5 µm	15 10, 15 15, 32 62	Nafcillin Nalidixic acid Naphthalene	100-5 C ₁₈ ec, C ₁₈ Pyramid, 5 µm PFP, 5 µm C ₁₈ Gravity, 5 µm C ₁₈ Gravity, C ₈ Gravity, C ₁₈ Pyramid Sphinx RP, 5 µm C ₁₈ HTec, 5 µm Sphinx RP, 5 µm 100-5 NH ₂ C ₁₈ Gravity, 1.8 vs. 3 µm C ₁₈ Gravity, 5 µm	40 41 42 40 41 39 30 6 15, 59
Linuron	100-3 C ₈ ec	60, 61			
Lorazepam	100-5 C ₁₈ ec	38			
n-Lupulone	100-5 C ₁₈ ec	51			
Lysine	100-5 C ₁₈ ec	46			
M					
Maleic acid	100-5 C ₁₈ ec 100-5 CN-RP C ₁₈ Gravity, 5 µm C ₁₈ Pyramid, 5 µm PFP, 5 µm PFP, C ₁₈ Gravity, 5 µm	28 28, 54 36 14 37 19			
Malic acid	C ₁₈ Pyramid, 5 µm	55	Naproxen	100-5 C ₈ ec C ₁₈ Gravity, 5 µm	
Maltose	100-5 NH ₂ -RP	30		33 34	
Mapenterol	C ₁₈ Pyramid, 5 µm	37	Nicardipine	100-5 CN-RP	
Maprotiline	C ₁₈ Isis, 5 µm	35	Nicotinamide	C ₁₈ Pyramid, 5 µm	
Marbofloxacin	Sphinx RP, 5 µm	39	Nicotinic acid	52	
MCPA	100-3 C ₈ ec	61	Nifedipine	HILIC, 3 µm	
Meclofenamic acid	100-5 C ₁₈ ec	34	Niflumic acid	100-5 CN-RP	
Mecoprop	100-3 C ₈ ec	61	Nimodipine	100-5 CN-RP	
Medrysone	100-5 CN/CN-RP	29	Nisoldipine	100-5 CN-RP	
Mefloquine	C ₁₈ Gravity, 5 µm	48	Nitrendipine	100-5 CN-RP	
Melamine	HILIC, 5 µm	45	Nitrofuran triacetic acid	C ₁₈ Pyramid, 5 µm	
Mephobarbital	100-5 C ₁₈ ec	34	Nitroaniline isomers	PFP, 5 µm	
Mepiquat	HILIC, 3 µm	27	Nitrobenzene	C ₁₈ Gravity, Sphinx RP, 5 µm	
Metalexyl	100-3 C ₈ ec	61		PFP, 3 µm	
Metamitron	100-3 C ₈ ec	60		PolarTec, 5 µm	
Metazachlor	100-3 C ₈ ec	60, 61	Nitrocresol isomers	C ₁₈ Gravity, 5 µm	
Methabenzthiazuron	100-3 C ₈ ec	60, 61	2-Nitrophenol	C ₁₈ Gravity, Sphinx RP, 5 µm	
Methacrylamide	HILIC, 5 µm	44	Nitrophenol isomers	C ₁₈ Gravity, 5 µm	
Methacrylic acid	HILIC, 5 µm	44		PFP, 5 µm	
Methionine	100-5 C ₁₈ ec C ₁₈ Gravity, 3 µm	46 46	Nitrotoluene isomers	PolarTec, 5 µm	
Methoxyphenol isomers	100-5 C ₈ ec 100-5 C ₁₈ ec	25 25, 56	Nizatidine	C ₁₈ Gravity, 3 µm	
2-Methyl-4,6-dinitrophenol	PFP, 5 µm	57	N-Methyl-N-2,4,6-tetrinitro-aniline (Tetryl)	N-Methyl-N-2,4,6-tetrinitro-aniline (Tetryl)	
o-, m-, p-Methylacetophenone	PFP, C ₁₈ Gravity, 5 µm	58	Norephedrine	PolarTec, 5 µm	
9-Methylanthracene	100-5 NH ₂	30		100-5 C ₁₈ ec/100-5 CN-RP	
4-Methylbenzylidene camphor	100-5 C ₁₈ ec C ₁₈ HTec, 5 µm	51 51	Norgestrel	C ₁₈ Gravity, 5 µm	
5-Methyl-2-furfurol	100-5 C ₁₈ ec	59	Nortriptyline	HILIC, 3 µm	
Methylhistidine isomers	PolarTec, 3 µm	17		100-5 CN/CN-RP	
Methyl-4-hydroxybenzoate	C ₁₈ Pyramid, 5 µm	32	Norvaline	100-5 CN-RP; C ₈ Gravity, C ₁₈ Gravity, C ₁₈ Isis, C ₁₈ Pyramid, 5 µm	
			Noscapine	100-5 C ₁₈ ec	
				C ₈ Gravity, 5 µm	
			NTA	C ₁₈ Gravity, 5 µm	
				C ₁₈ Pyramid, 5 µm	

Substance index

O

Octocrylene	100-5 C ₁₈ ec; C ₁₈ HTec, 5 µm	51
Octogen (HMX)	PolarTec, 5 µm	56
Ofloxacin	100-5 C ₁₈ ec	39
Ornithine	C ₁₈ Gravity, 3 µm	46
Oxacillin	PFP, 5 µm	41
Oxalic acid	HILIC, 3 µm	55
Oxazepam	100-5 C ₁₈ ec	38
Oxolinic acid	C ₈ Gravity, C ₁₈ Pyramid, 5 µm	40
	C ₁₈ Gravity, 5 µm	42
	C ₁₈ HTec, 5 µm	41
	Sphinx RP, 5 µm	39, 40

P

Panthothenic acid	PolarTec, 5 µm	52
Papaverine	C ₈ Gravity, 5 µm	48
	C ₁₈ Gravity, 5 µm	10
Paracetamol	100-5 C ₁₈ ec	28, 32
	100-5 CN-RP	28
	C ₈ Gravity, 5 µm	32
	C ₁₈ Gravity, 5 µm	36
	C ₁₈ Pyramid, 5 µm	32, 36
Paraquat	HILIC, 3 µm	62
Patulin	C ₁₈ Gravity, 5 µm	44
Penicillins G + V	100-5 C ₁₈ ec	39, 40
	C ₁₈ Pyramid, 5 µm	40
Penicillin V	C ₁₈ Gravity, 5 µm	42
	C ₁₈ HTec, PFP, 5 µm	41
Pentachlorophenol	PFP, 5 µm	57
Pentobarbital	100-5 C ₁₈ ec	34
Perfluorinated surfactants	Sphinx RP, 3 µm	62
Phenactin	C ₈ Gravity, 5 µm	32
Phenanthrene	C ₁₈ Gravity, 5 µm	59
Phenetole	100-5 C ₈ ec/C ₁₈ ec	25
	100-5 CN-RP	28, 29
Phenobarbital	100-5 C ₁₈ ec	34
Phenol	100-5 C ₈ ec	25
	100-5 C ₁₈ ec	25, 56
	C ₁₈ Gravity, 5 µm	56
	C ₁₈ HTec, 5 µm	22
	C ₁₈ HTec, 5 µm; EC vs. VP	23
	PFP, 3 µm	57
	PFP, 5 µm	57
	PolarTec, 5 µm	17
Phenoxyethanol	Sphinx RP, 5 µm	21, 56
Phenylalanine	PolarTec, 3 µm	43
	100-5 C ₁₈ ec; C ₁₈ Gravity, 3 µm	46
	HILIC, 3 µm	47
1-Phenyldodecane	100-5 NH ₂	30
Phenylglycine	HILIC, 3 µm	47
Phloroglucinol	PFP, 3 µm	57
Piperacillin	PFP, 5 µm	41
Pirenzepine hydrochloride	C ₁₈ Gravity, 3 µm	36
Piroxicam	100-5 C ₈ ec, 100-5 C ₁₈ ec	33
	C ₁₈ Gravity, 5 µm	34
Ponceau 4R and 6R	C ₁₈ Gravity, 5 µm	44
Prednisolone	100-5 C ₈ ec	49
	100-5 CN/CN-RP	29
	C ₁₈ Gravity, 5 µm	49
	C ₁₈ HTec, 5 µm	49
Procainamide	100-5 CN-RP	38
Progesterone	100-5 C ₈ ec; C ₁₈ Gravity, C ₁₈ HTec, 5 µm	49
Promazine	100-5 CN-RP	38
Promethazine	100-5 CN-RP	38
	PFP, 5 µm	37
	PFP, C ₁₈ Gravity, 5 µm	19

Propazine

Propiophenone	100-3 C ₈ ec	61
	100-5 C ₁₈ ec	59
Propyl 4-hydroxybenzoate	C ₁₈ Gravity, 1.8 vs. 3 µm	59
	C ₁₈ Gravity, Sphinx RP, 5 µm	56
Propyl paraben	PolarTec, 3 µm	43
Propyzamid	100-3 C ₈ ec	60
Protriptyline	C ₈ Gravity, C ₁₈ Gravity, C ₁₈ Isis, C ₁₈ Pyramid, 5 µm	35
Protryptiline	PolarTec, 5 µm	34
Pseudoephedrine	C ₁₈ Gravity, 5 µm	36
Pyrene	C ₁₈ Gravity, 5 µm	59
Pyridine	C ₁₈ Gravity, 5 µm	11
	C ₁₈ Isis, 5 µm	13, 58
	PolarTec, 5 µm	17
	Sphinx RP, 5 µm	21
	C ₁₈ Pyramid, 5 µm	53
	C ₁₈ Pyramid, 5 µm	53
	C ₁₈ Pyramid, 5 µm	53
	HILIC, 3 µm	52
	C ₁₈ Pyramid, 5 µm	52
	Pyrocatechol	25
	100-5 C ₈ ec	25, 56
	100-5 C ₁₈ ec	25, 56
	PFP, 3 µm	57

Q

Quercetin	Sphinx RP, C ₁₈ Gravity, C ₈ Gravity	21
Quinine	100-5 C ₈ ec	43
	C ₈ Gravity, C ₁₈ Gravity, 5 µm	48
Quinoline yellow	C ₁₈ Gravity, 5 µm	44

R

Ractopamine	C ₁₈ Pyramid, 5 µm	37
Reprotorel	C ₁₈ Pyramid, 5 µm	37
Resorcinol	100-5 C ₈ ec	25
	100-5 C ₁₈ ec	25, 56
	PFP, 3 µm	57
Riboflavin	C ₁₈ Pyramid, 5 µm	52, 53
Ritodrine	C ₁₈ Pyramid, 5 µm	37
Rutin	C ₁₈ Pyramid, 5 µm	52
	Sphinx RP, C ₁₈ Gravity, C ₈ Gravity	21

S

Saccharin	100-5 C ₈ ec; C ₁₈ HTec, 5 µm	43
Saccharose	100-5 NH ₂ -RP	30
Salicylic acid	100-5 C ₁₈ ec	32
	PolarTec, 3 µm	43
Salinomycin sodium	C ₁₈ Gravity, 3 µm	41
Sarafloxacin	Sphinx RP, 5 µm	39
Scopolamine	C ₈ Gravity, 5 µm	48
Sebutyhalazine	100-3 C ₈ ec	60, 61
Serine	100-5 C ₁₈ ec	46
	C ₁₈ Gravity, 3 µm	46
Shikimic acid	C ₁₈ Pyramid, 5 µm	55
Simazine	100-3 C ₈ ec	60, 61
Sorbitic acid	100-5 C ₈ ec; C ₁₈ HTec, 5 µm	43
Strychnine	C ₈ Gravity, 5 µm	48
Succinylsulfathiazole	C ₁₈ Gravity, 5 µm	42
Sulfadiazine	C ₁₈ Gravity, 5 µm	42
Sulfadimidine	C ₁₈ Gravity, 5 µm	42
Sulfamerazine	C ₁₈ Gravity, 5 µm	42
Sulfanilamide	C ₁₈ Gravity, 5 µm	42
Sulfathiazole	C ₁₈ Gravity, 5 µm	42
Sulindac	100-5 C ₁₈ ec; C ₁₈ Gravity, 5 µm	34
	C ₁₈ Gravity, 5 µm	44
Sunset yellow CFC	100-5 C ₈ ec, 100-5 C ₁₈ ec	33
Suprofen	C ₁₈ Gravity, 5 µm	34
	C ₁₈ Isis, 5 µm	54
Syringic acid	PolarTec, 5 µm	55

Alphabetical index of analytes

T

Tartaric acid	C ₁₈ Pyramid, 5 µm	14, 55
Tartrazine	C ₁₈ Gravity, 5 µm	44
Temazepam	100-5 C ₁₈ ec	38
Terbutylazine	100-3 C ₈ ec	60, 61
Terbutryl	100-3 C ₈ ec	60
o-, m-, p-Terphenyl	C ₁₈ Isis, 1.8 vs. 5 µm	7
o-Terphenyl	C ₁₈ Isis, 5 µm	12
Testosterone	C ₁₈ Isis, 5 µm	13
	100-5 C ₈ ec	49
	100-5 CN/CN-RP	29
	C ₁₈ HTec, 5 µm	49
Tetracaine	C ₁₈ Pyramid, 5 µm	32
Theobromine	Sphinx RP, 5 µm	42
Theophylline	100-5 C ₁₈ ec	24
	C ₁₈ Gravity, 5 µm	11
	C ₁₈ Isis, 5 µm	13
	Sphinx RP, 5 µm	20
Thiamine	C ₁₈ Pyramid, 5 µm	53
	HILIC, 3 µm	52
Thiamine hydrochloride	C ₁₈ Pyramid, 5 µm	52
Thiamylal	100-5 C ₁₈ ec	34
Threonine	C ₁₈ Gravity, 3 µm	46
Thymine	100-5 NH ₂ -RP	31
	C ₁₈ Pyramid, 5 µm	47
	HILIC, 5 µm	27, 47
	PolarTec, 3 µm	17
	PolarTec, 5 µm	47
α-Tocopherol	see Vitamin E	
Tocopherols	C ₁₈ Isis, 5 µm	53
Tolmetin	100-5 C ₈ ec	33
	100-5 C ₁₈ ec	34
	C ₁₈ Gravity, 5 µm	34
Toluene	100-5 C ₁₈ ec	24
	C ₁₈ Gravity, 5 µm	11, 56
	C ₁₈ HTec, 5 µm	22
	C ₁₈ Isis, 5 µm	13
	Sphinx RP, 5 µm	56
Tribunil	see Methabenzthiazuron	
2,4,6-Trichlorophenol	PFP, 5 µm	57
3,3,4-Triclocarbanilid	PolarTec, 3 µm	43
Trimipramine	100-5 CN-RP; C ₈ Gravity, C ₁₈ Gravity, 5 µm	35
	PolarTec, 5 µm	34
1,3,5-Trinitrobenzene	PolarTec, 5 µm	56
2,4,6-Trinitrotoluene	PolarTec, 5 µm	56
Triphenylene	C ₁₈ Isis, 1.8 vs. 5 µm	7
	C ₁₈ Isis, 5 µm	12, 13
Triprolidine	PFP, 5 µm	37
	PFP, C ₁₈ Gravity, 5 µm	19
Tryptophan	100-5 C ₁₈ ec	46
	C ₁₈ Gravity, 3 µm	46
Tyrosine	100-5 C ₁₈ ec	46
	C ₁₈ Gravity, 3 µm	46
	C ₁₈ Gravity, 5 µm	50
	HILIC, 3 µm	47

U

Uracil	100-5 C ₁₈ ec	24
	100-5 NH ₂ -RP	31
	C ₁₈ Gravity, 5 µm	56
	C ₁₈ HTec, 5 µm	22
	C ₁₈ Isis, 5 µm	58
	C ₁₈ Pyramid, 5 µm	47
	HILIC, 3 µm	27
	HILIC, 5 µm	27, 47
	PFP, 3 µm	57
	PolarTec, 3 µm	17
	PolarTec, 5 µm	47
	Sphinx RP, 5 µm	56, 62
Urea	C ₁₈ Pyramid, 5 µm	45
Uvinul Plus	C ₁₈ HTec, 5 µm	51
	V	
Valerophenone	100-5 C ₁₈ ec	59
	C ₁₈ Gravity, 1.8 vs. 3 µm	59
Valine	100-5 C ₁₈ ec	46
	C ₁₈ Gravity, 3 µm	46
	C ₁₈ Isis, 5 µm	54
	PolarTec, 5 µm	55
Vanillin	100-5 C ₈ ec	43
	100-5 C ₁₈ ec	58
	Sphinx RP, 5 µm	42
Veratrol	100-5 C ₈ ec	25
	100-5 C ₁₈ ec	24, 25
	Vitamin A + acetate	
	100-5 C ₁₈ ec; C ₁₈ Isis, 5 µm	53
	Vitamin B ₁ / B ₂ / B ₆	52
	Vitamin D ₂ / D ₃	
	100-5 C ₁₈ ec; C ₁₈ Isis, 5 µm	53
	Vitamin E + acetate	
	100-5 C ₁₈ ec; C ₁₈ Isis, 5 µm	53
	Vitamin K ₁	
	100-5 C ₁₈ ec; C ₁₈ Isis, 5 µm	53
	Vitamin K ₂	
	C ₁₈ Isis, 5 µm	53
	100-5 C ₁₈ ec	53
	Vitamins, water-soluble (B ₁ , B ₂ , B ₆ , B ₇ , B ₉ , B ₁₂ , C, H)	
	HILIC, 3 µm	52
	C ₁₈ Pyramid, 5 µm	52, 53
	X	
o-Xylene	100-5 CN-RP	28, 29
Xylene	C ₁₈ Gravity, Sphinx RP, 5 µm	56
Xylometazoline	100-5 CN-RP	37
	Y	
Yellow orange S	C ₁₈ Gravity, 5 µm	44
	Z	
Zearalenol	C ₁₈ Gravity, 5 µm	50
Zearalenone	C ₁₈ Gravity, 5 µm	50

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Appl. no. index / Trademarks

Index of application numbers

MN appl.	page								
117770	32	119160	32	120450	43	121662	59	123070	45
117800	35	119170	15	120460	40	121800	44	123090	45
117810	36	119200	35	120470	40	122130	33	123640	51
117820	34	119210	15	120481	61	122140	49	123710	49
117830	34	119220	15	120482	61	122160	30	123750	43
117850	38	119221	15	120483	61	122170	31	123760	41
117860	40	119271	29	120485	61	122180	30	124490	56
117870	39	119272	29	120490	60	122190	60	124531	19
117880	42	119280	35	120500	55	122450	53	124541	57
117890	53	119290	54	120740	59	122470	42	124562	55
117930	50	119300	38	120780	50	122500	44	124570	52
117960	48	119310	39	120810	9	122510	44	124590	43
117970	56	119320	38	120840	11	122520	36	124600	17
117980	59	119340	28	120850	11	122530	49	124610	17
117990	58	119350	29	120860	10	122550	34	124622	34
118010	10	119410	32	120870	14	122560	39	124672	47
118450	46	119760	37	120880	33	122580	41	124711	17
118470	38	119770	52	120881	33	122590	41	124720	19
118520	35	119780	54	120890	25	122650	56	124730	18
118540	48	119800	50	120891	25	122720	59	124740	57
118550	49	119810	42	120900	20	122730	59	124750	57
118560	43	119830	21	120910	21	122930	55	124761	58
118580	48	119840	56	121100	51	122940	54	124771	19
118590	33	119850	56	121160	53	122950	47	124800	58
118600	32	119870	40	121170	51	122970	52	124840	41
118630	63	119880	62	121180	54	122980	47	124851	37
118760	41	119890	60	121190	63	123000	45	124861	19
118980	46	119920	42	121200	58	123010	44	125140	17
119110	36	120380	28	121210	35	123030	50		
119120	36	120390	37	121500	51	123040	27		
119140	47	120400	39	121590	62	123050	62		
119150	40	120440	45	121612	36	123060	62		

Trademarks

MACHEREY-NAGEL trademarks

ALUGRAM	coated aluminium sheets for TLC
CHROMABOND	columns for solid phase extraction (SPE)
ChromCart	cartridge system for HPLC
NUCLEODUR	spherical high purity silica for HPLC

NUCLEOSIL OPTIMA

spherical standard silica for HPLC
high performance fused silica capillary columns with immobilized phases

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Ordering information

NUCLEODUR® C₁₈ Gravity

Pore size 110 Å; high density octadecyl phase, endcapped, 18% C;
eluent in column acetonitrile – water

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm						
NUCLEODUR® C₁₈ Gravity, 1.8 µm													
particle size 1.8 µm													
EC analytical columns													
2 mm ID	760078.20	760079.20	760071.20	760076.20		760075.20							
3 mm ID	760078.30	760079.30		760076.30									
4 mm ID	760078.40	760079.40		760076.40									
4.6 mm ID	760078.46	760079.46		760076.46									
EC guard columns*	4 x 2 mm: 761901.20		4 x 3 mm: 761901.30										
NUCLEODUR® C₁₈ Gravity, 3 µm													
particle size 3 µm													
Microbore analytical columns													
1 mm ID			717714.10	717715.10	717716.10	717717.10							
EC analytical columns													
2 mm ID		760080.20	760084.20	760081.20	760083.20	760082.20							
3 mm ID		760080.30	760084.30	760081.30	760083.30	760082.30							
4 mm ID		760080.40	760084.40	760081.40	760083.40	760082.40							
4.6 mm ID		760080.46	760086.46	760084.46	760081.46	760083.46	760082.46						
EC guard columns*	4 x 2 mm: 761902.20		4 x 3 mm: 761902.30										
CC guard columns**	8 x 3 mm: 761124.30		8 x 4 mm: 761124.40										
NUCLEODUR® C₁₈ Gravity, 5 µm													
particle size 5 µm													
Microbore analytical columns													
1 mm ID			717706.10	717707.10	717708.10	717705.10							
EC analytical columns													
2 mm ID		760102.20	760104.20	760100.20	760103.20	760101.20							
3 mm ID		760102.30	760104.30	760100.30	760103.30	760101.30							
4 mm ID		760102.40	760104.40	760100.40	760103.40	760101.40							
4.6 mm ID		760102.46	760106.46	760104.46	760100.46	760103.46	760101.46						
EC guard columns*	4 x 2 mm: 761903.20		4 x 3 mm: 761903.30										
CC guard columns**	8 x 3 mm: 761125.30		8 x 4 mm: 761125.40										
VarioPrep preparative columns													
10 mm ID		762103.100		762109.100		762113.100							
21 mm ID		762103.210		762109.210		762113.210							
32 mm ID						762113.320							
40 mm ID					762100.400	762113.400							
VP guard columns***	10 x 8 mm: 762160.80		10 x 16 mm: 762160.160		15 x 32 mm: 762163.320								
NUCLEODUR® C₁₈ Gravity, 10 µm													
particle size 10 µm													
VarioPrep preparative columns													
21 mm ID						762250.210							
40 mm ID						762250.400							
VP guard columns***	10 x 8 mm: 762160.80		10 x 16 mm: 762160.160		15 x 32 mm: 762163.320								
Microbore, EC, and VarioPrep columns in packs of 1, guard columns see page 73													



Packed columns

NUCLEODUR® C₈ Gravity

Pore size 110 Å; high density octyl phase, endcapped, 11% C;
eluent in column acetonitrile – water

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm		
NUCLEODUR® C₈ Gravity, 1.8 µm							particle size 1.8 µm		
EC analytical columns									
2 mm ID	760756.20	760755.20	760760.20	760757.20		760759.20			
3 mm ID	760756.30	760755.30		760757.30					
4 mm ID	760756.40	760755.40		760757.40					
4.6 mm ID	760756.46	760755.46		760757.46					
EC guard columns*	4 x 2 mm: 761905.20		4 x 3 mm: 761905.30						
NUCLEODUR® C₈ Gravity, 5 µm							particle size 5 µm		
EC analytical columns									
2 mm ID		760750.20		760754.20	760751.20	760752.20	760753.20		
3 mm ID		760750.30		760754.30	760751.30	760752.30	760753.30		
4 mm ID		760750.40		760754.40	760751.40	760752.40	760753.40		
4.6 mm ID		760750.46	760749.46	760754.46	760751.46	760752.46	760753.46		
EC guard columns*	4 x 2 mm: 761907.20		4 x 3 mm: 761907.30						
CC guard columns**	8 x 3 mm: 761754.30		8 x 4 mm: 761754.40						
VarioPrep preparative columns									
10 mm ID	762081.100					762071.100	762070.100		
21 mm ID	762081.210					762071.210	762082.210		
VP guard columns***	10 x 8 mm: 762097.80		10 x 16 mm: 762097.160						
EC and VarioPrep columns in packs of 1, guard columns see right, Microbore columns with NUCLEODUR® C ₈ Gravity on request!									

HPLC column systems from MACHEREY-NAGEL



microbore column



EC column



VarioPrep column

Microbore columns: On request available in lengths of 40, 60, 100, 125, 150, 200, 250 and 300 mm and with 0.05, 0.075, 0.1, 0.15, 0.3, 0.4, 0.5, 0.75, 1.0 and 1.5 mm ID.

EC columns: Analytical ready-to-use columns; available dimensions see page 85.

VarioPrep columns: Preparative columns with axially adjustable endfitting; available dimensions see page 88.

Ordering information

NUCLEODUR® C₁₈ Isis

Pore size 110 Å; octadecyl phase with high steric selectivity, polymer modification, 20% C; eluent in column acetonitrile – water

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm				
NUCLEODUR® C₁₈ Isis, 1.8 µm											
particle size 1.8 µm											
EC analytical columns											
2 mm ID	760406.20	760405.20	760396.20	760407.20		760409.20					
3 mm ID	760406.30	760405.30		760407.30							
4 mm ID	760406.40	760405.40		760407.40							
4.6 mm ID	760406.46	760405.46		760407.46							
EC guard columns*	4 x 2 mm: 761910.20		4 x 3 mm: 761910.30								
NUCLEODUR® C₁₈ Isis, 3 µm											
particle size 3 µm											
Microbore analytical columns											
1 mm ID	717760.10		717761.10	717762.10							
EC analytical columns											
2 mm ID	760400.20		760401.20	760402.20	760403.20	760404.20					
3 mm ID	760400.30		760401.30	760402.30	760403.30	760404.30					
4 mm ID	760400.40		760401.40	760402.40	760403.40	760404.40					
4.6 mm ID	760400.46		760397.46	760401.46	760402.46	760403.46	760404.46				
EC guard columns*	4 x 2 mm: 761911.20		4 x 3 mm: 761911.30								
CC guard columns**	8 x 3 mm: 761300.30		8 x 4 mm: 761300.40								
NUCLEODUR® C₁₈ Isis, 5 µm											
particle size 5 µm											
Microbore analytical columns											
1 mm ID	717770.10		717771.10	717772.10							
EC analytical columns											
2 mm ID	760410.20		760415.20	760412.20	760413.20	760414.20					
3 mm ID	760410.30		760415.30	760412.30	760413.30	760414.30					
4 mm ID	760410.40		760415.40	760412.40	760413.40	760414.40					
4.6 mm ID	760410.46		760416.46	760415.46	760412.46	760413.46	760414.46				
EC guard columns*	4 x 2 mm: 761912.20		4 x 3 mm: 761912.30								
CC guard columns**	8 x 3 mm: 761310.30		8 x 4 mm: 761310.40								
VarioPrep preparative columns											
10 mm ID	762404.100			762405.100		762403.100					
21 mm ID	762404.210			762405.210		762403.210					
32 mm ID						762403.320					
40 mm ID					762406.400	762403.400					
VP guard columns***	10 x 8 mm: 762420.80		10 x 16 mm: 762420.160		15 x 32 mm: 762422.320						
Microbore, EC, and VarioPrep columns in packs of 1, guard columns see below											

Guard column systems						Guard column holder
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
** ChromCart® guard columns (pack of)	CC	8/3 (3)	8/3 (3)	8/4 (3)	8/4 (3)	721359
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
*** VarioPrep guard columns (pack of)	VP	10/8 (2)	10/16 (2)	15/32 (1)	15/50 (1)	
VP guard column holder		718251	718256	718253	718255	



Packed columns

NUCLEODUR® C₁₈ Pyramid

Pore size 110 Å; octadecyl phase with hydrophilic endcapping, 14% C;
eluent in column acetonitrile – water

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm				
NUCLEODUR® C₁₈ Pyramid, 1.8 µm											
particle size 1.8 µm											
EC analytical columns											
2 mm ID	760271.20	760272.20	760275.20	760273.20		760274.20					
3 mm ID	760271.30	760272.30		760273.30							
4 mm ID	760271.40	760272.40		760273.40							
4.6 mm ID	760271.46	760272.46		760273.46							
EC guard columns*	4 x 2 mm: 761915.20		4 x 3 mm: 761915.30								
NUCLEODUR® C₁₈ Pyramid, 3 µm											
particle size 3 µm											
Microbore analytical columns											
1 mm ID	717740.10		717741.10	717742.10	717743.10	717744.10					
EC analytical columns											
2 mm ID	760263.20		760264.20	760260.20	760261.20	760262.20					
3 mm ID	760263.30		760264.30	760260.30	760261.30	760262.30					
4 mm ID	760263.40		760264.40	760260.40	760261.40	760262.40					
4.6 mm ID	760263.46		760259.46	760264.46	760260.46	760261.46	760262.46				
EC guard columns*	4 x 2 mm: 761916.20		4 x 3 mm: 761916.30								
CC guard columns**	8 x 3 mm: 761854.30		8 x 4 mm: 761854.40								
NUCLEODUR® C₁₈ Pyramid, 5 µm											
particle size 5 µm											
Microbore analytical columns											
1 mm ID	717722.10		717723.10	717724.10	717725.10						
EC analytical columns											
2 mm ID	760200.20		760204.20	760201.20	760203.20	760202.20					
3 mm ID	760200.30		760204.30	760201.30	760203.30	760202.30					
4 mm ID	760200.40		760204.40	760201.40	760203.40	760202.40					
4.6 mm ID	760200.46		760205.46	760204.46	760201.46	760203.46	760202.46				
EC guard columns*	4 x 2 mm: 761917.20		4 x 3 mm: 761917.30								
CC guard columns**	8 x 3 mm: 761800.30		8 x 4 mm: 761800.40								
VarioPrep preparative columns											
10 mm ID	762271.100		762273.100		762272.100						
21 mm ID	762271.210		762273.210		762272.210						
32 mm ID					762272.320						
40 mm ID					762269.400		762272.400				
VP guard columns***	10 x 8 mm: 762291.80		10 x 16 mm: 762291.160		15 x 32 mm: 762293.320						
Microbore, EC, and VarioPrep columns in packs of 1, guard columns see right											



Ordering information

NUCLEODUR® PolarTec

Pore size 110 Å; octadecyl phase with embedded polar group, endcapped, 17% C;
eluent in column acetonitrile – water

Length →	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm						
NUCLEODUR® PolarTec, 3 µm												
particle size 3 µm												
EC analytical columns												
2 mm ID	760473.20		760476.20	760477.20	760478.20	760479.20						
3 mm ID	760473.30		760476.30	760477.30	760478.30	760479.30						
4 mm ID	760473.40		760476.40	760477.40	760478.40	760479.40						
4.6 mm ID	760473.46	760475.46	760476.46	760477.46	760478.46	760479.46						
EC guard columns*	4 x 2 mm: 761981.20		4 x 3 mm: 761981.30									
CC guard columns**	8 x 3 mm: 761160.30		8 x 4 mm: 761160.40									
NUCLEODUR® PolarTec, 5 µm												
particle size 5 µm												
EC analytical columns												
2 mm ID	760483.20		760486.20	760487.20	760488.20	760489.20						
3 mm ID	760483.30		760486.30	760487.30	760488.30	760489.30						
4 mm ID	760483.40		760486.40	760487.40	760488.40	760489.40						
4.6 mm ID	760483.46	760485.46	760486.46	760487.46	760488.46	760489.46						
EC guard columns*	4 x 2 mm: 761982.20		4 x 3 mm: 761982.30									
CC guard columns**	8 x 3 mm: 761161.30		8 x 4 mm: 761161.40									
VarioPrep preparative columns												
10 mm ID	762220.100		762221.100		762223.100							
21 mm ID	762220.210		762221.210		762223.210							
32 mm ID					762223.320							
40 mm ID					762222.400	762223.400						
VP guard columns***	10 x 8 mm: 762224.80		10 x 16 mm: 762224.160		15 x 32 mm: 762226.320							
EC and VarioPrep columns in packs of 1, guard columns see below												

Guard column systems					
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)
** ChromCart® guard columns (pack of)	CC	8/3 (3)	8/3 (3)	8/4 (3)	8/4 (3)
Guard columns for VarioPrep columns with ID					
*** VarioPrep guard columns (pack of)		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm
VP guard column holder		10/8 (2)	10/16 (2)	15/32 (1)	15/50 (1)
		718251	718256	718253	718255
Guard column holder					



Packed columns

NUCLEODUR® PFP

Pore size 110 Å; pentafluorophenyl-propyl modification, multi-endcapped, 8% C;
eluent in column acetonitrile – water

Length →	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm		
NUCLEODUR® PFP, 3 µm						particle size 3 µm		
EC analytical columns								
2 mm ID	760443.20		760446.20	760447.20	760448.20	760449.20		
3 mm ID	760443.30		760446.30	760447.30	760448.30	760449.30		
4 mm ID	760443.40		760446.40	760447.40	760448.40	760449.40		
4.6 mm ID	760443.46	760445.46	760446.46	760447.46	760448.46	760449.46		
EC guard columns*		4 x 2 mm: 761976.20		4 x 3 mm: 761976.30				
CC guard columns**		8 x 3 mm: 761145.30		8 x 4 mm: 761145.40				
NUCLEODUR® PFP, 5 µm						particle size 5 µm		
EC analytical columns								
2 mm ID	760453.20		760456.20	760457.20	760458.20	760459.20		
3 mm ID	760453.30		760456.30	760457.30	760458.30	760459.30		
4 mm ID	760453.40		760456.40	760457.40	760458.40	760459.40		
4.6 mm ID	760453.46	760455.46	760456.46	760457.46	760458.46	760459.46		
EC guard columns*		4 x 2 mm: 761977.20		4 x 3 mm: 761977.30				
CC guard columns**		8 x 3 mm: 761146.30		8 x 4 mm: 761146.40				
VarioPrep preparative columns								
10 mm ID	762210.100			762211.100		762213.100		
21 mm ID	762210.210			762211.210		762213.210		
32 mm ID						762213.320		
40 mm ID					762212.400	762213.400		
VP guard columns***	10 x 8 mm: 762214.80		10 x 16 mm: 762214.160		15 x 32 mm: 762216.320			
EC and VarioPrep columns in packs of 1, guard columns see right								

Online Application Database • www.mn-net.com/apps

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Ordering information

NUCLEODUR® Sphinx RP

Pore size 110 Å; special bifunctional RP phase, 15% C;
eluent in column acetonitrile – water

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm				
NUCLEODUR® Sphinx RP, 1.8 µm											
EC analytical columns											
2 mm ID	760821.20	760822.20	760825.20	760823.20		760824.20					
3 mm ID	760821.30	760822.30		760823.30							
4 mm ID	760821.40	760822.40		760823.40							
4.6 mm ID	760821.46	760822.46		760823.46							
EC guard columns*	4 x 2 mm: 761920.20		4 x 3 mm: 761920.30								
NUCLEODUR® Sphinx RP, 3 µm											
EC analytical columns											
2 mm ID	760806.20		760812.20	760807.20	760805.20	760808.20					
3 mm ID	760806.30		760812.30	760807.30	760805.30	760808.30					
4 mm ID	760806.40		760812.40	760807.40	760805.40	760808.40					
4.6 mm ID	760806.46		760813.46	760812.46	760807.46	760805.46	760808.46				
EC guard columns*	4 x 2 mm: 761921.20		4 x 3 mm: 761921.30								
CC guard columns**	8 x 3 mm: 761557.30		8 x 4 mm: 761557.40								
NUCLEODUR® Sphinx RP, 5 µm											
Microbore analytical columns											
1 mm ID	717680.10		717681.10	717682.10	717683.10	717684.10					
EC analytical columns											
2 mm ID	760800.20		760809.20	760801.20	760802.20	760803.20					
3 mm ID	760800.30		760809.30	760801.30	760802.30	760803.30					
4 mm ID	760800.40		760809.40	760801.40	760802.40	760803.40					
4.6 mm ID	760800.46		760815.46	760809.46	760801.46	760802.46	760803.46				
EC guard columns*	4 x 2 mm: 761922.20		4 x 3 mm: 761922.30								
CC guard columns**	8 x 3 mm: 761550.30		8 x 4 mm: 761550.40								
VarioPrep preparative columns											
10 mm ID	762372.100		762375.100		762373.100						
21 mm ID	762372.210		762375.210		762373.210						
32 mm ID					762373.320						
40 mm ID			762371.400		762373.400						
VP guard columns***	10 x 8 mm: 762390.80		10 x 16 mm: 762390.160		15 x 32 mm: 762392.320						
Microbore, EC, and VarioPrep columns in packs of 1, guard columns see below											

Guard column systems						Guard column holder
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
** ChromCart® guard columns (pack of)	CC	8/3 (3)	8/3 (3)	8/4 (3)	8/4 (3)	721359
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
*** VarioPrep guard columns (pack of)	VP	10/8 (2)	10/16 (2)	15/32 (1)	15/50 (1)	
VP guard column holder		718251	718256	718253	718255	



Packed columns

NUCLEODUR® C₁₈ HTec

Pore size 110 Å; high density octadecyl phase, endcapped, 18% C;
eluent in column acetonitrile – water

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm		
NUCLEODUR® C₁₈ HTec, 1.8 µm							particle size 1.8 µm		
EC analytical columns									
2 mm ID	760301.20	760305.20	760304.20	760306.20			760308.20		
3 mm ID	760301.30	760305.30		760306.30					
4 mm ID	760301.40	760305.40		760306.40					
4.6 mm ID	760301.46	760305.46		760306.46					
EC guard columns*	4 x 2 mm: 761925.20		4 x 3 mm: 761925.30						
NUCLEODUR® C₁₈ HTec, 3 µm							particle size 3 µm		
EC analytical columns									
2 mm ID		760321.20		760323.20	760324.20	760325.20	760326.20		
3 mm ID		760321.30		760323.30	760324.30	760325.30	760326.30		
4 mm ID		760321.40		760323.40	760324.40	760325.40	760326.40		
4.6 mm ID		760321.46	760322.46	760323.46	760324.46	760325.46	760326.46		
EC guard columns*	4 x 2 mm: 761926.20		4 x 3 mm: 761926.30						
CC guard columns**	8 x 3 mm: 761120.30		8 x 4 mm: 761120.40						
NUCLEODUR® C₁₈ HTec, 5 µm							particle size 5 µm		
EC analytical columns									
2 mm ID		760311.20		760313.20	760314.20	760315.20	760316.20		
3 mm ID		760311.30		760313.30	760314.30	760315.30	760316.30		
4 mm ID		760311.40		760313.40	760314.40	760315.40	760316.40		
4.6 mm ID		760311.46	760312.46	760313.46	760314.46	760315.46	760316.46		
EC guard columns*	4 x 2 mm: 761927.20		4 x 3 mm: 761927.30						
CC guard columns**	8 x 3 mm: 761110.30		8 x 4 mm: 761110.40						
VarioPrep preparative columns									
10 mm ID		762551.100			762554.100		762556.100		
21 mm ID		762551.210		762553.210	762554.210		762556.210		
32 mm ID				762553.320		762555.320	762556.320		
40 mm ID						762555.400	762556.400		
50 mm ID				762553.500		762555.500	762556.500		
VP guard columns***	10 x 8 mm: 762591.80		10 x 16 mm: 762591.160		15 x 50 mm: 762592.500				
	15 x 32 mm: 762592.320								
NUCLEODUR® C₁₈ HTec, 7 µm							particle size 7 µm		
VarioPrep preparative columns									
10 mm ID		762561.100			762564.100		762566.100		
21 mm ID		762561.210		762563.210	762564.210		762566.210		
32 mm ID				762563.320		762565.320	762566.320		
40 mm ID						762565.400	762566.400		
50 mm ID				762563.500		762565.500	762566.100		
VP guard columns***	10 x 8 mm: 762591.80		10 x 16 mm: 762591.160		15 x 50 mm: 762592.500				
	15 x 32 mm: 762592.320								



Ordering information

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C₁₈ HTec, 10 µm							particle size 10 µm
VarioPrep preparative columns							
10 mm ID	762571.100			762574.100			762576.100
21 mm ID	762571.210		762573.210	762574.210			762576.210
32 mm ID			762573.320		762575.320		762576.320
40 mm ID					762575.400		762576.400
50 mm ID			762573.500		762575.500		762576.100
VP guard columns***	10 x 8 mm: 762591.80 15 x 32 mm: 762592.320		10 x 16 mm: 762591.160 15 x 50 mm: 762592.500				

EC and VarioPrep columns in packs of 1, guard columns see below

Guard column systems						Guard column holder
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
** ChromCart® guard columns (pack of)	CC	8/3 (3)	8/3 (3)	8/4 (3)	8/4 (3)	721359
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
*** VarioPrep guard columns (pack of)	VP	10/8 (2)	10/16 (2)	15/32 (1)	15/50 (1)	
VP guard column holder		718251	718256	718253	718255	

Scale up factors and parameters for typical MN column dimensions

	4 x 250	8 x 250	10 x 250	16 x 250	21 x 250	32 x 250	40 x 250	50 x 250	80 x 250
Linear scale-up factor	1	4	6.25	16	28	64	100	161	400
Typical sample mass* [mg]	0.02-2	0.08-8	0.13-13	0.3-35	0.6-60	1.3-130	2-210	3-350	10-850
Typical flow rate [ml/min]	0.5-1.5	2-6	3-9	8-24	14-40	32-96	50-150	80-250	200-600

* For RP material; the maximum amounts given here always depend on the separation problem and on the sample composition. In some cases even half of the amounts given can cause drastic overload, in other cases the maximum amounts can be even higher still giving acceptable separations.



Packed columns

NUCLEODUR® C₁₈ ec

Pore size 110 Å; octadecyl phase, endcapped, 17.5% C;
eluent in column acetonitrile – water

Length →	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm					
NUCLEODUR® 100-3 C₁₈ ec											
particle size 3 µm											
EC analytical columns											
2 mm ID	760050.20		760054.20	760051.20	760053.20	760052.20					
3 mm ID	760050.30		760054.30	760051.30	760053.30	760052.30					
4 mm ID	760050.40		760054.40	760051.40	760053.40	760052.40					
4.6 mm ID	760050.46	760046.46	760054.46	760051.46	760053.46	760052.46					
EC guard columns*		4 x 2 mm: 761931.20		4 x 3 mm: 761931.30							
CC guard columns**		8 x 3 mm: 761005.30		8 x 4 mm: 761005.40							
NUCLEODUR® 100-5 C₁₈ ec											
particle size 5 µm											
Microbore analytical columns											
1 mm ID			717701.10	717700.10	717702.10	717703.10					
EC analytical columns											
2 mm ID	760004.20		760013.20	760001.20	760008.20	760002.20					
3 mm ID	760004.30		760013.30	760001.30	760008.30	760002.30					
4 mm ID	760004.40		760013.40	760001.40	760008.40	760002.40					
4.6 mm ID	760004.46	760035.46	760013.46	760001.46	760008.46	760002.46					
EC guard columns*		4 x 2 mm: 761932.20		4 x 3 mm: 761932.30							
CC guard columns**		8 x 3 mm: 761100.30		8 x 4 mm: 761100.40							
VarioPrep preparative columns											
10 mm ID	762003.100			762029.100		762022.100					
21 mm ID	762003.210			762029.210		762022.210					
32 mm ID						762022.320					
40 mm ID					762027.400	762022.400					
VP guard columns***		10 x 8 mm: 762090.80		10 x 16 mm: 762090.160							
		15 x 32 mm: 762311.320		15 x 50 mm: 762311.500							
NUCLEODUR® 100-10 C₁₈ ec											
particle size 10 µm											
VarioPrep preparative columns											
10 mm ID	762011.100			762302.100		762010.100					
21 mm ID	762011.210			762302.210		762010.210					
32 mm ID						762010.320					
40 mm ID					762303.400	762010.400					
50 mm ID						762010.500					
VP guard columns***		10 x 8 mm: 762090.80		10 x 16 mm: 762090.160							
		15 x 32 mm: 762311.320		15 x 50 mm: 762311.500							
Microbore, EC, and VarioPrep columns in packs of 1, guard columns see right											



Ordering information

NUCLEODUR® C₈ ec

Pore size 110 Å; octyl phase, endcapped, 10.5 % C;
eluent in column acetonitrile – water

Length →	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® 100-3 C₈ ec						particle size 3 µm
EC analytical columns						
2 mm ID	760063.20		760059.20	760060.20		760062.20
3 mm ID	760063.30		760059.30	760060.30		760062.30
4 mm ID	760063.40		760059.40	760060.40		760062.40
4.6 mm ID	760063.46	760064.46	760059.46	760060.46	760061.46	760062.46
EC guard columns*		4 x 2 mm: 761936.20		4 x 3 mm: 761936.30		
CC guard columns**		8 x 3 mm: 761012.30		8 x 4 mm: 761012.40		
NUCLEODUR® 100-5 C₈ ec						particle size 5 µm
EC analytical columns						
2 mm ID	760700.20		760704.20	760701.20		760703.20
3 mm ID	760700.30		760704.30	760701.30		760703.30
4 mm ID	760700.40		760704.40	760701.40		760703.40
4.6 mm ID	760700.46	760706.46	760704.46	760701.46	760702.46	760703.46
EC guard columns*		4 x 2 mm: 761937.20		4 x 3 mm: 761937.30		
CC guard columns**		8 x 3 mm: 761704.30		8 x 4 mm: 761704.40		
VarioPrep preparative columns						
10 mm ID	762072.100			762061.100		762062.100
21 mm ID	762072.210			762061.210		762062.210
32 mm ID						762062.320
40 mm ID					762079.400	762062.400
VP guard columns***	10 x 8 mm: 762092.80		10 x 16 mm: 762092.160		15 x 32 mm: 762321.320	
EC and VarioPrep columns in packs of 1, guard columns see below, Microbore columns with NUCLEODUR® C ₈ ec on request!						

Guard column systems					Guard column holder
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)
** ChromCart® guard columns (pack of)	CC	8/3 (3)	8/3 (3)	8/4 (3)	8/4 (3)
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm
*** VarioPrep guard columns (pack of)	VP	10/8 (2)	10/16 (2)	15/32 (1)	15/50 (1)
VP guard column holder		718251	718256	718253	718255



Packed columns

NUCLEODUR® HILIC

Pore size 110 Å; zwitterionic phase for HILIC chromatography, 7% C; eluent in column acetonitrile – water 80:20

Length →	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm		
NUCLEODUR® HILIC, 1.8 µm							particle size 1.8 µm		
EC columns									
2 mm ID	760521.20	760523.20	760525.20	760526.20			760528.20		
3 mm ID	760521.30	760523.30		760526.30					
4 mm ID	760521.40	760523.40		760526.40					
4.6 mm ID	760521.46	760523.46		760526.46					
EC guard columns*	4 x 2 mm: 761960.20		4 x 3 mm: 761960.30						
NUCLEODUR® HILIC, 3 µm							particle size 3 µm		
EC columns									
2 mm ID	760532.20		760534.20	760531.20			760530.20		
3 mm ID	760532.30		760534.30	760531.30			760530.30		
4 mm ID	760532.40		760534.40	760531.40			760530.40		
4.6 mm ID	760532.46		760534.46	760531.46	760533.46		760530.46		
EC guard columns*	4 x 2 mm: 761961.20		4 x 3 mm: 761961.30						
CC guard columns**	8 x 3 mm: 761580.30		8 x 4 mm: 761580.40						
NUCLEODUR® HILIC, 5 µm							particle size 5 µm		
EC columns									
2 mm ID	760552.20		760554.20	760551.20			760550.20		
3 mm ID	760552.30		760554.30	760551.30			760550.30		
4 mm ID	760552.40		760554.40	760551.40			760550.40		
4.6 mm ID	760552.46		760554.46	760551.46	760553.46		760550.46		
EC guard columns*	4 x 2 mm: 761962.20		4 x 3 mm: 761962.30						
CC guard columns**	8 x 3 mm: 761590.30		8 x 4 mm: 761590.40						

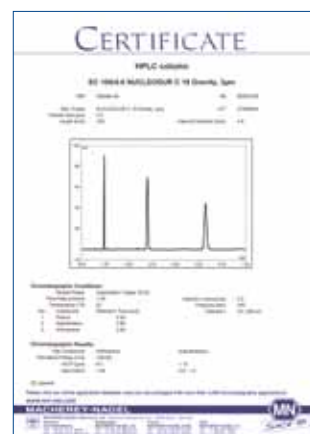
Columns in packs of 1, guard columns see right, Microbore columns and preparative columns with NUCLEODUR® HILIC on request!

Our HPLC QC policy

- **Highest production standard**
our facilities are EN ISO 9001:2008 certified
- **Strict quality specifications** for outstanding reliability
- **Perfect reproducibility** within each batch and from lot to lot
- Each column is individually tested and supplied with test chromatogram and test conditions

Test mixture for reversed phase columns

Designation	Pack of	REF
Test mixture for reversed phase columns in acetonitrile	1 ml	722394



Ordering information

NUCLEODUR® CN and CN-RP

Pore size 110 Å; cyano phase (nitrile), 7% C

Length →	50 mm	125 mm	150 mm	250 mm		
NUCLEODUR® 100-3 CN-RP		particle size 3 µm; eluent in column acetonitrile – water				
EC columns						
2 mm ID	760159.20	760157.20				
3 mm ID		760157.30				
4 mm ID			760156.40			
4.6 mm ID			760156.46			
EC guard columns*	4 x 2 mm: 761941.20		4 x 3 mm: 761941.30			
CC guard columns**	8 x 3 mm: 761430.30		8 x 4 mm: 761430.40			
NUCLEODUR® 100-5 CN-RP		particle size 5 µm; eluent in column acetonitrile – water				
EC columns						
4 mm ID		760153.40		760152.40		
4.6 mm ID		760153.46	760154.46	760152.46		
EC guard columns*	4 x 3 mm: 761944.30					
CC guard columns**	8 x 4 mm: 761420.40					
NUCLEODUR® 100-5 CN		particle size 5 µm; eluent in column n-heptane				
EC columns						
4 mm ID		760151.40	760149.40	760150.40		
4.6 mm ID		760151.46	760149.46	760150.46		
EC guard columns*	4 x 3 mm: 761943.30					
CC guard columns**	8 x 4 mm: 761419.40					
Columns in packs of 1, guard columns see below, Microbore columns and preparative columns with NUCLEODUR® CN / CN-RP on request!						

Guard column systems					Guard column holder
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)
** ChromCart® guard columns (pack of)	CC	8/3 (3)	8/3 (3)	8/4 (3)	8/4 (3)
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm
*** VarioPrep guard columns (pack of)	VP	10/8 (2)	10/16 (2)	15/32 (1)	15/50 (1)
VP guard column holder		718251	718256	718253	718255

Packed columns

NUCLEODUR® NH₂ and NH₂-RP

Pore size 110 Å; amino phase, 2.5 % C

Length →	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® 100-3 NH₂-RP particle size 3 µm; eluent in column acetonitrile – water				
EC columns				
2 mm ID	760740.20	760741.20		
4.6 mm ID			760742.46	760739.46
EC guard columns*	4 x 2 mm: 761951.20		4 x 3 mm: 761951.30	
CC guard columns**	8 x 3 mm: 761035.30		8 x 4 mm: 761035.40	
NUCLEODUR® 100-5 NH₂-RP particle size 5 µm; eluent in column acetonitrile – water				
EC columns				
2 mm ID		760730.20		760732.20
3 mm ID		760730.30		760732.30
4 mm ID		760730.40		760732.40
4.6 mm ID		760730.46	760731.46	760732.46
EC guard columns*	4 x 2 mm: 761953.20		4 x 3 mm: 761953.30	
CC guard columns**	8 x 3 mm: 761137.30		8 x 4 mm: 761137.40	
NUCLEODUR® 100-5 NH₂ particle size 5 µm; eluent in column n-heptane				
EC columns				
4 mm ID		760720.40		760722.40
4.6 mm ID		760720.46	760721.46	760722.46
EC guard columns*			4 x 3 mm: 761952.30	
CC guard columns**			8 x 4 mm: 761130.40	
Columns in packs of 1, guard columns see page 83, Microbore and preparative columns with NUCLEODUR® NH ₂ / NH ₂ -RP on request!				

Unmodified NUCLEODUR®

Pore size 110 Å; unmodified;
eluent in column n-heptane

Length →	50 mm	125 mm	150 mm	250 mm
NUCLEODUR® 100-3 particle size 3 µm				
EC analytical columns				
4.6 mm ID	760170.46		760172.46	760173.46
EC guard columns*			4 x 3 mm: 761966.30	
CC guard columns**			8 x 4 mm: 761007.40	
NUCLEODUR® 100-5 particle size 5 µm				
EC analytical columns				
4 mm ID				760007.40
4.6 mm ID	760023.46		760012.46	760007.46
EC guard columns*			4 x 3 mm: 761967.30	
CC guard columns**			8 x 4 mm: 761055.40	
VarioPrep preparative columns				
10 mm ID	762077.100	762078.100		762007.100
21 mm ID	762077.210	762078.210		762007.210
40 mm ID			762075.400	762007.400
VP guard columns***	10 x 8 mm: 762094.80	10 x 16 mm: 762094.160		15 x 32 mm: 762330.320
Columns in packs of 1, guard columns see page 83, Microbore columns with unmodified NUCLEODUR® on request!				



Ordering information

EC standard columns for analytical HPLC

Features

- Analytical column system manufactured from stainless steel
M 8 outer threads on both ends
Combination of sealing element and very fine-meshed stainless steel screen, PTFE ring and fitting adapter
Column heads SW 12, with inner threads M8 x 0.75 and UNF 10-32 (= 1/16" fitting)
- As screw-on guard column system we recommend the **Column Protection System** used with EC guard column cartridges with 4 mm length (see page 86).
- As built-in guard columns ChromCart® guard column cartridges with 8 mm length can be used with the guard column adapter EC (see page 87).
- Supplied with NUCLEODUR®, NUCLEOSHELL and NUCLEOSIL® spherical silicas



Available standard dimensions of EC columns

ID [mm]	20	30	50	75	Length [mm]					End fitting design
					100	125	150	200	250	
2	+	+	+	+	+	+	+	+	+	
3	+	+	+	+	+	+	+	+	+	
4	+	+	+	+	+	+	+	+	+	
4.6	+	+	+	+	+	+	+	+	+	

Please ask for availability of certain phases

Guard columns for EC columns

(pack of 3 each)	2 mm	EC columns with ID				Use guard column holder
		3 mm	4 mm	4.6 mm		
EC guard columns for Column Protection System guard column holder	EC	4/2	4/3	4/3	4/3	REF 718966
ChromCart® guard columns for EC guard column holder	CC	8/3	8/3	8/4	8/4	REF 721359

Guard column systems, accessories and replacement parts for EC columns • Ordering information

Description	Pack of	REF	
Column Protection System	1 kit	718966	see figures and description on page 86
Guard column adapter EC	1	721359	see figures and description on page 87
EC fitting adapter	1	718987	
EC column head (nut)	1	718988	
EC PTFE sealing ring	4	718992	
3-part sealing combination for EC columns	5 kits	718998	



Packed columns

Column Protection System

Innovative and universal screw-on guard column holder system
Suitable for all analytical HPLC columns with 1/16" fittings

Features

- Cartridges filled with specified NUCLEODUR®, NUCLEOSIL® and NUCLEOSHELL HPLC adsorbents
- Ideal protection for your analytical main column → significant increase in column lifetime
- Minimized void volume → suitable also for ultra fast HPLC
- Special ferrules → pressure stability up to 1034 bar (15 000 psi)
- Visual contamination check → in-time changing of the guard column
- Guard column length 4 mm, ID 2 mm (for main columns with 2 mm ID) or ID 3 mm (for main columns with 3, 4 and 4.6 mm ID)
- UNIVERSAL RP guard columns suitable for all HPLC columns under RP conditions



Content of the Column Protection System

Description	REF
Column Protection System	718966
Details	Content
Cartridge Holder	1
Capillaries	2
Ferrules	3
Wrenches	2
Manual	1

Replacement parts for the Column Protection System • Ordering information

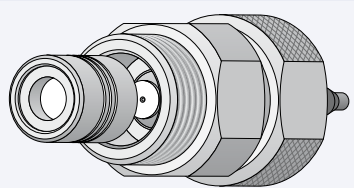
Description	Pack of	REF
Ferrules	5	718967
Replacement connector including O-ring	1	718968
Capillary tubes, nuts and metal ferrules	3	718969
Wrench (size 12 and 14 mm)	1	718970
EC 4/2 UNIVERSAL RP guard column (for main columns with 2 mm ID)	3	728777.20
EC 4/3 UNIVERSAL RP guard column (for main columns with 3, 4 and 4.6 mm ID)	3	728777.30

Visual Contamination Check

The cartridge is fitted with a white filter membrane:

A discoloration of the filter membrane usually indicates that the cartridge should be replaced.

In case of colorless contaminants, a rising back pressure and / or loss of chromatographic performance advise to change the guard column.

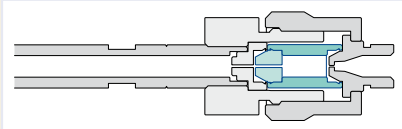


EC guard column adapter

Standard guard column built-in adapter system
Suitable for EC columns

Features

- Cartridges filled with specified NUCLEODUR® and NUCLEOSIL® HPLC adsorbents
- Ideal protection for your analytical EC main column → significant increase of column lifetime
- Guard column length 8 mm, ID 3 mm (for main columns with 2 and 3 mm ID) or ID 4 mm (for main columns with 4 and 4.6 mm ID)



EC guard column adapter • Ordering information

Description	Pack of	REF
EC guard column adapter	1	721359

Installation of the EC guard column adapter



1. Unscrew the column head
2. Remove the fitting
3. Unscrew the EC guard column adapter



4. Screw the adapter sleeve onto the column
5. Insert the CC guard column
6. Screw the nut of the guard column adapter in place



Packed columns

VarioPrep (VP) columns for preparative HPLC

Features

- Column system for preparative HPLC manufactured from stainless steel with two adjustable end fittings, e.g. for frequent use of back-flushing techniques
- Allows compensation of a dead volume, which could result at the column inlet after some time of operation, without need for opening the column
- Supplied with all NUCLEODUR® and NUCLEOSIL® spherical silicas

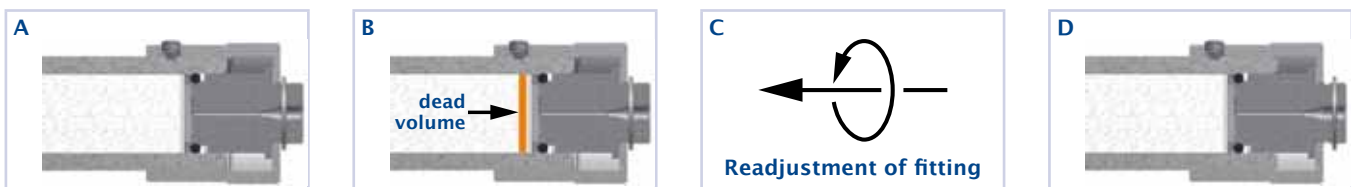


Available standard dimensions of VarioPrep columns with axially adjustable end fittings

ID [mm]	Length [mm]		Length [mm]						End fitting design
	10*	15*	50	75	100	125	150	250	
8	+		+		+	+	+	+	
10			+		+	+	+	+	
16	+		+		+	+	+	+	
21			+	+	+	+	+	+	
32		+			+		+	+	
40			+		+	+	+	+	+
50		+			+		+	+	
80							+	+	

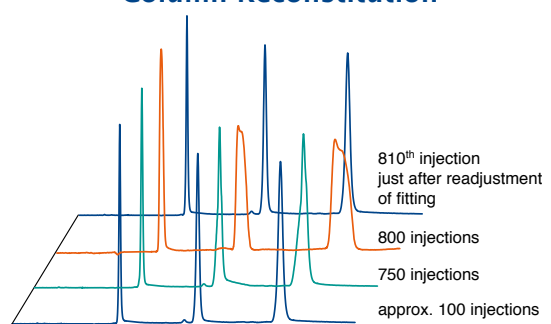
* 10 x 8, 10 x 16, 15 x 32 and 15 x 50 mm ID columns are used as guard columns and require adequate holders, see page 89.

The VarioPrep principle



Based on our special packing procedure VarioPrep columns are produced with highest packing quality and bed density (A). Due to intensively chemical and/or mechanical exposure of the column adsorbent, shrinking of the column bed can occur (B; orange gap). In this even unlikely case readjustment of the VarioPrep column fitting (C; turning the nut at column inlet clockwise) will eliminate the emerged dead volume (D). The performance of the VarioPrep column is completely reconstituted and column lifetime is significantly extended.

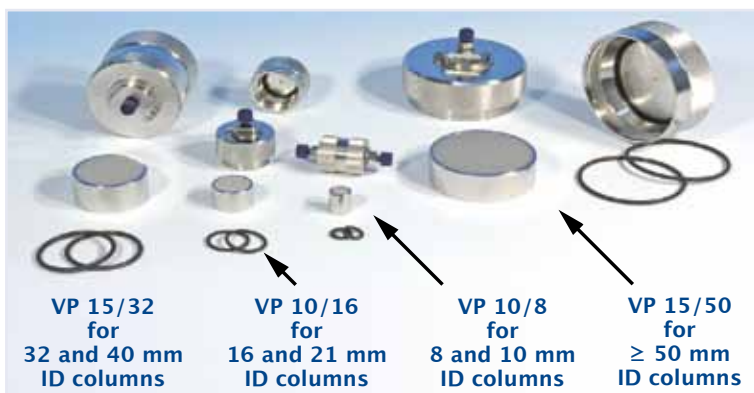
Column Reconstitution



Reconstitution of VarioPrep column performance

- Slight peak broadening and deformation after 800 injections under strongly demanding conditions (pH 11; 50 °C; sample in DMSO)
- Readjustment of the column fitting restores the column performance and prolongs column lifetime noticeably

Ordering information



The new guard column system for (semi-) preparative HPLC

- Easy handling and cartridge exchange
- Robust hardware
- Free rotary plunger fittings – low O-ring abrasion
- Cost-efficient cartridges
- Minimally invasive / no disturbance of the separation efficiency of main column
- Low back pressure
- Designed for pressures up to 400 bar

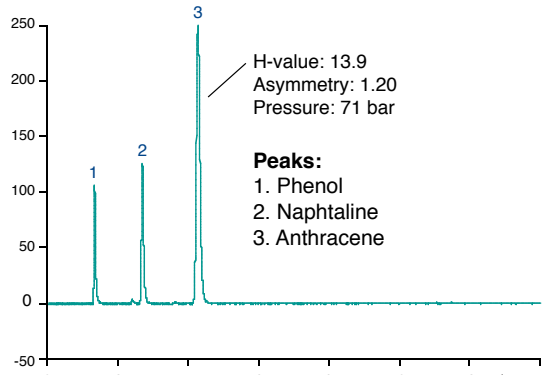
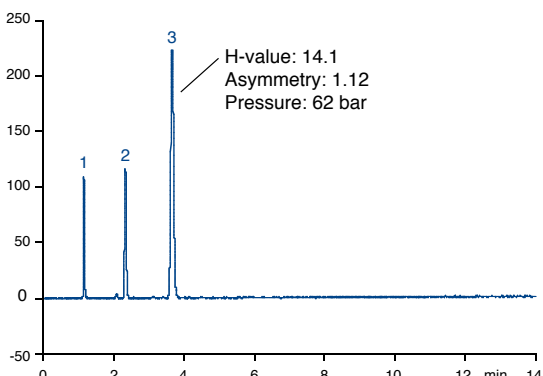
Performance of preparative column when used with VarioPrep guard column

Columns: 125 x 16 mm NUCLEODUR® C₁₈ HTec, 5 µm
125 x 16 mm NUCLEODUR® C₁₈ HTec, 5 µm + 10 x 16 mm NUCLEODUR® C₁₈ HTec guard column

Eluent: acetonitrile – water (80:20, v/v)

Flow rate: 16 mL/min

Temperature: 22 °C



Using VarioPrep guard columns provides ideal protection of your main column – symmetry, pressure and retention stay almost constant.

Technical data

	1/16" thread	free rotary plunger fittings – low O-ring abrasion	stainless steel			
Guard cartridge	Holder	Holder ID	Replacement O-ring (2 pcs)	Recommended for column ID	Preferred capillary ID	Typical flow rate
VP 10/8	REF 718251	8 mm	REF 718975	8 and 10 mm ID	0.17 and 0.25 mm	1-12 mL/min
VP 10/16	REF 718256	16 mm	REF 718976	16 and 21 mm ID	0.17, 0.25 and 0.5 mm	2-32 mL/min
VP 15/32	REF 718253	32 mm	REF 718977	32 and 40 mm ID	0.25, 0.5 and 1.0 mm	5-150 mL/min
VP 15/50	REF 718255	50 mm	REF 718978	≥ 50 mm ID	0.5 and 1.0 mm	20-250 mL/min

Guard columns for VarioPrep columns • Ordering information

		VP columns with ID		Use guard column holder	Holder ID		
		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm		
VP guard columns for VarioPrep guard column holder (pack of)	VP	10/8 (2)				REF 718251	8 mm
	VP		10/16 (2)			REF 718256	16 mm
	VP			15/32 (1)		REF 718253	32 mm
	VP				15/50 (1)	REF 718255	50 mm





HPLC



GC



TLC



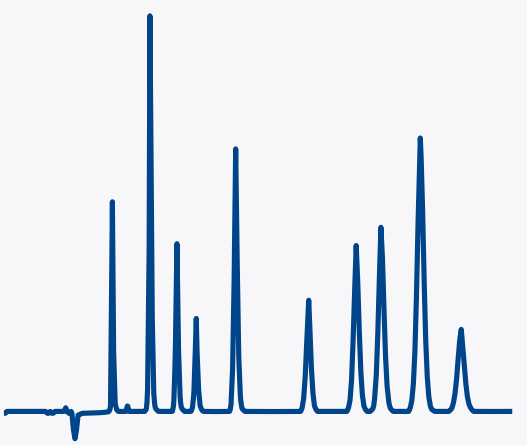
SPE & Flash



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MACHEREY-NAGEL GmbH & Co. KG · Neumann-Neander-Str. 6–8 · 52355 Düren · Germany

Germany

and international:

Tel.: +49 24 21 969-0

Fax: +49 24 21 969-199

E-mail: info@mn-net.com

Switzerland:

MACHEREY-NAGEL AG

Tel.: +41 62 388 55 00

Fax: +41 62 388 55 05

E-mail: sales-ch@mn-net.com

France:

MACHEREY-NAGEL EURL

Tel.: +33 388 68 22 68

Fax: +33 388 51 76 88

E-mail: sales-fr@mn-net.com

USA:

MACHEREY-NAGEL Inc.

Tel.: +1 484 821 0984

Fax: +1 484 821 1272

E-mail: sales-us@mn-net.com

