MACHEREY-NAGEL

















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High performance liquid chromatography (HPLC) is part of liquid chromatographic separating processes of substance mixtures and their analysis. At the beginning the technique was also called high pressure liquid chromatography due to the high back pressure of the column. HPLC offers qualitative (identification of substances) and quantitative (concentration determination) analysis by comparison with standard substances. The term HPLC was introduced in the 1970s, for the delineation of the high-performance method to the in the 1930s developed column liquid chromatography (column chromatography). At the beginning of the 21st century the HPLC was complemented by the even more efficient UHPLC (ultra high performance liquid chromatography). Hereby even higher pressures (> 400 bar) result in shorter analysis time and enhanced efficiency enabling a higher sample throughput with smaller sample volumes.

Application

HPLC/UHPLC is used additionally to gas chromatography (GC) for separation and determination of complex substance mixtures composed of low-volatile, polar and ionic, high-molecular or thermal instable substances. Therefore a sufficient solubility of the sample in a solvent or a solvent mixture is required. HPLC/UHPLC is used for purity control of chemicals and industrial products, determination of active agents for drug development, production and testing, environmental analytics, quality and purity control of foods, analysis of ingredients in cosmetics as well as for the isolation of biopolymers.

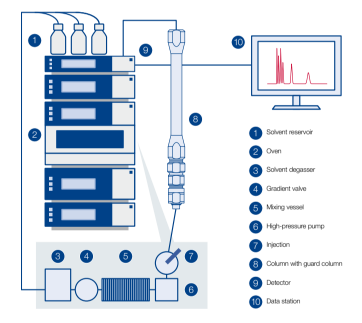
Basic principle

In liquid column chromatography a mobile phase (eluent) flows through a particle filled tube (separation column, stationary phase). In classic column chromatography this tube is a glass column with an inner diameter of several centimeters and a length up to 450 mm or even bigger. The filling material typically consists of coarse-grained particles like silica gel 60. The eluent is transported through the separation column either by hydrostatic pressure or a low-pressure pump with 1.5-2 bar.

In contrast HPLC columns consist of stainless steel with an inner diameter of 2-4.6 mm and a length of 20-300 mm. The column packing, mostly modified porous silica, has generally a particle size of 3, 5, 7 or 10 µm and a pore size of 50, 100, 120 (for low-molecular analytes) or 300-4000 Å (for high-molecular analytes). In UHPLC shorter columns in the range of 20-150 mm length with highly efficient particles of 1.8 µm size (sub-2 µm) are utilized. A guard column of a few millimeters length can be utilized and installed with a specific Column Protection System to increase the column lifetime. HPLC/UHPLC uses a high-pressure pump to transport the eluent from a storage vessel into the system with a column back pressure of up to 600/1200 bar.

Instrument

HPLC as well as UHPLC instruments have different building blocks. The storage vessel (eluent reservoir, 1) usually contains a deaerator unit (3) for the solvents. Followed by a gradient valve (4) with mixing chamber (5) in flow direction, which allows the usage of isocratic as well as gradient methods. A high-pressure pump (6) transports the sample into the system. The sample is injected via an injection valve (7). Usually this is operated automatically with a syringe by an autosampler. With the eluent flow the sample is transported to the guard and the separating column (8). For better reproducibility of the separation tempering with a column oven (2) should be performed. The separated substances are determined with a detector (9). In the resulting chromatogram each detector signal of a substance (peak), is related to the retention time of the column. With the data evaluation (10) these peaks can be identified and their concentration can be determined.



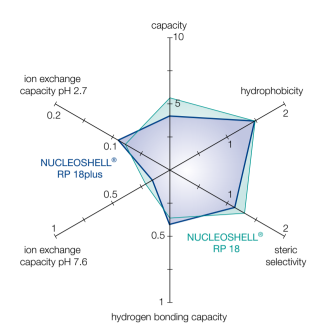


Separation mechanism

While flowing through the column each component of the solved mixture interacts differently with the stationary phase. According to the characteristics of the substance (hydrophobic, polar, ionic, aromatic, sterically hindered etc.) the strength of the interactions vary and thus the compounds are retained by the stationary phase in different ways. Essentially a distinction is drawn between normal phase (NP), reversed phase (RP) and ion exchange chromatography. Depending on the structure of the stationary phase diverse interactions e.g., van der Waals forces or π - π -stacking can occur and different polar mobile phases are required. For polar stationary normal phases (e.g., SiOH, CN, OH, NH₂) non-polar eluents like n-heptane, hexane, dichloromethane or 2-propanol are applicable. While for reversed phases (e.g., C₁₈, C₈, C₄, C₂, C₆H₅) typically polar RP eluents (e.g., acetonitrile or methanol with ultrapure water or buffer) and for ion exchange (e.g., SA, SB) aqueous buffers (e.g., phosphate, acetate, citric buffer) come to use.

Selectivity

The characteristic separation behavior of phases under certain conditions is also called selectivity. This is dependent on different parameters like structure and modifications of the base silica gel, nature of the chemical binding or the type of endcapping. In recent decades several methods have been developed to compare and distinguish the selectivity of various silica gels and their modifications. In this connection defined substances or substance classes are analyzed and the chromatographic parameters are graphically presented. A frequently applied model in specialist literature is e.g., the TANAKA plot, which allows a quick comparison of different HPLC phases. [4]



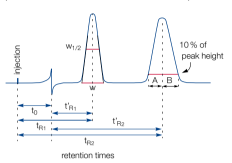
Parameter of the Tanaka diagram: Capacity = k' (pentylbenzene) Hydrophobicity = α (pentylbenzene, butylbenzene) Steric selectivity = α (triphenyl, o-terphenyl) Hydrogen bonding capacity (capacity of silanol) = α (caffeine, phenol) lon exchange capacity at pH 2.7 = α (benzylamine, phenol) lon exchange capacity at pH 7.6 = α (benzylamine, phenol)

The comparison of NUCLEOSHELL® RP 18 and NUCLEOSHELL RP® 18 plus for example shows a lower ion exchange capacity at pH 7.6 for the monomeric NUCLEOSHELL® RP 18 plus. The radar chart also reflects a more pronounced steric selectivity of NUCLEOSHELL® RP 18 due to a higher density of modifications with C_{18} chains.



Characteristic parameters

The success of a chromatographic separation depends apart from the stationary and mobile phase also on other characteristics like the quality of the separating column or the linear flow rate. The following schematic chromatogram illustrates the most important parameters which characterize a separation.



Schematic chromatogram

Peak width:	
W _{1/2}	peak width at half height
w	peak width of the peak (intersection point of the inflectional tangents with the zero line)
Peak symmetry:	
Α	peak front to peak maximum at 10 % of peak height
В	peak maximum to peak end at 10 % of peak height
Retention time::	
t _o	dead time of a column = retention time of a non-retarded substance
t _{R1} , t _{R2}	retention times of components 1 and 2
t' _{R1} , t' _{R2}	net retention times of components 1 and 2

In a chromatographic system the substances differ from each other in their retention time in or on the stationary phase. The time, which is needed by a sample component to migrate from column inlet (sample injection) to the column end (detector) is the retention time t_{R1} or t_{R2}. The dead time t₀ is the time required by an inert compound to migrate from column inlet to column end without any retardation by the stationary phase. Consequently, the dead time is identical with the retention time of the sample component remaining in the stationary phase. The difference of total retention time and dead time yields the net retention time t'_{B1} or t'_{B2} , which is the time a sample component remains in the stationary phase.

$$t'_{R1} = t_{R1} - t_0$$
 bzw. $t'_{R2} = t_{R2} - t_0$

To compare chromatograms that are recorded with columns of different lengths and internal diameters, as well as different flow rates, the retention time is converted into a dimensionless capacity factor k'.

$$k'_1 = \frac{t_{R1} - t_0}{t_0} \quad \text{bzw.} \quad k'_2 = \frac{t_{R2} - t_0}{t_0}$$

The relative retention a, also known as the separation factor, describes the ability of a chromatographic system (stationary and mobile phase) to distinguish between two compounds. This

is calculated from the rate of the capacity factors of the substances, where the figure in the denominator is the reference compound.

$$\alpha = \frac{k'_2}{k'_1}$$

The resolution R is a measure for the efficiency of the column to separate two substances. Besides the retention time t_B the peak width at half height $w_{1/2}$ is also included.

$$R = 1.18 \cdot \frac{t_{R2} - t_{R1}}{(W_{1/2})_2 + (W_{1/2})_1}$$

For practical reasons the peak symmetry is calculated at 10% of peak height. Ideally symmetry should be 1, i.e. A = B. Values > 1 indicate peak tailing, while values < 1 indicate peak fronting.

Peak symmetry
$$=\frac{B}{A}$$

Instead of the mobile phase volumetric flow rate [mL/min], which is controlled at the HPLC instrument, it is advantageous to use the linear velocity u [cm/sec]. The linear velocity is independent of the column cross section and proportional to the pressure drop in the column. The linear velocity can be calculated by means of the dead time, where L is the column length in cm and to the dead time in sec.

$$u = \frac{L}{t_0}$$

The quality of a column packing is determined through the number of theoretical plates N. High N values indicate a high capability to separate complex sample mixtures.

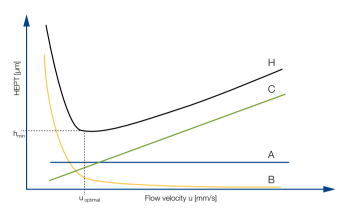
$$N = 5.54 \cdot \left(\frac{t_{R1}}{w_{1/2}}\right)^2$$

The value of the height equivalent to a theoretical plate HEPT is a criterion for the quality of a column. HEPT, is the length, in which the chromatographic equilibrium between mobile and stationary phase has been adjusted once. Its value depends on the particle size, the flow velocity, the mobile phase viscosity and especially on the packing quality. Small HEPT values, meaning a large number of theoretical plates N, facilitate the column to separate complex sample mixtures.

$$H = \frac{L}{N}$$

The Van Deemter equation shows the dependence of the HEPT on the velocity u.

$$H = A + \frac{B}{U} + C \cdot u$$



A term = eddy-diffusion, B term = longitudinal diffusion coefficient, C term = mass transfer coefficient, H = HEPT = height equivalent to a theoretical plate

The A term, also called eddy-diffusion, is a function of the particle size, the B term a function of the diffusion coefficient of the substance in the mobile phase and the C term the retardation

of a substance by the interface between stationary and mobile phase. In the point of intersection of h_{min} and u_{opt} the optimal separation efficiency for a column with high peak symmetry for the separated substances is obtained.

Column quality

Each HPLC/UHPLC column of MACHEREY-NAGEL is individually tested according to the most important characteristic parameters in quality control and the results are documented in a certificate of analysis.

Detailed information of the particular properties of the high-purity silica phases NUCLEODUR®, of the established standard silica NUCLEOSIL® and the modern Core-Shell material NUCLEOSHELL® as well as phases for special separations and the equivalent HPLC- and UHPLC-columns can be found on the following pages.



Strict quality specifications for outstanding reliability

- Highest production standard our facilities are EN ISO 9001:2008 certified
- · Perfect reproducibility from batch to batch and within each lot
- Each column is individually tested and supplied with test chromatogram and test conditions.

Test mixture* for reversed phase columns in acetonitrile, pack of 1 mL REF 722394



Furthermore custom-packed columns with different column types, dimensions and particle sizes are available on request.

^{*} This product (REF 722394) contains harmful substances which must be specially labeled as hazardous. For detailed information please see SDS.





USP spe	cification of MN HPLC phases		
Code	Specification	MN HPLC Phases	Page
		NUCLEODUR® C ₁₈ ec	181
		NUCLEODUR® C ₁₈ Gravity	158
		NUCLEODUR® C ₁₈ Gravity-SB	162
		NUCLEODUR® C ₁₈ HTec	178
		NUCLEODUR® C ₁₈ Isis	164
		NUCLEODUR® C ₁₈ PAH	227
		NUCLEODUR® C ₁₈ Pyramid	166
		NUCLEODUR® PolarTec	168
USP L1	octadecyl silane chemically bonded to porous silica particles 1.5 to 10 μm diameter, or monolithic silica gel	NUCLEODUR® Sphinx RP	176
		NUCLEOSHELL® RP 18	200
		NUCLEOSHELL® RP 18plus	202
		NUCLEOSIL® C ₁₈	214
		NUCLEOSIL® C ₁₈ AB	214
		NUCLEOSIL® C ₁₈ HD	214
		NUCLEOSIL® C ₁₈ MPN	243
		NUCLEOSIL® C ₁₈ PAH	229
		NUCLEOSIL® C ₁₈ PPN	244
		NUCLEODUR® SiOH	190
USP L3	porous silica particles, 1.5 to 10 μm diameter, or monolithic silica gel	NUCLEOSIL® SiOH	224
		NUCLEODUR® C ₈ ec	181
	octyl silane chemically bonded to totally porous silica particles,	NUCLEODUR® C ₈ Gravity	158
USP L7	1.8 to 10 μm diameter	NUCLEOSIL® C ₈	217
		NUCLEOSIL® C ₈ HD	217
		NUCLEODUR® NH ₂ /NH ₂ -RP	188
USP L8	an essentially monomolecular layer of aminopropyl silane chemically bonded to totally porous silica gel support, 1.5 to 10 µm diameter	NUCLEOSIL® Carbohydrate	246
	Since 30. Support, no to 10 pm diameter	NUCLEOSIL® NH ₂ /NH ₂ -RP	221
USP L9	irregular or spherical, totally porous silica gel having a chemically bonded, strongly acidic cation-exchange coating, 3 to 10 µm diameter	NUCLEOSIL® SA	223
		NUCLEODUR® CN/CN-RP	186
USP L10	nitrile groups chemically bonded to porous silica particles, 1.5 to 10 μ m diameter	NUCLEOSIL® CN/CN-RP	222



Code	specification of MN HPLC phases Specification	MN HPLC Phases	Page
Code	Specification	NUCLEODUR® Phenyl-Hexyl	170
		NUCL FODUR® π ²	172
1100144	phonyl graypo chamically handed to parago cilica particles 1.5 to 10 um diameter		-
USP L11	phenyl groups chemically bonded to porous silica particles, 1.5 to 10 μm diameter	NUCLEOSHELL® Phenyl-Hexyl	204
		NUCLEODUR® Sphinx RP	176
		NUCLEOSIL® C ₆ H ₅	220
USP L14	silica gel having a chemically bonded, strongly basic quaternary ammonium anion-exchange coating, 5 to 10 μm diameter	NUCLEOSIL® SB	223
USP L16	dimethylsilane chemically bonded to porous silica particles, 5 to 10 μm diameter	NUCLEOSIL® C ₂	219
LICD L 17	strong cation-exchange resin consisting of sulfonated cross-linked PS/DVB copolymer in the H	NUCLEOGEL® ION 300 OA	248
USP L17	form, 6 to 12 µm diameter	NUCLEOGEL® SUGAR 810 H	247
	strong cation-exchange resin consisting of sulfonated cross-linked PS/DVB copolymer in the Ca	NUCLEOGEL® SUGAR 810 Ca	247
USP L19	form, 5 to 15 µm particle size	NUCLEOGEL® SUGAR Ca	248
USP L20	dihydroxypropane groups chemically bonded to porous silica particles, 5 to 10 µm diameter	NUCLEOSIL® OH (Diol)	220
USP L21	a rigid, spherical styrene-divinylbenzene copolymer, 5 to 10 μm diameter	NUCLEOGEL® RP	245
USP L22	a cation-exchange resin made of porous polystyrene gel with sulfonic acid groups, about 10 μm NUCLEOGEL® SCX		240
USP L23	an anion-exchange resin made of porous polymethacrylate or polyacrylate gel with quaternary ammonium groups, about 10 µm in size	aternary NUCLEOGEL® SAX	
	butyl silane chemically bonded to totally porous silica particles, 5 to 10 µm diameter	NUCLEODUR® C ₄ ec	241
USP L26		NUCLEOSIL® C ₄	219
		NUCLEOSIL® C ₄ MPN	243
USP L32	a chiral ligand-exchange resin packing · L-proline copper complex covalently bonded to irregular shaped silica particles, 5 to 10 µm diameter	NUCLEOSIL® CHIRAL-1	235
USP L34	strong cation-exchange resin consisting of sulfonated cross-linked PS-DVB copolymer in the Pb form, 5 to 7 µm particle size	NUCLEOGEL® SUGAR Pb	248
USP L36	a 3,5-dinitrobenzoyl derivative of L-phenylglycine covalently bonded to 5 μm aminopropyl silica	NUCLEOSIL® CHIRAL-3	236
USP L40	cellulose tris-(3,5-dimethylphenylcarbamate) coated porous silica particles, 5 to 20 µm diameter	NUCLEOCEL DELTA	233
	pentafluorophenyl groups chemically bonded to silica particles by a propyl spacer, 1.5 to 10 µm	NUCLEODUR® PFP	174
USP L43	diameter	NUCLEOSHELL® PFP	206
USP L45	beta-cyclodextrin bonded to porous silica particles, R,S-hydroxypropyl ether derivative, 3 to 10 µm diameter	NUCLEODEX β-OH, β-PM	231
USP L58	strong cation-exchange resin consisting of sulfonated cross-linked PS/DVB copolymer in the Na form, 6 to 30 μ m diameter	NUCLEOGEL® SUGAR Na	248
LICD L CO	spherical porous silica gel, particle size of 10 µm diameter or smaller, the surface of which has	NUCLEODUR® PolarTec	168
USP L60	been covalently modified with alkyl amide groups and endcapped	NUCLEOSIL® C ₁₈ Nautilus	214
USP L75	A chiral-recognition protein, bovine serum albumin (BSA), chemically bonded to silica particles, about 7 μ m in diameter, with a pore size of 300 Angstrom	RESOLVOSIL BSA-7	234



NUCLEODUR® high purity silica for HPLC

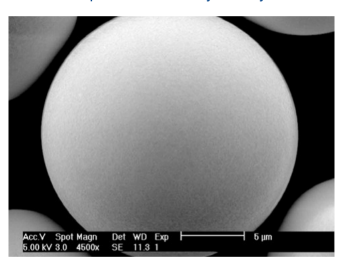


NUCLEODUR® is a fully synthetical type B silica (silica of 3rd generation) offering highly advanced physical properties like totally spherical particle shape, outstanding surface microstructure, high pressure stability and low metal content.

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.

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.

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				
Aluminum	< 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.

NUCLEODUR® silica is available with two pore sizes – 110 Å pore size as standard material and as 300 Å widepore material for the separation of biomolecules, like peptides and proteins.

Physical data of NUCLEODUR®					
	Standard	Widepore			
Pore size	110	300 Å			
Surface area (BET)	340 m²/g	100 m²/g			
Pore volume	0.9 mL/g	0.9 mL/g			
Density	0.47 g/mL	0.47 g/mL			

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.

For a summary of important properties of our NUCLEODUR® phases please see page 152.

1.8 µm particles for increased separation efficiency

Key feature

- · Decrease of analysis time (ultra fast HPLC)
- Shorter columns with high separation efficiency and significant improvement of resolution and detection sensitivity
- · Suitable for LC/MS due to low bleeding characteristics

Fractionation

 \cdot NUCLEODUR $^{\! B}$ 1.8 μm particles are fractionated to limit the increase in back pressure.

Availability

• The following NUCLEODUR® phases are available in 1.8 µm:

 $\rm C_{18}$ Gravity, $\rm C_8$ Gravity, $\rm C_{18}$ Gravity-SB, $\rm C_{18}$ Isis, $\rm C_{18}$ Pyramid, PolarTec, Phenyl-Hexyl, PFP, Sphinx RP, $\rm C_{18}$ HTec and HILIC

Advantages of 1.8 µm particle size

Miniaturization started in the early stage of HPLC with the reduction of particle size from 10 μm via 7 μm to standard 5 μm – still the most used particle diameter in analytical HPLC – to 3 μm spherical particles. With the introduction of 1.8 μm NUCLEODUR® particles researchers have turned over a new leaf in HPLC column technology, featuring extraordinary improvements in terms of plate numbers, column efficiency and resolution compared with 3 μm particles.

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

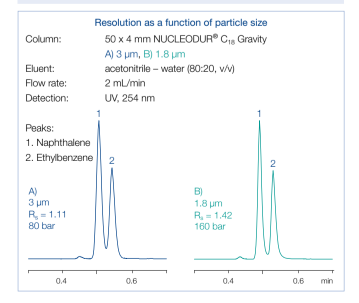
- · 50 x 4.6 mm NUCLEODUR® C₁₈ Gravity
- · 3 µm: N ≥ 100 000 plates/m (h-value≤ 10)
- 1.8 µm: N ≥ 166 667 plates/m (h-value≤ 6)

Increase of the plate number by $\sim 67\,\%$ offers the possibility of using shorter columns with equal plate number resulting in a decrease of analysis time.

Significant improvement in resolution

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

 R_s = resolution, α = selectivity (separation factor), k_i ' = retention N = plate number with $N \propto 1/d_P$, d_P = particle diameter



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

Column back pressure

Due to the smaller particles 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), LC = column length, η = viscosity, u = linear velocity, d_P = particle diameter

The high sphericity of the NUCLEODUR® particles and the very narrow particle size distribution allow to keep the back pressure on a moderate level.

Comparison of back pressures

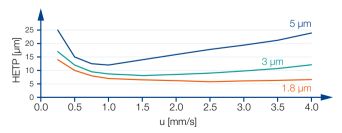
Eluent 100 % methanol, flow rate 1.5 mL/min temperature 22 °C, column dimensions 50 x 4.6 mm

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

Higher flow rates and shorter run times

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

Van Deemter curves



Column 50 x 4.6 mm, acetonitrile - water (50:50, v/v), analyte toluene

Technical requirements

To gain best results with 1.8 μ m particles certain technical demands must be met including pumps for flow rates of 2–3 mL with pressures of 250–1000 bar, minimized dead volume, and fast data recording.



ase	Specification	Page	Characteristic*	Stability	Structure
	octadecyl, high density coating, multi-endcapping 18 % C · USP L1	158	A •••••	pH 1–11, suitable for LC/MS	(Si-O ₂),
C ₁₈ Gravity			C •••		Z
	octadecyl (monomeric),	100	A ••••	 pH 1–9,	
₁₈ Gravity-SB	extensive endcapping 13 % C · USP L1	162	C -	suitable for LC/MS	NUCLEODUR®
	octyl, high density coating,		A •••		DUR®
C. Orașitu	multi-endcapping 11 % C · USP L7	158	B • C	pH 1–11, suitable for LC/MS	NUCLEODUR® (Si-O ₂) _n
C ₈ Gravity	octadecyl phase with specially crosslinked surface modification endcapping 20 % C · USP L1	164	A •••••• B •• C •••••	pH 1–10, suitable for LC/MS	NUCLEODUR® (Si-O ₂) _n
C ₁₈ Pyramid	octadecyl with polar endcapping 14 % C · USP L1	166	A • • • • B • • • C • • •	stable in 100 % aqueous eluent, pH 1–9, suitable for LC/MS	NUCLEODUR®
PolarTec	octadecyl with embedded polar group 17 % C · USP L1 and L60	168	A • • • • • B • • • • C • • • • • • • • •	stable in 100 % aqueous eluent, pH 1–9, suitable for LC/MS	NUCLEODUR® (Si-O ₂) (Si-O ₂) (Si(CH ^a) ^a)
Phenyl-Hexyl	phenylhexyl, multi-endcapping 10 % C · USP L11	170	A •• B ••• C •	pH 1–10, suitable for LC/MS	NUCLEODUR® (SI-O ₂) (SI-O ₂) (SI-O ₂)
π²	biphenylpropyl, multi-endcapping 17 % C · USP L11	172	A ••• B •••• C •••	pH 1.5–10	NUCLEODUR® (Si-O ₂), (Si-O ₂), (Si-O ₂),





Application	Similar phases**	Interactions · retention mecl	nanism
in general compounds with ionizable functional groups such as basic pharmaceuticals and pesticides	NUCLEOSIL® C ₁₈ HD Xterra® RP18 / MS C18; Luna® C18(2), Gemini®, Synergi® Max RP; Zorbax® Extend-C18; Inertsil® ODS III; Purospher® STAR RP-18; Hypersil™ BDS	hydrophobic (van der Waals interactions)	Si(CH ₃) ₃
overall sophisticated analytical separations, especially for polar compounds, e.g., antibiotics, water-soluble vitamins, organic acids	-	hydrophobic (van der Waals interactions) with additional polar inter- actions	Si-O-Si(CH ₃) ₃ H ₃ C
like C ₁₈ Gravity, however, generally shorter retention times for nonpolar compounds	NUCLEOSIL® C ₈ HD Xterra® RP8 / MS C8; Luna® C8; Zorbax® Eclipse XDB-C8	hydrophobic (van der Waals interactions)	OH CH ₃ OH CH ₃
high steric selectivity, thus suited for separation of positional and structural isomers, planar / nonplanar molecules	NUCLEOSIL® C ₁₈ AB Inertsil® ODS-P; Pro C18 RS	steric and hydrophobic	
basic pharmaceuticals, very polar compounds, organic acids	Aqua, Synergi® Hydro-RP; AQ; Atlantis® dC18; Polaris® C18-A	hydrophobic and polar (H bonds)	OH CH ₃ H ₃ C O
basic pharmaceuticals, organic acids, pesticides, amino acids, water-soluble vitamins	NUCLEOSIL® C ₁₈ Nautilus ProntoSIL® C18 AQ, Zorbax® Bonus-RP, Polaris® Amide-C18; Ascentis® RP Amide, SymmetryShield™ RP18; SUPELCOSIL™ LC-ABZ+; HyPURITY™ ADVANCE; ACCLAIM Polar AD.II	hydrophobic and polar (H bonds)	Si(CH ₃) HO
aromatic and unsaturated com- pounds, polar compounds like pharmaceuticals, antibiotics	Luna® Phenyl-Hexyl; Zorbax® Eclipse Plus Phenyl-Hexyl; Kromasil® Phenyl-Hexyl	π- $π$ and hydrophobic	O ₂ N
aromatic and unsaturated com- pounds, polar compounds like pharmaceuticals, antibiotics	Pinnacle® DB Biphenyl; Ultra Biphenyl	π-π and hydrophobic	O ₂ N
** phases which provide a similar	selectivity based on chemical and physical propertie	es	



е	Specification	Page	Characteristic*	Stability	Structure
	pentafluorophenylpropyl,		A ••	 nL 1 0	PD The second of
	multi-endcapping 8 % C · USP L43	174	В ••••	pH 1-9, suitable for LC/MS	(Si-O) (Si-O) (Si-O) (Si(CH ₃) ₃
PFP			C ••••		
	bifunctional, balanced ratio of		A •••	<u>.</u>	£
	propylphenyl and octadecyl, endcapping 15 % C · USP L1 and L11	176	B •••	pH 1–10, suitable for LC/MS	(Si-O ₂),
phinx RP	10 /0 0 001 ET and ETT		C		Z ₹
			A ••••		В
	octadecyl, high density coating, high capacity, multi-endcapping 18 % C · USP L1	178	В	pH 1-11, suitable for LC/MS	NUCLEODUR®
C ₁₈ HTec			C •••		N ,
	octadecyl, medium density,	endcapping Ivailable in 110 Å and 300 Å 181 B ● pH 1–9 Up pore size	® E		
	endcapping available in 110 Å and 300 Å pore size			•	(S)
C ₁₈ ec	17.5 % / 4 % C · USP L1		C ••••		Z
			A ••		e
	octyl, medium density, endcap- ping 10.5 % C · USP L7	181	B ••	pH 1–9	NUCLEODUR®
C ₈ ec			C •••		Ž
			Α •		ë E
	butyl, medium density, endcap- ping, 300 Å pore size 2.5 % C · USP L26	181	В ●●	pH 1–9	NUCLEODUR (Si-O) (Si-O) (Si-O) (Si(CH ^a) ^a
C ₄ ec			C ••		N Superings
			Α •		е Сн ₃ \
	zwitterionic ammonium – sulfonic acid phase 7 % C	184	В ••••	pH 2–8.5	NUCLEODURG (Si-OH) CH3 So.6 (CH3 So.6 (CH3 So.6 (CH3 So.6 (CH3 So.6 (CH3 So.6)
HILIC			C -	-	Z & H S - OH S 'S
		A •		B	
	cyano (nitrile) for NP and RP separations	186	В ••••	pH 1–8, stable towards highly	CEN CEN CEN CEN SI-OH SI-O-SI(CH ₃) ₃
N/CN-RP	7 % C · USP L10		C -	aqueous mobile phases	Sir ⁰ Si(CH ₃) ₃





Application	Similar phases**	Interactions · retention mech	hanism
aromatic and unsaturated com- pounds, halogen compounds, phenols, isomers, polar pharma- ceuticals, antibiotics	ACQUITY® CSH Fluoro-Phenyl; Hypersil™ GOLD PFP; Luna® PFP(2); Discovery® HS F5; Allure® PFP Propyl; Ultra II PFP Propyl	polar (H bond), dipole-dipole, π-π and hydrophobic	F F F
compounds with aromatic and multiple bond systems	no similar phases	π-π and hydrophobic	NO ₂
robust and well base deactivated C_{18} phase; all separation tasks with preparative potential	Xterra® RP18/MS C18/SunFire™ C18; Luna® C18(2), Gemini®, Synergi® Max RP; Zorbax® Extend-C18; Inertsil® ODS III; Purospher® STAR RP-18; Hypersil® BDS	hydrophobic (van der Waals interactions)	Si(CH ₃) ₃ H ₃ C O
robust C ₁₈ phase for routine analyses	NUCLEOSIL® C ₁₈ Spherisorb® ODS II; Symmetry® C18; Hypersil® ODS; Inertsil® ODS II; Kromasil® C18; LiChrospher® RP-18	hydrophobic (van der Waals interactions) some residual silanol interactions	SI(CH ₃) ₃ CH ₃
robust C_8 phase for routine analyses	NUCLEOSIL® C ₈ ec / C ₈ Spherisorb® C8; Symmetry® C8; Hypersil® MOS; Kromasil® C8; LiChrospher® RP-8	hydrophobic (van der Waals interactions) some residual silanol interactions	SIOH N CH ₃ SIOH CH ₃ SIOH CH ₃
biological macromolecules like proteins or peptides	Jupiter® C4; ACE® C4	hydrophobic (van der Waals interactions) some residual silanol interactions	SIOH R ₂
hydrophilic compounds such as polar organic acids and bases, polar natural compounds	Sequant™ ZIC®-HILIC; Obelisc™	ionic/ hydrophilic and electrost- atic	H ₃ C O CH ₃ O CH ₃ O CH ₃ NH NH ₂ SO ₃ O CH ₃ SO ₃
polar organic compounds (basic drugs), molecules containing π-electron systems	NUCLEOSIL® CN/CN-RP	π-π and polar (H bond), hydrophobic	C N HO



$ \rfloor \setminus $	V	U	L	

Phase	Specification	Page	Characteristic*	Stability	Structure
	aminopropyl for NP and RP separations 2.5 % C · USP L8		Α •		e E
		188	В ●●●●	pH 2–8, stable towards highly aqueous mobile phases	NUCLEODUR ₀ (Si-OH NH ² NH ²
NH ₂ /NH ₂ -RP			C -		OOZ
	unmodified high purity silica · USP L3		Α -	<u>.</u>	® ₩
		190	В -	pH 2–8	NUCLEODUR® (SI-O ₂) _n 90 - 9: 9: 9: 9: 9: 9: 9: 9: 9: 9: 9: 9: 9:
SiOH			C -		ON **





Application	Similar phases**	Interactions · retention mech	nanism
sugars, sugar alcohols and other hydroxy compounds, DNA ba- ses, polar compounds in general	NUCLEOSIL® NH ₂ /NH ₂ -RP	polar/ionic and hydro- phobic	NH ₃
polar compounds in general	NUCLEOSIL® SIOH	polar/ionic	SIOH \leftrightarrow O ₂ N $-$

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

NUCLEODUR® C₁₈ Gravity · C₈ Gravity nonpolar high density phase · USP L1 (C₁₈) · USP L7 (C₈)

Kev feature

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

Technical data

- Available as octadecyl (C_{18}) and octyl (C_{8}), multi-endcapped
- \cdot Pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm for $C_{18},$ 1.8 and 5 µm for $C_{8};$ 7, 10, 12 and 16 µm particles for preparative purposes on request
- \cdot Carbon content 18 % for $C_{18},\,11$ % for C_{8}

Recommended application

- Overall sophisticated analytical separations
- Compound classes separated include pharmaceuticals, e.g., analgesics, anti-inflammatory drugs, antidepressants; herbicides; phytopharmaceuticals; immunosuppressants

Base deactivation

NUCLEODUR® C_{18} Gravity and NUCLEODUR® C_{8} Gravity are based on the ultrapure NUCLEODUR® silica. Derivatization generates a homogeneous surface with a high density of bonded silanes (~18 % C for C_{18} , ~11 % C for C_{8}). 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. Even strongly basic pharmaceuticals like amitriptyline are eluted without tailing under isocratic conditions. For a discussion of the different retention behavior of C_{18} phases compared to C_{8} phases see page 182.

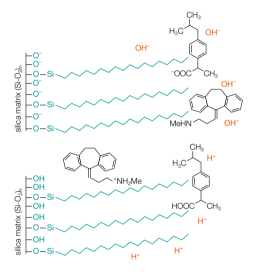
Enhanced pH stability

One major disadvantage of silica stationary phases is limited stability at strongly acidic or basic pH. Cleavage of the siloxane bonding by hydrolysis, or dissolution of the silica will rapidly lead to a considerable loss in column performance. 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® C_{18} and C_{8} Gravity allow for use at an expanded pH range from pH 1 to 11.

Benefits of enhanced pH stability

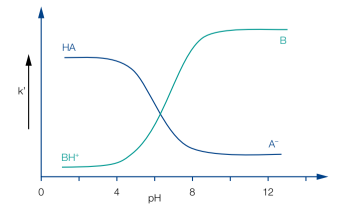
An expanded pH range is often required in method development. Many nitrogen containing compounds like basic drugs are protonated at acidic or neutral pH and exhibit poor retention on a standard C_{18} phase. The retention behavior can be improved by working at a higher pH, where the analyte is no longer 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



NUCLEODUR® columns

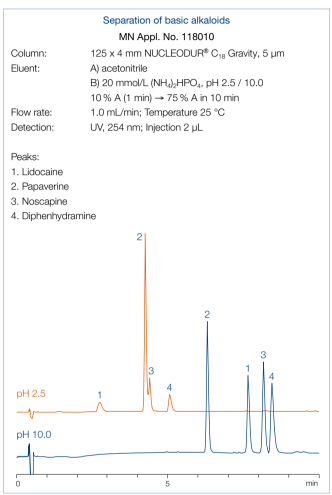


An example how selectivity can be controlled by pH is the separation of the acid ketoprofen, the base lidocaine and benzamide. Under acidic conditions the protonated lidocaine is eluted very fast due to lack of sufficiently strong hydrophobic interactions between analyte and C_{18} chains, while the formally neutral ketoprofen is eluted after about 3 min. However, at pH 10 a reversal of the elution order, with a visibly longer retention time for the basic lidocaine, is observed.

Influence of the pH value on selectivity MN Appl. No. 120860 Column: 125 x 4 mm NUCLEODUR® C_{18} Gravity, 5 μm Eluent: A) acetonitrile - 10 mmol/L ammonium formate, pH 3.0 (50:50, v/v); B) acetonitrile - 10 mmol/L ammonium bicarbonate, pH 10.0 (50:50, v/v) Flow rate: 1.0 mL/min 30 °C Temperature: UV, 230 nm Detection: Injection: 2 μL Peaks: 1. Lidocaine 2. Benzamide 3. Ketoprofen рН 3 pH 10 min

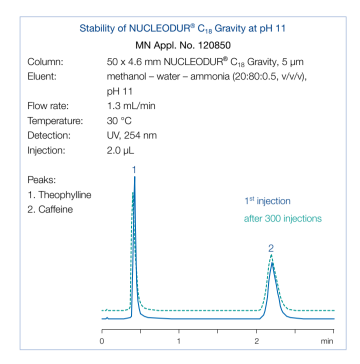
As mentioned above, pH stability of the stationary phase can be helpful for improving selectivity in method development. The following figure shows the separation of 4 basic drugs under acidic and basic conditions.

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.



The following chromatogram demonstrates the stability of NUCLEODUR® C_{18} Gravity under alkaline conditions. The ultrapure Gravity with its unique high density surface bonding technology withstands strong alkaline mobile phase conditions.



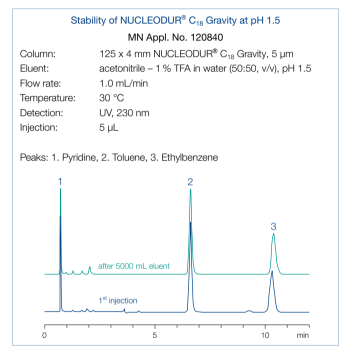


Even after 300 injections no loss of column efficiency – identified, e.g., by peak broadening or decrease in retention times – could be observed.

Under alkaline conditions dissolution of the silica support is possible, resulting in dead volume and thus peak broadening. 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® C_{18} Gravity in acidic conditions. Retention times of all three compounds in the column performance test remain 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.



Ordering informa	tion							
Eluent in column ace	etonitrile – w	ater						
	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C	3 ₁₈ Gravity	, 1.8 µm octa	decyl phase, part	icle size 1.8 µm,	18 % C · UHPLC			
Analytical EC column	S							
	2 mm	760078.20	760079.20	760071.20	760076.20		760075.20	
	3 mm	760078.30	760079.30		760076.30			
	4 mm	760078.40	760079.40		760076.40			
	4.6 mm	760078.46	760079.46		760076.46			
EC guard columns*			4 x 2 mm:	761901.20	4 x 3 mm:	761901.30		
NUCLEODUR® C	318 Gravity	, 3 µm octade	cyl phase, particl	e size 3 μm, 18 %	5 C			
Analytical EC column	S							
	2 mm		760080.20		760084.20	760081.20	760083.20	760082.20
	3 mm	•	760080.30	•	760084.30	760081.30	760083.30	760082.30
	4 mm		760080.40		760084.40	760081.40	760083.40	760082.40
	4.6 mm		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		



NUCLEODUR® columns



Eluent in column ac	ID	Length →						
	טו	20 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® (C ₁₈ Gravity	ν, 5 μm octade	cyl phase, particl	e size 5 µm, 18 9	% C			
nalytical EC colum	ns	•		·				
,	2 mm		760102.20		760104.20	760100.20	760103.20	760101.20
	3 mm	•	760102.30	•••••	760104.30	760100.30	760103.30	760101.30
	4 mm	••••••	760102.40	•••••	760104.40	760100.40	760103.40	760101.40
	4.6 mm	•	760102.46	760106.46	760104.46	760100.46	760103.46	760101.46
C guard columns*			4 x 2 mm:	761903.20	4 x 3 mm	: 761903.30		
reparative VarioPre	p columns							
	10 mm		762103.100			762109.100)	762113.100
	21 mm		762103.210			762109.210)	762113.210
	32 mm							762113.320
	40 mm	•					762100.400	762113.400
P guard columns	***************************************	•	10 x 8 mm:	762160.80	10 x 16 m	m: 762160.160	15 x 32 mm	n: 762163.320
*								
IUCLEODUR® (/, 10 μm octad	lecyl phase, parti	cle size 10 µm, 1	8 % C			
reparative VarioPre	•							760050 010
	21 mm			·····				762250.210
P guard columns *	40 mm	<u>.</u>			40 40	m: 762160.160	45 00	762250.400 n: 762163.320
luent in column ac	ation cetonitrile – w ID	/ater Length →						
luent in column ac	cetonitrile – w		50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
	cetonitrile – w ID	Length → 30 mm				125 mm	150 mm	250 mm
NUCLEODUR® (D ID C ₈ Gravity,	Length → 30 mm				125 mm	150 mm	250 mm
IUCLEODUR® (D ID C ₈ Gravity,	Length → 30 mm				125 mm	150 mm 760759.20	250 mm
NUCLEODUR® (ID C ₈ Gravity,	Length → 30 mm	phase, particle siz	ze 1.8 µm, 11 %	C · UHPLC	125 mm		250 mm
NUCLEODUR® (cetonitrile – w ID C ₈ Gravity, ns 2 mm	Length → 30 mm 1.8 µm octyl 760756.20	phase, particle siz	ze 1.8 µm, 11 %	C · UHPLC 760757.20	125 mm		250 mm
NUCLEODUR® (C ₈ Gravity, ns 2 mm 3 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30	phase, particle si: 760755.20 760755.30	ze 1.8 µm, 11 %	C · UHPLC 760757.20 760757.30	125 mm		250 mm
IUCLEODUR® (nalytical EC column	C ₈ Gravity, ns 2 mm 3 mm 4 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40	phase, particle six 760755.20 760755.30 760755.40 760755.46	ze 1.8 µm, 11 %	760757.20 760757.30 760757.40 760757.46	125 mm 125 mm		250 mm
IUCLEODUR® (nalytical EC column	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm:	ze 1.8 μm, 11 % 760760.20 761905.20	760757.20 760757.30 760757.40 760757.46			250 mm
IUCLEODUR® (nalytical EC column C guard columns*	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm:	ze 1.8 μm, 11 % 760760.20 761905.20	760757.20 760757.30 760757.40 760757.46			250 mm
NUCLEODUR® (nalytical EC column C guard columns*	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm:	ze 1.8 μm, 11 % 760760.20 761905.20	760757.20 760757.30 760757.40 760757.46			250 mm 760753.20
NUCLEODUR® (nalytical EC column C guard columns*	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity,	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm:	ze 1.8 μm, 11 % 760760.20 761905.20	C · UHPLC 760757.20 760757.30 760757.40 760757.46 4 x 3 mm	: 761905.30	760759.20	
NUCLEODUR® (nalytical EC column C guard columns*	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40	: 761905.30 760751.20 760751.30 760751.40	760759.20 760752.20	760753.20
NUCLEODUR® (analytical EC columns) C guard columns* NUCLEODUR® (analytical EC columns)	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 760750.46	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46	: 761905.30 760751.20 760751.30 760751.40 760751.46	760759.20 760752.20 760752.30	760753.20 760753.30
NUCLEODUR® (analytical EC columns* C guard columns* NUCLEODUR® (analytical EC columns*)	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 4 mm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 760750.46	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46	: 761905.30 760751.20 760751.30 760751.40	760759.20 760752.20 760752.30 760752.40	760753.20 760753.30 760753.40
NUCLEODUR® (nalytical EC column C guard columns* NUCLEODUR® (nalytical EC column C guard columns*	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 4 nm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 760750.46 4 x 2 mm:	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46	: 761905.30 760751.20 760751.30 760751.40 760751.46 : 761907.30	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
NUCLEODUR® (analytical EC columns* C guard columns* NUCLEODUR® (analytical EC columns*)	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 1 mm 4 mm 4.6 mm 2 mm 3 mm 4	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 760750.46 4 x 2 mm:	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46	760751.20 760751.30 760751.40 760751.46 : 761907.30	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
C guard columns* C guard columns* C guard columns* C guard columns*	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 4 nm 4.6 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 4 x 2 mm: 762081.100 762081.210	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.100 762071.210	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
IUCLEODUR® Inalytical EC column C guard columns* IUCLEODUR® Inalytical EC column C guard columns* Treparative VarioPre	Cancella Canada	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 4 x 2 mm: 762081.100 762081.210 10 x 8 mm:	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	760751.20 760751.30 760751.40 760751.46 : 761907.30	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
Analytical EC column EC guard columns* NUCLEODUR® Analytical EC column EC guard columns* Preparative VarioPre //P guard columns *	Cancella Canada	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 4 x 2 mm: 762081.100 762081.210 10 x 8 mm:	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.100 762071.210	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
NUCLEODUR® Analytical EC column CG guard columns* NUCLEODUR® Analytical EC column CG guard columns* Preparative VarioPre (P guard columns * CG and VarioPrep columns *	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 3 mm 4.6 mm 10 mm 21 mm 21 mm 21 mm 21 mm	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 4 x 2 mm: 762081.100 762081.210 10 x 8 mm:	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.100 762071.210	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
NUCLEODUR® Analytical EC column EC guard columns* NUCLEODUR® Analytical EC column EC guard columns* Preparative VarioPre EC and VarioPrep columns Guard columns *	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm 2 mm 3 mm 4	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 4 x 2 mm: 762081.100 762081.210 10 x 8 mm:	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	: 761905.30 760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.100 762097.160	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46
NUCLEODUR® Analytical EC column EC guard columns* NUCLEODUR® Analytical EC column EC guard columns* Preparative VarioPre EC and VarioPrep columns Guard columns for	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm 2 mm 3 mm 4	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 760750.46 4 x 2 mm: 762081.100 762081.210 10 x 8 mm: plumns see below.	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20 762097.80	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	: 761905.30 760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.210 m: 762097.160	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46 762070.100 762070.210
NUCLEODUR® Analytical EC column C guard columns* NUCLEODUR® Analytical EC column C guard columns* Preparative VarioPre C and VarioPrep columns Guard columns for Column Protection	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm 2 mm 3 mm 4	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph cks of 1, guard cc with ID k of)	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: ase, particle size 760750.20 760750.30 760750.40 760750.46 4 x 2 mm: 762081.100 762081.210 10 x 8 mm: plumns see below.	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20 762097.80	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm	: 761905.30 760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.210 n: 762097.160	760759.20 760752.20 760752.30 760752.40 760752.46	760753.20 760753.30 760753.40 760753.46 762070.100 762070.210
NUCLEODUR® Analytical EC columns* NUCLEODUR® Analytical EC columns* NUCLEODUR® Analytical EC columns* Preparative VarioPre Analytical EC columns* Columns for Column stor Column Protection Guard columns for VP guard colu	C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm C ₈ Gravity, ns 2 mm 3 mm 4 mm 4.6 mm 2 mm 3 mm 4	Length → 30 mm 1.8 µm octyl 760756.20 760756.30 760756.40 760756.46 5 µm octyl ph cks of 1, guard cc with ID k of)	760755.20 760755.30 760755.40 760755.46 4 x 2 mm: rase, particle size 760750.20 760750.30 760750.40 760750.40 760750.46 4 x 2 mm: 762081.100 762081.210 10 x 8 mm: plumns see below.	ze 1.8 μm, 11 % 760760.20 761905.20 5 μm, 11 % C 760749.46 761907.20 762097.80 3 m 3) 4/3 mm 16,	760757.20 760757.30 760757.40 760757.46 4 x 3 mm 760754.20 760754.30 760754.40 760754.46 4 x 3 mm 10 x 16 mm 10 x 16 mm (3) 4/3 21 mm 32	: 761905.30 760751.20 760751.30 760751.40 760751.46 : 761907.30 762071.100 762071.210 n: 762097.160 nm 4 3 (3) 4 40 mm ≥	760759.20 760752.20 760752.30 760752.40 760752.46) 762082.210 .6 mm Gu /3 (3) 71	760753.20 760753.30 760753.40 760753.46 762070.100 762070.210

For details of our column systems see page 250.

NUCLEODUR® C₁₈ Gravity-SB hydrophobic phase with polar selectivity · USP L1

Key feature

- Hydrophobic C₁₈ phase with distinct polar selectivity, ideal for method development, better retention of early eluting substances
- Excellent performance under highly aqueous conditions
- Suitable for LC/MS due to low bleeding characteristics

Technical data

- Monomeric octadecyl modification, extensive endcapping
- Pore size 110 Å; available particle sizes 1.8 μm, 3 μm and 5 μm; carbon content 13 %; pH stability 1–9

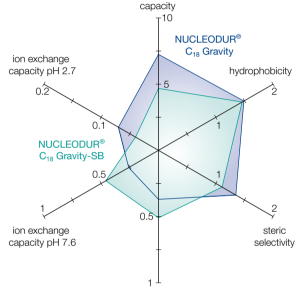
Recommended application

 Overall sophisticated analytical separations, especially for polar compounds, e.g., antibiotics, water-soluble vitamins, organic acids

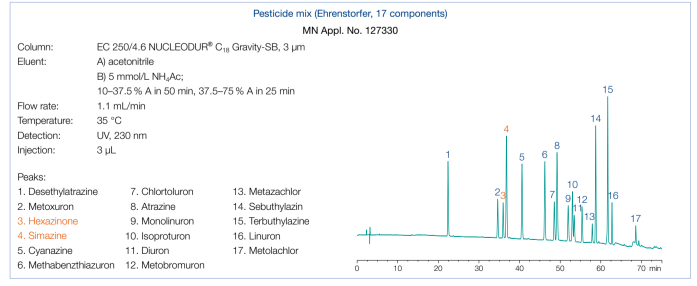
NUCLEODUR® C_{18} Gravity-SB excels with a relatively high hydrophobicity – similar to C_{18} Gravity – while simultaneously showing distinctive polar selectivity, without having polar embedded groups or polar endcapping. As a result the column displays better retention of early eluting analytes and high performance under strongly aqueous conditions. Additionally the column is suitable for LC/MS due to low bleeding characteristics. These features are achieved through side chains (isobutyl) of the monomeric C_{18} phase.

In the TANAKA plot the NUCLEODUR® Gravity-SB shows similar hydrophobicity than the Gravity, however with a reduced capacity. The ion exchange capacity under basic conditions (pH 7.6) is high, which favors good retention of early eluting, polar substances.

Due to the broad selectivity and stability the base deactivated NUCLEODUR® C_{18} Gravity-SB is versatile applicable, especially for polar analytes like nucleobases or pesticides the column shows good separation efficiency.



hydrogen bonding capacity



Good separation of the critical pair hexazinone/simazine





MN Appl. No. 127270

EC 150/4.6 mm Columns:

> NUCLEODUR® C_{18} Gravity-SB, 5 μm NUCLEODUR® C_{18} Gravity, 5 μm NUCLEODUR® C₁₈ Pyramid, 5 µm

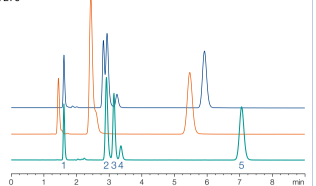
25 mmol/L KH_2PO_4 , pH 3 – methanol (95:5, v/v) Eluent:

Flow rate: 1.0 mL/min, Temperature: 20 °C Detection: UV, 220 nm, Injection: 2.5 µL (1 mg/mL)

Peaks:

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

3. Uracil



Better resolution of early eluting analyte

Ordering information

Eluent in column acetonitrile - water

	ID	Length →						
		30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C	C ₁₈ Gravity	/-SB, 1.8 μm	particle size 1.8 µ	m · UHPLC				
Analytical EC column	s							
	2 mm	760591.20	760593.20	760595.20	760596.20		760598.20	
	3 mm	760591.30	760593.30		760596.30			
	4 mm	760591.40	760593.40		760596.40			
	4.6 mm	760591.46	760593.46		760596.46			
EC guard columns*			4 x 2 mm:	761990.20	4 x 3 mm: 7	31990.30		
NUCLEODUR® C	C ₁₈ Gravity	/-SB, 3 μm pa	article size 3 µm					

NUCLEODUR®	C_{18} Gravity-SB, 3 μ m	particle size 3 µm
------------	--------------------------------	--------------------

Analytical EC columns

2 mm	760603.20	760606.20	760607.20	760608.20	760609.20
 3 mm	760603.30	760606.30	760607.30	760608.30	760609.30
4 mm	760603.40	760606.40	760607.40	760608.40	760609.40
4.6 mm	760603.46 760605.46	760606.46	760607.46	760608.46	760609.46
	•	•		······	· · · · · · · · · · · · · · · · · · ·

EC guard columns* 4 x 2 mm: 761991.20 4 x 3 mm: 761991.30

NUCLEODUR® C₁₈ Gravity-SB, 5 μm particle size 5 μm

Analytical EC columns

,							
	2 mm	760613.20		760616.20	760617.20	760618.20	760619.20
	3 mm	760613.30		760616.30	760617.30	760618.30	760619.30
	4 mm	760613.40		760616.40	760617.40	760618.40	760619.40
	4.6 mm	760613.46	760615.46	760616.46	760617.46	760618.46	760619.46
EC guard columns*	•	4 x 2 mm:	761992.20		761992.30		
Preparative VarioPrep	p columns						
	10 mm	762350.100			762351.100		762353.100
	21 mm	762350.210			762351.210		762353.210
	32 mm						762353.320
	10 mm	•	•		•	762352 400	762252 400

10 x 8 mm: 762354.80

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

Guard column systems

VP guard columns **

Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	Guard column holder
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
** VP 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	

10 x 16 mm: 762354.160

For details of our column systems see page 250.

15 x 32 mm: 762355.320

NUCLEODUR® C₁₈ Isis phase with high steric selectivity · USP L1

Key feature

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

Technical data

 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

Surface modification

By use of specific C_{18} silanes and polymeric bonding technologies a dense shield of alkyl chains protects the subjacent silica matrix. Elemental analysis of NUCLEODUR® C_{18} Isis shows a carbon load of 20%. The target crosslinking of the C_{18} 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.

Slot Model

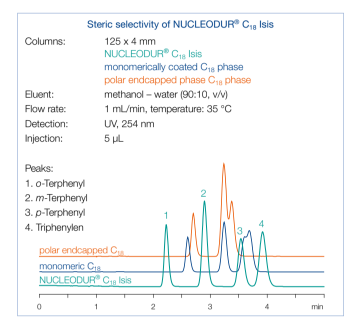
Sander and Wise [5] proposed a model for the retention of aromatic compounds based on molecular shape, which is referred to as "Slot Model". This model pictures the bonded C_{18} phase on the silica surface with slots which the analytes have to penetrate during retention. Planar molecules are able to penetrate these slots deeper than non-planar molecules of similar molecular weight and length-to-width ratio. Thus triphenylene (lower structure) is longer retained than o-terphenyl (upper structure).



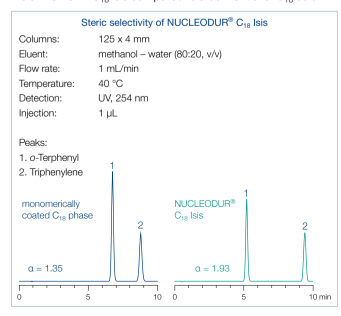


Steric selectivity

The following chromatograms reveal the improved resolution for positional isomers in a test mixture of aromatic compounds on NUCLEODUR® C_{18} Isis (green) in direct comparison with monomerically coated (blue) and polar endcapped (orange) C_{18} columns.



The separation of o-terphenyl and triphenylene is a good example to evaluate the selectivity of a RP column in terms of the shape of two molecules. The phenyl rings of o-terphenyl are twisted out of plane while triphenylene has a planar geometry. The separation factor α is a measure for the steric selectivity. As is shown below the α value is considerable larger on NUCLEODUR $^{\!8}$ C $_{\!18}$ Isis compared to a conventional C $_{\!18}$ column.





The surface bonding technology also provides improved stability features for the NUCLEODUR® C_{18} Isis phase.

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 application 121210 at www.mn-net.com/apps).

Ordering informa	tion							
Eluent in column ace	etonitrile – w	vater						
	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C	; ₁₈ Isis, 1.8	8 µm particle s	ize 1.8 μm · UHP	LC				
Analytical EC column	S							
	2 mm	760406.20	760405.20	760396.20	760407.20		760409.20	
	3 mm	760406.30	760405.30		760407.30			
	4 mm	760406.40	760405.40		760407.40			
	4.6 mm	760406.46	760405.46		760407.46	***************************************		
EC guard columns*	•	••••	4 x 2 mm:	761910.20	4 x 3 mm:	761910.30		
NUCLEODUR® C	; ₁₈ Isis, 3	µm particle size	e 3 µm					
Analytical EC columns								
•	2 mm		760400.20		760401.20	760402.20	760403.20	760404.20
	3 mm	•••••	760400.30	•••••	760401.30	760402.30	760403.30	760404.30
	4 mm		760400.40		760401.40	760402.40	760403.40	760404.40
	4.6 mm		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		
NUCLEODUR® C	3 ₁₈ Isis, 5	µm particle size	e 5 µm					
Analytical EC columns	S							
,	2 mm		760410.20		760415.20	760412.20	760413.20	760414.20
	3 mm		760410.30		760415.30	760412.30	760413.30	760414.30
	4 mm	····	760410.40		760415.40	760412.40	760413.40	760414.40
	4.6 mm	····•	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	•	
Preparative VarioPrep	columns							
	10 mm		762404.100			762405.100		762403.100
——————————————————————————————————————	21 mm		762404.210			762405.210		762403.210
	32 mm							762403.320
	40 mm						762406.400	762403.400
VP guard columns **			10 x 8 mm:	762420.80	10 x 16 mm	n: 762420.160	15 x 32 mm	: 762422.320
EC and VarioPrep col	umns in pac	cks of 1, guard co	olumns see below.					

Guard column systems						
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	Guard column holder
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
** VP 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	

For details of our column systems see page 250.

NUCLEODUR® C18 Pyramid phase for highly aqueous eluents · USP L1

Kev feature

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

Technical data

· Special phase with polar endcapping: pore size 110 Å: particle sizes 1.8 μ m, 3 μ m and 5 μ m (7 and 10 μ m 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

RP-HPLC with highly aqueous mobile phases

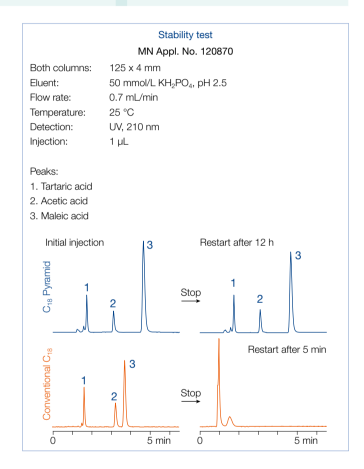
The efforts to neutralize unwanted silanol activity often results in well base-deactivated RP phases with high carbon load, but a limited scope of selectivity beyond non-polar interactions. Polar compounds like carboxylic acids or drug metabolites show only weak retention on densely bonded RP columns 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 [6].

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.

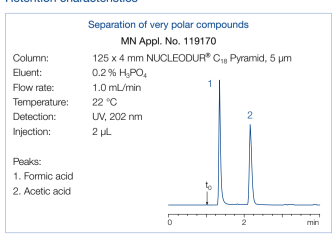
Stability features

NUCLEODUR® C₁₈ Pyramid is a silica phase with hydrophilic endcapping, designed especially for use in eluent systems of up to 100% water. The upper figure shows the retention behavior of tartaric, acetic and maleic acid under purely aqueous conditions on NUCLEODUR® C18 Pyramid in comparison with a conventionally bonded C₁₈ 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 h, whilst the performance of the conventional RP column already collapsed totally after 5 min.



Retention characteristics





NUCLEODUR® columns



The polar surface exhibits retention characteristics different from conventional C_{18} phases. Application 119170 shows the improved retention behavior of the very polar short chain organic acids, which are insufficiently retained on RP columns with predominantly hydrophobic surface properties. In addition to the exceptional polar selectivity NUCLEODUR® C_{18} Pyramid also provides adequate hydrophobic retention (see applicati-

on No. 19190 at www.mn-net.com). 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 at www.mn-net.com/apps).

	ID	Length →						
	טו	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C	C ₁₈ Pyrami	id, 1.8 μm par	ticle size 1.8 µm	· UHPLC				
Analytical EC column	S							
	2 mm	760271.20	760272.20	760275.20	760273.20		760274.20	
	3 mm	760271.30	760272.30		760273.30			
	4 mm	760271.40	760272.40		760273.40			
	4.6 mm	760271.46	760272.46	•••••	760273.46	•••••	•••••	
EC guard columns*	•	•	4 x 2 mm:	761915.20	4 x 3 mm:	761915.30		
NUCLEODUR® C	C ₁₈ Pyrami	d, 3 µm partic	le size 3 µm					
Analytical EC column	S							
	2 mm		760263.20		760264.20	760260.20	760261.20	760262.20
	3 mm	•	760263.30		760264.30	760260.30	760261.30	760262.30
	4 mm	•	760263.40		760264.40	760260.40	760261.40	760262.40
	4.6 mm	•	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		
NUCLEODUR® C	C ₁₈ Pyrami	d, 5 µm partic	le size 5 µm					
Analytical EC column	S	-						
	2 mm		760200.20		760204.20	760201.20	760203.20	760202.20
	3 mm		760200.30		760204.30	760201.30	760203.30	760202.30
	4 mm	***************************************	760200.40	••••••	760204.40	760201.40	760203.40	760202.40
	4.6 mm	•	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		
Preparative VarioPrep	columns							
	10 mm		762271.100			762273.100		762272.100
	21 mm		762271.210			762273.210		762272.210
	32 mm							762272.320
	40 mm						762269.400	762272.400
VP guard columns **			10 x 8 mm:	762291.80	10 x 16 mm	n: 762291.160	15 x 32 mm	: 762293.320

Guard column systems						
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	Guard column holder
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
** VP 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	•

For details of our column systems see page 250.

NUCLEODUR® PolarTec RP phase with embedded polar group · USP L1 and L60

Kev feature

- · Excellent base deactivation
- Suitable for LC/MS and 100 % aqueous eluents
- · Pronounced steric selectivity

Technical data

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

✓ Recommended application

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

RP-HPLC under 100 % aqueous conditions

The dominant form of interactions of conventional C_{18} phases are nonpolar London dispersion forces. Besides nonpolar interactions phases with embedded polar groups possess the ability to show polar interactions (dipole-dipole, hydrogen bonds, π - π , etc.). These interactions enhance retention and selectivity for polar compounds like carboxylic acids, phenols and nitrogen containing compounds.

Separation of histidines

MN Appl. No. 125140

Column: 150 x 3 mm NUCLEODUR® PolarTec, 3 µm
Eluent: 1.0 mmol/L perfluoropentanoic acid in water –

0.5 mmol/L perfluoropentanoic acid in acetonitrile

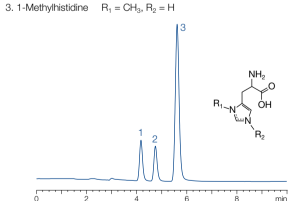
(99.5:0.5, v/v)

Flow rate: 0.4 mL/min
Temperature: 20 °C
Detection: UV, 230 nm

Peaks:

1. 3-Methylhistidine $R_1 = H$, $R_2 = CH_3$

2. Histidine $R_1 = R_2 = H$

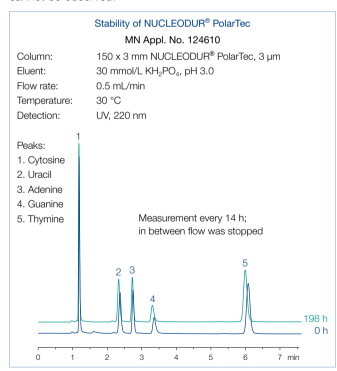


In order to increase retention for polar compounds it is often necessary to decrease the organic ratio of the mobile phase to zero. Under these conditions many conventional C_{18} phases display the so-called dewetting effect which means that the mobile phase is expelled from the pores. This phenomenon 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.

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.

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

Even after days or weeks of operation in purely aqueous eluents the C_{18} chains of NUCLEODUR® PolarTec are neither folded nor show any collapsing. A significant reduction of retention time cannot be observed.



In spite of the polar character of the embedded functional group NUCLEODUR® PolarTec exhibits sufficient hydrophobic properties and is very well suited for analyzing basic compounds.



NUCLEODUR® columns



Ordering informa										
Eluent in column ac	etonitrile – w ID	/ater Length →								
	ID	20 mm	50 mm	75	mm	100 mm	125 mm	150 m	nm	250 mm
NUCLEODUR® F	PolarTec, 1	.8 µm particle	size 1.8 µ	m · UHPLC						
Analytical EC column	ns	-								
•	2 mm	760461.20	760463	3.20 760	0465.20	760466.20)	76046	88.20	
	3 mm	760461.30	760463	3.30	•••••	760466.30)			
	4 mm	760461.40	760463	3.40		760466.40)			
	4.6 mm	760461.46	760463	3.46		760466.46	3	•		
EC guard columns*			4 x	2 mm: 76198	30.20	4 x 3 n	nm: 761980.30	-		
NUCLEODUR® F	PolarTec, 3	β μm particle si	ze 3 µm							
Analytical EC column	ns									
	2 mm		760473	3.20		760476.20	760477.	20 76047	'8.20	760479.20
	3 mm		760473	3.30		760476.30	760477.	30 76047	'8.30	760479.30
	4 mm		760473	3.40	•••••	760476.40	760477.	40 76047	'8.40	760479.40
	4.6 mm		760473	3.46 760	0475.46	760476.46	760477.	46 76047	'8.46	760479.46
EC guard columns*	•	••••	4 x	2 mm: 76198	31.20	4 x 3 n	nm: 761981.30			
NUCLEODUR® F	PolarTec, 5	µm particle si	ze 5 µm							
Analytical EC column										
, , , , , , , , , , , , , , , , , , , ,	2 mm		760483	3.20		760486.20	760487.	20 76048	88.20	760489.20
	3 mm	••••••	760483	3.30		760486.30	760487.	30 76048	38.30	760489.30
	4 mm	••••••	760483	3.40		760486.40	760487.	40 76048	38.40	760489.40
	4.6 mm		760483	3.46 760	0485.46	760486.46	760487.	46 76048	38.46	760489.46
EC guard columns*	····		4 x	2 mm: 76198	32.20	4 x 3 n	nm: 761982.30			
Preparative VarioPrep	o columns									
	10 mm	<u>.</u>	762220).100			762221.	100		762223.100
	21 mm	····	762220).210			762221.:	210		762223.210
——~L\\\\\\\\\\	32 mm	····						·····		762223.320
	40 mm	···•					·····	76222	22.400	762223.400
VP guard columns **	:			8 mm: 76222	24.80	10 x 16	mm: 762224.16	30 15	x 32 mm	762226.320
EC and VarioPrep co	lumns in pad	cks of 1, guard co	olumns see	below.						
Guard column sy	ystems									
Guard columns for E	EC columns	with ID		2 mm	3 mm	۱ 4	4 mm	4.6 mm	Gua	ard column holder
* Column Protection			EC	4/2 (3)	4/3 (3	<u> </u>	4/3 (3)	4/3 (3)	718	966
Guard columns for \		lumns with ID		8, 10 mm			32, 40 mm	≥ 50 mm		
** VP guard columns			VP	10/8 (2)	10/16		15/32 (1)	15/50 (1)		
VP guard column ho	lder			718251	7182	56	718253	718255		

For details of our column systems see page 250.

NUCLEODUR® Phenyl-Hexyl productive for polar/aromatic compunds · USP L11

Kev feature

- · Hydrophobic phase with alternative selectivity compared to classical C₁₈ modifications
- · Separation principle based on 2 retention mechanisms: π - π interactions and hydrophobic interactions
- · Suitable for LC/MS due to low bleeding characteristics

Technical data

· Phase with phenyl-hexyl modification and multi-endcapping; pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm; carbon content 10 %; pH stability 1-10

Recommended application

· Aromatic and unsaturated compounds, polar compounds like pharmaceuticals, antibiotics

Phenylhexyl modified phases are an interesting alternative to classical C₁₈ phases due to an excellent separation of aromatic and unsaturated compounds especially with electron withdrawing groups.

The combination of hydrophobic and polar π - π interactions result in an interesting and alternate selectivity in comparison to C₁₈ and C₈ modified phases.

Through short phenylhexyl chains the NUCLEODUR® Phenyl-Hexyl is more polar than the bifunctional modified NUCLEODUR® Sphinx RP. Therefore shorter analysis times can be achieved with mixtures of structural similar aromatic and aliphatic unsaturated compounds.

With NUCLEODUR® Phenyl-Hexyl e.g., tricyclic antidepressants or water soluble vitamins can be separated in good resolution.

Separation of water-soluble vitamins on NUCLEODUR® Phenyl-Hexyl

MN Appl. No. 125920

Column: 100 x 3 mm NUCLEODUR® Phenyl-Hexyl, 3 µm

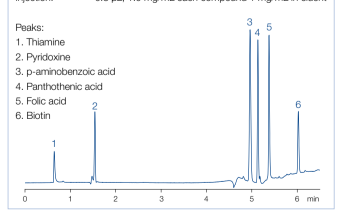
Eluent: A) 0.1 % phosphoric acid in water

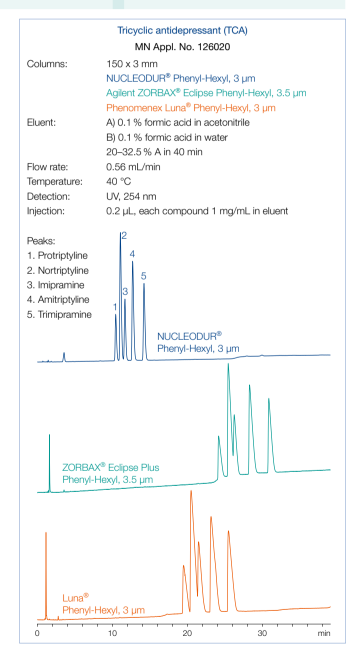
B) 0.1 % phosphoric acid in acetonitrile

0 % B for 2 min, then to 60 % B in 7 min

Flow rate: 0.56 mL/min Temperature: 35 °C Detection: UV, 215 nm

0.8 µL, 1.0 mg/mL each compound 1 mg/mL in eluent Injection:







NUCLEODUR® columns



Ordering inform										
Eluent in column ac										
	ID	Length → 30 mm	50 mm	75 mm	100	mm	125 mm	150	mm	250 mm
NUCLEODUR®	Phenyl-He	xyl, 1.8 µm pa	article size 1.8	µm · UHPLC						
Analytical EC colum	ns									
	2 mm	760561.20	760563.20	760565.2	20 760	566.20		7605	568.20	
	3 mm	760561.30	760563.30		760	566.30		***************************************		
	4 mm	760561.40	760563.40		760	566.40		•••••		
	4.6 mm	760561.46	760563.46		760	566.46		•		
EC guard columns*			4 x 2 m	m: 761985.20		4 x 3 mm:	761985.30			
NUCLEODUR®	Phenyl-He	xyl, 3 µm part	icle size 3 µm							
Analytical EC colum	ns									
	2 mm		760573.20		760	576.20	760577.20	7605	78.20	760579.20
	3 mm	•••••	760573.30	••••••	760	576.30	760577.30	7605	78.30	760579.30
	4 mm	•••••••	760573.40		760	576.40	760577.40	7605	78.40	760579.40
	4.6 mm	***************************************	760573.46	760575.4	46 760	576.46	760577.46	6 7605	78.46	760579.46
EC guard columns*	•	•	4 x 2 m	m: 761986.20		4 x 3 mm:	761986.30	•		
NUCLEODUR®	Phenyl-He	xyl, 5 µm part	icle size 5 µm							
Analytical EC colum	ns									
	2 mm		760583.20		760	586.20	760587.20	7605	588.20	760589.20
	3 mm		760583.30		760	586.30	760587.30	7605	588.30	760589.30
	4 mm	•	760583.40		760	586.40	760587.40	7605	588.40	760589.40
	4.6 mm	····	760583.46	760585.4	46 760	586.46	760587.46	7605	88.46	760589.46
EC guard columns*			4 x 2 m	m: 761987.20		4 x 3 mm:	761987.30			
Preparative VarioPre	p columns									
	10 mm		762210.100)			762211.10	00		762213.100
	21 mm		762210.210)			762211.21	10		762213.210
	32 mm									762213.320
	40 mm							7622	212.400	762213.400
VP guard columns *	*		10 x 8 m	m: 762234.80	1	0 x 16 mn	n: 762234.160	15	5 x 32 mm	: 762236.320
EC and VarioPrep co	olumns in pac	cks of 1, guard co	olumns see belo	DW.						
Guard column s	systems									
Guard columns for	EC columns	with ID	2 r	nm	3 mm	4 m	ım	4.6 mm	Gu	ard column holder
* Column Protection	System (pac	k of)	EC 4/2	2 (3)	4/3 (3)	4/3	(3)	4/3 (3)	718	8966
Guard columns for	VarioPrep co	lumns with ID		10 mm	16, 21 mm	32,	40 mm	≥ 50 mm		
** VP guard columns	s (pack of)		VP 10	/8 (2)	10/16 (2)	15/	32 (1)	15/50 (1)		

718251

718256

718253

718255

For details of our column systems see page 250.

VP guard column holder

$NUCLEODUR^{®} \pi^{2}$ hydrophobic biphenylpropyl phase · USP L11

Key feature

- Hydrophobic phase with alternative selectivity compared to classical C₁₈ modifications
- Separation principle based on 2 retention mechanisms (π - π interactions and hydrophobic interactions)
- Better retention of aromatic and unsaturated substances
- Excellent performance under highly aqueous conditions

Technical data

Phase with biphenylpropyl modification and multi-endcapping; pore size 110 Å; particle size 5 µm; carbon content 17 %; pH stability 1.5–10

Recommended application

 Overall sophisticated analytical separations, especially aromatic and unsaturated compounds, polar compounds like pharmaceuticals, antibiotics, steroids

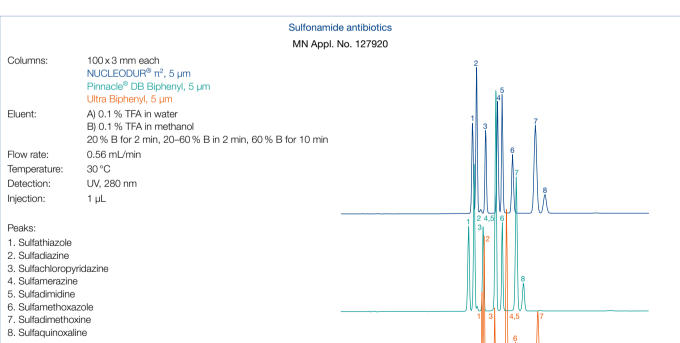
Stationary HPLC phases with biphenyl ligands like NUCLEODUR® π^2 provide an interesting alternative to classical alkyl modified C_{18} and C_{8} HPLC phases due to their remarkable orthogonal selectivity.

Furthermore the NUCLEODUR[®] π^2 provides an excellent separation performance for aromatic and unsaturated analytes by combination of hydrophobic and π - π interactions.

A unique feature is the predominant separation mechanism $(\pi$ - π or hydrophobic interactions) and thus the selectivity can be controlled by selection of the eluent. In acetonitrile/water

NUCLEODUR® π^2 shows similar retention strength then C_{18} modified phases and thereby displays a significantly stronger retention than phenyl phases. These interactions are even further enhanced in a methanol / water eluent.

NUCLEODUR® π^2 exceeds other aryl phases in terms of stability under strongly aqueous conditions. Therefore i.a. steroids, sulfonamides and acidic pharmaceuticals are separated in good resolution with NUCLEODUR® π^2 . NUCLEODUR® π^2 is the stationary phase with the highest aromatic analyte selectivity.



20

4.0

6.0

8 0



Columns: 125 x 4 mm each

 $NUCLEODUR^{\text{(8)}}\,\pi^{2},\,5\;\mu\text{m}$

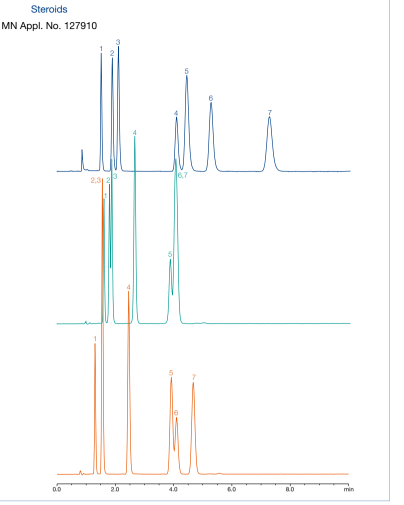
NUCLEODUR® Phenyl-Hexyl, 5 µm NUCLEODUR® C₁₈ Gravity, 5 µm

Eluent: acetonitrile - water (45:55, v/v)

Injection: 1 µL 1 mL/min Flow rate: Temperature: 25 °C Detection: UV, 230 nm

Peaks:

- 1. Estriol
- 2. Hydrocortisone
- 3. Prednisone
- 4. β-Estradiol
- 5. Corticosterone
- 6. Cortisonacetate
- 7. Testosterone



Ordering information Eluent in column acetonitrile - water ID Length → 50 mm 75 mm 100 mm 125 mm 150 mm 250 mm NUCLEODUR[®] π^2 , 5 μ m particle size 5 μ m Analytical EC columns 2 mm 760620.20 760621.20 760622.20 760623.20 760624.20 760625.20 760622.30 760620.30 760621.30 760623.30 760624.30 760625.30 3 mm 4 mm 760625.40 760620.40 760621.40 760622.40 760623.40 760624.40 4.6 mm 760620.46 760621.46 760622.46 760623.46 760624.46 760625.46 4 x 2 mm: 761810.20 4 x 3 mm: 761810.30

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

EC guard columns*

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

For details of our column systems see page 250.

NUCLEODUR® PFP hydrophobic pentafluorophenyl phase · USP L43

Key feature

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

Technical data

 Phase with pentafluorophenyl-propyl modification and multi-endcapping; pore size 110 Å; particle sizes 1.8 µm, 3 µm and 5 µm; carbon content 8 %; pH stability 1–9

Recommended application

 Aromatic and unsaturated compounds, phenols, halogen compounds, isomers, polar compounds like pharmaceuticals, antibiotics; strong retention of basic compounds

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 F_5). 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.

While a typical C_{18} phase just provides hydrophobic interactions between stationary phase and analyte NUCLEODUR® PFP offers four different retention mechanisms: polar interactions (H bonds), dipole-dipole, π - π , and hydrophobic interactions. Especially the pronounced ion exchange capacity and distinct steric selectivity are typical for fluorinated phases.

Due to low bleeding characteristics NUCLEODUR® PFP is also suitable for LC/MS. Based on a special surface modification procedure NUCLEODUR® PFP offers highest stability also at low pH values.

NUCLEODUR® PFP offers a completely different retention behavior compared to alkyl modified silica and is often used for separations which provide insufficient results on traditional C_{18} phases.

Applications in the areas of (bio-)pharma, natural compounds and environment show the broad applicability of this phase.

Separation of antihistamines MN Appl. No. 124861

Columns: 250 x 3 mm NUCLEODUR® PFP, 5 μm

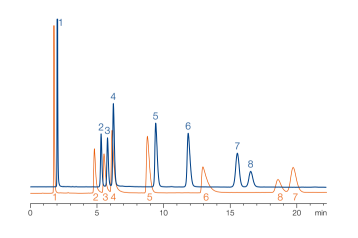
250 x 3 mm NUCLEODUR® C₁₈ Gravity, 5 μm

Eluent: acetonitrile – 20 mmol/L KH₂PO₄ (30:70, v/v)

Flow rate: 1.3 mL/min
Temperature: 30 °C
Detection: UV, 210 nm

Peaks:

- 1. Maleic acid
- 2. Chlorpheniramine
- 3. Brompheniramine
- 4. Triprolidine
- 5. Diphenhydramine
- 6. Promethazine
- 7. Cetirizine
- 8. Hydroxyzine





Separation of phenol isomers

125 x 4 mm NUCLEODUR® PFP, 5 µm

125 x 4 mm NUCLEODUR $^{\rm @}$ C₁₈ HTec, 5 μ m

Eluent: acetonitrile, 0.1 % formic acid – water, 0.1 %

formic acid (35:65, v/v)

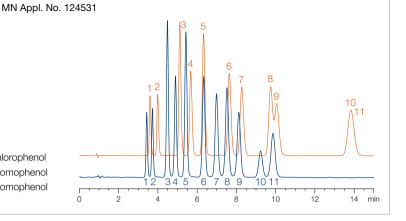
Flow rate: 1 mL/min
Temperature: 35 °C
Detection: UV, 280 nm

Peaks:

Column:

1. o-Kresol5. 2,5-Dimethylphenol9. 3,4-Dichlorophenol2. m-Kresol6. 2,6-Dichlorophenol10. 2,4-Dibromophenol3. 3,4-Dimethylphenol7. 2,3-Dichlorophenol11. 3,5-Dibromophenol

4. 3,5-Dimethylphenol 8. 2,4-Dichlorophenol



Ordering information Eluent in column acetonitrile - water Length → 100 mm 250 mm 30 mm 50 mm 75 mm 125 mm 150 mm NUCLEODUR® PFP, 1.8 μm particle size 1.8 μm · UHPLC Analytical EC columns 2 mm 760431.20 760433.20 760435.20 760436.20 760438.20 760431.30 760433.30 760436.30 3 mm 4 mm 760431.40 760433.40 760436.40 4.6 mm 760431.46 760433.46 760436.46 EC guard columns* 4 x 2 mm: 761975.20 4 x 3 mm: 761975.30 NUCLEODUR® PFP, 3 μm particle size 3 μm Analytical EC columns 760449.20 760443.20 760446.20 760447.20 760448.20 2 mm 760447.30 760448.30 760449.30 3 mm 760443.30 760446.30 4 mm 760443.40 760446.40 760447.40 760448.40 760449.40 4.6 mm 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 NUCLEODUR® PFP. 5 um particle size 5 um

VP guard columns **		10 x 8 mm:	762214.80	10 x 16 mm	n: 762214.160	15 x 32 mm	: 762216.320
	40 mm			.	-	762212.400	762213.400
——————————————————————————————————————	32 mm						762213.320
	21 mm	762210.210			762211.210		762213.210
	10 mm	762210.100		<u> </u>	762211.100		762213.100
Preparative VarioPrep	columns						
EC guard columns*			761977.20		761977.30		
	4.6 mm	760453.46	760455.46	760456.46	760457.46	760458.46	760459.46
	4 mm	760453.40		760456.40	760457.40	760458.40	760459.40
————	3 mm	760453.30		760456.30	760457.30	760458.30	760459.30
	2 mm	760453.20		760456.20	760457.20	760458.20	760459.20
Analytical EC columns	3						
	, .	•					

Guard	column	cyctome	

Gradier Cordinario, Cyclorico						
Guard columns for EC columns with ID		2 mm	3 mm	4 mm	4.6 mm	Guard column holder
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
** VP 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	

For details of our column systems see page 250.

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

NUCLEODUR® Sphinx RP bifunctional RP phase · USP L1 and L11

Kev feature

- · 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 data

· 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

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.

Stability of NUCLEODUR® Sphinx RP at pH 10 MN Appl. No. 120900 50 x 4.6 mm NUCLEODUR® Sphinx RP, 5 µm Column: Eluent: methanol - dil. NH₃, pH 10 (20:80, v/v) Flow rate: 1.0 mL/min, temperature 30 °C UV. 275 nm Detection: Injection: 3 μL Peaks: 1. Theophylline 2. Caffeine after 300 injections (with 5 L eluent) 1st injection

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.

Separation of flavonoids on three different NUCLEODUR® phases

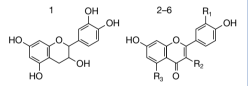
MN Appl. No. 119830

Columns: 150 x 4.6 mm

> NUCLEODUR® Sphinx RP, 5 µm NUCLEODUR® C₁₈ Gravity, 5 µm NUCLEODUR® C₈ Gravity, 5 µm

water - methanol (40:60, v/v) Fluent:

Flow rate: 1 mL/min 30 °C Temperature: Detection: UV, 270 nm Injection: 3 μL



Peaks:

1. Catechin

2. Rutin

 $R_1 = R_3 = OH$, $R_2 = O$ -Rutinose 3. Fisetin

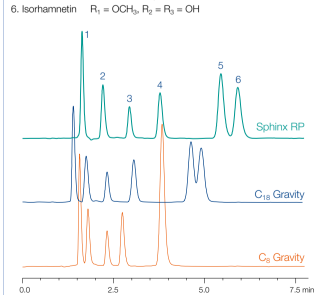
4. Quercetin

 $R_1 = R_2 = OH, R_3 = H$

 $R_1 = R_2 = R_3 = OH$

5. Kaempferol

 $R_1 = H, R_2 = R_3 = OH$





NUCLEODUR® columns



Ordering informa	ation									
Eluent in column ac	etonitrile – w	vater								
	ID	Length →								
		30 mm	50 mm	75 n	nm	100 mm	125 mm	150 m	ım	250 mm
NUCLEODUR® S	•	, 1.8 µm partio	cle size 1.8	µm · UHPLC						
Analytical EC column	าร									
	2 mm	760821.20	760822		825.20	760823.20	·····	76082	24.20	
	3 mm	760821.30	760822			760823.30	·····	·····		.
	4 mm	760821.40	760822	.40		760823.40	·····			.
	4.6 mm	760821.46	760822	.46		760823.46				
EC guard columns*			4 x	2 mm: 761920).20	4 x 3 mr	n: 761920.30			
NUCLEODUR® S	Sphinx RP,	, 3 µm particle	size 3 µm							
Analytical EC column	าร									
	2 mm		760806	.20		760812.20	760807.2	20 76080	5.20	760808.20
————	3 mm		760806	.30		760812.30	760807.3	30 76080	5.30	760808.30
	4 mm		760806	.40		760812.40	760807.4	10 76080	5.40	760808.40
	4.6 mm		760806	.46 7608	813.46	760812.46	760807.4	16 76080	5.46	760808.46
EC guard columns*	•	•	4 x	2 mm: 761921	.20	4 x 3 mr	m: 761921.30			
NUCLEODUR® S	Sphinx RP,	, 5 µm particle	size 5 µm							
Analytical EC column	าร									
	2 mm		760800	.20		760809.20	760801.2	20 76080	2.20	760803.20
	3 mm	•	760800	.30		760809.30	760801.3	30 76080	2.30	760803.30
	4 mm	····	760800	.40		760809.40	760801.4	10 76080	2.40	760803.40
	4.6 mm	••••	760800	.46 7608	815.46	760809.46	760801.4	16 76080	2.46	760803.46
EC guard columns*			4 x	2 mm: 761922	2.20	4 x 3 mr	m: 761922.30	•••••		
Preparative VarioPrep	o columns									
	10 mm		762372	.100			762375.1	100		762373.100
	21 mm		762372	.210			762375.2	210		762373.210
	32 mm									762373.320
	40 mm							76237	1.400	762373.400
VP guard columns **	f	***************************************	10 x	8 mm: 762390	0.80	10 x 16 m	nm: 762390.16	0 15:	x 32 mn	n: 762392.320
EC and VarioPrep co	olumns in pac	cks of 1, guard co	olumns see	below.						
Guard column s	ystems									
Guard columns for I	EC columns	with ID		2 mm	3 mm	4	mm	4.6 mm	Gı	uard column holder
* Column Protection	System (pac	k of)	EC	4/2 (3)	4/3 (3	3) 4/	(3 (3)	4/3 (3)	71	8966
Guard columns for	VarioPrep co	lumns with ID		8, 10 mm	16, 2	1 mm 32	2, 40 mm	≥ 50 mm		
** VP guard columns	(pack of)		VP	10/8 (2)	10/16	5 (2) 15	5/32 (1)	15/50 (1)		
VP guard column ho	lder			718251	7182	56 7 ⁻	18253	718255		

For details of our column systems see page 250.

NUCLEODUR® C₁₈ HTec base-deactivated preparative octadecyl phase · USP L1

Key feature

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

Technical data

• High density octadecyl modification (C₁₈); pore size 110 Å; particle sizes 1.8 μ m, 3 μ m, 5 μ m, 7 μ m and 10 μ m 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

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.

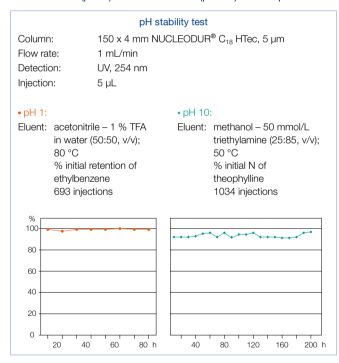
Selectivity and base deactivation

The innovative 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.

Engelhardt test MN Appl. No. 123580 250 x 4 mm NUCLEODUR® C₁₈ HTec, 5 µm Column: methanol - water (49:51, v/v) Eluent: 1 mL/min Flow rate: 40 °C Temperature: Detection: UV, 254 nm Injection: 5 µL Peaks: 5. N,N-Dimethylaniline 1. Uracil 2. Aniline 6. Toluene 3. Phenol 7. Ethylbenzene 4. p-Ethylaniline 20 10 30

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 results 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.



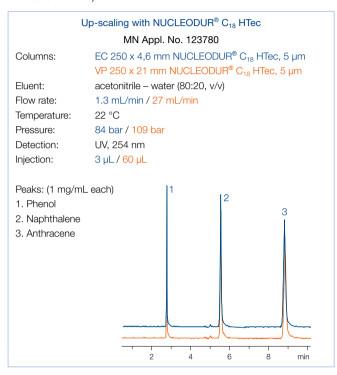
Due to innovative surface coating procedures NUCLEODUR $^{\$}$ C₁₈ HTec offers excellent analytical separation properties and is the first choice for up-scaling to preparative column dimensions.

NUCLEODUR® columns



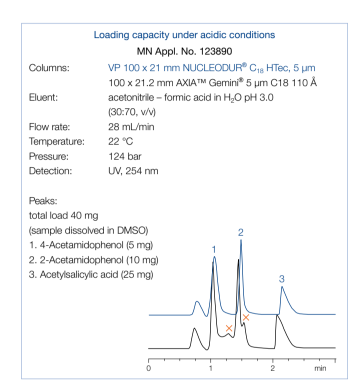
Up-scaling

Due to highest quality standards in silica production and phase chemistry combined with optimized packing technology, NUCLEODUR® C_{18} HTec allows exceptional transferability from analytical to preparative scale with respect to different particle sizes (e.g., 5, 7 or 10 μm) as well as column dimensions (e.g., ID 4.6 to 21 mm).



Capacity

A vital criterion for efficiency in preparative HPLC is the capacity of the separation medium. NUCLEODUR® C_{18} HTec is characterized by a notably high loading capacity under both basic and acidic conditions, while competitor columns show overload effects even at lower loads (x).



Ordering informa	ıtion							
Eluent in column ace	etonitrile – w	vater						
	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C	C ₁₈ HTec,	1.8 µm particle	size 1.8 µm · UF	IPLC				
Analytical EC column	S							
	2 mm	760301.20	760305.20	760304.20	760306.20		760308.20	
————	3 mm	760301.30	760305.30	•	760306.30			
	4 mm	760301.40	760305.40		760306.40			
	4.6 mm	760301.46	760305.46		760306.46			
EC guard columns*			4 x 2 mm:	761925.20	4 x 3 mm:	761925.30		
NUCLEODUR® C	C ₁₈ HTec,	3 µm particle s	ize 3 µm					
Analytical EC column	S							
	2 mm		760321.20		760323.20	760324.20	760325.20	760326.20
	3 mm	••••••	760321.30		760323.30	760324.30	760325.30	760326.30
	4 mm	•••••	760321.40		760323.40	760324.40	760325.40	760326.40
	4.6 mm	-	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		





	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® (C ₁ , HTec.					.20		200
nalytical EC column								
,	2 mm		760311.20		760313.20	760314.20	760315.20	760316.20
	3 mm	*	760311.30	·····	760313.30	760314.30	760315.30	760316.30
	4 mm	···•	760311.40		760313.40	760314.40	760315.40	760316.40
	4.6 mm	···•	760311.46	760312.46	760313.46	760314.46	760315.46	760316.46
C guard columns*			4 x 2 mm:	761927.20	4 x 3 mm:	761927.30		
reparative VarioPrep	columns							
	10 mm		762551.100			762554.100		762556.100
	21 mm		762551.210		762553.210	762554.210		762556.210
	32 mm				762553.320		762555.320	762556.320
	40 mm						762555.400	762556.400
	50 mm				762553.500		762555.500	762556.500
P guard columns **			10 x 8 mm:	762591.80	10 x 16 mm	: 762591.160		
		-	15 x 32 mm:	762592.320	15 x 50 mm	762592.500		
		7 μm particle s						
	columns 10 mm	7 µm particle s	762561.100		762563 210	762564.100 762564.210		762566.100 762566.210
	columns	7 μm particle s			762563.210 762563.320	762564.100 762564.210	762565.320	762566.100 762566.210 762566.320
	columns 10 mm 21 mm	7 μm particle s	762561.100		·····•	· · · · · · · · · · · · · · · · · · ·	762565.320 762565.400	762566.210
	10 mm 21 mm 32 mm	7 μm particle s	762561.100		·····•	· · · · · · · · · · · · · · · · · · ·	*	762566.210 762566.320
reparative VarioPrep	10 mm 21 mm 32 mm 40 mm	7 μm particle s	762561.100		762563.320 762563.500	· · · · · · · · · · · · · · · · · · ·	762565.400	762566.210 762566.320 762566.400
Preparative VarioPrep	10 mm 21 mm 32 mm 40 mm	7 μm particle s	762561.100 762561.210	762591.80	762563.320 762563.500 10 x 16 mm	762564.210	762565.400	762566.210 762566.320 762566.400
reparative VarioPrep	20 columns 10 mm 21 mm 32 mm 40 mm 50 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.320 762563.500 10 x 16 mm	762564.210 - 762591.160	762565.400	762566.210 762566.320 762566.400
P guard columns **	o columns 10 mm 21 mm 32 mm 40 mm 50 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.320 762563.500 10 x 16 mm	762564.210 - 762591.160	762565.400	762566.210 762566.320 762566.400
P guard columns **	o columns 10 mm 21 mm 32 mm 40 mm 50 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.320 762563.500 10 x 16 mm	762564.210 - 762591.160	762565.400	762566.210 762566.320 762566.400
reparative VarioPreparative VarioPrepara	o columns 10 mm 21 mm 32 mm 40 mm 50 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm	762591.80	762563.320 762563.500 10 x 16 mm	762564.210 : 762591.160 : 762592.500	762565.400	762566.210 762566.320 762566.400 762566.500
reparative VarioPreparative VarioPrepara	o columns 10 mm 21 mm 32 mm 40 mm 50 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm	762591.80	762563.320 762563.500 10 x 16 mm 15 x 50 mm	762564.210 : 762591.160 : 762592.500 762574.100	762565.400	762566.210 762566.320 762566.400 762566.500
P guard columns **	20 columns 10 mm 21 mm 32 mm 40 mm 50 mm Columns 10 mm 21 mm 21 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm	762591.80	762563.320 762563.500 10 x 16 mm 15 x 50 mm	762564.210 : 762591.160 : 762592.500 762574.100	762565.400 762565.500	762566.210 762566.320 762566.400 762566.500 762576.100 762576.210
P guard columns **	20 columns 10 mm 21 mm 32 mm 40 mm 50 mm Columns 10 mm 10 mm 21 mm 32 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm	762591.80	762563.320 762563.500 10 x 16 mm 15 x 50 mm	762564.210 : 762591.160 : 762592.500 762574.100	762565.400 762565.500 762575.320	762566.210 762566.320 762566.400 762566.500 762576.100 762576.210 762576.320
Preparative VarioPreparative VarioPrepar	20 columns 10 mm 21 mm 32 mm 40 mm 50 mm C ₁₈ HTec, 0 columns 10 mm 21 mm 32 mm 40 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm	762591.80 762592.320	762563.320 762563.500 10 x 16 mm 15 x 50 mm 762573.210 762573.320 762573.500	762564.210 : 762591.160 : 762592.500 762574.100	762565.400 762565.500 762575.320 762575.400	762566.210 762566.320 762566.400 762566.500 762576.100 762576.210 762576.320 762576.400
P guard columns ** IUCLEODUR® (reparative VarioPrep	20 columns 10 mm 21 mm 32 mm 40 mm 50 mm Columns 10 mm 21 mm 21 mm 21 mm 32 mm 40 mm		762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm 762571.100 762571.210	762591.80 762592.320 762591.80	762563.320 762563.500 10 x 16 mm 15 x 50 mm 762573.210 762573.320 762573.500 10 x 16 mm	762564.210 : 762591.160 : 762592.500 762574.100 762574.210	762565.400 762565.500 762575.320 762575.400	762566.210 762566.320 762566.400 762566.500 762576.100 762576.210 762576.320 762576.400
Preparative VarioPreparative VarioPrepar	20 columns 10 mm 21 mm 32 mm 40 mm 50 mm Columns 10 mm 21 mm 32 mm 40 mm 21 mm 32 mm 40 mm 50 mm	10 μm particle	762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm 762571.100 762571.210 10 x 8 mm: 15 x 32 mm:	762591.80 762592.320 762591.80	762563.320 762563.500 10 x 16 mm 15 x 50 mm 762573.210 762573.320 762573.500 10 x 16 mm	762564.210 : 762591.160 : 762592.500 762574.100 762574.210	762565.400 762565.500 762575.320 762575.400	762566.210 762566.320 762566.400 762566.500 762576.100 762576.210 762576.320 762576.400

For details of our column systems see page 250.

Guard columns for VarioPrep columns with ID

* Column Protection System (pack of)

** VP guard columns (pack of)

VP guard column holder

NUCLEODUR® C₁₈ HTec bulk material in 7 and 10 µm for self-packing of preparative columns see page 256.

4/2 (3)

8, 10 mm

10/8 (2)

718251

4/3 (3)

16, 21 mm

10/16 (2)

718256

4/3 (3)

32, 40 mm

15/32 (1)

718253

4/3 (3)

≥ 50 mm

15/50 (1)

718255

NUCLEODUR® columns



$NUCLEODUR^{\circledR} \ C_{18} \ ec \cdot C_{8} \ ec \cdot C_{4} \ ec \quad \text{nonpolar phases for routine analysis} \cdot \ \text{USP L1 } (C_{18}) \cdot \ \text{L26 } (C_{4}) \cdot \$

Key feature

- Ideal and reliable standard RP phase for daily routine analysis and up-scaling for preparative HPLC
- Medium density Octadecyl (C₁₈) and octyl (C₈) with pore size of 110 Å with exhaustive endcapping for a wide range of applications
- Octadecyl (C₁₈) and butyl (C₄) with pore size of 300 Å for the separation of biomolecules

Technical data

- \cdot 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 C_{18} , 10.5 % for C_{8} ; pH stability 1–9; high reproducibility from lot to lot
- Pore size 300 Å: technical data and applications in chapter "HPLC column for biochemical separations" (see page 241)

Recommended application

• 110 Å:

basic, neutral or acidic drugs; derivatized amino acids; pesticides; fat-soluble vitamins; aldehydes and ketones; phenolic compounds

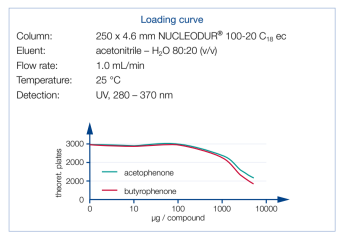
 300 Å: biomolecular macromolecules, like proteins and peptides

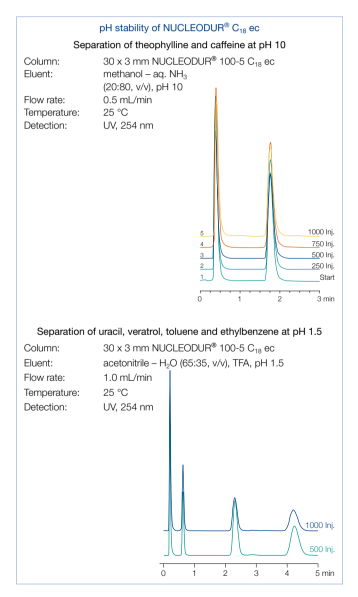
NUCLEODUR® C₁₈ ec for daily routine analysis

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® C_{18} ec is available in 9 different particle sizes (3, 5, 7, 10, 12, 16, 20, 30 and 50 μ m) which cover the whole range from high speed analytical HPLC up to medium and low pressure prep LC. NUCLEODUR® C_{18} ec is also an ideal tool for scale-up purposes.

Loading capacity

Loading capacity, 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® 100-20 C_{18} ec column describes the correlation between the increase of column loading and the decrease of separation efficiency.





Chemical stability

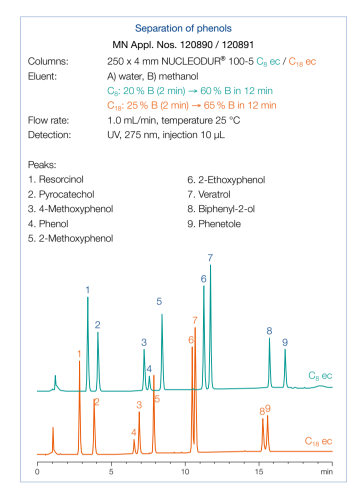
The utmost purity of the base silica and the exceptional silane bonding chemistry minimize the risk of dissolution, or hydrolysis at pH extremes.

The chromatograms show the retention behavior at pH values of 1.5 and 10.0 for NUCLEODUR® 100-5 C_{18} ec.

NUCLEODUR® octyl phases

In addition to NUCLEODUR® C_{18} phases MACHEREY-NAGEL offers octyl modified NUCLEODUR® C_8 Gravity and NUCLEODUR® C_8 ec columns to expand the RP tool box. Based on the same spherical high purity silica the C_8 phases exhibit the same chemical and mechanical stability as the C_{18} counterparts. Indeed NUCLEODUR® C_8 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 C_{18} phases). NUCLEODUR® C_8 ec and NUCLEODUR® C_8 Gravity are most suitable for the development of new methods but also for robust routine analyses.

There are no general guidelines which could make the choice between C_8 and C_{18} phases easier but it will always be beneficial to add both phases to the existing pool of RP columns in the laboratory. Comparative studies reveal some different selectivity patterns of NUCLEODUR® C_8 ec and C_{18} ec. The separation of phenols at right 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.



NUCLEODUR® phases for biochromatography

A description and applications for C_{18} and C_4 modified 300 Å NUCLEODUR[®] widepore materials for the separation of biopolymers, like peptids and proteins can be found in chapter "HPLC column for biochemical separations" (see page 241).

$C_{18} \mbox{ or } C_8 \cdot \mbox{the best of both worlds}$

- · Octyl phases (C₈) show superior polar selectivity.
- · Octadecyl phases (C₁₈) show superior hydrophobic selectivity.
- · Hydrophobic compounds show shorter retention times on C₈ phases.

Ordering informa	tion						
Eluent in column ace	tonitrile – wa	ter					
	ID	Length → 50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® 1	00-3 C ₁₈ ed	octadecyl phase,	particle size 3 µm,	17.5 % C			
Analytical EC columns	S						
	2 mm	760050.20		760054.20	760051.20	760053.20	760052.20
————	3 mm	760050.30	•	760054.30	760051.30	760053.30	760052.30
	4 mm	760050.40		760054.40	760051.40	760053.40	760052.40
	4.6 mm	760050.46	760046.46	760054.46	760051.46	760053.46	760052.46
EC guard columns*			4 x 2 mm: 7	761931.20	4 x 3 mm: 7	761931.30	



NUCLEODUR® columns



Eluent in column ac	cetonitrile – wat	ter					
	ID	Length → 50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR®	100-5 C ₁₈ ec	c octadecyl phase,	particle size 5 µm,	17.5 % C			
Analytical EC colum			•				
,	2 mm	760004.20		760013.20	760001.20	760008.20	760002.20
	3 mm	760004.30		760013.30	760001.30	760008.30	760002.30
	4 mm	760004.40	·····	760013.40	760001.40	760008.40	760002.40
	4.6 mm	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: 7		
Preparative VarioPre	ep columns						
•	10 mm	762003.100			762029.100		762022.100
	21 mm	762003.210	•••••		762029.210		762022.210
	32 mm		······				762022.320
	40 mm					762027.400	762022.400
VP guard columns *	*		10 x 8 mm:	762090.80	10 x 16 mm:	· · · · · · · · · · · · · · · · · · ·	
J	•		15 x 32 mm:		15 x 50 mm:	· · · · · · · · · · · · · · · · · · ·	
NLICLEODUR®	100-10 C	C octadecyl phase					
Preparative VarioPre		octadecyi priase	, particle Size 10 μ	111, 17.5 /0 0			
reparative varioPre		760011 100			760000 100		760010 100
	10 mm	762011.100	·····		762302.100		762010.100
	21 mm	762011.210			762302.210	<u>.</u>	762010.210
	32 mm						762010.320
	40 mm				······	762303.400	762010.400
	50 mm	······					762010.500
VP guard columns *	•	······	10 x 8 mm:		10 x 16 mm:	· · · · · · · · · · · · · · · · · · ·	
			15 x 32 mm:	102311.320	15 x 50 mm:	702311.500	
Oud avia a infava	_*:		15 x 32 mm:	702311.320	15 X 50 mm:	762311.500	
			15 x 32 mm:	702311.320	15 X 50 mm:	762311.500	
	cetonitrile – wat		15 x 32 mm:	702311.320	15 X 50 mm:	762311.500	
		Length →					250 mm
Eluent in column ad	cetonitrile – wat ID	Length → 50 mm	75 mm	100 mm	15 X 50 mm:	150 mm	250 mm
Eluent in column ac	ID 100-3 C ₈ ec	Length →	75 mm	100 mm			250 mm
Eluent in column ac	ID 100-3 C ₈ ec	Length → 50 mm	75 mm	100 mm			250 mm
Eluent in column ac	ID 100-3 C ₈ ec	Length → 50 mm	75 mm	100 mm			250 mm 760062.20
Eluent in column ac	D 100-3 C ₈ ec	Length → 50 mm octyl phase, particl	75 mm	100 mm % C	125 mm		
Eluent in column ac	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm	Length → 50 mm octyl phase, particl 760063.20	75 mm	100 mm % C 760059.20	125 mm 760060.20		760062.20
Eluent in column ac	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30	75 mm	100 mm % C 760059.20 760059.30	125 mm 760060.20 760060.30		760062.20 760062.30
NUCLEODUR® Analytical EC colum	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40	75 mm e size 3 μm, 10.5 9	100 mm % C 760059.20 760059.30 760059.40 760059.46	760060.20 760060.30 760060.40	150 mm 760061.46	760062.20 760062.30 760062.40
NUCLEODUR® Analytical EC colum	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm:	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20	760060.20 760060.30 760060.40 760060.46	150 mm 760061.46	760062.20 760062.30 760062.40
NUCLEODUR® Analytical EC colum EC guard columns*	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm:	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20	760060.20 760060.30 760060.40 760060.46	150 mm 760061.46	760062.20 760062.30 760062.40
NUCLEODUR® Analytical EC colum EC guard columns*	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm:	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7	150 mm 760061.46	760062.20 760062.30 760062.40 760062.46
NUCLEODUR® Analytical EC colum EC guard columns*	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm:	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20 % C 760704.20	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7	150 mm 760061.46	760062.20 760062.30 760062.40 760062.46
NUCLEODUR® Analytical EC colum EC guard columns*	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm:	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7	150 mm 760061.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30
NUCLEODUR® Analytical EC colum EC guard columns*	2 mm 3 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40
NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20 % C 760704.20 760704.30 760704.40 760704.46	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.46	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30
NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum	2 mm 3 mm 4.6 mm 2 mm 3 mm 4.6 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40
NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum	cetonitrile – wat ID 100-3 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40 760700.46	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20 % C 760704.20 760704.30 760704.40 760704.46	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.46 4 x 3 mm: 7	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46
NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum	2 mm 3 mm 4 mm 4.6 mm 2 mm 3 mm 4 mm 4.6 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40 760700.46	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20 % C 760704.20 760704.30 760704.40 760704.46	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.46 4 x 3 mm: 7	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46
Analytical EC colum	2 mm 3 mm 4 mm 4.6 mm 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm 4	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40 760700.46	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20 % C 760704.20 760704.30 760704.40 760704.46	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.46 4 x 3 mm: 7	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46
NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum	2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec ns 2 mm 3 mm 4 mm 2 mm 4 mm 4 mm 4 mm 4 mm 4 mm 4	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40 760700.46	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9	100 mm % C 760059.20 760059.30 760059.40 760059.46 761936.20 % C 760704.20 760704.30 760704.40 760704.46	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.46 4 x 3 mm: 7	760061.46 61936.30 760702.46 61937.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.46 762062.100 762062.210 762062.320
NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum EC guard columns*	2 mm 3 mm 4 mm 4.6 mm 100-5 C ₈ ec 100-5 C ₈ ec	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.30 760700.40 760700.46	75 mm e size 3 μm, 10.5 9 760064.46 4 x 2 mm: e size 5 μm, 10.5 9 760706.46 4 x 2 mm:	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46 761937.20	760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.46 4 x 3 mm: 7	760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.46 762062.100 762062.210 762062.320 762062.400

Guard column systems see previous NUCLEODUR® phases. For details of our column systems see page 250.

NUCLEODUR® C₁₈ ec bulk material with 10–50 μm for self-packing of preparative columns see page 256.

The ordering information for C_{18} and C_4 modified 300 \mathring{A} NUCLEODUR® widepore materials for the separation of biopolymers can be found in the chapter "HPLC" column for biochemical separations" (see page 241).

^{*} and ** for corresponding guard column systems see page 180.

NUCLEODUR® HILIC zwitterionic phase

Key feature

- Ideal for reproducible and stable chromatography of highly polar analytes
- Suitable for analytical and preparative applications
- · Very short column conditioning period

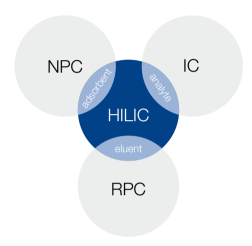
Technical data

 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

Hydrophilic interaction chromatography



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.

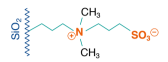
The expression HILIC (Hydrophilic Interaction Chromatography) was firstly published by Andrew Alpert in 1990 – since then it took quite some efforts to develop robust and reproducible hydrophilic HPLC phases for HILIC chromatography [7].

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, NH_2 , 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.

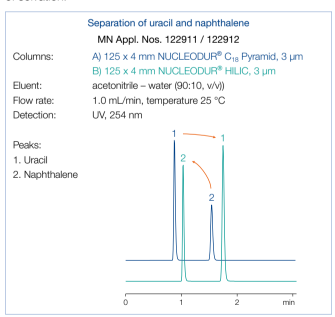
Summarized: "HILIC is NP chromatography of polar and ionic compounds under RP conditions."

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 mobile and stationary phases. 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 compounds exhibit faster elution profiles due to minor hydrophobic interactions. In the separation of uracil and naphthalene the elution order is quite often inverse on HILIC columns compared to RP columns.

NUCLEODUR® columns

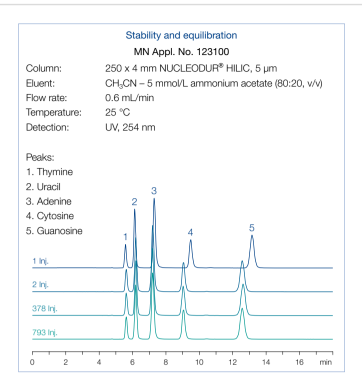


Stability features

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. Due to its high loading capacity 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.



Eluent in column ace	etonitrile – w	vater (80:20, v/v)						
	ID	Length →						
		30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® H	HILIC, 1.8	µm particle size	e 1.8 μm · UHPLC	;				
Analytical EC column	IS							
	2 mm	760521.20	760523.20	760525.20	760526.20		760528.20	
	3 mm	760521.30	760523.30		760526.30			
	4 mm	760521.40	760523.40		760526.40			
	4.6 mm	760521.46	760523.46		760526.46			
EC guard columns*			4 x 2 mm:	761960.20	4 x 3 mm:	761960.30		
NUCLEODUR® H	HILIC, 3 µr	n particle size 3	μm					
Analytical EC column	IS							
	2 mm	····	760532.20		760534.20	760531.20	760533.20	760530.20
	3 mm		760532.30		760534.30	760531.30	760533.30	760530.30
	4 mm	····	760532.40		760534.40	760531.40	760533.40	760530.40
	4.6 mm	····	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		
NUCLEODUR® H	HLIC, 5 µr	n particle size 5	μm					
Analytical EC column	IS							
	2 mm		760552.20		760554.20	760551.20	760553.20	760550.20
	3 mm		760552.30		760554.30	760551.30	760553.30	760550.30
	4 mm		760552.40		760554.40	760551.40	760553.40	760550.40
	4.6 mm		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		

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

For details of our column systems see page 250.

NUCLEODUR® CN/CN-RP cyano-modified high purity silica phase · USP L10

Kev feature

- · 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 data

- · 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

Alternative bonded-phase functionality

In reversed phase HPLC it is fairly common to start with C_{18} 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 NUCLEODUR® 100-5 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 below).

Separation of cold medicine ingredients on two different NUCLEODUR® phases

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

250 x 4 mm NUCLEODUR® 100-5 CN-RP

MN Appl. No. 119340

Eluent: acetonitrile - 100 mmol/L sodium citrate pH 2.5

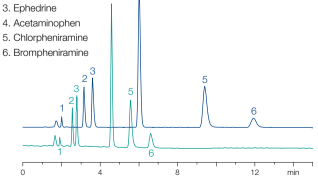
(15:85, v/v)

Flow rate: 1.0 mL/min, temperature 25 °C Detection: UV, 254 nm, injection 10 µL

Peaks:

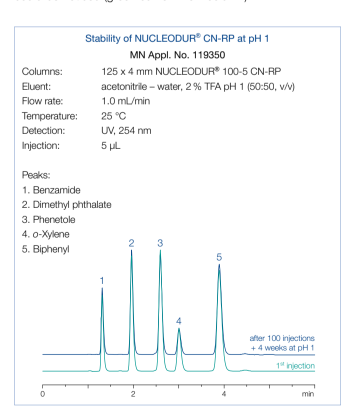
1. Maleic acid

- 2. Norephedrine
- 4. Acetaminophen
- 6. Brompheniramine



The polarity of NUCLEODUR® 100-5 CN-RP can be classified as intermediate based on multiple retention mechanisms such as dipole-dipole, π - π , and also hydrophobic interactions [8]. 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) [9].

Short-chain bonded phases are sometimes suspected of revealing shortcomings in stability towards hydrolysis at low pH [10]. Application 119350 shows 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)

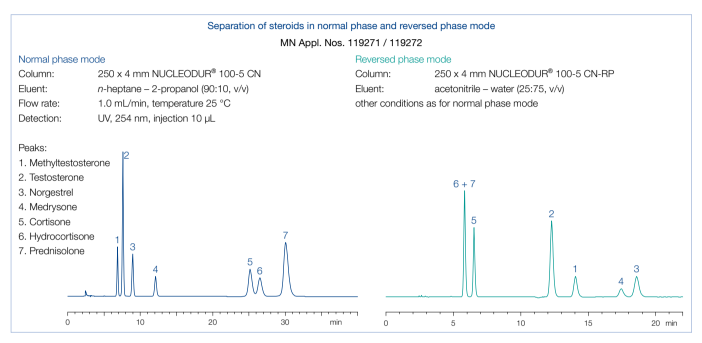




Multi-mode columns

Due to its polarity the cyano phase can also be run in normal phase mode. NUCLEODUR® CN columns for NP applications are shipped in *n*-heptane. The change in selectivity and order of elution for a mixture of various steroids in NP and RP mode is

displayed below. The high coverage combined with a thorough endcapping makes NUCLEODUR® 100-5 CN-RP suitable for separation of ionizable compounds such as basic drugs.



	ation				
	ID	Length → 50 mm	125 mm	150 mm	250 mm
					230 111111
NUCLEODUR® 1	00-3 CN-RP	particle size 3 µm; eluent ir	n column acetonitrile - water	r	
Analytical EC column	is .				
	2 mm	760159.20	760157.20		
	3 mm		760157.30		
	4 mm			760156.40	
	4.6 mm			760156.46	
EC guard columns*		4 x 2 mm: 7619	41.20	4 x 3 mm: 7619	941.30
NUCLEODUR® 1	00-5 CN-RP	particle size 5 µm; eluent ir	n column acetonitrile – water	r	
Analytical EC column					
ni iaiyildal EO GOlultili.					
- Trialytical EC Column	4 mm		760153.40		760152.40
A laytical EC Column			760153.40 760153.46	760154.46	760152.40 760152.46
	4 mm			760154.46 4 x 3 mm: 7619	760152.46
EC guard columns*	4 mm 4.6 mm	cle size 5 µm; eluent in colu	760153.46		760152.46
EC guard columns*	4 mm 4.6 mm	cle size 5 μm; eluent in colι	760153.46		760152.46
EC guard columns*	4 mm 4.6 mm	cle size 5 μm; eluent in colι	760153.46		760152.46
EC guard columns*	4 mm 4.6 mm	cle size 5 μm; eluent in colι	760153.46 umn <i>n-</i> heptane	4 x 3 mm: 7619	760152.46 944.30

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

For details of our column systems see page 250.

Guard column system

NUCLEODUR® NH₂ / NH₂-RP amino-modified high purity silica · USP L8

Key feature

- 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 data

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

✓ Recommended application

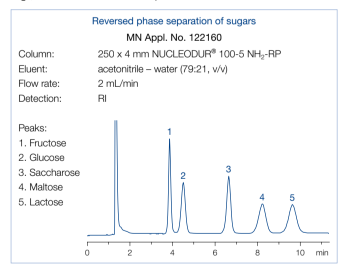
Polar compounds under RP conditions (sugars, DNA bases), hydrocarbons under NP conditions

- 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 C_{18} phases. Polar-modified silica phases offer alternative selectivities thus expanding the spectrum of analytical HPLC into the polar range.

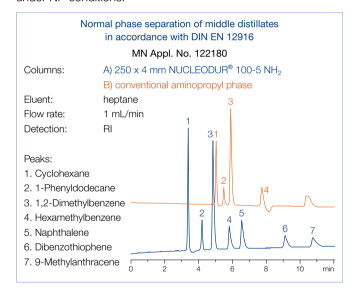
Multi-mode columns

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® NH_2 , too, belongs to the so-called multimode columns. It can be used for RP chromatography of polar compounds such as sugars in aqueous-organic eluent systems, for NP chromatography 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® NH_2 is the separation of simple and complex sugars, sugar alcohols and other hydroxy compounds under RP conditions as well as hydrocarbons under NP conditions.

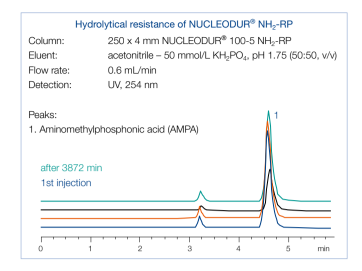


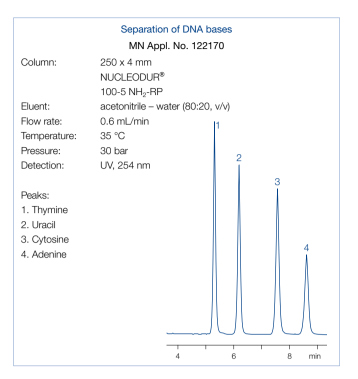
Due to the special method of surface modification NUCLEODUR® NH_2 features a pronounced stability at higher as well as at lower pH values. The following figure shows, that even after several days of exposure of the column material at pH 1.75 good separation efficiency and peak symmetry are maintained. The resulting high column life allows cost reduction due to lower column consumption.

This example shows the enhanced pH stability of NUCLEODUR® NH₂ and the outstanding suitability for the separation of total herbicides (AMPA, glyphosate, glufonisate, ...) - see application 122190 in our online data base at www.mn-net.com/apps.

NUCLEODUR® columns







Based on superspherical NUCLEODUR® this phase features a high pressure stability, which makes it the perfect choice for preparative separations as well as for LC/MS. Additionally, the high batch-to-batch reproducibility of NUCLEODUR® NH₂ enables reliable analyses especially for routine work.

	ID	Length →			
		100 mm	125 mm	150 mm	250 mm
NUCLEODUR® -	100-3 NH ₂ -RP	particle size 3 µm; eluent	in column acetonitrile – wate	er	
Analytical EC columr	าร				
	2 mm	760740.20	760741.20		
	4.6 mm			760742.46	760739.46
EC guard columns*		4 x 2 mm: 76	1951.20	4 x 3 mm: 76	1951.30
NUCLEODUR® -	100-5 NH ₂ -RP	particle size 5 µm; eluent	in column acetonitrile – wate	er	
Analytical EC columr					
	2 mm		760730.20		760732.20
	3 mm	•••••••••••••••••••••••••••••••••••••••	760730.30		760732.30
	4 mm		760730.40		760732.40
	4.6 mm		760730.46	760731.46	760732.46
EC guard columns*		4 x 2 mm: 76	1953.20	4 x 3 mm: 76	1953.30
NUCLEODUR® -	100-5 NH ₂ par	ticle size 5 µm; eluent in co	lumn <i>n</i> -heptane		
Analytical EC columr					
	4 mm		760720.40		760722.40
	4.6 mm	•	760720.46	760721.46	760722.46
EC guard columns*		•		4 x 3 mm: 76	1952.30
EC columns in packs	s of 1, quard colu	mns in packs of 3.			

3 mm

4/3 (3)

4 mm

4/3 (3)

2 mm

4/2 (3)

EC

For details of our column systems see page 250.

Guard columns for EC columns with ID

* Column Protection System (pack of)

Guard column holder

718966

4.6 mm

4/3 (3)



NUCLEODUR® SIOH unmodified silica for normal phase · USP L3

Key feature

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

Technical data

 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 page 150)

Recommended application

 Polar and midpolar compounds under normal phase conditions

Ordering information

Eluent in column *n*-heptane

ID Length →

50 mm 125 mm 150 mm 250 mm

NUCLEODUR® 100-3 particle size 3 µm

Analytical EC columns

4.6 mm 760170.46 760172.46 760173.46

EC guard columns* 4 x 3 mm: 761966.30

NUCLEODUR® 100-5 particle size 5 µm

Analytical EC columns

, and factor and						
————	4 mm				760007.40	
	4.6 mm	760023.46		760012.46	760007.46	
EC guard columns*				4 x 3 mm: 76		
Preparative VarioPre	ep columns					
	10 mm	762077.100	762078.100		762007.100	

21 mm

VP guard columns

columns					
10 mm	762077.100	762078.100		762007.100	
21 mm	762077.210	762078.210		762007.210	
40 mm			762075.400	762007.400	
	10 x 8 mm: 70		10 x 16 mm: 7	762094.160	
	15 x 32 mm:				

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	Guard column holder
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966
Guard columns for VarioPrep columns with ID		8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	
** VP 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	

For details of our column systems see page 250.

Unmodified NUCLEODUR® bulk material in 10-50 µm for self-packing of preparative columns see page 256.





MACHEREY-NAGEL your partner in HPLC · also online

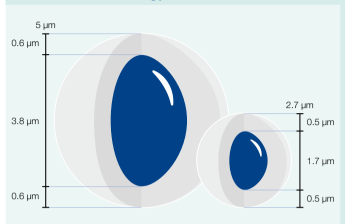
Besides to this catalog our website provides useful information

- Applications Database without registration, with more than 3000 free chromatography applications for your separation task.
- · Instruction manuals General advises for column care and individual column cleaning are available in the attached instruction manual or online.
- · HPLC troubleshooting Sometimes during chromatographic separation unexpected effects occur. We give advise of possible reasons and how to avoid or remedy these.
- · Flyers, brochures, catalogs Our product information is available online as PDF file at any time.



NUCLEOSHELL® core-shell silica for HPLC

Core-shell technology

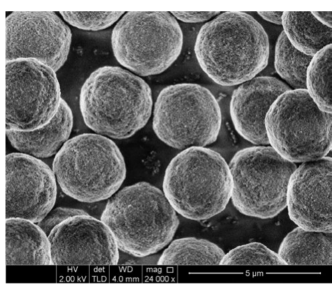


Key feature

- · Solid core of silicon dioxide, homogeneous shell of porous
- · Highest efficiency compared to traditional totally porous materials
- · Pore size 90 Å; particle size 2.7 μm (core 1.7 μm) and 5 µm (core 3.8 µm); specific surface 130 (2.7 µm) and 90 (5 µm) m²/g lower back pressure enables use on conventional LC systems
- · Pressure stability 600 bar

Demands on HPLC separations are constantly increasing with respect to separation efficiency, detection limits, and the time requirements for each analysis.

Several approaches have been made to achieve fast separations without losing chromatographic performance. HPLC columns packed with particles < 2 µm show very high efficiencies (plates/meter) and allow the use of smaller column sizes with the positive side effect of significant solvent saving. However they generate a high back pressure of the mobile phase during column runs which requires specifically designed equipment.



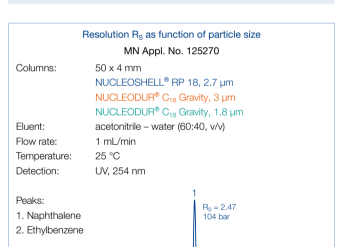
Electron microscopic image of NUCLEOSHELL®

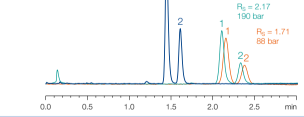
NUCLEOSHELL® silica particles consist of a non-porous solid core of 1.7 µm diameter and a porous outer shell of 0.5 µm thickness. Accordingly the total diameter of the particle is 2.7 µm.

Utilizing a proprietary process of synthesis, NUCLEOSHELL® particles exhibit a distinct narrow particle size distribution (d90/ d10 ~ 1.1). Columns packed with NUCLEOSHELL core shell particles feature exceptional separation efficiencies with theoretical plate numbers easily comparable to totally porous sub 2 micron particles.

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

 R_s = resolution, α = selectivity (separation factor), k_i = retention N = plate number with N \propto 1/d_P, d_P = particle diameter







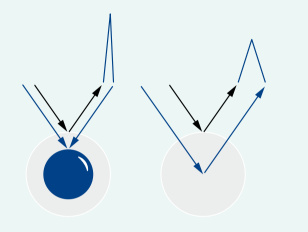
NUCLEOSHELL® core-shell silica for HPLC



Theoretical colu	mn efficienc	y (optimal cor	nditions)					
Silica	d _p [µm]	L [m]	HETP [µm]	Efficiency [plates/m]	L [mm]	N	R_s	Analysis time
NULCU ECOLUEUL®	2.7	1	4	250 000	100	25 000	112 %	40 %
NUCLEUSHELL ³	5	1	6.5	154 000	150	23 000	115%	60 %
	1.8	1	4.5	222 222	100	22 000	105 %	40 %
NUCLEODUR®	3	1	7.5	133 333	150	20 000	100 %	60 %
	5	1	12.5	80 000	250	20 000	100 %	100 %

Benefits of core-shell technology

Core-shell particles vs. totally porous silica



Short diffusion paths

- · Fast mass transfer (term C of Van Deemter equation)
- · High flow velocity without peak broadening for fast LC

Narrow particle size distribution $(d_{90}/d_{10} \sim 1.1)$

· Stable packing

High heat transfer

- · Minimized influence of frictional heat
- Efficiency of NUCLEOSHELL® ~ 250 000 m⁻¹ (HETP $\sim 4 \mu m$)

With conventional fully porous particles the mass transfer between stationary and mobile phase usually results in peak broadening at higher flow rates (C-term in van Deemter equation). The short diffusion paths in the core-shell particles reduce the

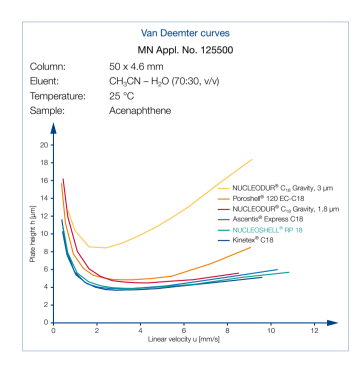
dwell time of the analyte molecules in the stationary phase, so that even at high flow velocities of the mobile phase, optimal separation results can be obtained.

The van Deemter plots demonstrate how efficiency is affected by flow rate.

In comparison with fully porous silicas, core-shell particles from various manufacturers maintain the efficiency optimum (max. plates/m) over a long range of increasing linear mobile phase velocity.

$$H = A + \frac{B}{II} + C \cdot u$$

A term = eddy-diffusion, B term = longitudinal diffusion coefficient, C term = mass transfer coefficient

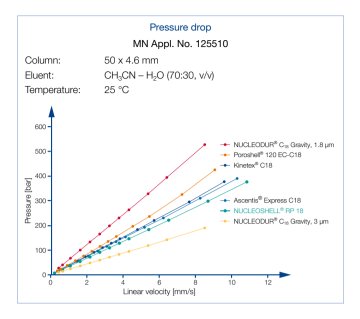


NUCLEOSHELL® core-shell silica for HPLC

In direct comparison with conventional sub 2 micron phases, NUCLEOSHELL® columns only generate about 60% of the back pressure and can be operated with the majority of conventional HPLC systems. In order to develop the maximum performance of NUCLEOSHELL® columns, we recommend reducing extra column voids by using suitable capillaries (< 0.15 mm inner diameter) and specially adapted detector cells. Moreover detector settings should be optimized by increasing the measuring rate or by decrease of the time constant.

$$\Delta_{p} = \frac{\Phi \cdot L_{C} \cdot \eta \cdot \iota}{d_{p}^{2}}$$

 $\Delta_P = pressure \; drop, \; \Phi = flow \; resistance \; (nondimensional), \; LC = column \\ length, \; \eta = viscosity, \; u = linear \; velocity, \; d_P = particle \; diameter$

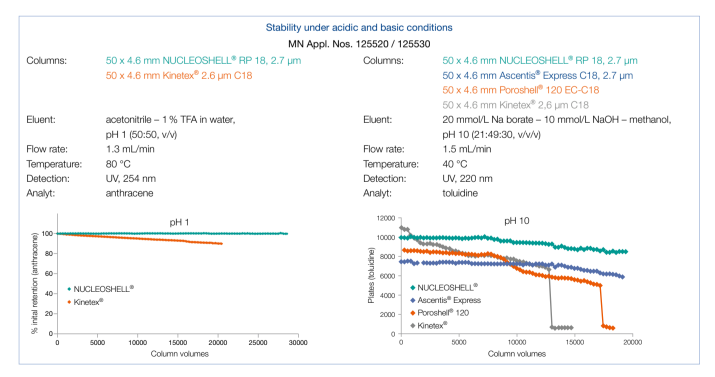


Core-shell particle technology from MACHEREY-NAGEL is an alternate route to gain highest column efficiency and resolution in HPLC at short run time, but with moderate back pressure.

Features of NUCLEOSHELL® particles

A criterion for the long-term stability of the column at pH extremes is the percentage decrease of initial retention and initial plates, respectively.

The following figure shows a column stability test of NUCLEOSHELL® RP 18 at mobile phase levels pH 1 and pH 10 compared with three competing phases.



End (t = 40 h)

NUCLEOSHELL® core-shell silica for HPLC



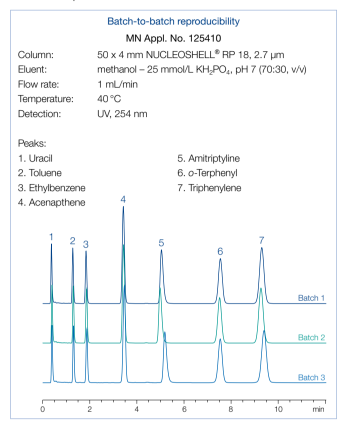
Columns can be operated at elevated temperatures without loss in retention, efficiency or peak symmetry.

Temperature stability MN Appl. No. 125400 Stability test: Column: 50 x 2 mm NUCLEOSHELL® RP 18, 2.7 µm Eluent: A) 10 mmol/L ammonium formate - methanol $(9:1, v/v) + 120 \mu L$ formic acid, ~ pH 4 B) 10 mmol/L ammonium formate - methanol $(1:9, v/v) + 120 \mu L$ formic acid, ~ pH 4 0-100 % B in 7 min Flow rate: 0.5 mL/min, Temperature: 100°C Detection: UV. 220 nm Peaks: 1. Phenol 2. Naphthalene 38 h 30 h 26 h 22 h min Efficiency test: Eluent: Acetonitrile - water (60:40, v/v) 0.33 mL/min; Flow rate: 25°C Temperature: UV, 254 nm Detection: Analyte: Anthracene HETP [µm] Asymmetry Start (t = 0)0.98 5.2

5.2

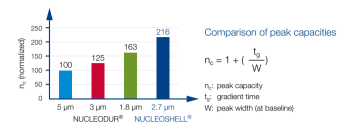
1.01

Uniformly shaped NUCLEOSHELL® particles combined with optimized bonding technology safeguard tightly packed columns for 100 % reproducible results.



Peak capacity

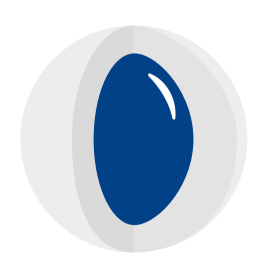
The peak capacity is a measure for the number of sample analytes that can be separated on HPLC columns per time unit. Narrow peaks increase the peak capacity and thus the efficiency of the analytical column.



The example shows, that in comparison with totally porous NUCLEODUR® silica (1.8 µm) NUCLEOSHELL® provides 33 % higher peak capacity.

Peak capacity MN Appl. No. 125540 100 x 4.6 mm each Columns: NUCLEOSHELL® RP 18, 2.7 µm NUCLEODUR® C₁₈ Gravity, 1.8 µm NUCLEODUR® C₁₈ Gravity, 3 µm A) acetonitrile, B) water, 40-100 % A in 4 min Eluent: Flow rate: 1.5 mL/min Temperature: 25°C Detection: UV. 230 nm Peaks: 1. Acetophenone 2. Benzoin 3. Propiophenone 4. Butyrophenone 5. Benzophenone 6. Valerophenone Max. pressure [bar] Resolution (4.5) NUCLEOSHELL®, 2.7 µm 255 5.45 NUCLEODUR®, 1.8 µm 450 4.14 NUCLEODUR®, 3 µm 214 2.97 NUCLEODUR®, 5 µm 142 2.30





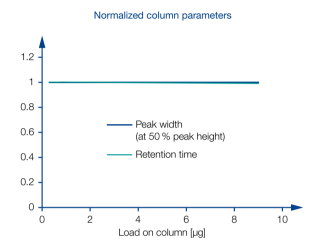
1,1

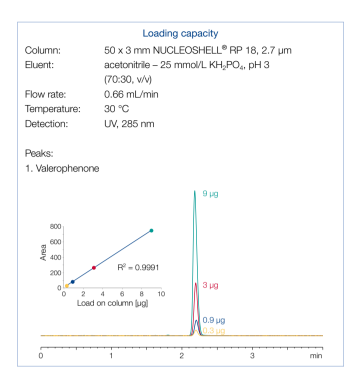
NUCLEOSHELL® core-shell silica for HPLC



Loading capacity

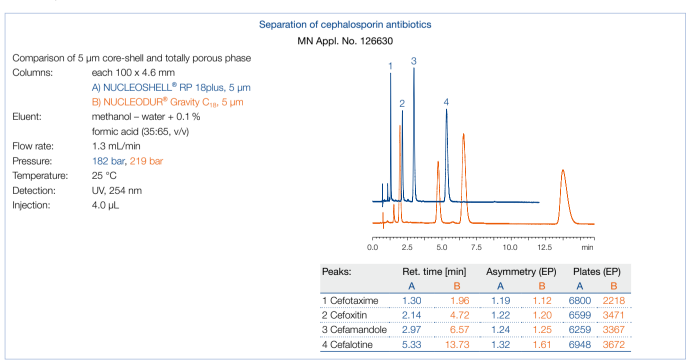
NUCLEOSHELL® columns allow reliable quantification in a wide analytical detection range. Retention time and peak width at 50% height remain constant with increasing columns load although core-shell particles are suspected of showing a slightly lower loading capacity compared to fully porous silica materials.





Method transfer of 5 µm particle columns

NUCLEOSHELL $^{\$}$ is also available in 5 μm particle size to offer all benefits of core-shell technology to all applications which are bound to particle size.





NUCLEOSHELL® phase overview



ase	Specification	Page	Ch	naracteristic*	Stability	Structure		
	octadecyl, multi-endcapping		Α	••••		©		
	7.8 % C (2.7 μm particles) 6.1 % C (5 μm particles)	200	В	•	pH 1-11, suitable for LC/MS	NUCLEOSHELL® (SI-O ₂),		
RP 18	USP L1		С	•••		NO.		
	octadecyl (monomeric), multi-endcapping 5.7 % C (2.7 µm particles) 4.4 % C (5 µm particles)		Α	••••		ELL®		
		202	В	••(pH 2–9, suitable for LC/MS	NUCLEOSHELL® (Si-O ₂) Si(CH ₃		
RP 18plus	USP L1		С	-				
	phenylhexyl, multi-endcapping 4.5 % C (2.7 µm particles)	phenylhexyl,		A	••		©	
		204	В	•••	pH 1–10, suitable for LC/MS	NUCLEOSHELL®		
Phenyl-Hexyl	USP L11		С	•		N Section 1		
	pentafluorophenyl,		Α	••		⊜ □ □ _ ⊁si- <mark>OH</mark>		
	multi-endcapping ~ 3 % C (2.7 μm particles)	206	В	••••	pH 1–9, suitable for LC/MS	NUCLEOSHELL®		
PFP	USP L43		С	••••				
			A •			© CH ₅		
	zwitterionic ammonium – sulfonic acid 1.3 % C (2.7 µm particles)	208	В	••••	pH 2-8.5, suitable for LC/MS	NUCLEOSHELL® (Si-OH) (CH ² SO ² (CH ²		
HILIC	, , , ,		С	-		Z SI-OH3		



NUCLEOSHELL® phase overview



Application	Similar phases**	Interactions · retention mecl	hanism
overall sophisticated analytical separations, e.g., analgesics, anti-inflammatory drugs, antidepressants; herbicides; phytopharmaceuticals; immunosuppressants	Kinetex® C18; Cortecs® C18; Raptor® C18; Accucore® C18; Ascentis® Express C18	hydrophobic (van der Waals interactions)	Si(CH ₃) ₃
overall sophisticated analytical separations, especially for polar compounds, e.g., pharmaceuti- cals like antibiotics, water-solub- le vitamins, organic acids	Kinetex® XB-C18; Bonshell® ASB-C18; Raptor® ARC-C18;	hydrophobic (van der Waals interactions)	Si Si(CH ₃) ₃ H ₄ C
aromatic and unsaturated com- pounds, polar compounds like pharmaceuticals, antibiotics	Ascentis® Express Phenyl-Hexyl; Kinetex® Phenyl-Hexyl; Accucore® Phenyl-Hexyl; Ultracore® Phenyl-Hexyl; Poroshell® Phenyl-Hexyl	π-π and hydrophobic	O ₂ N
aromatic and unsaturated com- pounds, phenols, halogenated hydrocarbons, isomers, polar compounds like pharmaceuti- cals, antibiotics	Kinetex® PFP; Ascentis® Express F5; Accucore® PFP	polar (H bond), dipole-dipole, π-π and hydrophobic	F F H
hydrophilic compounds such as organic polar acids and bases, polar natural compounds	-	ionic/ hydrophilic and electro- static	H ₃ C O CH ₃ O CH ₃ O CH ₃ NH NH NH ₂ NCH ₃ SO ₃ O NH ₂
** phases which provide a similar	selectivity based on chemical and physical propertie	es	

NUCLEOSHELL® RP 18 nonpolar high density phase · USP L1

Key feature

- · Core-shell technology for fast and efficient HPLC
- · Suitable for LC/MS and HPLC at pH extremes (pH 1-11)
- · Superior base deactivation, ideal for method development

Technical data

· Octadecyl modification, multi-endcapped; pore size 90 Å, particle size 2.7 and 5 µm, carbon content 7.8 % for 2.7 µm, 6.1 % for 5 µm; pH stability 1-11; suitable for LC/MS

Recommended application

· Overall sophisticated analytical separations, e.g., analgesics, anti-inflammatory drugs, antidepressants; herbicides; phytopharmaceuticals; immunosuppressants

NUCLEOSHELL® RP 18 is based on core-shell silica. A unique derivatization process generates a homogeneous surface with a high density of bonded silanes. The following thorough endcapping suppresses any unwanted polar interactions between the silica surface and the sample, which makes NUCLEOSHELL® RP 18 particularly suitable for the separation of basic and other ionizable analytes. The extremely reduced silanol activity of the phase can be demonstrated by applying basic analytes, such as tricyclic antidepressants. The chromatogram below shows a sharp elution profile (superior resolution!) of these highly polar compounds with an excellent asymmetry value for amitriptyline of 1.12.

Tricyclic antidepressants · comparison of selectivity and resolution MN Appl. No. 124960 Columns: 50 x 4.6 mm each Peaks: NUCLEOSHELL® RP 18, 2.7 µm 1. Protriptyline Ascentis® Express C18 2. Desipramine Kinetex® 2.6 µm C18 3. Maprotiline Poroshell® 120 EC-C18 4. Nortriptyline Eluent: methanol – acetonitrile – 25 mmol/L KH₂PO₄, pH 7 5. Doxepin (22.5:22.5:55, v/v/v) 6. Imipramine Flow rate: 2 mL/min 7. Amitriptyline Pressure: 224 bar, 239 bar, 248 bar, 212 bar 8. Clomipramine Temperature: 40 °C 9. Trimipramine Detection: UV, 220 nm Asymmetry Resolution (amitriptyline) (8, 9)**NUCLEOSHELL®** 1.12 3.35 Ascentis® Express 2.07 1.91 Kinetex® 1.33 n.a. 1.05 1.95 Poroshell® **NUCLEOSHELL®** Ascentis® Kinetex® Poroshell® 0 18

NUCLEOSHELL® columns



NUCLEOSHELL® RP 18 combines innovative silica technology and excellent surface deactivation, that outperforms conventional C₁₈ silicas in terms of efficiency, resolution and speed.

Due to the applied core-shell particle design the back pressure at elevated flow rates remains at a moderate level and in many cases permits the use of existing HPLC equipment. NUCLEOSHELL® RP 18 with extended pH stability, low bleed characteristics in LC/MS applications, and overall robustness is an ideal tool for method development and routine analyses in modern HPLC.

The separation of 13 β-lactam antibiotics illustrates how time of analysis can be shortened to a fractional part by using core-shell particles without loss of resolution at moderate back pressure.

13 β-lactam antibiotics in less than 3 min

MN Appl. No. 124940

50 x 4 mm NUCLEOSHELL® RP 18, 2.7 µm Columns:

150 x 4 mm NUCLEODUR® C₁₈ Gravity, 5 µm

A) acetonitrile B) 20 mmol/L KH₂PO₄, pH 3.5 Eluent:

 $10 \% A (0.5 min) \rightarrow 50 \% A in 1.5 min (0.5 min 50 % A)$

 $10\% A (3 min) \rightarrow 50\% A in 9 min (3 min 50\% A)$

Flow rate: 2 mL/min, 1 mL/min Pressure:

270 bar, 110 bar Temperature: 25 °C

Detection: UV. 220 nm

Peaks:

1. Amoxicillin 9. Penicillin V 2. Ampicillin 10. Oxacillin 3. Cephalexin 11. Cloxacillin 4. Cefotaxime 12. Nafcillin 5. Cefoxitin 13. Dicloxacillin

6. Cefamandole 7. Cephalothin 8. Piperacillin

10 12 10 11 13 4 6 7 8 9 5 5 5 5 6 7 8 9
0 2 4 6 8 10 12 min
2.5 min 270 bar 6 7 8 9 12 23 4 5
0.0 0.4 0.8 1.2 1.6 2.0 min

Ordering information

Eluent in column acetonitrile - water

	ID	Length →								
		50 mm	100 mm	150 mm	250 mm	EC guard columns*				
NUCLEOSHELL [®] RP 18, 2.7 μm particle size 2.7 μm										
Analytical EC column	ns									
	2 mm	763132.20	763134.20	763136.20		763138.20				
	3 mm	763132.30	763134.30	763136.30		763138.30				
	4 mm	763132.40	763134.40	763136.40		763138.30				
	4.6 mm	763132.46	763134.46	763136.46		763138.30				
NUCLEOSHELL	.® RP 18, 5 μ	m particle size 5 μm								
Analytical EC column	ns									
	2 mm	763152.20	763154.20	763156.20	763157.20	763158.20				
	3 mm	763152.30	763154.30	763156.30	763157.30	763158.30				
	4 mm	763152.40	763154.40	763156.40	763157.40	763158.30				

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

763152.46

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

763156.46

763157.46

763154.46

For details of the EC column system please see page 250.

763158.30

NUCLEOSHELL® RP 18plus C₁₈ phase with polar selectivity · USP L1

Key feature

- Based on core-shell particle technology for fast and efficient HPLC
- Hydrophobic C₁₈ phase with distinct polar selectivity, ideal for method development
- Excellent performance under highly aqueous conditions

Technical data

 Monomeric octadecyl modification, multi-endcapped; pore size 90 Å, available particle sizes 2.7 µm and 5 µm, carbon content 5.7 % for 2.7 µm, 4.4 % for 5 µm; pH stability 2–9; suitable for LC/MS

Recommended application

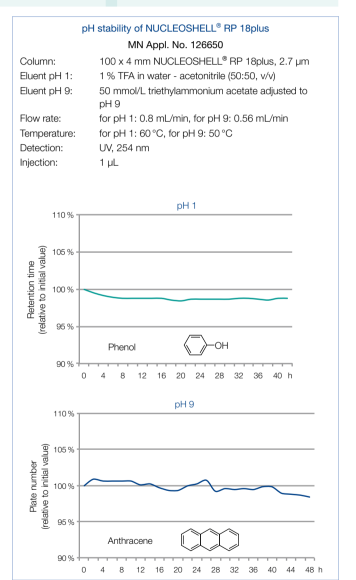
 Overall sophisticated analytical separations, especially for polar compounds, e.g., pharmaceuticals like antibiotics, water-soluble vitamins, organic acids

NUCLEOSHELL® RP 18 plus is a C_{18} modified core-shell silica. Due to a monomeric bonding chemistry this HPLC phase offers hydrophobic characteristics with distinct polar selectivity. A special derivatization process generates a medium density of bonded silanes with reduced steric selectivity compared to NUCLEOSHELL® RP 18.

Bleeding characterisitics MN Appl. No. 126640 50 x 2 mm NUCLEOSHELL® RP 18plus, 2.7 µm Column: Eluent: A) 0.1 % formic acid in water B) 0.1 % formic acid in acetonitrile $95 \% A \rightarrow 5 \% A \text{ in } 4.5 \text{ min } (0.5 \text{ min}) \rightarrow 95 \% A \text{ in }$ 0.5 min (4.5 min) Flow rate: 0.5 mL/min 25 °C Temperature: Detection: MS NUCLEOSHELL® RP 18 plus NUCLEOSHELL® RP 18 Poroshell® C18 m/z 50-1000 — Kinetex® XB-C18 ion chromatogram (TIC), Total 6

NUCLEOSHELL® RP 18plus combines superbly hydrophobic and polar selectivity – so it is a useful tool for method development in RP chromatography. Good pH stability and low bleeding characteristics make it ideal especially for LC/MS applications.

Retention time [min]



Also a comparison of retention of the glycopeptide antibiotic vancomycin on several octadecyl modified core-shell phases underlines the polar selectivity of NUCLEOSHELL® RP 18plus.

NUCLEOSHELL® columns



Polar selectivity shown for vancomycin

MN Appl. No. 126660

Columns: 50 x 3 mm each

NUCLEOSHELL® RP 18 plus, 2.7 μ m NUCLEOSHELL® RP 18, 2.7 μ m

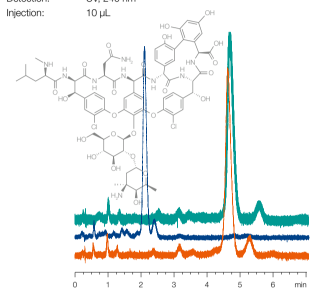
Kinetex® 2.6 µm C18

Eluent: water - methanol - acetonitrile - glacial acetic acid

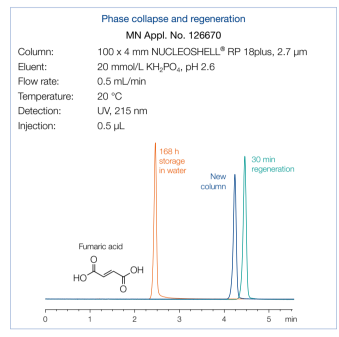
(100:8:2:0.3, v/v/v/v) adjusted to pH 3.2 with sodium

hydroxide solution

Flow rate: 0.9 mL/min
Temperature: 35 °C
Detection: UV, 240 nm
Injection: 10 µL



In addition NUCEOSHELL® RP 18plus provides a good stability under highly aqueous conditions. Even by long term usage or storage of the phase phase collapse and loss of retention are hardly observed. The original performance can be regained after a short regeneration procedure.



Eluent in column ad	cetonitrile – wat	er				
	ID	Length →				
		50 mm	100 mm	150 mm	250 mm	EC guard columns*
NUCLEOSHELL	® RP 18plus	, 2.7 µm particle siz	ze 2.7 µm			
Analytical EC colum	ns					
	2 mm	763232.20	763234.20	763236.20		763238.20
	3 mm	763232.30	763234.30	763236.30		763238.30
	4 mm	763232.40	763234.40	763236.40		763238.30
	4.6 mm	763232.46	763234.46	763236.46		763238.30
NUCLEOSHELL	® RP 18plus	, 5 µm particle size	5 μm			
Analytical EC colum	ns					
	2 mm	763252.20	763254.20	763256.20	763257.20	763258.20
	3 mm	763252.30	763254.30	763256.30	763257.30	763258.30
	4 mm	763252.40	763254.40	763256.40	763257.40	763258.30
	4.6 mm	763252.46	763254.46	763256.46	763257.46	763258.30

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

For details of the EC column system please see page 250.

NUCLEOSHELL® Phenyl-Hexyl nonpolar high density phase · USP L11

Kev feature

- Based on core-shell particle technology for fast and efficient HPLC
- Hydrophobic phase with alternative selectivity compared to classical C₁₈ modifications
- Separation principle based on 2 retention mechanisms: π - π interactions and hydrophobic interactions

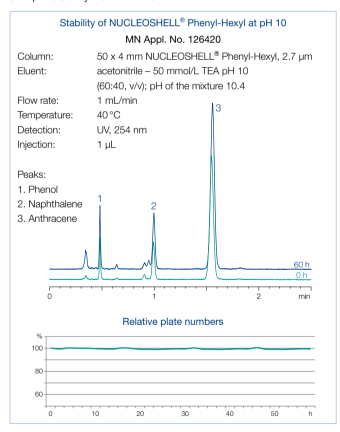
Technical data

 Phenyl-Hexyl modification, multi-endcapped; pore size 90 Å, particle size 2.7 µm; carbon content 4.5 %; pH stability 1–10; suitable for LC/MS

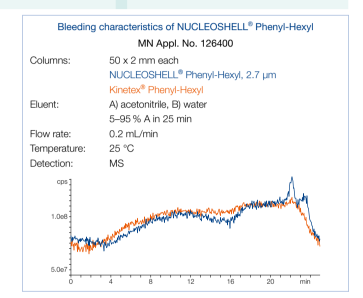
Recommended application

 Aromatic and unsaturated compounds, polar compounds like pharmaceuticals, antibiotics

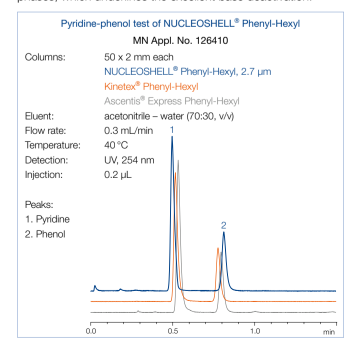
Phenyl-Hexyl modified phases offer an excellent separation efficiency especially for aromatic and unsaturated compounds with electron-withdrawing groups. The combination of hydrophobic and $\pi\text{-}\pi$ interactions results in an alternative and interesting selectivity profile compared to C_{18} or C_{8} modifications. NUCLEOSHELL® Phenyl-Hexyl is based on a unique surface bonding chemistry - therefore it is suitable for LC/MS due to low bleeding characteristics and offers high temperature stability and pH stability from 1 to 10.



NUCLEOSHELL® Phenyl-Hexyl is a robust phase with an alternative RP selectivity for aromatic and unsaturated analytes compared to classical $\rm C_{18}$ / $\rm C_{8}$ phases – it is an additional and useful tool for all chromatography users.



The pyridine-phenol test shows that NUCLEOSHELL® Phenyl-Hexyl provides a symmetrical peak for pyridine and higher resolution in comparison to other core-shell based Phenyl-Hexyl phases, which underlines the excellent base deactivation.







MN Appl. No. 125860

Columns: 150 x 3 mm each

> NUCLEOSHELL® Phenyl-Hexyl, 2.7 µm NUCLEODUR® Phenyl-Hexyl, 1.8 µm NUCLEODUR® Phenyl-Hexyl, 3 µm NUCLEODUR® Phenyl-Hexyl, 5 μm

Eluent: A) methanol

B) 0.1 % formic acid in water

20-80 % A in 10 min

Flow rate: 0.56 mL/min Temperature: 40°C Detection: UV, 254 nm Injection: 0.5 µL

Peaks:

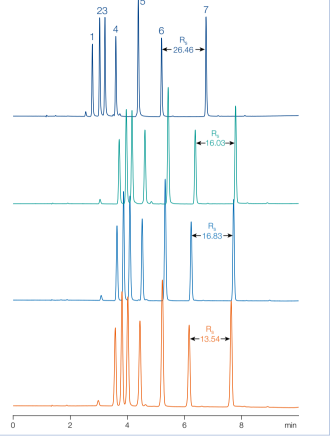
On NUCLEOSHELL® Phenyl-Hexyl 1. Sulfadiazine the resolution of the last two peaks is 2. Sulfachlorpyridazine higher than on the fully porous 1.8 µm NUCLEODUR® Phenyl-Hexyl.

3. Sulfapyridine 4. Sulfamerazine

5. Sulfadimidine

6. Sulfathiazole

7. Sulfadimethoxine



The separation of sulfonamides proves the scalability from ful-Iv porous NUCLEODUR® to NUCLEOSHELL® Phenyl-Hexyl.

Thus, method transferability between NUCLEODUR® and NUCLEOSHELL® is guaranteed, either for speeding up your methods or scaling up for preparative requirements.

ly porodo reocleobort to reocleobries i mony moxy.
Hereby the core-shell silica exhibits identical selectivity, narrower
peaks and slightly shorter retention under the same conditions.
Ordering information

Ordering information	ation										
Eluent in column acetonitrile – water											
	ID	Length → 50 mm	100 mm	150 mm	EC guard columns*						
NUCLEOSHELL® Phenyl-Hexyl, 2.7 μm particle size 2.7 μm											
Analytical EC colum	ns										
	2 mm	763732.20	763734.20	763736.20	763738.20						
	3 mm	763732.30	763734.30	763736.30	763738.30						
	4 mm	763732.40	763734.40	763736.40	763738.30						
	4.6 mm	763732.46	763734.46	763736.46	763738.30						

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

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

For details of the EC column system please see page 250.

NUCLEOSHELL® PFP hydrophobic pentafluorophenyl phase · USP L43

Kev feature

- Core-shell technology for fast and efficient HPLC
- Hydrophobic phase with alternative selectivity in comparison to classical C₁₈ modifications
- Separation principle based on 4 retention mechanisms (polar interactions (H bonds), dipole-dipole, π - π , hydrophobic interactions)

Technical data

 Phase with pentafluorophenylpropyl modification, multi-endcapping; pore size 90 Å, particle size 2.7 µm; carbon content ~ 3 %; pH stability 1–9; suitable for LC/MS

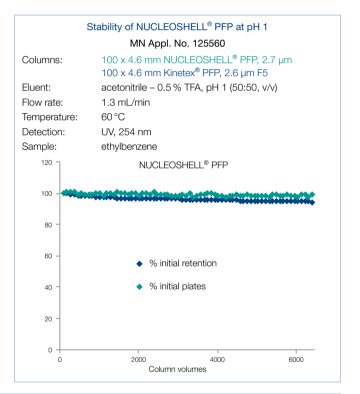
Recommended application

 Aromatic and unsaturated compounds, phenols, halogen compounds, isomers, polar compounds like pharmaceuticals, antibiotics; strong retention of basic compounds

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 F_5). Especially the orthogonal selectivity compared to traditional alkyl phases widens the scope in analytical HPLC. Thus NUCLEOSHELL® PFP offers an excellent selectivity especially for highly polar analytes, aromatic and unsaturated compounds, phenols or halogenated hydrocarbons.

While a typical C_{18} phase just provides hydrophobic interactions between stationary phase and analyte NUCLEOSHELL® 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.





Columns: 100 x 4.6 mm

NUCLEOSHELL® RP 18, 2.7 µm NUCLEOSHELL® PFP, 2.7 µm

Eluent: A) acetonitrile + 0.1 % formic acid

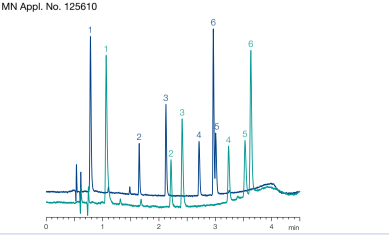
B) 0.1 % formic acid

10-35 % A in 2.5 min, 35-50 % A in 2 min

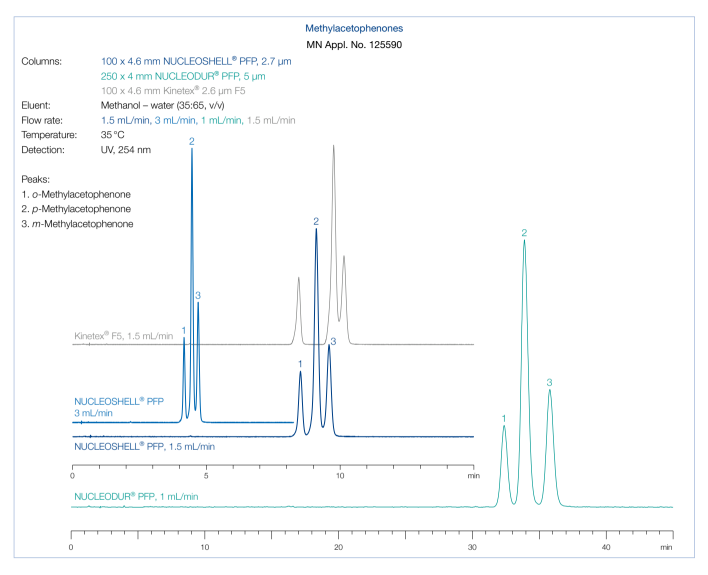
Flow rate: 1.7 mL/min
Temperature: 25 °C
Detection: UV, 280 nm

Peaks:

Atenolol
 Pindolol
 Metroprolol
 Alprenolol
 Propranolol







NUCLEOSHELL® PFP combines the benefits of core-shell technology, high stability, and orthogonal selectivity. Thus it is a useful complementary tool for highly efficient separations especially of isomers, halogenated, aromatic and / or polar compounds.

luent in column a	acetonitrile – water				
	ID	Length → 50 mm	100 mm	150 mm	EC guard columns*
NUCLEOSHEL	L [®] PFP, 2.7 µm	particle size 2.7 µm			
nalytical EC colur	mns				
	2 mm	763532.20	763534.20	763536.20	763538.20
———	3 mm	763532.30	763534.30	763536.30	763538.30
	4 mm	763532.40	763534.40	763536.40	763538.30
	4.6 mm	763532.46	763534.46	763536.46	763538.30

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

For details of the EC column system please see page 250.

NUCLEOSHELL® HILIC zwitterionic phase

Key feature

- Core-shell technology for fast and efficient HPLC
- Ideal for reproducible and stable chromatography of highly polar analytes
- · Very short column equilibration times

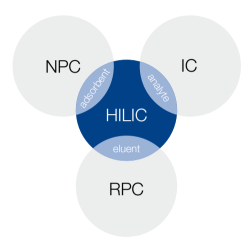
Technical data

 Ammonium - sulfonic acid modified silica; pore size 90 Å, particle size 2.7 µm; carbon content 1.3 %; pH stability 2–8.5; suitable for LC/MS

Recommended application

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

Hydrophilic interaction chromatography

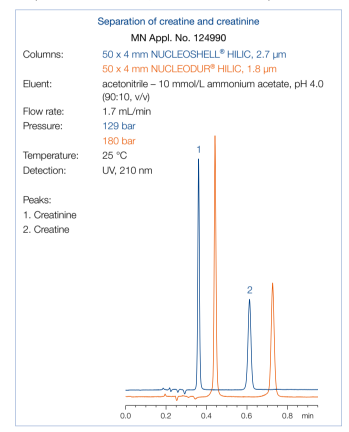


Hydrophilic interaction chromatography (HILIC) is a separation technique using polar stationary phases and organic-aqueous mobile phases. A minimum water content of at least $2\,\%$ is indispensable to provide a permanent water layer between the adsorbent surface and the organic fraction of the mobile phase. The sample molecules become separated in a partition chromatography, in which polar analytes are more strongly retained than neutral, less hydrophilic compounds. Consequently, increasing the aqueous part in the mobile phase will diminish retention of the polar sample constituents. In this way HILIC behaves inverse to classical RP chromatography. The particular retention profile of HILIC enables the chromatography of very polar and often small molecules, which won't show any retention on C_8 or C_{18} reversed phases.

Ultra-fast separations at moderate back pressure

NUCLEOSHELL® HILIC is a core-shell technology based stationary phase with a covalently bonded 3-N,N-dimethylamino-propane sulfonic acid ligand (pat. p nd.). The betaine character of the strong ion-exchanger results in full charge balancing and facilitates fast equilibration times.

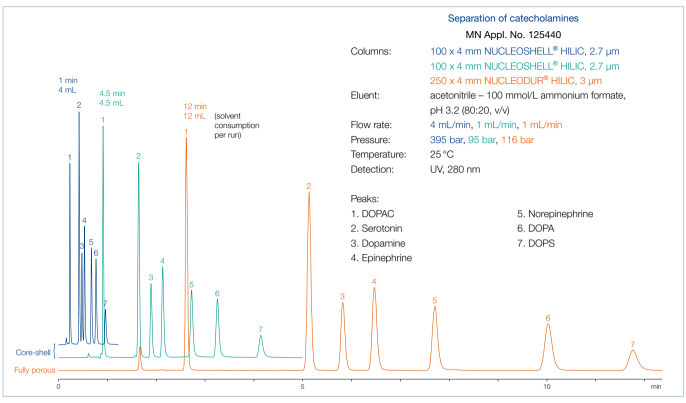
Good separation of polar compounds like the physiologically important substances creatine and creatinine can be achieved on NUCLEOSHELL® HILIC as well as on NUCLEODUR® HILIC, 1.8 µm at similar retention, but much lower back pressure.



The following chromatograms show the method transfer from a fully porous 3 μ m HILIC phase to 2.7 μ m core-shell silica with equal selectivity features.

Run time has been cut down to 1 min. Column back pressure remains modest < 400 bar, while solvent demand is reduced to less than 35 %.





Core-shell silica: separation in 1 min pressure < 400 bar

NUCLEOSHELL® HILIC provides stable and reproducible chromatography, comprising all the benefits of a state-of-the-art core-shell silica.

uent in column	acetonitrile – water				
	ID	Length → 50 mm	100 mm	150 mm	EC guard columns*
IUCLEOSHEL	_L [®] HILIC, 2.7 μr	n particle size 2.7 µm			
nalytical EC colu	mns				
	2 mm	763332.20	763334.20	763336.20	763338.20
	3 mm	763332.30	763334.30	763336.30	763338.30
	4 mm	763332.40	763334.40	763336.40	763338.30
	4.6 mm	763332.46	763334.46	763336.46	763338.30

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

For details of the EC column system please see page 250.





The guard column system for HPLC / UHPLC from MN

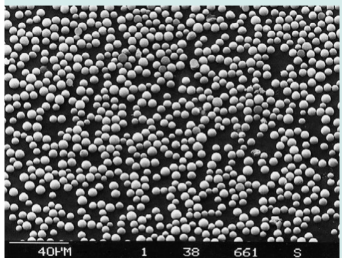
- · Ideal protection for your analytical main column: significant increase in column lifetime
- · Minimized void volume: suitable also for ultra fast HPLC (UHPLC)
- · Special ferrules: pressure stability up to 1300 bar (18850 psi)
- · Cartridges filled with NUCLEODUR®, NUCLEOSIL® and NUCLEOSHELL® HPLC adsorbents.
- · Universal screw-on guard column holder system
- · Suitable for all analytical HPLC columns with 1/16" fittings Further information on page 251.



NUCLEOSIL® standard silica for HPLC







Kev feature

- NUCLEOSIL® is a family of totally porous spherical silicas.
 They feature a very pure and uniform SiO₂ structure and have gained wide acceptance as routine chromatographic packings for very different fields of modern chromatography.
- · One of the first spherical silicas used in HPLC
- Developed in the early seventies, it became a worldrenowned HPLC packing
- · Absolutely reliable choice for routine analyses
- · Largest variety of modified HPLC silicas available
- pH stability 2-8 (for NUCLEOSIL® 100-5 C₁₈ AB 1-9)
- Due to its particle sizes NUCLEOSIL[®] finds application in analytical as well as in preparative columns.

Benefits of NUCLEOSIL® silica

- · High efficiency due to narrow particle size distribution
- High separation performance due to optimized binding techniques
- · High chemical and mechanical stability
- · High load capacity and recovery rates
- · High reproducibility from lot to lot

Physical properties

NUCLEOSIL® is manufactured with different pore diameters (50, 100, 120, 300, 500, 1000 and 4000 Å) and particle sizes from 3 μ m (only NUCLEOSIL® 50, 100 and 120) to 10 μ m with very narrow fractionation. All narrow-pore NUCLEOSIL® packings are stable up to 500 bar (7 250 psi), the wide-pore NUCLEOSIL® silicas are stable up to 300 or 400 bar (4 200 or 5 600 psi).

Phase	Pore size	Pore volume	Surface (BET)	Density	Pressure stability*
	016 3126		. ,	_ = =,	1 1633ule Stability
NUCLEOSIL® 50	50 A	0.8 mL/g	420 m²/g	0.45 g/mL	500 bar
NUCLEOSIL® 100	100 Å	1 mL/g	350 m²/g	0.36 g/mL	500 bar
NUCLEOSIL® 120	120 Å	0.65 mL/g	200 m²/g	0.55 g/mL	500 bar
NUCLEOSIL® 300	300 Å	0.8 mL/g	100 m²/g	0.45 g/mL	400 bar
NUCLEOSIL® 500	500 Å	0.8 mL/g	35 m²/g	0.45 g/mL	400 bar
NUCLEOSIL® 1000	1000 Å	0.8 mL/g	25 m²/g	0.45 g/mL	300 bar
NUCLEOSIL® 4000	4000 Å	0.7 mL/g	10 m ² /g	0.48 g/mL	300 bar

NUCLEOSIL® modifications

- NUCLEOSIL® packings are available as unmodified silica or with numerous chemically bonded phases: RP phases like C₁₈ AB, C₁₈ HD, C₁₈ Nautilus, C₁₈, C₁₈ ec, Protect I, C₈ HD, C₈ ec, C₈, C₄, C₂ and C₆H₅ separate mainly by hydrophobic interactions (van der Waals forces). The less polar the sample molecules, the more they are retained the more polar the sample, the weaker are the hydrophobic interactions and consequently the retention times are shorter.
- Phases with chemically bonded polar groups such as CN, NH₂, N(CH₃)₂, OH show selective separation properties.
 Due to the availability of different functional groups it is pos-
- sible to vary the chemical characteristics of the surface and consequently the adsorption characteristics of the stationary phase.
- Silica-based ion exchangers (NUCLEOSIL® SA and SB) are stable from pH 2 to 8 and do not swell. Compared to resin-based ion exchangers they offer the advantage of constant permeability, even when the ionic strength and/or pH of the eluent are changed. The separation can be influenced by
- the type of buffer
- the ionic strength and
- the pH value.

A tabular overview of NUCLEOSIL® phases can be found on page 212.

NUCLEOSIL® phase overview



Phase	NUCLEOSIL® HPLC phases Specification	Page	Stability	Interactions	Structu	ra
NUCLEOSIL® RF		1 age	Glability	Interactions	Ollucial	
C ₁₈	octadecyl phase, medium density modification, endcapping 15 % C · USP L1	214	pH 2–8	hydrophobic (van der Waals) interactions slight residual silanol interactions	NUCLEOSIL® (Si-O ₂) _n	Si O Si(CH ₃) ₃
C ₁₈ HD	octadecyl phase, high density monomeric modification, end- capping 20 % C · USP L1	214	pH 2-9	hydrophobic (van der Waals) interactions	NUCLEOSIL® (Si-O ₂) _n	
C ₁₈ AB	octadecyl phase, special crosslinked modification, endcapping 25 % C · USP L1	214	pH 1–9	steric and hydrophobic interactions	NUCLEOSIL® (Si-O ₂),	
C ₁₈ Nautilus	octadecyl phase, embedded polar group, endcapping 16 % C · USP L60	214	pH 2–8 up to 100 % H ₂ O	hydrophobic and polar interactions	NUCLEOSIL® (Si-O ₂) _n	Pol Si-OH Si-O Si(CH ₃) ₃
Protect I	special RP phase, protective polar group, monomeric modi- fication, endcapping 11 % C	216	pH 2–8 up to 100 % H ₂ O	hydrophobic and polar interactions	NUCLEOSIL® (Si-O ₂),	Si-O-Si(CH ₃) ₃
C ₈ ec	octyl phase, medium density modification, endcapping 9 % C · USP L7	217	pH 2–8	hydrophobic (van der Waals) interactions slight residual silanol interactions	NUCLEOSIL® (Si-O ₂) _n	Si-OH Si-OSi(CH ₃) ₃
C ₈	octyl phase, no endcapping 8.5 % C · USP L7	217	pH 2–8	hydrophobic (van der Waals) interactions noticeable residual silanol interac- tions	NUCLEOSIL® (Si-O ₂) _h	Si-OH
C ₈ HD	octyl phase, high density modification, endcapping 13 % C · USP L7	218	pH 2–8	hydrophobic (van der Waals) interactions	NUCLEOSIL® (Si-O ₂) _n	
C ₄	butyl phase, medium density modification, endcapping ~ 2 % C · USP L26	219	pH 2-8	hydrophobic (van der Waals) interactions residual silanol interac- tions	NUCLEOSIL® (Si-O ₂) _n	Si-OH Si-O Si(CH ₃) ₃

NUCLEOSIL® phase overview



Phase	IUCLEOSIL® HPLC phases Specification	Page	Stability	Interactions	Structu	re
C ₂	dimethyl phase 3.5 % C · USP L16	219	pH 2-8	hydrophobic (van der Waals) interactions noticeable residual silanol interactions	NUCLEOSIL® (Si-O ₂) _n	Si-OH Si-O-Si(CH ₃) ₂ Si-OH
C_6H_5	phenyl phase, no endcapping 8 % C · USP L11	220	pH 2-8	π-π interactions and hydrophobic interactions noticeable residual silanol interactions	NUCLEOSIL® (Si-O ₂) _n	Si-OH
	_® phases and NUCLEOSIL® ion e	xchange	rs			
	cyano (nitrile) phase USP L10	222	pH 2-8	π– $π$, polar and hydrophobic interactions	NUCLEOSIL® (Si-O ₂) _n	C=N Si-OH C=N
CN/CN-RP						
OH (Dial)	diol · USP L20	220	pH 2-8	polar interactions (hydro- gen bonds)	NUCLEOSIL® (Si-O ₂) _n	Si-OH OH
OH (Diol)						
	amino · USP L8	221	pH 2–8	polar and hydrophobic interactions, weak ion exchange interactions	NUCLEOSIL® (Si-O ₂) _n	NH ₂ Si-OH NH ₂ Si-OH
NH ₂ / NH ₂ -RP						
N/CH.).	dimethylamino	221	pH 2-8	polar and hydrophobic interactions, weak ion exchange interactions	NUCLEOSIL® (Si-O ₂) _n	Si-OH CH ₃
N(CH ₃) ₂						
20	sulfonic acid, strongly acid cation exchanger (SCX) USP L9	223	pH 2-8	strong ion exchange interactions	NUCLEOSIL® (Si-O ₂) _n	Si-OH SO ₃ Na Si-OH SO ₃ Na
SA						
	quaternary ammonium, strongly basic anion exchanger (SAX) USP L14	223	pH 2-8	strong ion exchange interactions	NUCLEOSIL® (Si-O ₂) _n	Si-OH CH ₃ Cr NCH ₃ CH ₃ CH ₃
SB						
	unmodified spherical silica USP L3	224	pH 2-8	polar	NUCLEOSIL® (Si-O ₂) _n	Si-OH
SiOH						



NUCLEOSIL® octadecyl phases (C₁₈)

NUCLEOSIL® standard octadecyl phases · USP L1

Technical data

- · Nonpolar phases
- · pH stability at 20 °C: 2-8
- · carbon content depending on pore size (see table)
- · Corresponding NUCLEODUR® phases see C₁₈ ec page 181

NUCLEOSIL® C18 HD · USP L1

-(CH₂)₁₇-CH₃

-(CH₂)₁₇-CH₃

Technical data

- · Nonpolar hydrophobic high density phases; monomeric modification
- · pH stability 2-9

- · Carbon content 20 %
- · Corresponding NUCLEODUR® phases see C₁₈ Gravity page 158

NUCLEOSIL® C₁₈ AB · USP L1

-(CH₂)₁₇-CH₃

-(CH₂)₁₇-CH₃

Technical data

- · Crosslinked hydrophobic phase; polymeric modification; inert towards acidic and basic substances with high affinity for silica
- · pH stability 1-9

- · Carbon content 25 %; distinct steric selec-
- · Corresponding NUCLEODUR® phases see C₁₈ Isis page 164

NUCLEOSIL® C₁₈ Nautilus · USP L60

Technical data

- · Stable in 100 % aqueous eluents
- · Carbon content 16 %
- · Interesting polar selectivity features; very good base deactivation

· Corresponding NUCLEODUR® phases see C₁₈ PolarTec page 168

All NUCLEOSIL® octadecyl phases are endcapped.

Custom-packed columns with different column dimensions are available on request.

Ordering information

Eluent in column acetonitrile - water

	100 mm	125 mm	150 mm	250 mm	EC guard columns*
NUCLEOSIL® 50-5 C ₁₈ ec	particle size 5 µm, por	re size 50 Å, endcappe	ed, 14.5 % C		

Analytical EC columns



$NUCLEOSIL^{\$}$ 100-3 C_{18} particle size 3 μm , pore size 100 Å, endcapped, 15 % C

Lenath →

Analytical EC columns

4 mm		720150.40		720133.40	721022.30
4.6 mm	720841.46	720150.46	720949.46	720133.46	721022.30

NUCLEOSIL[®] 100-5 C₁₈ particle size 5 μm, pore size 100 Å, endcapped, 15 % C

Ana

nalytical EC columi	ns						
	2 mm		720002.20		720014.20	721074.20	
	3 mm		720002.30		720014.30	721074.30	
	4 mm	720141.40	720002.40	720120.40	720014.40	721074.30	
	4.6 mm	720141.46	720002.46	720120.46	720014.46	721074.30	



NUCLEOSIL® columns



Ordering information						
Eluent in column acetonit	rile – water					
ID	Length - 100 mm		150 m	m	250 mm	EC guard columns*
NUCL FOSII ® 100-7		n, pore size 100 Å, endcap			200	20 guara corannio
Analytical EC columns	ο 18 ραιτισίο σί2ο / μπ	ι, ροιο 6120 100 / ι, οπασαργ	300, 10 70 0			
•	mm				720018.40	
4.6	mm	720951.46	72011	0.46	720018.46	
NUCLEOSIL® 100-10	C ₁₈ particle size 10	μm, pore size 100 Å, endc	apped, 15 % C			
Analytical EC columns					=======================================	
[]	· mm · mm	720701.46	72014	0.46	720023.40 720023.46	
4.0	· 11 11	720701.40	72014	0.40	720023.40	
NUCLEOSIL® 120-3	$C_{18}\;$ particle size 3 µm	n, pore size 120 Å, endcap _l	oed, 11 % C			
Analytical EC columns						
()	mm 720149. mm 720149.	·····	72074	0.46	720055.40	721075.30
				0.46	720055.46	721075.30
Analytical EC columns	U ₁₈ particle size 5 μm	n, pore size 120 Å, endcapp	bed, 11% C			
•	· mm	720051.40			720041.40	721070.30
[[]	mm	720051.46	72073	0.46	720041.46	721070.30
NUCLEOSIL® 120-7	C ₁₈ particle size 7 μm	n, pore size 120 Å, endcap	oed, 11 % C			
Analytical EC columns						
	mm				720042.40	
	C ₁₈ particle size 10	μm, pore size 120 Å, endo	apped, 11 % C			
Analytical EC columns					700040 40	
4 1	mm				720043.40 720043.46	
	C ₁₈ HD particle size	3 μm, pore size 100 Å, 20 °	% C			
Analytical EC columns	nm	720191.40				721196.30
()	mm	720191.46	72019	3.46	···•	721196.30
NUCL FOSII ® 100-5	C ₁₀ HD particle size	5 μm, pore size 100 Å, 20 °		-		
Analytical EC columns	0 18 · · · 2 par iloro oizo	о рин, рего оде тост, де	,,,			
•	mm	720296.40			720280.40	721072.30
4.6	mm	720296.46	72029	4.46	720280.46	721072.30
NUCLEOSII ® 100-5	Cus AB particle size	5 μm, pore size 100 Å, 25 9	% C			
Analytical EC columns	O ₁₈ / (D) partiolo size	o pm, pore size 10071, 207	,			
•	mm	720935.40			720936.40	721073.30
4.6	mm	720935.46	72030	5.46	720936.46	721073.30
NUCLEOSII® 100-3	C Nautilus partiol	e size 3 μm, pore size 100	Å 16%C			
Analytical EC columns	O ₁₈ reactifus partici	e size o piri, pore size 100	л, 10 /0 С			
•	mm	720472.40				721649.30
[]	mm	720472.46	72047	1.46		721649.30
NUCLEOSIL® 100-5	C ₁₈ Nautilus particl	e size 5 μm, pore size 100	Å, 16 % C			
Analytical EC columns						
[[]	mm	720430.40			720431.40	721133.30
4.6	mm	720430.46	72043	2.46	720431.46	721133.30
Guard column system	n					
Guard columns for EC co		2 mm	3 mm	4 mm	4.6 mm	Guard column holde
* Column Protection Syste	m (pack of)	EC 4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966

NUCLEOSIL® octadecyl phases (C₁₈) wide pore octadecyl phases · USP L1

Technical data

-(CH₂)₁₇-CH₃

- · Many biologically interesting molecules can not be separated using conventional narrow pore silicas with pore sizes of about 100 Å. This is why MACHEREY-NAGEL offers a complete line of wide pore packings with pore sizes of 300, 500, 1000 and 4000 Å.
- · These materials can also be used for size exclusion chromatography (SEC).

All NUCLEOSIL® octadecyl phases are endcapped.

Custom-packed columns with different column dimensions are available on request.

Ordering information Eluent in column acetonitrile - water Lenath → 250 mm EC guard columns* NUCLEOSIL[®] 300-5 C₁₈ particle size 5 μm, pore size 300 Å, endcapped, 6.5 % C Analytical EC columns 720065.40 721085.30 720065.46 721085.30 NUCLEOSIL 8 500-7 C_{18} particle size 7 μm , pore size 500 Å, endcapped, 2 % CAnalytical EC columns 720074.46 4.6 mm $NUCLEOSIL^{\circledR}$ 1000-7 $C_{18}~$ particle size 7 $\mu m,$ pore size 1000 Å, endcapped, \sim 1 % C Analytical EC columns 4.6 mm 720077.46

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

VarioPrep preparative HPLC columns with NUCLEOSIL® packing material on request.

NUCLEOSIL® 100 Protect I special RP phase with protective polar group

Technical data

- · RP phase with pronounced hydrophilic properties
- Endcapped

· Monomeric coating

· Carbon content 11 %

Ordering information

Eluent in column acetonitrile - water

	ID	Length →			
		125 mm	150 mm	250 mm	EC guard columns*
NUCLEOSIL® 10	0-5 Protect	particle size 5 μm, pore size 100 Å			
Analytical EC column	ns				
	4 mm	720175.40		720170.40	721157.30
	4.6 mm	720175.46	720174.46	720170.46	721157.30

Guard co	lumn	sys	tem
O			

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



NUCLEOSIL® octyl phases (C₈) NUCLEOSIL® standard octyl phases · USP L7

-(CH₂)₇-CH₃

Technical data

- Nonpolar phases for RP and ion-pairing chromatography
- Endcapped and non-endcapped modifications available; pH stability at 20 °C: 2–8
- · Carbon content depending on pore size (see table)

✓ Recommended application

- Separation of moderately to highly polar (water-soluble) compounds: steroids, nucleosides, cyclodextrins, pharmacological plant constituents
- Corresponding NUCLEODUR® phases see C₈ ec page 183

Ordering information						
Eluent in column acetonitrile – water ID		Langth .				
U		Length → 125 mm	150 mm		250 mm	EC guard columns*
NUCLEOSIL® 100-5 C ₈ ec parti	icle size 5 µm, pore	size 100 Å, endca	oped, 9 % C			
Analytical EC columns						
4.6 mm					720165.46	721096.30
NUCLEOSIL® 100-5 C ₈ particle	size 5 µm, pore size	e 100 Å, not endca	pped, 8.5 % C			
Analytical EC columns						
4 mm		720001.40			720013.40	721194.30
4.6 mm		720001.46	720990.4	16	720013.46	721194.30
NUCLEOSIL® 100-7 C ₈ particle	size 7 µm, pore size	e 100 Å, not endca	pped, 8.5 % C			
Analytical EC columns						
4.6 mm					720017.46	
NUCLEOSIL® 100-10 C ₈ particle	e size 10 µm, pore s	size 100 Å, not end	capped, 8.5 % C			
Analytical EC columns						
4 mm					720022.40	·····
4.6 mm					720022.46	
NUCLEOSIL® 120-3 C ₈ particle	size 3 um. pore size	e 120 Å. not endca	pped. 6.5 % C			
Analytical EC columns		, , , , , , , , , , , , , , , , , , , ,				
4 mm		720071.40				721093.30
4.6 mm		720071.46	720214.4	16	····	721093.30
NUCLEOSIL® 120-5 C ₈ particle	size 5 µm, pore size	e 120 Å, not endca	pped, 6.5 % C			
Analytical EC columns			,			
4 mm		720050.40			720052.40	721095.30
4.6 mm		720050.46	720735.4	16	720052.46	721095.30
		9				
NUCLEOSIL® 300-5 C ₈ particle	size 5 µm, pore size	e 300 A, not endca	pped, ~ 3 % C			
Analytical EC columns					700000 10	704004.63
4.6 mm					720062.46	721061.30
EC columns in packs of 1, guard colum	nns in packs of 3.					
Custom-packed columns with different	column dimensions	are available on red	quest.			
Guard column system						
Guard columns for EC columns with I	D	2 mm	3 mm	4 mm	4.6 mm	Guard column holde
* Column Protection System (pack of)	EC	4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966



NUCLEOSIL® octyl phases (C8) NUCLEOSIL® C8 HD · USP L7

-(CH₂)₇-CH₃

Technical data

- · Nonpolar high density phases; monomeric modification; endcapped; carbon content 13%
- · Corresponding NUCLEODUR® phases see C₈ Gravity page 158

Recommended application

· Separation of moderate to strong polar (water soluble) analytes like steroids, cyclodextrines, pharmalogical plant ingredients

Ordering information

Fluent in column acetonitrile – water

Lident in Coldini ac	betoritine – water				
	ID	Length →			
		125 mm	150 mm	250 mm	EC guard columns*
NUCLEOSIL® 1	00-5 C ₈ HD particle size 5 μm, pore s	size 100 Å			
Analytical EC colum	ns				
	4 mm			720196.40	721071.30
	4.6 mm		720194.46	720196.46	721071.30
CO and mana in model	f 1				

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

Custom-packed columns with different column dimensions are available on request.

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

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



Beside analytical HPLC columns we also produce VarioPrep columns (see page 252) for preparative applications.



NUCLEOSIL® columns



NUCLEOSIL® butyl phases (C₄) · USP L26

-(CH₂)₃-CH₃

Technical data

- · Endcapped phases for RP and ion-pairing chromatography
- · pH stability at 20 °C: 2-8; carbon content ~
- \cdot Retention times are shorter than on C_8 and C₁₈ phases

Recommended application

- · For separation of macromolecules and hydrophobic substances
- · For butyl phases for biochemical separations please refer to page 241

Ordering informa	ation		
Eluent in column ac	eetonitrile – water		
	ID	Length → 250 mm	EC guard columns*
NUCLEOSIL® 12	$205~C_4~$ particle size 5 µm, pore size 120 Å		
Analytical EC column	ns		
	4.6 mm	720096.46	721083.30
NUCLEOSIL® 30	$00-5$ C_4 particle size 5 μm, pore size 300 Å		
Analytical EC column	ns		
	4 mm	720059.40	721916.30
	4.6 mm	720059.46	721916.30
EC columns in packs	s of 1, guard columns in packs of 3.		
Guard column s	ystem		

3 mm

4/3 (3)

4 mm

4/3 (3)

NUCLEOSIL® dimethyl phase (C2) · USP L16



Guard columns for EC columns with ID

* Column Protection System (pack of)



· Non-endcapped phase for RP and ion-pairing chromatography

2 mm

4/2 (3)

- · pH stability at 20 °C: 2-8; carbon content 3.5%
- · Retention times are much shorter than for the other RP phases

4.6 mm

4/3 (3)

Guard column holder

718966

Ordering information

Eluent in column acetonitrile – water		
ID	Length →	
	250 mm	EC guard columns*
NUCLEOSIL [®] 100-7 C ₂ particle size 7 μm, pore size 100 Å		
Analytical EC columns		
4.6 mm	720089.46	721030.30



NUCLEOSIL® phenyl phases (C₆H₅) · USP L11

Technical data

- · Relatively nonpolar, non-endcapped phases for RP and ion pairing chromatography
- · Polarity similar to C₈, but with different selectivity for PAHs, polar aromatics, fatty acids
- · pH stability at 20 °C: 2-8; carbon content 8%

Recommended application

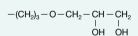
· Separation of moderately polar compounds

Ordering information

Eluent in column acetonitrile - water

ID	Length → 250 mm	EC guard columns*
NUCLEOSIL $^{\$}$ 100-5 C_6H_5 particle size 5 μ m, pore size 100 Å, not endcapped		
Analytical EC columns		
4.6 mm	720956.46	721137.30
NUCLEOSIL [®] 100-7 C ₆ H ₅ particle size 7 μm, pore size 100 Å, not endcapped		
Analytical EC columns		
4 mm	720019 40	

NUCLEOSIL® diol phases · USP L20





- · Dihydroxypropyl modified silica for RP and NP chromatography
- · Less polar than unmodified silica, very easily wettable with water
- · pH stability at 20 °C: 2-8; carbon content 5%

720019.46

Ordering information

Eluent in column is n-heptane. When using an eluent which is not miscible with n-heptane (e.g., water), it is necessary to rinse the column with THF first.

ID	Length → 250 mm	EC guard columns*
NUCLEOSIL® 100-5 OH (Diol) particle size 5 μm, pore size 100 Å		
Analytical EC columns		
4.6 mm	720143.46	721142.30

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



NUCLEOSIL® amino phases · USP L8

Technical data

- \cdot Aminopropyl modified polar silica phase; pH stability at 20 °C: 2–8; carbon content 3.5 %
- Corresponding NUCLEODUR® phases see page 188

Н

Recommended application

Multi-mode chromatography

- NP chromatography with hexane, dichloromethane or 2-propanol as mobile phase for polar compounds such as substituted anilines, esters, chlorinated pesticides
- RP chromatography of polar compounds like carbohydrates in aqueous-organic eluent systems
- Anion exchange chromatography of anions and organic acids using common buffers (e.g., acetate or phosphate) in conjunction with organic modifiers (e.g., acetonitrile)

Ordering information

-(CH₂)₃-NH₂

Eluent in column is *n*-heptane (except for NH₂ RP). When using an eluent which is not miscible with *n*-heptane (e.g., water), it is necessary to rinse the column with THF first.

COIGITITI WILLT TITLE III	J.,		
	ID	Length →	
		250 mm	EC guard columns*
NUCLEOSIL® 10	00-5 NH ₂ particle size 5 μm, pore size 100 Å; eluent in column <i>n</i> -heptane		
Analytical EC column	ns		
	4.6 mm	720095.46	721020.30
NUCLEOSIL® 10	00-5 NH ₂ -RP particle size 5 μm, pore size 100 Å; eluent in column acetonitrile – w	vater (80:20)	
Analytical EC column	ns		
	4.6 mm	720095.46RP	721155.30
NUCLEOSIL® 10	00-10 NH ₂ particle size 10 μm, pore size 100 Å; eluent in column <i>n</i> -heptane		
Analytical EC column	ns		
	4.6 mm	720025.46	

NUCLEOSIL® dimethylamino phase

-(CH₂)₃-N(CH₃)₂



- Weakly basic anion exchanger, pH stability at 20 °C: 2–8; carbon content 4 %
- Recommended application
- · Separation of many anions; can also be used in a similar way as the NH₂ phase

Ordering information

Eluent in column is n-heptane. When using an eluent which is not miscible with n-heptane (e.g., water), it is necessary to rinse the column with THF first.

ID	Length →	
	250 mm	EC guard columns*
$NUCLEOSIL^{\oplus}$ 100-5 $N(CH_3)_2$ particle size 5 μm , pore size 100 Å		
Analytical EC columns		
4.6 mm	720994.46	721158.30

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



NUCLEOSIL® cyano phases · USP L10

Technical data

- · Polar to midpolar cyano (nitrile) modified silica
- \cdot pH stability at 20 °C: 2–8; carbon content 5 % for 100 Å pores, \sim 3 % for 120 Å pores
- Corresponding NUCLEODUR® phases see page 186

Recommended application

Reversed phase and normal phase chromatography

- Normal phase: with low-polarity solvents for many compounds, which can also be separated on unmodified silica, however, due to the rapid equilibration much more suitable for gradient separations
- Reversed phase:
 with different selectivity than C₁₈, C₈ or phenyl modified packings

Ordering information

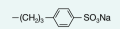
-(CH₂)₃-CN

Eluent in column (except for NUCLEOSIL® 100-5 CN-RP) is *n*-heptane. When using an eluent which is not miscible with *n*-heptane (e.g., water), it is necessary to rinse the column with THE first

necessary to rinse	the column with THF first.		
, , , , , , , , , , , , , , , , , , , ,	ID	Length →	
		250 mm	EC guard columns*
NUCLEOSIL® 1	00-5 CN particle size 5 μm, pore size 100 Å; eluent in column <i>n</i> -heptane		
Analytical EC colum	ns		
	4 mm	720090.40	721078.30
	4.6 mm	720090.46	721078.30
NUCLEOSIL® 10 Analytical EC colum	00-5 CN-RP particle size 5 μ m, pore size 100 Å; eluent in column acetonitrile -	- water	
	4 mm	720205.40	721039.30
	4.6 mm	720205.46	721039.30
NUCLEOSIL® 10	00-10 CN particle size 10 μm, pore size 100 Å; eluent in column <i>n</i> -heptane		
Analytical EC colum	ns		
	4 mm	720024.40	
	4.6 mm	720024.46	
NUCLEOSIL® 12	20-7 CN particle size 7 μm, pore size 120 Å; eluent in column <i>n</i> -heptane		
Analytical EC colum	ins		
	4 mm	720057.40	
	4.6 mm	720057.46	



NUCLEOSIL® SA phases · USP L9

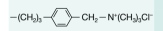


Technical data

- · Strongly acidic cation exchanger (SCX) with benzenesulfonic acid modification
- · Capacity ~ 1 meq/g; pH stability at 20 °C: 2-8; carbon content 6.5 %

Ordering informa	ation				
Eluent in column 0.	15 mol/L (NH ₄) ₂ HPO ₄ , pH 5				
	ID	Length →			
		125 mm	150 mm	250 mm	EC guard columns*
NUCLEOSIL® 10	00-5 SA particle size 5 μm, pore size	100 Å			
Analytical EC column	าร				
	4 mm			720097.40	721024.30
	4.6 mm	720709.46	720182.46	720097.46	721024.30
NUCLEOSIL® 10	00-10 SA particle size 10 μm, pore si.	ze 100 Å			
Analytical EC column	าร				
	4.6 mm			720028.46	

NUCLEOSIL® SB phases · USP L14



- Technical data
- · Strongly basic anion exchanger (SAX) with quaternary ammonium modification
- · Capacity ~ 1 meq/g; pH stability at 20 °C: 2-8; carbon content 10 %

		.,			-,	
Ordering informa	ation					
Eluent in column 0.1	15 mol/L (NH ₄)₂HPO₄, pH 5				
	ID		Length → 125 mm	150 mm	250 mm	EC guard columns*
NUCLEOSIL® 10	00-5 SB par	rticle size 5 µm, pore size	100 Å			
Analytical EC column	ns					
	4 mm				720996.40	721025.30
	4.6 mm		720989.46	720183.46	720996.46	721025.30
NUCLEOSIL® 10	00-10 SB p	article size 10 µm, pore s	ize 100 Å			
Analytical EC column	าร					
	4.6 mm				720029.46	



EC guard columns*

NUCLEOSIL® SiOH unmodified silica · USP L3

Technical data

- · Spherical silica, pH stability 2-8
- For physical properties of unmodified NUCLEOSIL® materials please see page 211.
- Maximum working pressure for the EC columns listed below is 400 bar.

Length → 250 mm

Ordering information

Eluent in column is n-heptane. When using an eluent which is not miscible with n-heptane (e.g., water), it is necessary to rinse the column with THF first.

)

 $NUCLEOSIL^{\$}$ 50-5 particle size 5 µm, pore size 50 Å

Analytical EC columns

4.6 mm 720093.46 721167.30

 $NUCLEOSIL^{\circledR}$ 100-5 particle size 5 µm, pore size 100 Å

Analytical EC columns

4.6 mm 720099.46 721518.30

Guard column system

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



Analytical columns with LiChrospher®



LiChrospher® packings manufactured by E. Merck (D)

Phase	USP	Particle size	Pore size	Modification	Endcapped	Carbon content
LiChrospher® 100 RP 18, 5 µm	L1	nom. 5 µm	100 Å	Octadecyl	_	21 %
LiChrospher® 100 RP 18 ec, 5 µm	L1	nom. 5 µm	100 Å	Octadecyl	+	21 %
LiChrospher® 60 RP select B, 5 µm	L7	nom. 5 µm	60 Å	Octyl	+	12 %
All phases as packed ChromCart® carti	ridges					
ChromCart® columns require the CC or	onnocting kit	(DEE 701600)				

Ordering information

Eluent in column acetonitrile - water

ID	Length →			
	125 mm	150 mm	250 mm	Guard columns*
LiChrospher [®] 100 RP 18, 5 μm	η particle size 5 μm, pore size 100 Å			
2 mm	728031.20		728032.20	728053.30
3 mm	728031.30	•	728032.30	728053.30
4 mm	728031.40		728032.40	728053.40
4.6 mm	728031.46	728033.46	728032.46	728053.40
LiChrospher® 100 RP 18 ec, 5	μm particle size 5 μm, pore size 100 Å			
2 mm	728034.20		728035.20	728054.30
3 mm	728034.30		728035.30	728054.30
4 mm	728034.40		728035.40	728054.40
4.6 mm	728034.46	728036.46	728035.46	728054.40
LiChrospher® 60 RP select B,	5 μm particle size 5 μm, pore size 100 Å			
2 mm	728037.20		728038.20	728055.30
3 mm	728037.30	•	728038.30	728055.30
4 mm	728037.40		728038.40	728055.40
4.6 mm	728037.46	728039.46	728038.46	728055.40

8 mm ChromCart® guard column cartridges in packs of 3, all other columns in packs of 1.



Phase overview for special separations



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

11/1

HPLC columns for environmental analyses



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

Technical data

Guard columns for EC columns with ID

* Column Protection System (pack of)

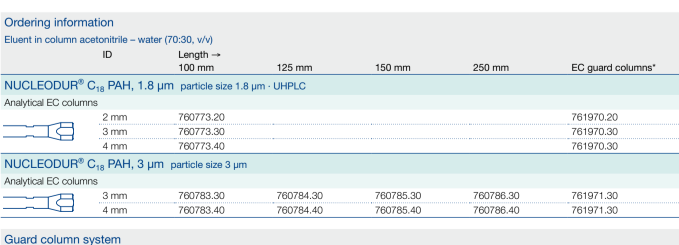
 \cdot Base material NUCLEODUR® silica, particle sizes 1.8 and 3 µm, pore size 110 Å; polymeric coating

Recommended application

Allows efficient gradient separation of the 16 PAHs according to EPA

Analysis of 16 EPA PAHs with or without acetonitrile MN Appl. Nos. 123820/123830 Separation with acetonitrile Separation without acetonitrile Peaks: Column: 100 x 4 mm Column: 125 x 4 mm 1. Naphthalene NUCLEODUR® C18 PAH, 3 µm NUCLEODUR® C18 PAH, 3 µm 2. Acenaphthylene (not detectable by Eluent: A) methanol – water (80:20, v/v) Eluent: fluorescence) B) acetonitrile 2-20 % B in 1.2 min, B) methanol 65-97 % B in 6 min, 3. Acenaphthene 20-100 % B in 0.5 min. 100 % B 97 % B for 5 min. 97-65 % B in 4. Fluorene for 2.5 min, 100-2 % B in 0.4 min 0.5 min 5. Phenantrene Flow rate: 2.5 mL/min, temperature 35 °C Flow rate: 2 mL/min, temperature 35 °C 6. Anthracene Detection: UV. 254 nm fluorescence (see chromatogram) Detection: 7. Fluoranthene fluorescence (see chromatogram) 8. Pyrene 9. Benz[a]anthracene 10. Chrysene 10 11. Benzo[b]fluoranthene 12. Benzo[k]fluoranthene 13. Benzo[a]pyrene 14. Dibenz[ah]anthracene 15. Benzo[ghi]perylene 16. Indeno[1,2,3-cd]pyrene 16 10 375 425 330 420 335 440 315 330 375 345 405 420 460 420 315

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



3 mm

4/3 (3)

4 mm

4/3 (3)

2 mm

4/2 (3)

EC

Guard column holder

718966

4.6 mm

4/3 (3)

Column:

Eluent:

Flow rate:

Injection:

Detection:

Peaks:

Temperature:

Fluorescence:

1.-16. see page 227

1-me-n: 1-methylnaphthalene 2-me-n: 2-methylnaphthalene

125 x 4 mm

A) methanol - water (70:30, v/v); B) acetonitrile 0-20 % B in 1.5 min. 20-50 % B in 1.5 min,

50-100 % B in 1.0 min,

100 % B for 3 min,

1.5 ml /min

35 °C

0.5 µL

(concentrations 10 ng/µL per compound)

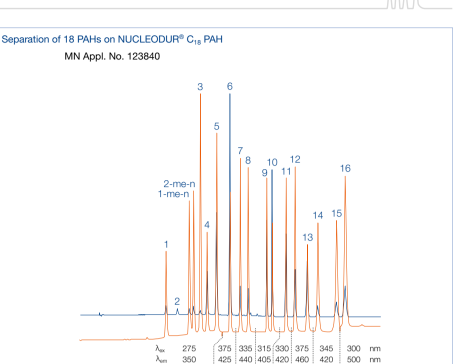
UV: 1 μL,

UV. 254 nm fluorescence (see chromatogram)

100-0 % B in 0.5 min

NUCLEODUR® C₁₈ PAH, 3 µm

HPLC columns for environmental analyses



Analysis of polycyclic aromatic hydrocarbons (PAHs) by HPLC

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

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

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



500 nm

HPLC columns for PAH analysis

For PAH analyses we have developed specially modified C_{18} phases based on NUCLEODUR® and NUCLEOSIL® which allow efficient gradient separation of 16 PAHs according to EPA. Detection of the separated PAHs can be achieved by UV (250-280 nm), with diode array or with fluorescence detection at different wavelengths for excitation and emission. Acenaphthylene cannot be analyzed with fluorescence detection. For cost-effective routine PAH analysis we recommend applications using methanol instead of acetonitrile as eluent. For rapid analysis NUCLEODUR® C₁₈ PAH (3 μm) in short columns (100 mm) provides excellent results at high flow rates. Hereby separation of 16 PAHs according to EPA can be achieved in less than 3 min.

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

HPLC columns for environmental analyses



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

Technical data

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

Recommended application

· Efficient gradient separation of the 16 PAHs according to **EPA**

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

MN Appl. No. 115040

150 x 4 mm NUCLEOSIL® 100-5 C₁₈ PAH Column:

A) methanol - water (80:20) Eluent:

> B) acetonitrile - tetrahydrofuran (93:7) 0-100 % B in 10 min, 5 min 100 % B

Flow rate: 1 mL/min Pressure: 140 bar 20 °C Temperature: Detection: UV, 260 nm

Peaks: (10 µg/mL each in acetonitrile)

1. Naphthalene

10. Chrysene

2. Acenaphthylene

11. Benzo[b]fluoranthene

3. Acenaphthene

12. Benzo[k]fluoranthene

4. Fluorene

13. Benzo[a]pyrene

5. Phenanthrene

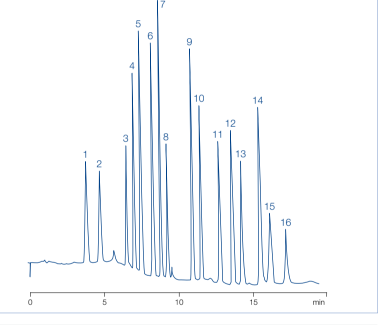
14. Dibenz[ah]anthracene

6. Anthracene

15. Benzo[ghi]perylene 16. Indeno[1,2,3-cd]pyrene

7. Fluoranthene 8. Pyrene

9. Benz[a]anthracene



Ordering information

Eluent in column acetonitrile - water 70:30

Length → 150 mm 250 mm EC guard columns* NUCLEOSIL® 100-5 C₁₈ PAH particle size 5 µm, pore size 100 Å

Analytical EC columns

licai Lo colui i	1110				
	2 mm		720117.20	721168.20	
	3 mm	720923.30	720117.30	721168.30	
	4 mm	720923.40	720117.40	721168.30	
	4 6 mm	•	720117 46	721168.30	

PAH standard according to EPA for HPLC

Analytical EC columns

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

Cuard column avetem

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

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

HPLC columns for environmental analyses

Anion columns for analysis of inorganic anions

NUCLEOGEL® Anion I

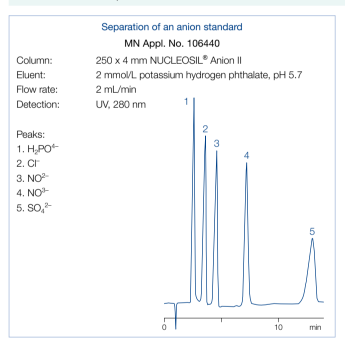
Technical data

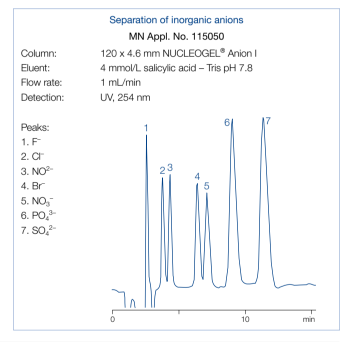
- \cdot Strongly basic polymer-based anion exchanger, particle size 10 μ m; pH stability 1–14
- · Eluent in column 4 mmol/L salicylate buffer pH 7.8
- Contrary to the silica-based phase also suited for fluoride analysis

NUCLEOSIL® Anion II

Technical data

- \cdot Base material NUCLEOSIL® silica, particle size 10 µm, pore size 300 Å strongly basic anion exchanger, exchange capacity 50 µeq/g, pH stability 2–7.5
- Eluent in column 0.15 mol/L (NH₄)₂HPO₄ buffer pH 5.2 recommended buffer concentration for separation of inorganic anions: 2 mmol/L phthalate
- Preferred method of detection: conductivity or negative UV detection





Ordering information			
ID	Length → 120 mm	250 mm	Guard columns*
NUCLEOGEL® Anion I eluent 4 mmol/L salicylate buffer pH 7.8			
Analytical Valco type columns			
4.6 mm	719533		719543
NUCLEOSIL® Anion II eluent 0.15 mol/L (NH ₄) ₂ HPO ₄ buffer pH 5.2			
Analytical EC columns			
4 mm		720094.40	721169.30

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

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



NUCLEODEX columns enantiomer separation based on cyclodextrins

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

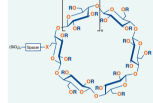
Technical data

- Base material NUCLEOSIL[®] silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Separation based on hydrogen bonds and dipole interactions between functional groups of the analyte and hydroxyl groups of the cyclodextrin
- Examples for successful enantiomer separations: chlorthalidone and other compounds, which require free hydroxyl groups for enantioselective interactions
- Eluent in column CH₃OH 0.1 % TEAA pH 4 (55:45)

NUCLEODEX α -PM permethylated α -cyclodextrin (R = CH₃; n = 1)

Technical data

- Base material NUCLEOSIL[®] silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Examples for successful enantiomer separations: mecoprop and dichlorprop as free carboxylic acids, trans-stilbene oxide, styrene oxide
- Eluent in column CH₃OH 50 mmol/L phosphate pH 3 (70:30)



NUCLEODEX β-PM permethylated β-cyclodextrin (R = CH₃; n = 2) · USP L45

Technical data

- Base material NUCLEOSIL® silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Examples for successful enantiomer separations: mephobarbital (prominal), pesticide derivatives mecoprop methyl and dichlorprop methyl
- Eluent in column CH₃OH 0.1 % TEAA pH 4 (65:35)

NUCLEODEX γ -PM permethylated γ -cyclodextrin (R = CH₃; n = 3)

Technical data

- Base material NUCLEOSIL[®] silica, particle size 5 µm, pore size 100 Å modified cyclodextrins as chiral selectors
- Examples for successful enantiomer separations: steroids or other larger molecules
- Eluent in column CH₃OH 0.1 % TEAA pH 4 (55:45)

Recommended application

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





Separation of the positional isomers of nitroaniline

MN Appl. No. 101420

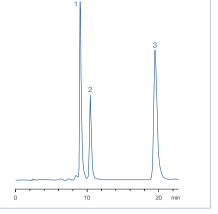
Column: 200 x 4 mm NUCLEODEX β-OH

Eluent: methanol – 0.1 % triethylammonium acetate pH 4.0 (50:50, v/v)

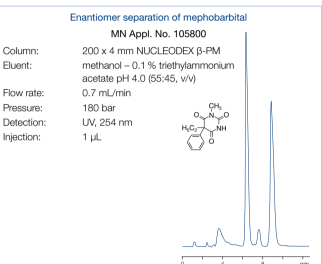
Flow rate: 0.7 mL/min
Pressure: 180 bar
Detection: UV, 254 nm
Injection: 1 µL

Peaks:

m-Nitroaniline
 o-Nitroaniline
 p-Nitroaniline



Enantiomer separation of styrene oxide MN Appl. No. 106160 Column: 200 x 4 mm NUCLEODEX α-PM Eluent: methanol – 0.1 % triethylammonium acetate pH 4.0 (60:40, v/v) Flow rate: 0.7 mL/min Pressure: 160 bar Detection: UV, 230 nm Injection: 2 μL

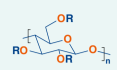


Ordering information		
ID	Length → 200 mm	EC guard columns*
NUCLEODEX β-OH eluent methanol – 0.1 % TEAA pH 4 (55:45)		
Analytical EC columns		
4 mm	720124.40	721171.30
NUCLEODEX α-PM eluent methanol – 50 mmol/L phosphate pH 3 (70:30)		
Analytical EC columns		
4 mm	720127.40	721469.30
NUCLEODEX β-PM eluent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

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



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



Technical data

• Base material silica, chiral selector cellulose tris-(3,5-dimethylphenylcarbamate)
High resolution type (S) with 5 µm particle size, allows use of shorter columns (150 mm) for faster separations, pressure stability up to ~150 bar (2000 psi), pH stability 1–9
NUCLEOCEL DELTA for normal phase applications: eluent in column n-heptane – 2-propanol (90:10, v/v) typical eluents are heptane – propanol mixtures

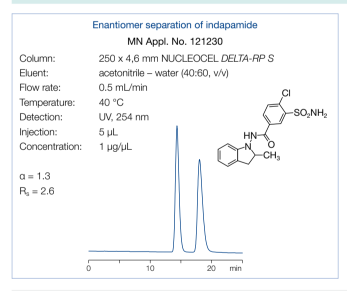
$$\mathbf{R} = \bigvee_{\mathbf{C}}^{\mathbf{H}} \bigvee_{\mathbf{CH}_3}^{\mathbf{CH}_3}$$

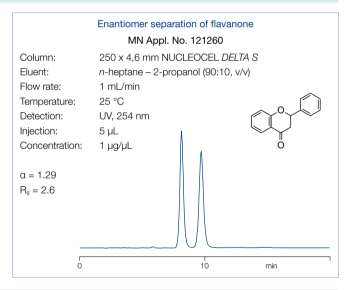
NUCLEOCEL DELTA-RP for reversed phase applications: eluent in column acetonitrile – water (40:60, v/v) designed for use either in polar organic mode or with eluents containing high concentrations of chaotropic salts such as perchlorate

Recommended application

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

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





Ordering information			
ID	Length →		
	150 mm	250 mm	EC guard columns*
NUCLEOCEL DELTA S, 5 µm eluent n-heptane – 2-propanol (90:10, v/v)			
Analytical EC columns			
4.6 mm		720445.46	721185.30
NUCLEOCEL DELTA-RP S, 5 µm eluent acetonitrile – water (40:60, v/v)			
Analytical EC columns			
4.6 mm	720451.46	720450.46	721186.30

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

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

Technical data

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

Recommended application

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

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

MN Appl. No. 105450

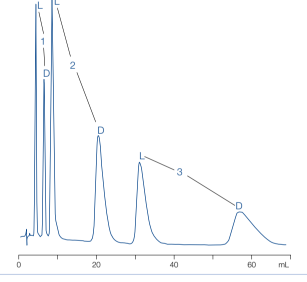
S. Allenmark et al. in "Affinity chromatography and biological recognition" (I. Chaiken, M. Wilchek, and I. Parikh. Eds.), Academic Press, New York, 1983, 259–260

Column: 150 x 4 mm RESOLVOSIL BSA-7 Eluent: 50 mmol/L phosphate buffer pH 6.5

+ 1 % 1-propanol

Flow rate: 0.70 mL/min Detection: UV, 225 nm

Peaks:
1. Serine
2. Alanine
3. Phenylalanine

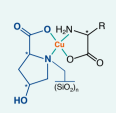


Ordering information Eluent in column 0.1 mol/L phosphate buffer pH 7.5, 2 %	-propanol	
ID	Length → 150 mm	EC guard columns*
RESOLVOSIL BSA-7		
Analytical EC columns		
4 mm	720046.40	721402.30

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



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



Technical data

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

Recommended application

Column:

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

D,L-alanine enantiomers

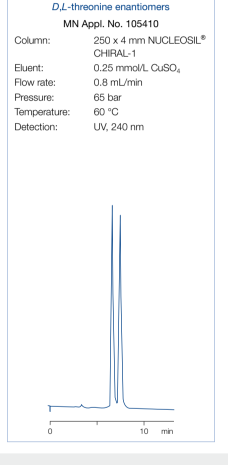
MN Appl. No. 105410

Column: 250 x 4 mm NUCLEOSIL®

CHIRAL-1

Eluent: 0.5 mmol/L CuSO₄

Flow rate: 1 mL/min
Pressure: 60 bar
Temperature: 60 °C
Detection: UV, 250 nm



Eluent: 0.5 mmol/L CuSO₄ Flow rate: 0.8 mL/min Temperature: 60 °C Detection: UV, 240 nm Injection: 1 µL

Lactic acid enantiomers

MN Appl. No. 105560

CHIRAL-1

250 x 4 mm NUCLEOSIL®

Ordering information

Eluent in column 0.5 mmol/L copper sulfate solution

10 min

ID Length → 250 mm EC quard columns*

NUCLEOSIL® CHIRAL-1

Analytical EC columns

4 mm 720081.40 721188.30

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

10 min

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

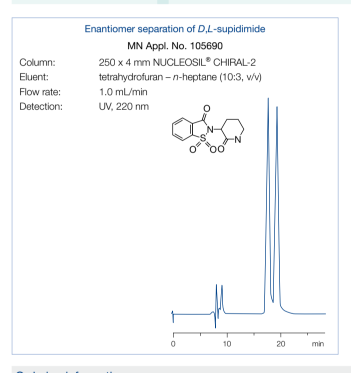
O₂N X Spacer (SiO₃)_n

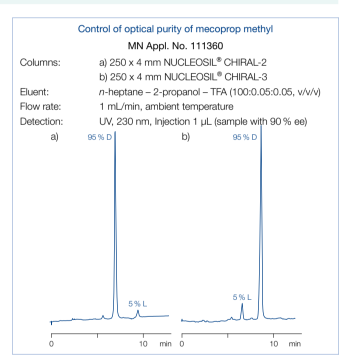
Technical data

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

Recommended application

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





Ordering information		
Eluent in column <i>n</i> -heptane – 2-propanol – TFAA (100:0.05:0.05, v/v/v)		
ID	Length → 250 mm	EC guard columns*
NUCLEOSIL® CHIRAL-2		
Analytical EC columns		
4 mm	720088.40	721190.30
NUCLEOSIL® CHIRAL-3		
Analytical EC columns		
4 mm	720350.40	721190.30

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

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



NUCLEOGEN® columns anion exchange chromatography of nucleic acids

NUCLEOGEN® 60-7 DEAE pore size 60 Å

Technical data

- \cdot Base material silica, particle size 7 $\mu m;$ DEAE anion exchanger
- \cdot For the separation of oligonucleotides up to chain lengths of 40 bases with recoveries $>95\,\%$ capacity 200 $\rm A_{260}/mL$ ($\sim300\,A_{260}$ for a 125 x 4 mm ID column, 1875 $\rm A_{260}$ for a 125 x 10 mm ID column)
- Preparative separations possible when using higher flow rates and longer gradient times

NUCLEOGEN® 500-7 DEAE pore size 500 Å



Technical data

- \cdot Base material silica, particle size 7 $\mu m;$ DEAE anion exchanger
- For the separation of tRNA, 5S RNA, viroids and messenger RNA in the intermediate molecular weight range (25–1 000 kDa) with recoveries > 95 %
- \cdot Capacity 730 $\rm A_{260}$ for a 125 x 6 mm ID column, 1940 $\rm A_{260}$ for a 125 x 10 mm ID column

NUCLEOGEN® 4000-7 DEAE pore size 4000 Å

Technical data

- \cdot Base material silica, particle size 7 $\mu m;$ DEAE anion exchanger
- For the separation of plasmids, DNA restriction fragments, ribosomal RNA, messenger RNA and viral RNA, i.e. very high molecular weight nucleic acids (e.g., 1–50 MDa)
- Capacity 120 A_{260} for a 125 x 6 mm ID column, 350 A_{260} for a 125 x 10 mm ID column

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

Separation of plasmid pBR 322

MN Appl. No. 107480

M. Colpan, D. Riesner, private communication A) isolation of plasmid DNA from a crude cell lysate

5 µg plasmid pBR 322 containing cleared lysate from Sample:

E. coli

Column: 125 x 6 mm NUCLEOGEN® 4000-7 DEAE

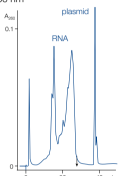
Eluent: A) 20 mmol/L K phosphate buffer pH 6.9; 5 mol/L urea

> B) eluent A + 1.5 mol/L KCl 20-100 % B in 50 min:

arrow = ionic strength of 850 mmol/L

Flow rate: 1.0 mL/min, 70 bar, ambient temperature

Detection: UV, 260 nm



B) separation of supercoiled plasmid from relaxed and linear forms

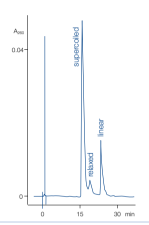
plasmid pBR 322, supercoiled, relaxed and linear Sample:

125 x 6 mm NUCLEOGEN® 4000-7 DEAE Column:

Eluent: A) 20 mmol/L K phosphate buffer pH 6.8; 6 mol/L urea

> B) eluent A + 2 mol/L KCl 42-100 % B in 230 min

Flow rate: 1.5 mL/min, 45 bar, ambient temperature



Separation of oligo(rA)_n

MN Appl. No. 115180

Column: 125 x 4 mm NUCLEOGEN® 60-7 DEAE

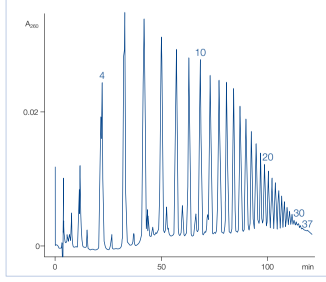
A) 20 mmol/L phosphate buffer, pH 5.5,

5 mol/L urea

B) buffer A + 1 mol/L KCl 0-100 % B in 200 min

Flow rate: 2 mL/min Pressure: 110 bar Temperature: ambient UV, 260 nm Detection:

Eluent:



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

MN Appl. No. 107490

D. Riesner, BioEngineering 1 (1988) 42-48

Column: 125 x 6 mm NUCLEOGEN® 500-7 DEAE

Eluent: A) `250 mmol/L KCl, 20 mmol/L phosphate buffer,

pH 6.6, 5 mol/L urea

B) 1 mol/L KCl, 20 mmol/L phosphate buffer, pH 6.6,

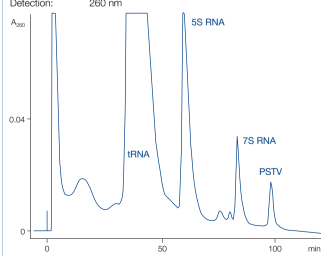
5 mol/L urea

0-50 % B in 120 min, 50-100 % B in 250 min

Flow rate: 3 mL/min

Pressure: 40 bar, ambient temperature

Detection: 260 nm







Ordering informa	ation		
Eluent in column me	ethanol		
	ID	Length → 125 mm	Guard columns*
NUCLEOGEN® (60-7 DEAE particle size 7 μm, pore size 60 Å	Å	
Analytical EC column	ns		
	4 mm	736596.40	736400.40
Preparative VarioPre	p columns		
	10 mm	736597.100	736400.40
NUCLEOGEN® 5	500-7 DEAE particle size 7 μm, pore size 50	00 Å	
Analytical Valco type	columns		
	6 mm	736598	736400.40
Preparative VarioPre	p columns		
	10 mm	736599.100	736400.40
NUCLEOGEN® 4	4000-7 DEAE particle size 7 μm, pore size 4	4000 Å	
Analytical Valco type	columns		
	6 mm	736601	736400.40
Preparative VarioPre	p columns		
	10 mm	736602.100	736400.40
	ard columns are 30 mm long and require the CC f 1, guard columns in packs of 2.	column holder 30 mm (REF 721823).	



NUCLEOGEL® SAX anion exchange of biological macromolecules · USP L23

Technical data

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

Recommended application

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

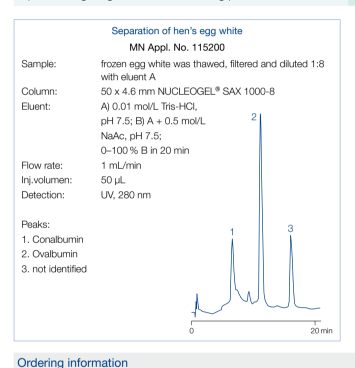
NUCLEOGEL® SCX cation exchange of biological macromolecules · USP L22

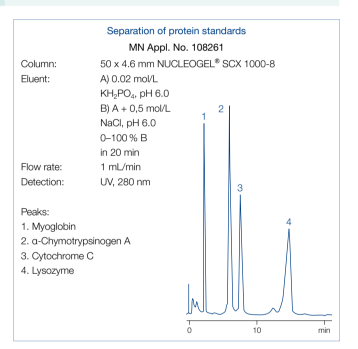
Technical data

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

Recommended application

· Proteins, peptides and carbohydrates with high isoelectric point





Eluent in column 0.1 mol/L Na ₂ SO ₄ + 0.2 % NaN ₃		
ID	Length →	
	50 mm	Guard columns*
NUCLEOGEL® SAX pore size 1000 Å		
Analytical Valco type columns		
4.6 mm	719469	719600

NUCLEOGEL® S	CX pore size 1000 Å		
Analytical Valco type	columns		
	4.6 mm	719475	719540

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



$NUCLEODUR^{\circledR}~300~C_{18}~ec~\cdot~C_{4}~ec~~\text{wide pore silica for biochromatography}~\cdot~\text{USP L1}~(C_{18})~\cdot~\text{USP L26}~(C_{4})~$

Key feature

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

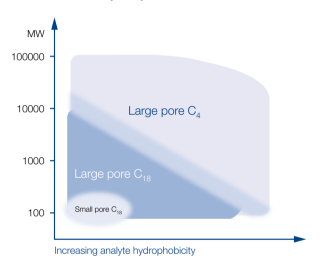
Technical data

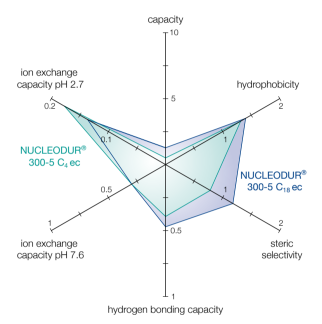
• Pore size 300 Å; particle size 5 μ m, carbon content 4 % for C₁₈, 2.5 % for C₄; pH stability 1–9; high reproducibility from lot to lot

Recommended application

Biological macromolecules like proteins or peptides

Column selection by analyte characteristics



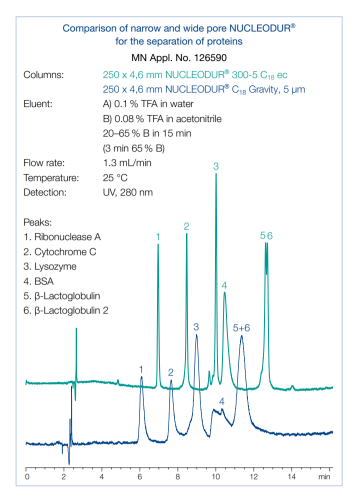


Tanaka plots of NUCLEODUR® wide pore phases

Batch-to-batch reproducibility of NUCLEODUR® 300-5 C₄ ec and NUCLEODUR® 300-5 C₁₈ ec MN Appl. Nos. 126551 / 126552 NUCLEODUR® 300-5 C₁₈ ec Columns: 250 x 4 mm NUCLEODUR® 300-5 C₄ ec Eluent: A) 0.1 % TFA in water B) 0.08 % TFA in acetonitrilel 20-60 % B in 15 min 56 Flow rate: 1 mL/min Temperature: 25 °C Detection: UV, 280 nm Peaks: 1. Ribonuclease A 2. Cytochrome C 3. Lysozyme 4. BSA 5. β-Lactoglobulin 6. β-Lactoglobulin 2 10 10 12 12 14







Tryptic digest of cytochrome C MN Appl. No. 126600 250 x 4.6 mm NUCLEODUR® 300-5 C₁₈ ec Columns: 250 x 4.6 mm Jupiter® C₁₈, 5 μm A) 0.1 % TFA in water Eluent: B) 0.08 % TFA in acetonitrile 5-40 % B in 15 min (1 min 40 % B) Flow rate: 1.3 mL/min 30 °C Temperature: Detection: UV, 280 nm

Sharper peaks of larger molecules on wide pore material

Less tailing and better separation on NUCLEODUR® 300 C₁₈ ec

Ordering informa	ation					
Eluent in column ac	cetonitrile – wat	er				
	ID	Length →				
		100 mm	125 mm	150 mm	250 mm	EC guard columns*
NUCLEODUR [®] 300-5 C ₁₈ eC octadecyl phase, particle size 5 μm, pore size 300 Å, endcapped, 4 % C						
Analytical EC column	ns					
	2 mm	760183.20	760184.20	760185.20	760186.20	761988.20
	3 mm	760183.30	760184.30	760185.30	760186.30	761988.30
	4 mm	760183.40	760184.40	760185.40	760186.40	761988.30
	4.6 mm	760183.46	760184.46	760185.46	760186.46	761988.30
NUCLEODUR [®] 300-5 C ₄ ec butyl phase, particle size 5 μm, pore size 300 Å, endcapped, 2.5 % C						
Analytical EC columns						
	2 mm	760193.20	760194.20	760195.20	760196.20	761989.20
	3 mm	760193.30	760194.30	760195.30	760196.30	761989.30
	4 mm	760193.40	760194.40	760195.40	760196.40	761989.30
	4.6 mm	760193.46	760194.46	760195.46	760196.46	761989.30

^{*} EC guard columns require the Column Protection System guard column holder (REF 718966, see page 251). EC columns in packs of 1, guard columns in packs of 3.



NUCLEOSIL® MPN RP chromatography of biological macromolecules

NUCLEOSIL® 100-5 C₁₈ MPN · USP L1

Key feature

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

Technical data

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

NUCLEOSIL® 300-5 C4 MPN · USP L26

Key feature

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

Technical data

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

Separation of haemoglobin chains

MN Appl. No. 108240

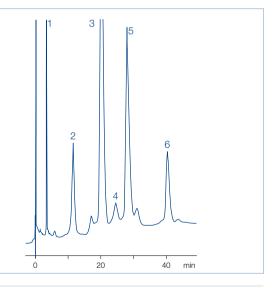
B) 60 % acetonitrile, 40 % water, 0.1 % TFA

 $40\text{--}60\,\%$ B in 60 min

Flow rate: 1 mL/min
Detection: UV, 220 nm

Peaks: 1. Hem 2. β -globin 3. α -globin 4. $^{A}\gamma^{T}$ -globin 5. $^{G}\gamma$ -globin

6. $^{A}\gamma^{I}$ -globin



Ordering information

Eluent in column methanol

250 mm EC guard columns*

Length →

NUCLEOSIL® 100-5 C₁₈ MPN

Analytical EC columns

4 mm 720231.40

NUCLEOSIL® 300-5 C₄ MPN

Analytical EC columns

4 mm 720245.40 721119.30

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



NUCLEOSIL® PPN RP chromatography of biological macromolecules

NUCLEOSIL® 100-5 C₁₈ PPN · USP L1

Kev feature

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

Technical data

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

NUCLEOSIL® 500-5 C₁₈ PPN · USP L1

Key feature

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

Technical data

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

Separation of a protein standard MN Appl. No. 108220 125 x 4 mm NUCLEOSIL® 100-5 C₁₈ PPN

Column:

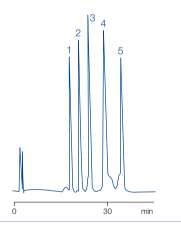
A) 0.1 % TFA in H₂O Eluent: B) 0.08 % TFA in CH₃CN

20-60 % B in 10 min

Flow rate: 1.0 mL/min Detection: UV, 280 nm

Peaks:

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



Separation of pancreatic secretion of piglets MN Appl. No. 108280 125 x 4 mm NUCLEOSIL® 500-5 C₁₈ PPN Column: A) 0.1 % TFA in H₂O Eluent: B) 0.08 % TFA in CH₃CN 30-50 % B in 14 min, then 50-65 % B in 6 min Flow rate: 1 mL/min Detection: UV, 215 nm Peaks: 1. Trypsin + trypsinogen 2. Proelastase 3. Lipase + α-Chymotrypsin 4. Chymotrypsinogen 5. α-Amylase 6., 7. Procarboxypeptidase

Ordering information

Eluent in column methanol

Length → 250 mm

EC guard columns*

20 min

$NUCLEOSIL^{\$}$ 100-5 C_{18} PPN particle size 5 μm , pore size 100 Å

Analytical EC columns



720252.40

721567.30

$NUCLEOSIL^{\$}$ 500-5 C_{18} PPN particle size 5 μm , pore size 500 Å

Analytical EC columns

720258.40 721924.30

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

11/1

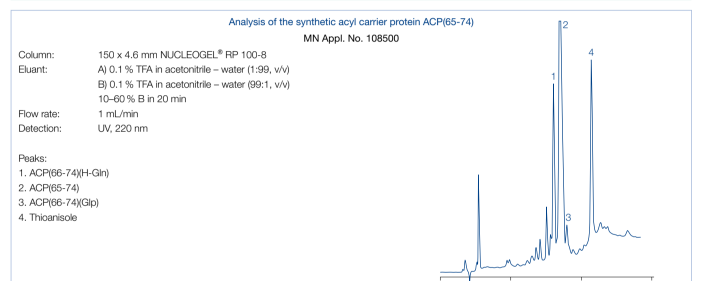
HPLC columns for biochemical separations



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

Technical data

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



Ordering information				
Eluent in column acetonitrile -	water			
ID	Length → 50 mm	→ 150 mm	250 mm	Guard columns*
		150 11111	250 111111	Guard Columns
NUCLEOGEL® RP 100-5	particle size 5 μ m, pore size 100 Å			
Analytical Valco type columns				
4.6 mm		719454	719455	719542
NUCLEOGEL® RP 100-8	particle size 8 µm, pore size 100 Å			
Analytical Valco type columns				
4.6 mm		719456	719520	719542
NUCLEOGEL® RP 300-5	particle size 5 µm, pore size 300 Å			
Analytical Valco type columns				
4.6 mm	719459			719542
NUCLEOGEL® RP 300-8	particle size 8 µm, pore size 300 Å			
Analytical Valco type columns				
4.6 mm	719460			719542

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

HPLC columns for sugar analyses

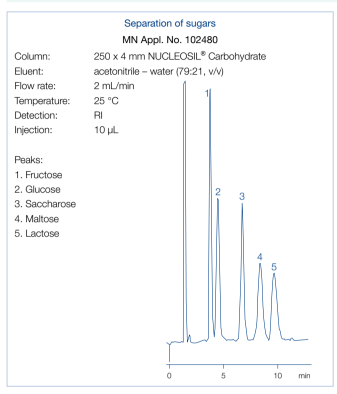
NUCLEOSIL® Carbohydrate separation of mono- and disaccharides · USP L8

Technical data

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

Recommended application

· RP separation of mono- and disaccharides



Ordering information		
Eluent in column acetonitrile – water (79:21, v/v)		
ID	Length →	
	250 mm	EC guard columns*
NUCLEOSIL® Carbohydrate		
Analytical EC columns		
4 mm	720905.40	721170.30

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

HPLC columns for sugar analyses



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

Technical data

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

Recommended application

· H⁺ form:

Separation of sugars, sugar alcohols and organic acids; eluent in column 5 mmol/L H_2SO_4

· Ca²⁺ form:

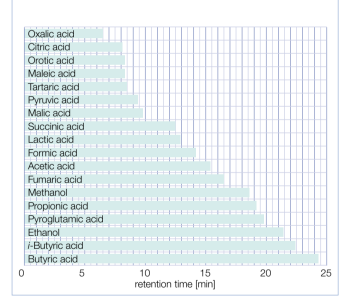
Separation of mono-, di- and oligosaccharides; eluent in column water

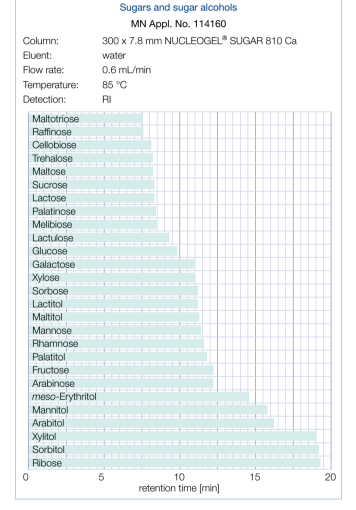
Organic acids and alcohols

MN Appl. No. 113870

Column: 300 x 7.8 mm NUCLEOGEL® SUGAR 810 H

 $\begin{tabular}{lll} Eluent: & 5 mmol/L H_2SO_4$ \\ Flow rate: & 0.6 mL/min \\ Temperature: & 35 °C \\ Detection: & RI \\ Injection: & 5 μL \\ \end{tabular}$





Ordering information		
ID	Length →	
	300 mm	Guard columns*
NUCLEOGEL® SUGAR 810 H eluent in column 5 mmol/L H ₂ SO ₄		
Analytical Valco type columns		
7.8 mm	719574	719575
NUCLEOGEL® SUGAR 810 Ca eluent in column water		
Analytical Valco type columns		
7.8 mm	719570	719571

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



HPLC columns for sugar analyses



NUCLEOGEL® ION 300 OA / SUGAR

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

Technical data

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

Recommended application

NUCLEOGEL® ION 300 OA:

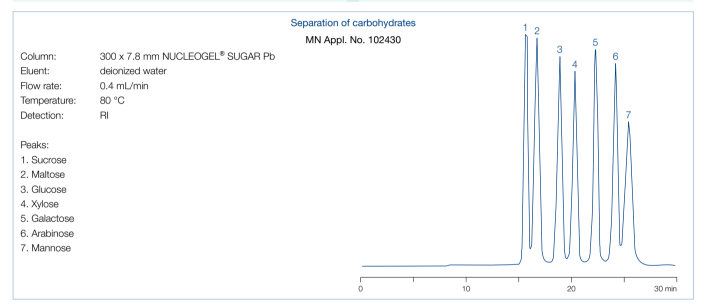
 $\ensuremath{H^{\scriptscriptstyle{+}}}$ form for separation of sugars, alcohols and organic acids

NUCLEOGEL® SUGAR:

Ca²⁺ form: separation of mono- and oligosaccharides, sugar alcohols

Pb²⁺ form: separation of mono- and disaccharides from food and biological samples

Na⁺ form: separation of oligosaccharides from starch hydrolysates and food



Ordering information				
ID	Length → 300 mm	Guard columns*		
${\color{red} NUCLEOGEL}^{\tiny{\$}} \ {\color{blue} ION \ 300 \ OA} \ \ {\color{blue} eluent \ in \ column \ 5 \ mmol/L \ H_2SO_4 \ 5 \ mmol/L \ H_2SO_5 \ 6 \ mmol/L \ H_2SO_6 \ $				
Analytical Valco type columns				
7.8 mm	719501	719537		
NUCLEOGEL® SUGAR Ca eluent in column water + 0.02 % azide				
Analytical Valco type columns				
6.5 mm	719531	719535		
NUCLEOGEL® SUGAR Pb eluent in column water + 0.02 % azide				
Analytical Valco type columns				
7.8 mm	719530	719534		
NUCLEOGEL® SUGAR Na eluent in column water + 0.02 % azide				
Analytical Valco type columns				
7.8 mm	719532	719536		
* Valco Type guard columns measure 21 x 4 mm and require the guard column holder C. REF 719538, see page 250.				

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



Columns for gel permeation chromatography

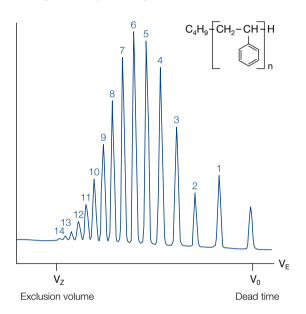


NUCLEOGEL® GPC for GPC of water-insoluble substances

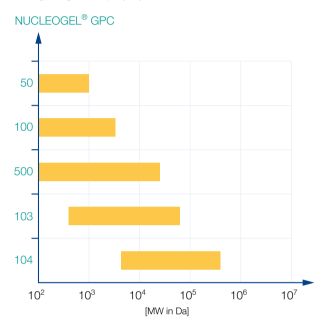
Technical data

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

Chromatogram of styrene oligomers



Working ranges for polystyrene



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

Columns and guard columns in packs of 1.



EC standard columns for analytical HPLC / UHPLC



- Analytical column system manufactured from stainless steel M8 outer threads on both ends combination of sealing element and very fine-meshed stainless steel screen, PTFE ring and fitting adaptor column heads SW 12, with inner threads M8 x 0.75 and UNF 10-32 (= 1/16" connection)
- EC column hardware guarantees pressure stability of 1200 bar - hereby EC columns are suitable for UHPLC applications (ultra fast HPLC) and all modern HPLC systems.
- As screw-on guard column system we recommend the Column Protection System used with EC guard column cartridges with 4 mm length.
- EC guard columns supplied with NUCLEODUR®, NUCLEOSIL® spherical silicas and NUCLEOSHELL® spherical core shell silica particles

Available standard dimensions of EC columns

ID	Length →										
	20 mm	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	200 mm	250 mm	300 mm	
2 mm	+	+	+	+	+	+	+	+	+	+	
3 mm	+	+	+	+	+	+	+	+	+	+	
4 mm	+	+	+	+	+	+	+	+	+	+	
4.6 mm	+	+	+	+	+	+	+	+	+	+	
Please ask	Please ask for availability of certain phases.										

Note: NUCLEODUR® and NUCLEOSHELL® column head must not be removed!

Guard columns for EC columns								
EC column with ID	EC guard column*							
2 mm	4/2							
3 mm	4/3							
3 mm	4/3							
3 mm	4/3							
Packs of 3 cartridges								
* Information about the Colum	nn Protection System on page 251.							

For preparative applications MN offers the so-called VarioPrep® hardware system, which is described from page 252 on.

Valco type columns



- Analytical column system manufactured from stainless steel
- Available inner diameters:
 4.6 mm ID (1/4" OD) and 7.7 mm (3/8" OD)
- Mainly used for NUCLEOGEN[®] and NUCLEOGEL[®] (see page 226)

Ordering information

Description	Pack of	REF
Accessories for Valco type columns		
Guard column holder B for VA columns 5 x 3 mm	1	719539
Guard column holder C for VA guard columns 21 x 4 mm	1	719538

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MN column systems



Column Protection System

Innovative and universal guard column holder system



- Suitable for all analytical HPLC columns with 1/16" fittings
- Cartridges filled with special NUCLEODUR[®], NUCLEOSIL[®] and NUCLEOSHELL[®] HPLC adsorbents
- · Ideal protection for your analytical main column
- → significant increase in column lifetime
- Minimized dead volume → suitable also for ultra-fast HPLC
- Special ferrules → pressure stability up to 1300 bar (18 850 psi)

- Visual contamination check
 → in-time changing of the guard
 column
- Suitable guard columns with 4 mm length, 2 mm ID (for main columns with 2 mm ID); 3 mm ID (for main columns with 3, 4 and 4.6 mm), respectively
- UNIVERSAL RP guard columns suitable for all HPLC columns under RP conditions

Content of the Column Protection System



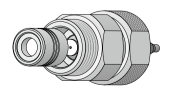
Description	Pack of	REF
Guard column holder	1	
Capillaries (0.12 mm ID)	2	
Ferrules	3	718966
Wrenches	2	
Manual	1	

Ordering information		
Description	Pack of	REF
Replacement parts for the Column Protection System		
Special ferrules made of PEEK	5	718967
Replacement connector including O-ring	1	718968
Stainless steel capillaries 0.12 mm ID, nuts and metal ferrules	3	718969
Stainless steel capillaries 0.18 mm ID (for higher flow rates), nuts and metal ferrules	3	718971
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 4/2 UNIVERSAL RP guard column (for main columns with 2 mm ID), value pack	9	728778.20
EC 4/3 UNIVERSAL RP guard column (for main columns with 3, 4 and 4.6 mm ID)	3	728777.30
EC 4/3 UNIVERSAL BP guard column (for main columns with 3, 4 and 4.6 mm ID), value pack	9	728778.30

Visual contamination check

The cartridge is fitted with a special filter membrane:

- If this silver membrane is contaminated (bright or dark discoloration), it is advisable to replace the cartridge.
- If the contaminants are colorless, replace the cartridge if the pressure rises or the chromatographic performance decreases.



VarioPrep (VP) columns for preparative HPLC



- Column system for preparative HPLC, manufactured from stainless steel with two adjustable end fittings, suitable for frequent use of back-flushing techniques
- Allows compensation of a dead volume, which could occur at the column inlet after some time of operation, without need for opening the column
- Can be packed with all NUCLEODUR[®] and NUCLEOSIL[®] spherical silicas

Available standard dimensions of VarioPrep columns with axially adjustable end fittings

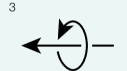
	ID	Length →		Length →						
End fitting design		10* mm	15* mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm	500 mm
	8	+		+		+	+	+	+	
	10			+		+	+	+	+	
	16	+		+		+	+	+	+	
	21			+	+	+	+	+	+	
	32		+			+		+	+	
	40			+		+	+	+	+	+
	50		+			+		+	+	
	80		***************************************						+	+

^{* 10} x 8, 10 x 16, 15 x 32 and 15 x 50 mm ID columns are used as guard columns and require the respective holders, see page 253.

The VarioPrep principle







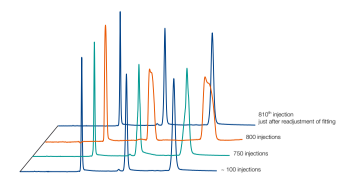


Readjustment of fitting

VarioPrep columns are produced with highest packing quality and bed density (1). Due to intensive chemical and/ or mechanical exposure of the column adsorbent, shrinking of the column bed can occur (2; orange gap). in this even unlikely case readjustment of the VarioPrep

column fitting (3; turning the nut at the column inlet clockwise) will eliminate the emerged dead volume (4). 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 column performance and prolongs column lifetime noticeably.

MN column systems

ange



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



- (1) VP 15/32 for 32 and 40 mm ID columns
- ③ VP 10/8 for 8 and 10 mm ID columns
- ② VP 10/16 for 16 and 21 mm ID columns
- ④ VP 15/50 for ≥ 50 mm ID columns

· Robust hardware

· Free rotary plunger fittings - low O-ring abrasion

· Easy handling and cartridge exch-

- · Cost-efficient cartridges
- · Minimally invasive / no disturbance of the separation efficiency of main column
- · Low back pressure
- · Designed for pressures up to 400 bar

Column performance without and with guard column

125 x 16 mm NUCLEODUR® C₁₈ HTec, 5 µm Columns:

 $125 \times 16 \text{ mm NUCLEODUR}^{\otimes} \text{ C}_{18} \text{ HTec}, 5 \mu\text{m} + 10 \times 16 \text{ mm NUCLEODUR}^{\otimes} \text{ C}_{18} \text{ HTec guard column}$

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

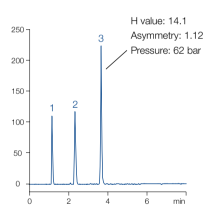
Flow rate: 16 mL/min Temperature: 22 °C

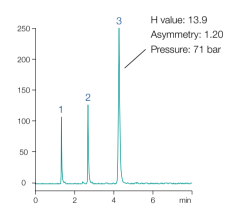
Peaks:

1. Phenol

2. Naphthalene

3. Anthracene





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

Technical data

· free rotary plunger fittings – low O-ring abrasion

· 1/16 triread	· free rotary pluriger	illings – low	O-ring abrasion · stainless ste	30 1		
Guard cartridge	Holder REF	Holder ID	Recommended for column ID	Preferred capillary ID	Typical flow rate	
VP 10/8	718251	8 mm	8 and 10 mm ID	0.17 and 0.25 mm	1–12 mL/min	
VP 10/16	718256	16 mm	16 and 21 mm ID	0.17, 0.25 and 0.5 mm	2–32 mL/min	
VP 15/32	718253	32 mm	32 and 40 mm ID	0.25, 0.5 and 1.0 mm	5–150 mL/min	
VP 15/50	718255	50 mm	≥ 50 mm ID	0.5 and 1.0 mm	20-250 mL/min	

Ordering information

Guard column holders for VarioPrep columns

Guarc	data obtainin holders for varior rep obtaining										
	VP Guard col	umns for VarioPre	p columns with ID) →	Pack of	Replacement O-ring	Holder				
	8, 10 mm	16, 21 mm	32, 40 mm	≥ 50 mm	guard columns	(pack of 2)	ID	REF			
VP	10/8				2	718975	8 mm	718251			
VP		10/16			2	718976	16 mm	718256			
VP			15/32		1	718977	32 mm	718253			
VP	•		•	15/50	1	718978	50 mm	718255			

For REF numbers of individual VP guard column cartridges see respective NUCLEODUR® and NUCLEOSIL® phases.





Accessories for stainless steel HPLC columns



- · Stainless steel columns are most frequently used in HPLC.
- · The material is corrosion resistant, pressure stable and easy to work mechanically.

Ordering information		
Description	Pack of	REF
Capillary accessories		
1/16" column end caps (plastic)	4	718582
1/16" nut for connecting 1/16" capillaries	5	718583
1/16" ferrule	5	718584
Capillary unions		
Typ 1: 100 mm x 1/16" x 0.25 mm	1	718637
Typ 2: 100 mm x 1/16" x 0.12 mm	1	719489
Cutter for 1/16" capillary tubing	1	706290

For accessories and replacement parts for EC columns see page 251, for accessories and replacement parts for VarioPrep columns see page 253.



SPE accessories for sample preparation, like e.g., CHROMABOND® vacuum manifolds can be found on page 65.

PEEK accessories

· PEEK (= polyether ether ketone) is a high performance polymer belonging to the group of polyarylether ketones (PAEK), which meets all requirements of HPLC columns with respect to chemical resistance and mechanical stability. In some fields of application in HPLC like, e.g., in ion chromatography and chromatography of biopolymers, PEEK fulfils the requirements for a nonmetallic material.

· All fittings can be tightened by hand.

Ordering in	formation				
Description			Pack of	REF	
PEEK fitting	gs				
	ngertight fitting,		1	718770	718770
	nation nut + ferrule			710771	718771
1/16" PEEK fir	errule for REF 718771		1	718771 718772	
1/16" PEEK d	·····		1	718775	
			<u>:</u>		718775
					718772
					1012
1/10" DEEL					
	nion, both sides inner thr double ferrules	eads, equipped with 2 finger-	1	718766	
1/16" PEEK u and without fe		eads, however without nuts	1	718767	
1/16" PEEK union, both sides outer threads		1	718768		
AD	ID [mm]	Length	Pack of	REF	
	dard capillaries				
1/16"	0.13	1 m	1	718765	
1/16"	0.17	1 m	1	718760	
1/16"	0.25 0.5	1 m	1	718761 718762	
1/16"	0.75	1 m	! 1	718763	
Description	0.70	1 111	Pack of	REF	
	EEK capillaries				
	2.04				
Guillotine cutte	er for PEEK and PTFE ca	pillaries	1	718769	
		J	•		
					/
Clean-Cut cut	ter for different capillary o	outer diameters	1	718755	

NUCLEODUR® high purity silica for HPLC



Basics of preparative HPLC

In principal for preparative HPLC the same rules apply than for analytic HPLC. However both differ significantly in their aim. The aim of analytic HPLC is a preferably complete separation of the single components of a mixture with subsequent peak identification. In contrast the goal of preparative HPLC is isolation of the desired product in defined purity, maximum amount while having a cost effective method of operating.

Demand of a preparative separation

- Throughput
- Purity
- Yield

Upscaling table for current MN column dimensions

	•	•	0	0	0	0	0	0	
ID x Length [mm]	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	27.6	64	100	156.3	400
Typical amount of sample* [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

^{*} based on RP material; the herein stated maximum amounts of sample are dependent on the separation problem and the sample. In some cases half the maximum amount of sample can already lead to a drastic overload of the column, in other cases the maximum amount of sample still leads to an acceptable separation.

NUCLEODUR® bulk packings

· Fully spherical high purity silica

- · Bigger particles for preparative application
- Pore size 110 Å; pore volume 0.9 mL/g; surface (BET) 340 m²/g; density 0.47 g/mL; pressure stable up to 600 bar

Ordering information						
Phase	Endcapped	Carbon content	Particle size	Pack of 100 g	Pack of 1000 g	
NUCLEODUR® C ₁₈ HTec premiu	um octadecyl phase	e (see page 178)				
NUCLEODUR® C ₁₈ HTec, 7 µm	yes	18 % C	7 μm	713831.0100	713831.1	
NUCLEODUR [®] C ₁₈ HTec, 10 μm	yes	18 % C	10 μm	713832.0100	713832.1	
NUCLEODUR® C ₁₈ ec standard	octadecyl phase (s	ee page 181)				
NUCLEODUR® 100-10 C ₁₈ ec	yes	17.5 % C	10 μm	713611.0100	713611.1	
NUCLEODUR® 100-12 C ₁₈ ec	yes	17.5 % C	12 µm	713618.0100	713618.1	
NUCLEODUR® 100-16 C ₁₈ ec	yes	17.5 % C	16 µm	713621.0100	713621.1	
NUCLEODUR® 100-20 C ₁₈ ec	yes	17.5 % C	20 μm	713601.0100	713601.1	
NUCLEODUR® 100-30 C ₁₈ ec	yes	17.5 % C	30 µm	713631.0100	713631.1	
NUCLEODUR® 100-50 C ₁₈ ec	yes	17.5 % C	50 μm	713550.0100	713550.1	
Unmodifiziertes NUCLEODUR®	SiOH silica (see pa	ge 190)				
NUCLEODUR® 100-10			10 μm	713610.0100	713610.1	
NUCLEODUR® 100-12			12 µm	713615.0100	713615.1	
NUCLEODUR® 100-16			16 µm	713620.0100	713620.1	
NUCLEODUR® 100-20			20 μm	713600.0100	713600.1	
NUCLEODUR® 100-30			30 µm	713630.0100	713630.1	
NUCLEODUR® 100-50			50 μm	713551.0100	713551.1	



POLYGOSIL® irregular silica for HPLC



POLYGOSIL® bulk packings

- · Irregular silica for analytical applications
- · pH stability 2–8

Physical properties of unmo	dified POLYGOSIL® materials
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, , ,					
Phase	Pore size	Pore volume	Surface (BET)	Density	Pressure stability
POLYGOSIL® 60	60 Å	0.75 mL/g	350 m ² /g	0.45 g/mL	600 bar
POLYGOSIL® 100	100 Å	1 mL/g	280 m²/g	0.35 g/mL	400 bar
POLYGOSIL® 300	300 Å	0.8 mL/g	100 m²/g	0.45 g/mL	400 bar
POLYGOSIL® 1000	1000 Å	0.8 mL/g	25 m²/g	0.45 g/mL	300 bar

Modification of POLYGOSIL® follows the same processes as for NUCLEOSIL® silica.

Ordering information			_			
Phase	Endcapped	Carbon content	Pore size	Particle size	Pack of 10 g	Pack of 100 g
Octadecyl phases -(CH ₂) ₁₇	-CH ₃					
POLYGOSIL® 60-5 C ₁₈	yes	12 % C	60 Å	5 μm	711330.10	711330.100
POLYGOSIL® 60-7 C ₁₈	yes	12 % C	60 Å	7 μm	711340.10	711340.100
POLYGOSIL® 60-10 C ₁₈	yes	12 % C	60 Å	10 μm	711350.10	711350.100
POLYGOSIL® 100-5 C ₁₈	yes	14 % C	100 Å	5 μm	711560.10	711560.100
POLYGOSIL® 100-7 C ₁₈	yes	14 % C	100 Å	7 μm	711570.10	711570.100
POLYGOSIL® 100-10 C ₁₈	yes	14 % C	100 Å	10 μm	711580.10	711580.100
POLYGOSIL® 300-7 C ₁₈	yes	4 % C	300 Å	7 μm	711710.10	711710.100
POLYGOSIL® 1000-7 C ₁₈	yes	~ 1 % C	1000 Å	7 μm	711992.10	711992.100
Octyl phases -(CH ₂) ₇ -CH ₃						
POLYGOSIL® 60-5 C ₈	no	7 % C	60 Å	5 µm	711300.10	711300.100
POLYGOSIL® 60-7 C ₈	no	7 % C	60 Å	7 μm	711310.10	711310.100
POLYGOSIL® 60-10 C ₈	no	7 % C	60 Å	10 μm	711320.10	711320.100
Butyl phases -(CH ₂) ₃ -CH ₃						
POLYGOSIL® 300-7 C ₄	yes	~ 1 % C	300 Å	7 µm	711680.10	711680.100
POLYGOSIL® 1000-7 C ₄	yes	< 1 % C	1000 Å	7 μm	711991.10	711991.100
Cyano phases (nitrile) -(CI	H ₂) ₃ – CN					
POLYGOSIL® 60-5 CN		~ 5 % C	60 Å	5 µm	711380.10	711380.100
POLYGOSIL® 60-10 CN	······································	~ 5 % C	60 Å	10 µm	711390.10	711390.100
Amino phases -(CH ₂) ₃ -NH	2					
POLYGOSIL® 60-5 NH ₂		~ 3 % C	60 Å	5 µm	711360.10	711360.100
POLYGOSIL® 60-10 NH ₂	······································	~ 3 % C	60 Å	 10 μm	711370.10	711370.100
Dimethylamino phases -(0	CH ₂) ₃ – N(CH ₃) ₂			•		
POLYGOSIL® 60-5 N(CH ₃) ₂	2.0 (0.2	~ 3.5 % C	60 Å	5 μm	711420.10	711420.100
POLYGOSIL® 60-10 N(CH ₃) ₂	•	~ 3.5 % C	60 Å	10 µm	711430.10	711430.100
Jnmodified silica SiOH						
POLYGOSIL® 60-5			60 Å	5 µm	711010.10	711010.100
POLYGOSIL® 60-7	······································		60 Å	7 µm	711280.10	711280.100
POLYGOSIL® 60-10			60 Å	10 µm	711020.10	711020.100
POLYGOSIL® 100-5			100 Å	5 μm	711510.10	711510.100
POLYGOSIL® 100-7			100 Å	7 μm	711520.10	711520.100
POLYGOSIL® 100-10			100 Å	10 μm	711530.10	711530.100
POLYGOSIL® 300-7			300 Å	7 μm	711600.10	711600.100
POLYGOSIL® 1000-7			1000 Å	7 μm	711890.10	711890.100

POLYGOPREP irregular silica for HPLC

POLYGOPREP bulk packings

- · Irregular silica for preparative applications
- · pH stability 2–8

	,						
Phase	Pore size	Pore volume	Surface (BET)	Density	Pressure stability		
POLYGOPREP 60	60 Å	0.75 mL/g	350 m²/g	0.45 g/mL	600 bar		
POLYGOPREP 100	100 Å	1 mL/g	280 m²/g	0.35 g/mL	400 bar		
POLYGOPREP 300	300 Å	0.8 mL/g	100 m²/g	0.45 g/mL	400 bar		
POLYGOPREP 1000	1000 Å	0.8 mL/g	35 m²/g	0.45 g/mL	300 bar		
Modification of POLYGOPREP follows the same processes as for NUCLEOSIL® silica.							

Ordering information						
Phase	Endcapped	Carbon content	Pore size	Particle size	Pack of 100 g	Pack of 1 kg
Octadecyl phases -(CH ₂) ₁₇	-CH ₃					
POLYGOPREP 60-12 C ₁₈	no*	12 % C	60 Å	10–15 μm	711009.100	711009.1000
POLYGOPREP 60-20 C ₁₈	no*	12 % C	60 Å	15–25 µm	711031.100	711031.1000
POLYGOPREP 60-30 C ₁₈	no*	12 % C	60 Å	25–40 µm	711480.100	711480.1000
POLYGOPREP 60-50 C ₁₈	no*	12 % C	60 Å	40–63 µm	711500.100	711500.1000
POLYGOPREP 60-80 C ₁₈	no*	12 % C	60 Å	63–100 µm	711011.100	711011.1000
POLYGOPREP 60-130 C ₁₈	no*	12 % C	60 Å	63–200 µm	711590.100	711590.1000
POLYGOPREP 100-12 C ₁₈	no*	14 % C	100 Å	10–15 µm	711018.100	711018.1000
POLYGOPREP 100-20 C ₁₈	no*	14 % C	100 Å	15–25 µm	711019.100	711019.1000
POLYGOPREP 100-30 C ₁₈	no*	14 % C	100 Å	25–40 µm	711032.100	711032.1000
POLYGOPREP 100-50 C ₁₈	no*	14 % C	100 Å	40–63 µm	711021.100	711021.1000
POLYGOPREP 300-12 C ₁₈	yes	4 % C	300 Å	10–15 µm	711024.100	711024.1000
POLYGOPREP 300-20 C ₁₈	yes	4 % C	300 Å	15–25 µm	711025.100	711025.1000
POLYGOPREP 300-30 C ₁₈	yes	4 % C	300 Å	25–40 µm	711720.100	711720.1000
POLYGOPREP 300-50 C ₁₈	yes	4 % C	300 Å	40–63 µm	711730.100	711730.1000
POLYGOPREP 1000-30 C ₁₈	yes	~ 1 % C	1000 Å	25–40 µm	711028.100	711028.1000
POLYGOPREP 1000-50 C ₁₈	yes	~ 1 % C	1000 Å	40–63 µm	711029.100	711029.1000
Octyl phases -(CH ₂) ₇ -CH ₃						
POLYGOPREP 60-12 C ₈	no*	7 % C	60 Å	10–15 µm	711007.100	711007.1000
POLYGOPREP 60-20 C ₈	no*	7 % C	60 Å	15–25 μm	711008.100	711008.1000
POLYGOPREP 60-30 C ₈	no*	7 % C	60 Å	25–40 µm	711470.100	711470.1000
POLYGOPREP 60-50 C ₈	no*	7 % C	60 Å	40–63 µm	711490.100	711490.1000
On request, these POLYGOPREF	P RP phases can be e	ndcapped at surcharg	e.			
Butyl phases -(CH ₂) ₃ -CH ₃						
POLYGOPREP 300-12 C ₄	yes	~ 1 % C	300 Å	10–15 μm	711022.100	711022.1000
POLYGOPREP 300-20 C ₄	yes	~ 1 % C	300 Å	15–25 μm	711023.100	711023.1000
POLYGOPREP 300-30 C ₄	yes	~ 1 % C	300 Å	25–40 µm	711690.100	711690.1000
POLYGOPREP 300-50 C ₄	yes	~ 1 % C	300 Å	40–63 μm	711700.100	711700.1000
POLYGOPREP 1000-30 C ₄	yes	< 1 % C	1000 Å	25–40 μm	711026.100	711026.1000
POLYGOPREP 1000-50 C ₄	yes	< 1 % C	1000 Å	40–63 µm	711027.100	711027.1000
Cyano phases (nitrile) -(CF	H ₂) ₃ – CN					
POLYGOPREP 60-12 CN		~ 4.5 % C	60 Å	10–15 µm	711015.100	711015.1000
POLYGOPREP 60-20 CN		~ 4.5 % C	60 Å	15–25 µm	711016.100	711016.1000
POLYGOPREP 60-30 CN		~ 4.5 % C	60 Å	25–40 μm	711017.100	711017.1000
Amino phases -(CH ₂) ₃ -NH ₂						
POLYGOPREP 60-12 NH ₂		~ 3 % C	60 Å	10–15 µm	711012.100	711012.1000
POLYGOPREP 60-20 NH ₂	••••••	~ 3 % C	60 Å	15–25 µm	711013.100	711013.1000
POLYGOPREP 60-30 NH ₂		~ 3 % C	60 Å	25–40 µm	711014.100	711014.1000



POLYGOPREP irregular silica for HPLC



Ordering information	า				
Phase	Pore size	Particle size	Pack of 100 g	Pack of 1 kg	Pack of 5 kg
Unmodified POLYG	OPREP sil	ica SiOH			
POLYGOPREP 60-12	60 Å	10–15 μm		711001.1000	711001.5000
POLYGOPREP 60-20	60 Å	15–25 μm	•	711240.1000	711240.5000
POLYGOPREP 60-30	60 Å	25–40 μm		711250.1000	711250.5000
POLYGOPREP 60-50	60 Å	40–63 μm		711260.1000	711260.5000
POLYGOPREP 60-80	60 Å	63–100 µm		711270.1000	711270.5000
POLYGOPREP 60-130	60 Å	63–200 µm		711037.1000	711037.5000
POLYGOPREP 100-12	100 Å	10–15 μm		711002.1000	711002.5000
POLYGOPREP 100-20	100 Å	15–25 μm		711003.1000	711003.5000
POLYGOPREP 100-30	100 Å	25–40 μm		711540.1000	711540.5000
POLYGOPREP 100-50	100 Å	40–63 μm		711550.1000	711550.5000
POLYGOPREP 100-80	100 Å	63–100 µm		711033.1000	711033.5000
POLYGOPREP 100-130	100 Å	63–200 µm		711034.1000	711034.5000
POLYGOPREP 300-12	300 Å	10–15 μm	711004.100	711004.1000	
POLYGOPREP 300-20	300 Å	15–25 μm	711610.100	711610.1000	
POLYGOPREP 300-30	300 Å	25–40 μm	711620.100	711620.1000	
POLYGOPREP 300-50	300 Å	40–63 μm	711630.100	711630.1000	
POLYGOPREP 1000-12	1000 Å	10–15 μm	711035.100	711035.1000	
POLYGOPREP 1000-20	1000 Å	15–25 μm	711036.100	711036.1000	
POLYGOPREP 1000-30	1000 Å	25–40 μm	711005.100	711005.1000	
POLYGOPREP 1000-50	1000 Å	40–63 μm	711006.100	711006.1000	

Adsorbents for column chromatography



Silica adsorbents for low pressure column chromatography



- · Silica 60; pore size ~ 60 Å; pore volume ~ 0.75 mL/g; spec. surface BET ~ 500 m²/g highly porous, amorphous silicic acid in the form of hard, opalescent particles, prepared by precipitation of water glass with sulfuric acid
- · For higher demands on the performance of column packings we recommend our high-purity irregular POLYGOPREP silicas (see before).
- · Silica FIA for the fluorescence indicator adsorption procedure for the determination of hydrocarbon groups in the testing of liquid fuels in accordance with DIN 51791 and ASTM D 1319-58T
- · The FIA method determines saturated hydrocarbons, olefins and aromatic hydrocarbons of a sample chromatographically by adsorption and desorption in a column filled with FIA silica, in the presence of a fluorescent dye mixture.

Ordering information				
Description	Particle size	1 kg	5 kg	25 kg
Silica 60, 0.015-0.04 mm	_	815650.1	815650.5	815650.25
Silica 60, 0.025-0.04 mm	_	815300.1	815300.5	815300.25
Silica 60, 0.04-0.063 mm	230–400 mesh	815380.1	815380.5	815380.25
Silica 60 M, 0.04-0.063 mm	230–400 mesh	815381.1	815381.5	815381.25
Silica 60, 0.05-0.1 mm	130–270 mesh	815390.1	815390.5	815390.25
Silica 60, 0.05-0.2 mm	70–270 mesh	815320.1	815320.5	815320.25
Silica 60, 0.063-0.2 mm	70–230 mesh	815330.1	815330.5	815330.25
Silica 60, < 0.063 mm	+230 mesh	815400.1	815400.5	815400.25
Silica 60, < 0.08 mm	+190 mesh	815310.1	815310.5	815310.25
Silica 60, 0.1-0.2 mm	70–130 mesh	815340.1	815340.5	815340.25
Silica 60, 0.2–0.5 mm	35–70 mesh	815350.1	815350.5	815350.25
Silica 60, 0.5–1.0 mm	18-35 mesh	815360.1	815360.5	815360.25
Silica FIA fine	0.071-0.16 mm	815410.1		
Silica FIA coarse	0.071-0.63 mm	815430.1		

Aluminum oxide

- · Aluminum oxides produced by dehydration of different aluminum hydroxides, e.g., hydrargillite between 400 and 500 °C.
- · Activity grade I, particle size 50-200 µm, specific surface (BET) $\sim 130 \text{ m}^2/\text{g}$

Ordering information

Description	рН	1 kg	5 kg	25 kg
Aluminum oxide 90 basic	pH 9.5 ± 0.3	815010.1	815010.5	815010.25
Aluminum oxide 90 neutral	pH 7 ± 0.5	815020.1	815020.5	815020.25
Aluminum oxide 90 acidic	pH 4 ± 0.3	815030.1	815030.5	815030.25

Adsorbents for column chromatography



Kieselguhr

- Naturally occurring amorphous silicic acids of fossil origin, also known as diatomaceous earth or diatomite purified for chromatographic applications
- Compared to silica, kieselguhr has a small surface of low activity → application in partition chromatography; impregnated with various substances (paraffin, silicone oil, undecane) it can be used for reversed phase chromatography
- The following grades of kieselguhr are manufactured by Johns-Manville. They are narrowly classified with homogeneous particle size distributions and high purity.
- For columns packed with kieselguhr please see CHROMABOND[®] XTR for liquid-liquid extraction, page 63.

Ordering information

3				
Description	Rel. purification factor	Rel. flow rate	1 kg	5 kg
Filter-Cel®	100	100	815510.1	815510.5
Hyflo [®] Super-Cel [®]	58	534	815530.1	815530.5
Celite® 503	42	910	815540.1	815540.5
Celite® 535	35	1269	815550.1	815550.5
Celite® 545	32	1830	815560.1	815560.5

Florisil[®]

- \cdot Hard granular magnesia silica gel: MgO 15.5 \pm 0.5 % \cdot SiO $_2$ 84.0 \pm 0.5 % \cdot Na $_2$ SO $_4$ \leq 1.0 %; 60/100 mesh
- Recommended application
 Sample preparation (see chapter "Solid phase extraction", page 16)
- Clean-up of pesticide residues, separation of chlorinated pesticides, extraction of steroids, sex hormones, antibiotics, lipids etc.

Ordering information

•				
Description	Particle size	1 kg	5 kg	
Florisil standard 60/100 mesh	0.15/0.25 mm	815710.1	815710.5	

Adsorbents for column chromatography

Polyamide

- · Polyamide 6 = ε-polycaprolactam
- · The separation mechanism mainly based on hydrogen
- · Recommended application Separation of phenolic compounds (e.g., isolation of natural products) carboxylic acids, aromatic nitro compounds
- · For SPE columns packed with polyamide see CHROMABOND® PA page 44.

Ordering information					
Description	Particle size	1 kg	5 kg		
Polyamide SC 6, < 0.07 mm	< 0,07 mm	815610.1	815610.5		
Polyamide SC 6, 0.05-0.16 mm	0.05–0.16 mm	815620.1	815620.5		
Polyamide SC 6, 0.10-0.30 mm	0.10–0.30 mm	815600.1	815600.5	••••••	

Unmodified cellulose

- · Cellulose MN 100: native fibrous cellulose, standard grade average degree of polymerization 620–680, fiber length (85 %) 20–100 μm, specific surface acc. to Blaine ~ 6500 cm²/g; residue on ignition at 850 °C < 10000 ppm, < 20 ppm Fe, < 5 ppm Cu, < 7 ppm P, CH₂Cl₂ extract < 0.20 %
- · Cellulose MN 2100: native fibrous cellulose, purified grade (washed with different eluents) average degree of polymerization 620-680, fiber length (85 %) 20-75 µm, specific surface acc. to Blaine ~ 5500 cm²/g residue on ignition at 850 °C < 1000 ppm,
- < 2 ppm Fe, < 1 ppm Cu, < 2 ppm P, CH₂Cl₂ extract < 0.15 %
- · Grade MN 2100ff is a defatted cellulose MN 2100 with a CH₂Cl₂ extract < 0.02 %

Ordering information						
Description	1 kg	5 kg	25 kg			
Cellulose MN 100	815050.1	815050.5	815050.25			
Cellulose MN 2100	815060.1	815060.5	815060.25			
Cellulose MN 2100ff (Cellulose MN 2100 defatted)	815070.1					





MACHEREY-NAGEL optimal autosampler vials for your sample

Vials and closures

For reliable and reproducible analysis the correct storage of sample solutions is important. MACHEREY-NAGEL offers diverse vials and suitable closures.

Our product range includes

- · Different vial types from N 8 to N 24
- Crimp neck
- Screw neck
- Snap ring
- · Clear glass, amber glass and polypropylene vials, with or without scale and label
- · Diverse inserts for small sample volumes
- · Variety of closures and septa of different material
- · Suitable accessories like crimping tools and vial contain-
- · Compatibility with different autosamplers from page 136 onwards



Our broad range of vials and closures can be found from page 97 onwards.

Also use our VialFinder on www.mn-net.com/VialFinder

For specific technical questons on LC columns, please contact:

Authorised Distributor:

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local distributor Instrument Solutions Technology for Your Success!

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