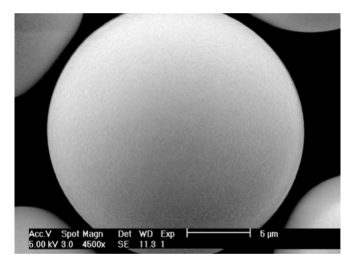


NUCLEODUR<sup>®</sup> is a fully synthetical type B silica (silica of 3rd generation) offering highly advanced physical properties like totally spherical particle shape, outstanding surface micro-structure, high pressure stability and low metal content.

NUCLEODUR<sup>®</sup> 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<sup>®</sup> 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<sup>®</sup> 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<sup>®</sup> is virtually free of metal impurities and low acidic surface silanols. Elemental analysis data of NUCLEODUR<sup>®</sup> 5  $\mu$ m measured by AAS are listed below.

Elementary ar	alysis (metal ions)	of NUCLEODUR <sup>®</sup> 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.

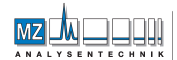
NUCLEODUR<sup>®</sup> silica is available with two pore sizes – 110 Å pore size as standard material and as 300 Å widepore material for the separation of biomolecules, like peptides and proteins.

Physical data of N	NUCLEODUR <sup>®</sup>	
	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<sup>®</sup> 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<sup>®</sup> phases please see page 152.



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AUTHORIZED DISTRIBUTOR



### 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

 NUCLEODUR<sup>®</sup> 1.8 µm particles are fractionated to limit the increase in back pressure.

### Advantages of 1.8 µm particle size

Miniaturization started in the early stage of HPLC with the reduction of particle size from 10  $\mu$ m via 7  $\mu$ m to standard 5  $\mu$ m – still the most used particle diameter in analytical HPLC – to 3  $\mu$ m spherical particles. With the introduction of 1.8  $\mu$ m 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  $\mu$ m particles.

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

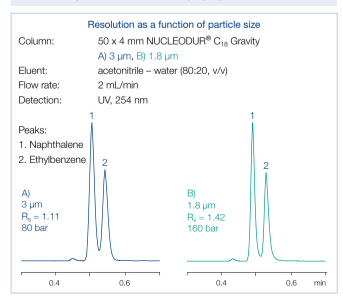
- 50 x 4.6 mm NUCLEODUR® C<sub>18</sub> Gravity
- $\cdot$  3 µm: N ≥ 100 000 plates/m (h-value≤ 10)
- 1.8  $\mu$ m: N  $\geq$  166 667 plates/m (h-value $\leq$  6)

Increase of the plate number by ~ 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,  $\alpha$  = selectivity (separation factor),  $k'_i$  = retention N = plate number with N  $\propto$  1/d<sub>P</sub>, d<sub>P</sub> = particle diameter



### Availability

- $\cdot$  The following NUCLEODUR® phases are available in 1.8  $\mu m:$
- C<sub>18</sub> Gravity, C<sub>8</sub> Gravity, C<sub>18</sub> Gravity-SB, C<sub>18</sub> Isis,
- C18 Pyramid, PolarTec, Phenyl-Hexyl, PFP, Sphinx RP,
- C<sub>18</sub> HTec and HILIC

Use of 1.8  $\mu$ m instead of 3  $\mu$ 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

$$\Delta_{\rm P} = \frac{\Phi \cdot L_{\rm C} \cdot \eta \cdot u}{d_{\rm P}^2}$$

$$\label{eq:DeltaP} \begin{split} \Delta_{\!P} &= \text{pressure drop, } \Phi = \text{flow resistance (nondimensional), } LC = \text{column} \\ & \text{length, } \eta = \text{viscosity, } u = \text{linear velocity, } d_{\!P} = \text{particle diameter} \end{split}$$

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

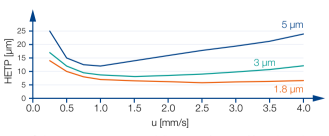
#### Comparison of back pressures

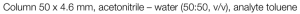
Eluent 100 % methanol, flow rate 1.5 mL/min

lemperature 22	0, column umensions 30 x 4.0	
	NUCLEODUR <sup>®</sup> C <sub>18</sub> 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  $\mu m$  particles is higher than for 3 and 5  $\mu m$  particles (see figure – the flow rate should be at the van Deemter minimum).





#### **Technical requirements**

Van Deemter curves

To gain best results with 1.8  $\mu$ 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.



nase	Specification	Page	Characteristic*	Stability	Structure		
			A •••••		e C		
	octadecyl, high density coating, multi-endcapping 18 % C · USP L1	158	В	pH 1–11, suitable for LC/MS	NUCCLEODUR® (Si-Og)n		
C <sub>18</sub> Gravity			C •••				
			A ••••		e I		
	octadecyl (monomeric), extensive endcapping 13 % C · USP L1	162	B ●●€	pH 1–9, suitable for LC/MS			
G <sub>18</sub> Gravity-SB			C -				
			A ●●€		e L		
	octyl, high density coating, multi-endcapping 11 % C · USP L7	158	В	pH 1–11, suitable for LC/MS	NUCLEODUR®		
C <sub>8</sub> Gravity			C •(				
	octadecyl phase with specially crosslinked surface modification endcapping 20 % C · USP L1		A ••••		۳ ۲		
		164	В●●	pH 1–10, suitable for LC/MS	(SI-0_2) (SI-0_2)		
C <sub>18</sub> Isis			C •••••				
			A ●●●●	eluent, pH 1–9, suitable for LC/MS	 ۲		
	octadecyl with polar endcapping 14 $\%~C\cdot USP~L1$	166	B ●●€		NUCCLEODUR®		
C <sub>18</sub> Pyramid			C ••				
			A ••••	stable in 100 % aqueous			
	octadecyl with embedded polar group 17 % C · USP L1 and L60	168	B ●●●	eluent, pH 1–9,	CLEODUR® Second Second		
PolarTec			C ••••	suitable for LC/MS	€ € Si <sup>-O*</sup> Si(CH <sub>3</sub> ) <sub>3</sub>		
			A ••				
	phenylhexyl, multi-endcapping 10 % C · USP L11	170	В ●●●	pH 1–10, suitable for LC/MS	NUCLEODUR (Si-O.2) (Si-O.2) (Si-O.2)		
Phenyl-Hexyl			C 🔴		D 5		
	histopylarocyl		A ••	pH 1.5–10			
	biphenylpropyl, multi-endcapping 17 % C · USP L11	172	B ●●●●		Sichally Sic		
$\pi^2$			С •••				





Application	Similar phases**	Interactions · retention mecl	hanism
in general compounds with ionizable functional groups such as basic pharmaceuticals and pesticides	NUCLEOSIL <sup>®</sup> C <sub>18</sub> HD Xterra <sup>®</sup> RP18 / MS C18; Luna <sup>®</sup> C18(2), Gemini <sup>®</sup> , Synergi <sup>®</sup> Max RP; Zorbax <sup>®</sup> Extend-C18; Inertsil <sup>®</sup> ODS III; Purospher <sup>®</sup> STAR RP-18; Hypersil <sup>™</sup> BDS	hydrophobic (van der Waals interactions)	Si(CH <sub>3</sub> ) <sub>3</sub> H <sub>5</sub> C
 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(CHa)a Hac-N-N-Si(CHa)a Hac-N-N-Si(CHa)a Hac-N-N-Si(CHa)a Hac-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N
like C <sub>18</sub> Gravity, however, gene- rally shorter retention times for nonpolar compounds	NUCLEOSIL <sup>®</sup> C <sub>8</sub> HD Xterra <sup>®</sup> RP8 / MS C8; Luna <sup>®</sup> C8; Zorbax <sup>®</sup> Eclipse XDB-C8	hydrophobic (van der Waals interactions)	OH SI(CH <sub>3</sub> ) <sub>3</sub> OH OH OH OH OH
high steric selectivity, thus suited for separation of positional and structural isomers, planar / nonplanar molecules	NUCLEOSIL <sup>®</sup> C <sub>18</sub> AB Inertsil <sup>®</sup> ODS-P; Pro C18 RS	steric and hydrophobic	
basic pharmaceuticals, very polar compounds, organic acids	Aqua, Synergi <sup>®</sup> Hydro-RP; AQ; Atlantis <sup>®</sup> dC18; Polaris <sup>®</sup> C18-A	hydrophobic and polar (H bonds)	
basic pharmaceuticals, organic acids, pesticides, amino acids, water-soluble vitamins	NUCLEOSIL <sup>®</sup> C <sub>18</sub> Nautilus ProntoSIL <sup>®</sup> C18 AQ, Zorbax <sup>®</sup> Bonus-RP, Polaris <sup>®</sup> Amide-C18; Ascentis <sup>®</sup> RP Amide, SymmetryShield <sup>™</sup> RP18; SUPELCOSIL <sup>™</sup> LC-ABZ <sup>+</sup> ; HyPURITY <sup>™</sup> ADVANCE; ACCLAIM Polar AD.II	hydrophobic and polar (H bonds)	
aromatic and unsaturated com- pounds, polar compounds like pharmaceuticals, antibiotics	Luna <sup>®</sup> Phenyl-Hexyl; Zorbax <sup>®</sup> Eclipse Plus Phenyl-Hexyl; Kromasil <sup>®</sup> Phenyl-Hexyl	π-π and hydrophobic	O <sub>2</sub> N
aromatic and unsaturated com- pounds, polar compounds like pharmaceuticals, antibiotics	Pinnacle® DB Biphenyl; Ultra Biphenyl	π-π and hydrophobic	

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

# NUCLEODUR<sup>®</sup> phase overview



se	Specification	Page	Characteristic*	Stability	Structure
			A ••		® └└ ───── ३-Si-OH
	pentafluorophenylpropyl, multi-endcapping 8 % C · USP L43	174	B ●●●●	pH 1–9, suitable for LC/MS	Si-OH Si(CH <sub>abs</sub> )
PFP			C ••••		
	bifunctional, balanced ratio of		A •••		e ۲
	propylphenyl and octadecyl, endcapping	176	В ●●◀	pH 1–10, suitable for LC/MS	NUCLEODUR® (Si-O <sub>2</sub> )n
Sphinx RP	15 % C · USP L1 and L11		C 🔴		NUN
			A •••••		e C
	octadecyl, high density coating, high capacity, multi-endcapping 18 % C · USP L1	178	В	pH 1–11, suitable for LC/MS	NUCLEODUR <sup>®</sup> (Si-O <sub>2</sub> ), (Si-O <sub>2</sub> ),
C <sub>18</sub> HTec			C •••		
	octadecyl, medium density, endcapping available in 110 Å and 300 Å pore size 17.5 % / 4 % C · USP L1		A ••••		
		181	В	pH 1–9	NUCLEODU <sub>(S)</sub>
C <sub>18</sub> ec			C ••••		
			A ••	рН 1–9	
	octyl, medium density, endcap- ping 10.5 % C · USP L7	181	В●€		NUCLEOUR (S) (S) (S) (S) (CH <sup>9</sup> ) (S) (CH <sup>9</sup> )
C <sub>8</sub> ec			C •••		
			A ●		
	butyl, medium density, endcap- ping, 300 Å pore size 2.5 % C · USP L26	181	В●●	pH 1–9	SI-OH SI-OH SI-OH SI-OH SI-OH
C <sub>4</sub> ec			C •(		
	zwitterionic ammonium – sulfonic		A •		
	acid phase 7 % C	184	B ●●●●●	pH 2–8.5	NUCLEODUR o'o'o o'o'o'o'
HILIC			C -		Z <sup>\$</sup>
	cyano (nitrile) for NP and RP		A •	<sup></sup> рН 1–8,	
	separations 7 % C · USP L10	186	В ●●●●	pH 1–8, stable towards highly aqueous mobile phases	
CN/CN-RP			C -		





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

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

## NUCLEODUR<sup>®</sup> phase overview



#### Overview of NUCLEODUR® HPLC phases Phase Specification Page Characteristic\* Stability Structure А NUCLEODUR<sup>®</sup> (Si-O<sub>2</sub>)<sub>n</sub> aminopropyl for NP and RP pH 2–8, Si-OH separations 188 В stable towards highly .... 2.5 % C · USP L8 aqueous mobile phases Si-OH С -NH<sub>2</sub>/NH<sub>2</sub>-RP A -NUCLEODUR<sup>®</sup> (Si-O<sub>2</sub>)<sub>n</sub> -Si-<mark>OH</mark> unmodified high purity silica · 190 в pH 2–8 USP L3 Si-OH С \_ SiOH \* A = • hydrophobic selectivity, B = • polar / ionic selectivity, C = • steric selectivity

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Application	Similar phases**	Interactions · retention mech	nanism
sugars, sugar alcohols and other hydroxy compounds, DNA ba- ses, polar compounds in general	NUCLEOSIL® NH2/NH2-RP	polar/ionic and hydro- phobic	
polar compounds in general	NUCLEOSIL® SIOH	polar/ionic	SIOH ← → O₂N
** phases which provide a similar	selectivity based on chemical and physical propertie	S	



### NUCLEODUR<sup>®</sup> C<sub>18</sub> Gravity · C<sub>8</sub> Gravity nonpolar high density phase · USP L1 (C<sub>18</sub>) · USP L7 (C<sub>8</sub>)

### 📩 Key feature

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

### 🖊 Technical data

- $\cdot$  Available as octadecyl (C\_{18}) and octyl (C\_8), multi-endcapped
- Pore size 110 Å; particle sizes 1.8  $\mu$ m, 3  $\mu$ m and 5  $\mu$ m for C<sub>18</sub>, 1.8 and 5  $\mu$ m for C<sub>8</sub>; 7, 10, 12 and 16  $\mu$ m 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**

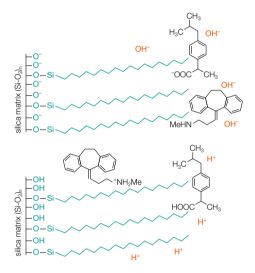
NUCLEODUR<sup>®</sup> C<sub>18</sub> Gravity and NUCLEODUR<sup>®</sup> C<sub>8</sub> Gravity are based on the ultrapure NUCLEODUR<sup>®</sup> silica. Derivatization generates a homogeneous surface with a high density of bonded silanes (~18 % C for C<sub>18</sub>, ~11 % C for C<sub>8</sub>). Thorough 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<sub>18</sub> phases compared to C<sub>8</sub> 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<sup>®</sup> C<sub>18</sub> and C<sub>8</sub> Gravity allow for use at an expanded pH range from pH 1 to 11.

### Benefits of enhanced pH stability

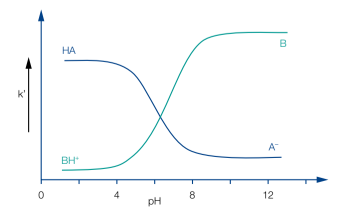
An expanded pH range is often required in method development. Many nitrogen containing compounds like basic drugs are protonated at acidic or neutral pH and exhibit poor retention on a standard  $C_{18}$  phase. The retention behavior can be improved by working at a higher pH, where the analyte is no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

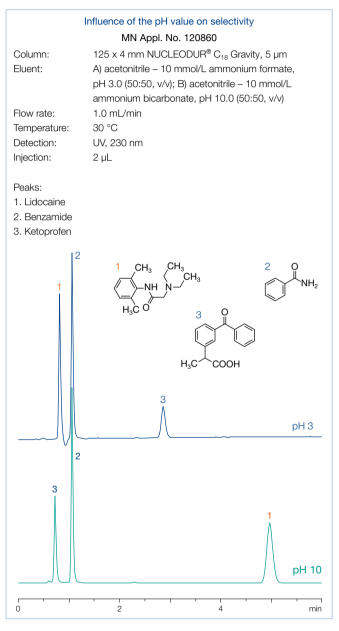
The figure above shows the extent of protonation of surface silanols and of two exemplary analytes at acidic and alkaline pH. The following graph explains the general correlation between retention and pH.

Correlation between retention and pH for basic and acidic compounds

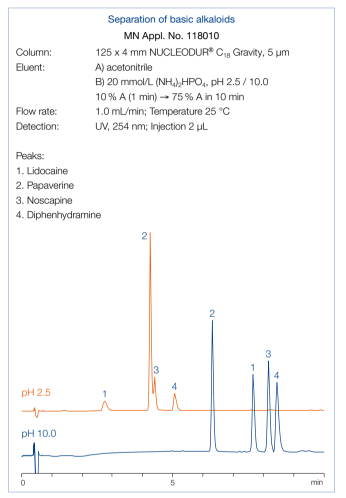




An example how selectivity can be controlled by pH is the separation of the acid ketoprofen, the base lidocaine and benzamide. Under acidic conditions the protonated lidocaine is eluted very fast due to lack of sufficiently strong hydrophobic interactions between analyte and  $C_{18}$  chains, while 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.

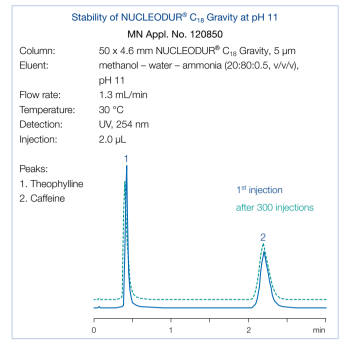


As mentioned above, pH stability of the stationary phase can be helpful for improving selectivity in method development. The following figure shows the separation of 4 basic drugs under acidic and basic conditions. At pH 2.5 the protonated analytes exhibit poor retention (early elution) and in addition 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<sup>®</sup> C<sub>18</sub> Gravity under alkaline conditions. The ultrapure Gravity with its unique high density surface bonding technology withstands strong alkaline mobile phase conditions.

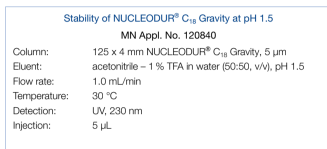




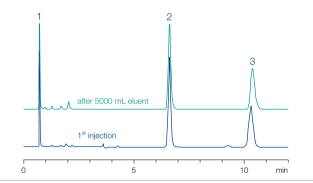
Even after 300 injections no loss of column efficiency – identified, e.g., by peak broadening or decrease in retention times – could be observed.

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<sup>®</sup> C<sub>18</sub> 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 <sup>®</sup> C	C18 Gravity	, <b>1.8 μm</b> 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 <sup>®</sup> C	C18 Gravity	, 3 µm octade	cyl phase, particl	e size 3 µm, 18 %	6 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		





### Ordering information

Eluent in column acetonitrile - water	
---------------------------------------	--

	ID	Length →						
		30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR <sup>®</sup> C	318 Gravity	, 5 µm octadec	yl phase, particle	e size 5 µm, 18 %	C			
Analytical EC column	S							
	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
EC guard columns*			4 x 2 mm:	761903.20	4 x 3 mm:	761903.30		
Preparative VarioPrep	columns							
	10 mm		762103.100			762109.100		762113.100
	21 mm		762103.210			762109.210		762113.210
L	32 mm							762113.320
	40 mm						762100.400	762113.400
VP guard columns			10 x 8 mm:	762160.80	10 x 16 mm	: 762160.160	15 x 32 mm	: 762163.320
NUCLEODUR® C	Gravity	, 10 µm octade	ecyl phase, partic	cle size 10 µm, 18	3 % C			
Preparative VarioPrep	columns							
	21 mm							762250.210
L	40 mm							762250.400
VP guard columns **					10 x 16 mm	: 762160.160	15 x 32 mm	: 762163.320

### Ordering information

Ordering informa								
Eluent in column ace	etonitrile – w	vater						
	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR <sup>®</sup> C	<sub>8</sub> Gravity,	1.8 µm octyl	phase, particle siz	e 1.8 μm, 11 % 0				
Analytical EC column	s							
	2 mm	760756.20	760755.20	760760.20	760757.20		760759.20	
	3 mm	760756.30	760755.30		760757.30			
	4 mm	760756.40	760755.40		760757.40			
	4.6 mm	760756.46	760755.46		760757.46			
EC guard columns*			4 x 2 mm:	761905.20	4 x 3 mm:	761905.30		
NUCLEODUR <sup>®</sup> C	S <sub>8</sub> Gravity,	5 µm octyl ph	ase, particle size	5 µm, 11 % C				
Analytical EC column								
	2 mm		760750.20		760754.20	760751.20	760752.20	760753.20
	3 mm		760750.30	•	760754.30	760751.30	760752.30	760753.30
	4 mm		760750.40		760754.40	760751.40	760752.40	760753.40
	4.6 mm	-	760750.46	760749.46	760754.46	760751.46	760752.46	760753.46
EC guard columns*			4 x 2 mm:	761907.20	4 x 3 mm:	761907.30		
Preparative VarioPrep	columns							
	10 mm		762081.100			762071.100		762070.100
L	21 mm		762081.210			762071.210	762082.210	762070.210
VP guard columns **			10 x 8 mm:	762097.80	10 x 16 mn	n: 762097.160		
EC and VarioPrep col	umns in pac	cks of 1, guard co	olumns see below.					
Guard column sy	vstems							
Guard columns for E	C columns	with ID	2 mm	3 mn	n 4 m	im 4.6	mm G	uard column hole

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	

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

### 🔀 Key feature

- Hydrophobic C<sub>18</sub> 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

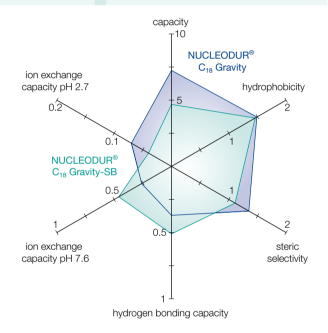
### Recommended application

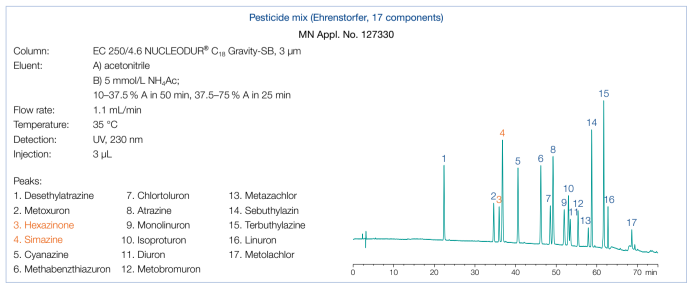
 Overall sophisticated analytical separations, especially for polar compounds, e.g., antibiotics, water-soluble vitamins, organic acids

NUCLEODUR<sup>®</sup> C<sub>18</sub> Gravity-SB excels with a relatively high hydrophobicity – similar to C<sub>18</sub> 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<sub>18</sub> phase.

In the TANAKA plot the NUCLEODUR<sup>®</sup> 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<sup>®</sup> C<sub>18</sub> Gravity-SB is versatile applicable, especially for polar analytes like nucleobases or pesticides the column shows good separation efficiency.





Good separation of the critical pair hexazinone/simazine





#### Comparing of selectivity for nucleobases MN Appl. No. 127270 Columns: EC 150/4.6 mm NUCLEODUR® C<sub>18</sub> Gravity-SB, 5 µm NUCLEODUR® C<sub>18</sub> Gravity, 5 µm NUCLEODUR® C18 Pyramid, 5 µm Eluent: 25 mmol/L KH<sub>2</sub>PO<sub>4</sub>, pH 3 – methanol (95:5, v/v) 1.0 mL/min, Temperature: 20 °C Flow rate: Detection: UV, 220 nm, Injection: 2.5 µL (1 mg/mL) Peaks: 4. Guanine 1. Cytosine VV 2. Adenine 5. Thymine 234 5 3. Uracil 3 7 2 0 5 i 4 6 8 min

Better resolution of early eluting analyte

	ID	Length →								
	10	30 mm	50 mm	75 mm	1(	00 mm	125 mm	150 ו	mm	250 mm
NUCLEODUR® C	Gravity	-SB, 1.8 μm	particle size	1.8 µm · UHPL	0					
Analytical EC column										
,	2 mm	760591.20	760593.2	20 760595	.20 70	60596.20		7605	98.20	
	3 mm	760591.30	760593.3	30	76	60596.30		•		•••••
	4 mm	760591.40	760593.4	40	76	60596.40		•••••		•••••
	4.6 mm	760591.46	760593.4	46	70	60596.46	•••••	••••••		
EC guard columns*	•	•	4 x 2	mm: 761990.20	)	4 x 3 mm:	761990.30	•••••		
NUCLEODUR <sup>®</sup> (	Gravity	-SB, 3 µm pa	rticle size 3	μm						
Analytical EC column										
	2 mm		760603.2	20	70	60606.20	760607.2	0 7606	08.20	760609.20
	3 mm	••••	760603.3	••••••••••••••••	••••••	50606.30	760607.3	• • • • • • • • • • • • • • • • • • • •	08.30	760609.30
	4 mm		760603.4	40	76	50606.40	760607.4	0 7606	08.40	760609.40
	4.6 mm	•	760603.4	46 760605	.46 76	50606.46	760607.4	6 7606	08.46	760609.46
EC guard columns*	•••••	•	4 x 2	mm: 761991.20	)	4 x 3 mm:	761991.30	•••••		•••••
NUCLEODUR <sup>®</sup> (	Gravity	-SB. 5 um pa	rticle size 5	um						
Analytical EC column		- A - F - F-								
	2 mm		760613.2	20	76	50616.20	760617.2	0 7606	18.20	760619.20
	3 mm		760613.3		•••••••	50616.30	760617.3	••••••	18.30	760619.30
	4 mm		760613.4	••••••••••••••••	••••••	50616.40	760617.4	••••••	18.40	760619.40
	4.6 mm	•••••	760613.4	•••••••••••••••••••••••••••••••••••••••	••••••	50616.46	760617.4	• • • • • • • • • • • • • • • • • • • •	18.46	760619.46
EC guard columns*		····	• • • • • • • • • • • • • • • • • • • •	mm: 761992.2	•••••••••••••••••••••••••••••••••••••••	•••••	761992.30			
Preparative VarioPrep	columns									
	10 mm		762350.	100			762351.1	00		762353.100
	21 mm	•	762350.2	210	•••••		762351.2	10		762353.210
	32 mm	•						•••••		762353.320
	40 mm		•••••	•	•••••		•••••	7623	52.400	762353.400
VP guard columns **			10 x 8	mm: 762354.80	)	10 x 16 mm: 762354.160			x 32 mm	: 762355.320
EC and VarioPrep co	lumns in pac	ks of 1, guard co	olumns see b	elow.						
Guard column sy	/stems									
Guard columns for E	C columns	with ID		2 mm	3 mm	4 m	m	4.6 mm	Gu	ard column holde
* Column Protection	System (pac	k of)	EC	4/2 (3)	4/3 (3)	4/3	(3)	4/3 (3)	71	3966
Guard columns for V	/arioPrep co	lumns with ID		8, 10 mm	16, 21 mr	n 32,	40 mm	≥ 50 mm		
	•			10/0 (0)	10/10/0)		0 (1)	15/50 (1)		

10/8 (2)

718251

10/16 (2)

718256

VP

For details of our column systems see page 250.

\*\* VP guard columns (pack of)

VP guard column holder

15/50 (1)

718255

15/32 (1)

718253



### $NUCLEODUR^{\circledast} \ C_{18} \ Isis$ phase with high steric selectivity $\cdot \ \text{USP L1}$

### 🔽 Key feature

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

### 꾿 Technical data

 C<sub>18</sub> 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

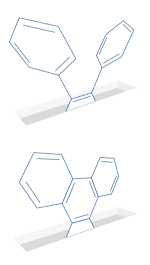
• Steroids, (*o*,*p*,*m*-)substituted aromatics, fat-soluble vitamins

### Surface modification

By use of specific C<sub>18</sub> silanes and polymeric bonding technologies a dense shield of alkyl chains protects the subjacent silica matrix. Elemental analysis of NUCLEODUR<sup>®</sup> C<sub>18</sub> Isis shows a carbon load of 20 %. The target crosslinking of the C<sub>18</sub> 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.

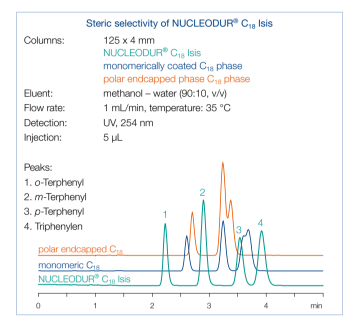
### Slot Model

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).

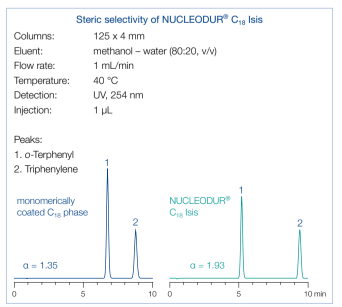


### Steric selectivity

The following chromatograms reveal the improved resolution for positional isomers in a test mixture of aromatic compounds on NUCLEODUR<sup>®</sup> C<sub>18</sub> Isis (green) in direct comparison with monomerically coated (blue) and polar endcapped (orange) C<sub>18</sub> 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  $\alpha$  is a measure for the steric selectivity. As is shown below the  $\alpha$  value is considerable larger on NUCLEODUR® C18 lsis compared to a conventional C18 column.







The surface bonding technology also provides improved stability features for the NUCLEODUR  $^{\circledast}$  C\_{18} Isis phase.

### Surface deactivation

The chromatography of basic analytes requires a high density of surface-bonded  $C_{18}$  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*).

Eluent in column ac		ater								
	ID	Length → 30 mm	50 mm	75 m	m	100 mm	125 mm	ı 1	50 mm	250 mm
NUCLEODUR® (	C₁₀ Isis. 1.8	3 um particle s	ize 1.8 um	·UHPLC						
Analytical EC column		- F F								
	2 mm	760406.20	760405	5.20 7603	96.20	760407.20	)	70	60409.20	
	3 mm	760406.30	760405		••••••	760407.30	·····			
	4 mm	760406.40	760405		•••••••	760407.40	· · · · · · · · · · · · · · · · · · ·	•••••		
	4.6 mm	760406.46	760405	••••••	•••••••••••••••••••••••••••••••••••••••	760407.46	· · · · · · · · · · · · · · · · · · ·	•••••		
EC guard columns*			4 x	2 mm: 761910.	20	4 x 3 r	nm: 761910.30	•••••		
NUCLEODUR <sup>®</sup> (	Cas Isis, 3 (	Im particle size	a 3 um							
Analytical EC column			, o p							
	2 mm		760400	0.20		760401.20	760402.	.20 70	60403.20	760404.20
	3 mm	•••••	760400	••••••	•••••••••••••••••••••••••••••••••••••••	760401.30		· · · · · · · · · · · · · · · · · · ·	60403.30	760404.30
	4 mm		760400	••••••	•••••••	760401.40	· · · · · · · · · · · · · · · · · · ·		60403.40	760404.40
	4.6 mm		760400	••••••	••••••	760401.46	••••••	·····	60403.46	760404.46
EC guard columns*	••••••	•••••••••••••••••••••••••••••••••••••••	4 x	2 mm: 761911.	20	4 x 3 r	nm: 761911.30	•••••		
NUCLEODUR <sup>®</sup> (	Cas Isis, 5 (	Im particle size	e 5 um							
Analytical EC column										
	2 mm		760410	).20		760415.20	760412.	.20 70	60413.20	760414.20
	3 mm	••••	760410	).30	•••••	760415.30	) 760412.	.30 70	60413.30	760414.30
	4 mm		760410	).40	•••••	760415.40	) 760412.	.40 70	60413.40	760414.40
	4.6 mm	•	760410	0.46 7604	16.46	760415.46	6 760412.	.46 70	60413.46	760414.46
EC guard columns*	•••••		4 x	2 mm: 761912.	20	4 x 3 r	nm: 761912.30	•••••		
Preparative VarioPrep	o columns									
	10 mm		762404	.100			762405.	100		762403.100
	21 mm		762404	.210			762405.	.210		762403.210
L	32 mm									762403.320
	40 mm							7(	62406.400	762403.400
VP guard columns **			10 x	8 mm: 762420.	80	10 x 16	mm: 762420.1	60	15 x 32 mm	: 762422.320
EC and VarioPrep co	lumns in pac	ks of 1, guard co	olumns see	below.						
Guard column sy	/stems									
Guard columns for E	C columns	with ID		2 mm	3 mm		4 mm	4.6 mm	Gu	ard column holde
* Column Protection	System (pac	< of)	EC	4/2 (3)	4/3 (3)		4/3 (3)	4/3 (3)	718	3966
Guard columns for \	/arioPrep co	lumns with ID		8, 10 mm	16, 21 n		32, 40 mm	≥ 50 mm	า	
** VP guard columns	(pack of)		VP	10/8 (2)	10/16 (2	)	15/32 (1)	15/50 (1	)	
VP guard column ho	der			718251	718256	•••••	718253	718255	•••••	

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

### 📩 Key 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 μm, 3 μm and 5 μm (7 and 10 μm particles for preparative purposes on request); carbon content 14 %; pH stability 1–9

### Recommended application

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

### RP-HPLC with highly aqueous mobile phases

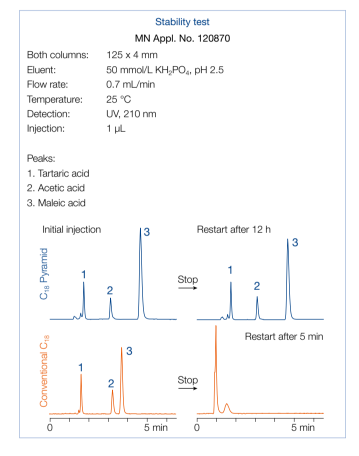
The efforts to neutralize unwanted silanol activity often results in well base-deactivated RP phases with high carbon load, but a limited scope of selectivity beyond non-polar interactions. Polar compounds like carboxylic acids or drug metabolites show only weak retention on densely bonded RP columns 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<sup>®</sup> PolarTec may be taken as an example for the embedded polar group strategy, in which a C<sub>18</sub> silane with a polar function is successfully linked to the silica surface.

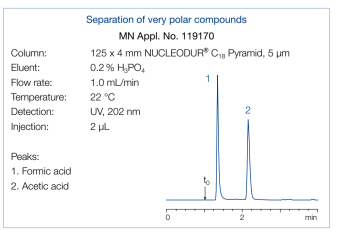
### Stability features

NUCLEODUR<sup>®</sup> C<sub>18</sub> 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<sup>®</sup> C<sub>18</sub> Pyramid in comparison with a conventionally bonded C<sub>18</sub> phase.

It can be shown that the retention times for NUCLEODUR<sup>®</sup> C<sub>18</sub> 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 to-tally after 5 min.



### **Retention characteristics**







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*).

Ordering information	ation									
Eluent in column ac	etonitrile – w	ater								
	ID	Length $\rightarrow$								
		30 mm	50 mm	ו	75 mm	100 mm	125 mm	150 mm	n 250 mm	1
NUCLEODUR® (	C <sub>18</sub> Pyrami	d, 1.8 μm pa	rticle size	1.8 µm · Uŀ	HPLC					
Analytical EC colum	ns									
	2 mm	760271.20	76027	2.20	760275.20	760273.20	)	760274	.20	
	3 mm	760271.30	76027	2.30		760273.30	)			
	4 mm	760271.40	76027	2.40		760273.40	)			
	4.6 mm	760271.46	76027	2.46		760273.46	;	•		
EC guard columns*			4 x	( 2 mm: 76 <sup>-</sup>	1915.20	4 x 3 n	nm: 761915.30			
NUCLEODUR <sup>®</sup> (	C <sub>18</sub> Pyrami	d, 3 µm partio	cle size 3 µ	ım						
Analytical EC colum	ns									
	2 mm		76026	3.20		760264.20	760260.2	20 760261	.20 760262.3	.20
	3 mm	•	76026	3.30		760264.30	760260.3	30 760261	.30 760262.3	.30
	4 mm	•	76026	3.40		760264.40	760260.4	10 760261	.40 760262.4	.40
	4.6 mm	•••••	76026	3.46	760259.46	760264.46	760260.4	16 760261	.46 760262.4	.46
EC guard columns*	•••••		4 x	2 mm: 76 <sup>-</sup>	1916.20	4 x 3 n	nm: 761916.30	•••••		
NUCLEODUR <sup>®</sup>	C₁₀ Pvrami	d. 5 um partio	cle size 5 i	ım						
Analytical EC colum										
	2 mm		76020	0.20		760204.20	760201.2	20 760203	.20 760202.3	.20
	3 mm	•••••••••••••••••••••••••••••••••••••••	76020	0.30		760204.30	760201.3	30 760203	.30 760202.3	.30
	4 mm	•••	76020	0.40		760204.40	760201.4	10 760203	.40 760202.4	.40
	4.6 mm	•	76020	0.46	760205.46	760204.46	760201.4	16 760203	.46 760202.4	.46
EC guard columns*	•		4 x	2 mm: 76	1917.20	4 x 3 n	nm: 761917.30	•	•••••	
Preparative VarioPre	p columns									
	10 mm		76227	1.100			762273.1	100	762272.	.100
	21 mm		76227	1.210			762273.2	210	762272.	.210
L	32 mm								762272.3	.320
	40 mm						•	762269	.400 762272.4	.400
VP guard columns *	*		10 ×	( 8 mm: 762	2291.80	10 x 16	mm: 762291.16	0 15 x	32 mm: 762293.32	20
EC and VarioPrep co	olumns in pac	ks of 1, guard co	olumns see	e below.						
Guard column s	ystems									
Guard columns for	EC columns	with ID		2 mm	3 mn	n 4	1 mm	4.6 mm	Guard column	1 holde
* Column Protection	System (pac	k of)	EC	4/2 (3)	4/3 (	3) 4	4/3 (3)	4/3 (3)	718966	
	VarioPrep co	lumns with ID		8, 10 mn	n 16, 2	1 mm 3	32, 40 mm	≥ 50 mm		
Guard columns for	ranor rop oo									
<pre>Guard columns for ** VP guard columns</pre>	-		VP	10/8 (2)	10/1	6 (2) ·	15/32 (1)	15/50 (1)		

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

### 📩 Key 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,  $\pi$ - $\pi$ , etc.). These interactions enhance retention and selectivity for polar compounds like carboxylic acids, phenols and nitrogen containing compounds.

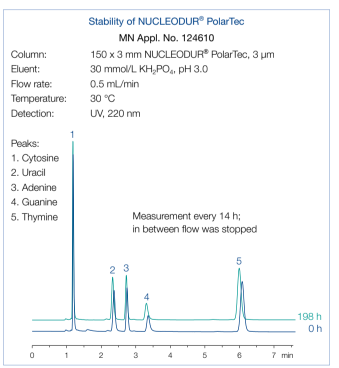
Separation of histidines							
MN Appl. No. 125140							
Column:	Column: 150 x 3 mm NUCLEODUR <sup>®</sup> PolarTec, 3 µm						
Eluent:	1.0 mmol/L perfluoropentanoic acid in water –						
	0.5 mmol/L perfluoropentanoic acid in acetonitrile						
	(99.5:0.5, v/v)						
Flow rate:	0.4 mL/min						
Temperature:	20 °C						
Detection:	UV, 230 nm						
Peaks:							
1. 3-Methylhistid	line $R_1 = H, R_2 = CH_3$						
2. Histidine	$R_1=R_2=H$						
3. 1-Methylhistid	line $R_1 = CH_3, R_2 = H$						
	13						
	Ŭ						
	NH <sub>2</sub>						
	B <sub>1</sub> , J, OH						
	R <sub>1</sub> ,						
	2						
	· · · · · · · · · · · · · · · · · · ·						
0	2 4 6 8 min						

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<sup>®</sup> 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<sup>®</sup> 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  $\ensuremath{\mathsf{NUCLEODUR}}^{\ensuremath{\$}}$  PolarTec is also suitable for LC/MS.

Even after days or weeks of operation in purely aqueous eluents the  $C_{18}$  chains of NUCLEODUR<sup>®</sup> 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<sup>®</sup> PolarTec exhibits sufficient hydrophobic properties and is very well suited for analyzing basic compounds.





### Ordering information

Eluent in columr	n acetonitrile – water
------------------	------------------------

		Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
	PolarTec, 1	I.8 µm particle	size 1.8 µm · UH	PLC				
Analytical EC column								
	2 mm	760461.20	760463.20	760465.20	760466.20		760468.20	
	3 mm	760461.30	760463.30	•••••	760466.30	••••	•••••	•••••
	4 mm	760461.40	760463.40	•	760466.40		•••••	•
	4.6 mm	760461.46	760463.46	•	760466.46			
EC guard columns*			4 x 2 mm:	761980.20	4 x 3 mm:	761980.30		
NUCLEODUR <sup>®</sup> F	PolarTec, 3	βμm particle si	ize 3 µm					
Analytical EC column	S							
	2 mm		760473.20		760476.20	760477.20	760478.20	760479.20
	3 mm		760473.30	•••••	760476.30	760477.30	760478.30	760479.30
	4 mm		760473.40	•	760476.40	760477.40	760478.40	760479.40
	4.6 mm	•	760473.46	760475.46	760476.46	760477.46	760478.46	760479.46
EC quard columns*		•	4 x 2 mm:	761981.20	4 x 3 mm:	761981.30		
	PolarTec, 5	5 µm particle si	ize 5 µm					
		δμm particle si	ze 5 µm					
		ο μm particle si	ize 5 μm 760483.20		760486.20	760487.20	760488.20	760489.20
	S	ο μm particle si					760488.20 760488.30	760489.20 760489.30
	s 2 mm	5 μm particle si	760483.20		760486.20	760487.20		
	s 2 mm 3 mm	5 μm particle si	760483.20 760483.30	760485.46	760486.20 760486.30	760487.20 760487.30	760488.30	760489.30
Analytical EC column	s 2 mm 3 mm 4 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46		760486.20 760486.30 760486.40 760486.46	760487.20 760487.30 760487.40	760488.30 760488.40	760489.30 760489.40
NUCLEODUR <sup>®</sup> F Analytical EC column	s 2 mm 3 mm 4 mm 4.6 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46	760485.46	760486.20 760486.30 760486.40 760486.46	760487.20 760487.30 760487.40 760487.46	760488.30 760488.40	760489.30 760489.40
NUCLEODUR® F Analytical EC column	s 2 mm 3 mm 4 mm 4.6 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46	760485.46	760486.20 760486.30 760486.40 760486.46	760487.20 760487.30 760487.40 760487.46	760488.30 760488.40	760489.30 760489.40 760489.46
NUCLEODUR <sup>®</sup> F Analytical EC column	s 2 mm 3 mm 4 mm 4.6 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46 4 x 2 mm:	760485.46	760486.20 760486.30 760486.40 760486.46	760487.20 760487.30 760487.40 760487.46 761982.30	760488.30 760488.40	760489.30 760489.40 760489.46 762223.100
NUCLEODUR® F Analytical EC column	s 2 mm 3 mm 4 mm 4.6 mm 0 columns 10 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46 4 x 2 mm: 762220.100	760485.46	760486.20 760486.30 760486.40 760486.46	760487.20 760487.30 760487.40 760487.46 761982.30 762221.100	760488.30 760488.40	760489.30 760489.40 760489.46 762223.100 762223.210
NUCLEODUR <sup>®</sup> F Analytical EC column	s 2 mm 3 mm 4 mm 4.6 mm 0 columns 10 mm 21 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46 4 x 2 mm: 762220.100	760485.46	760486.20 760486.30 760486.40 760486.46	760487.20 760487.30 760487.40 760487.46 761982.30 762221.100	760488.30 760488.40	760489.30 760489.40
NUCLEODUR® F Analytical EC column EC guard columns* Preparative VarioPrep	s 2 mm 3 mm 4 mm 4.6 mm 0 columns 10 mm 21 mm 32 mm	5 μm particle si	760483.20 760483.30 760483.40 760483.46 4 x 2 mm: 762220.100	760485.46 761982.20	760486.20 760486.30 760486.40 760486.46 4 x 3 mm:	760487.20 760487.30 760487.40 760487.46 761982.30 762221.100	760488.30 760488.40 760488.46 760488.46	760489.30 760489.40 760489.46 762223.100 762223.210 762223.320

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	

### NUCLEODUR<sup>®</sup> Phenyl-Hexyl productive for polar/aromatic compunds · USP L11

### 📩 Key feature

- Hydrophobic phase with alternative selectivity compared to classical C<sub>18</sub> 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  $\mu$ m, 3  $\mu$ m and 5  $\mu$ m; carbon content 10 %; pH stability 1–10

### Recommended application

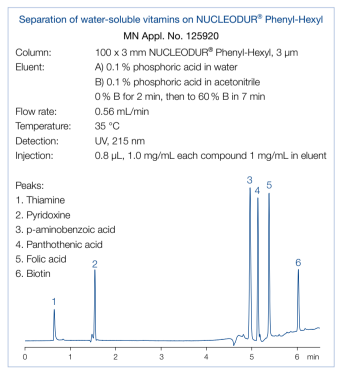
 Aromatic and unsaturated compounds, polar compounds like pharmaceuticals, antibiotics

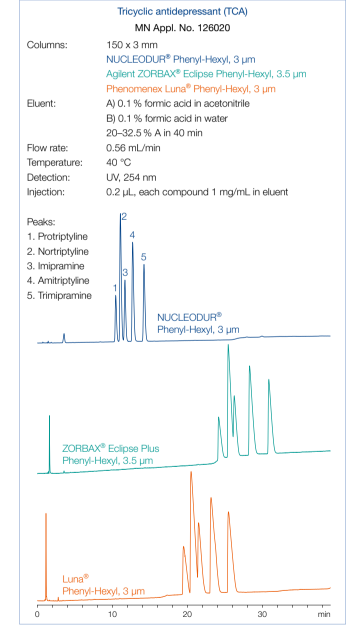
Phenylhexyl modified phases are an interesting alternative to classical  $C_{18}$  phases due to an excellent separation of aromatic and unsaturated compounds especially with electron withdrawing groups.

The combination of hydrophobic and polar  $\pi$ - $\pi$  interactions result in an interesting and alternate selectivity in comparison to C<sub>18</sub> and C<sub>8</sub> modified phases.

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

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









### Ordering information

Eluent in column acetonitrile - water

	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
	henyl-He	xyl, 1.8 µm p	article size 1.8 µm	· UHPLC				
Analytical EC column		<b>9</b> 7 - 1						
	2 mm	760561.20	760563.20	760565.20	760566.20		760568.20	
	3 mm	760561.30	760563.30		760566.30	••••	•••••	
	4 mm	760561.40	760563.40		760566.40			
	4.6 mm	760561.46	760563.46		760566.46			•
EC guard columns*			4 x 2 mm:	761985.20	4 x 3 mm:	761985.30		
NUCLEODUR <sup>®</sup> F	henyl-He	xyl, 3 µm par	ticle size 3 µm					
Analytical EC column	s							
	2 mm		760573.20		760576.20	760577.20	760578.20	760579.20
	3 mm		760573.30		760576.30	760577.30	760578.30	760579.30
	4 mm		760573.40		760576.40	760577.40	760578.40	760579.40
	4.6 mm		760573.46	760575.46	760576.46	760577.46	760578.46	760579.46
EC guard columns*			4 x 2 mm:	761986.20	4 x 3 mm:	761986.30		
NUCLEODUR <sup>®</sup> F	henyl-He	xyl, 5 µm par	ticle size 5 µm					
Analytical EC column	s							
	2 mm		760583.20		760586.20	760587.20	760588.20	760589.20
	3 mm		760583.30		760586.30	760587.30	760588.30	760589.30
	4 mm		760583.40		760586.40	760587.40	760588.40	760589.40
	4.6 mm		760583.46	760585.46	760586.46	760587.46	760588.46	760589.46
EC guard columns*			4 x 2 mm:	761987.20	4 x 3 mm:	761987.30		
Preparative VarioPrep	columns							
	10 mm		762210.100			762211.100		762213.100
	21 mm		762210.210		<b>.</b>	762211.210		762213.210
L <u></u>	32 mm		· · · · · · · · · · · · · · · · · · ·		<b>.</b>		·····	762213.320
	40 mm	····					762212.400	762213.400
VP guard columns **			10 x 8 mm:	762234.80	10 x 16 mm	n: 762234.160	15 x 32 mm	: 762236.320
			olumns see below.					

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	





### $NUCLEODUR^{\textcircled{8}}$ $\pi^2$ hydrophobic biphenylpropyl phase $\cdot$ USP L11

### 🔀 Key feature

- Hydrophobic phase with alternative selectivity compared to classical C<sub>18</sub> modifications
- Separation principle based on 2 retention mechanisms ( $\pi$ - $\pi$  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

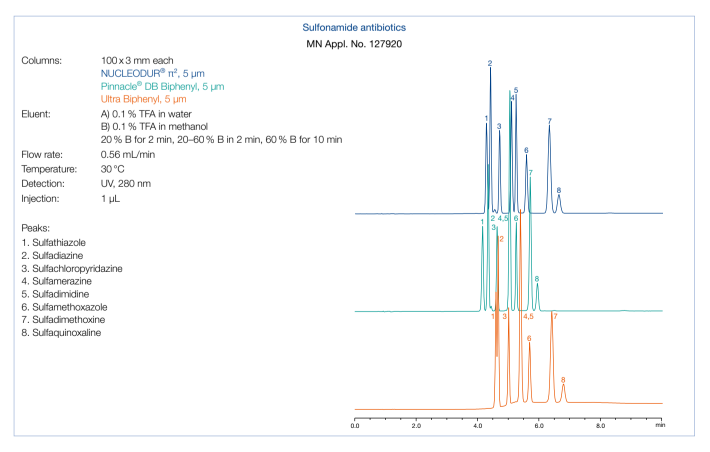
Stationary HPLC phases with biphenyl ligands like NUCLEODUR<sup>®</sup>  $\pi^2$  provide an interesting alternative to classical alkyl modified C<sub>18</sub> and C<sub>8</sub> HPLC phases due to their remarkable orthogonal selectivity.

Furthermore the NUCLEODUR<sup>®</sup>  $\pi^2$  provides an excellent separation performance for aromatic and unsaturated analytes by combination of hydrophobic and  $\pi$ - $\pi$  interactions.

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

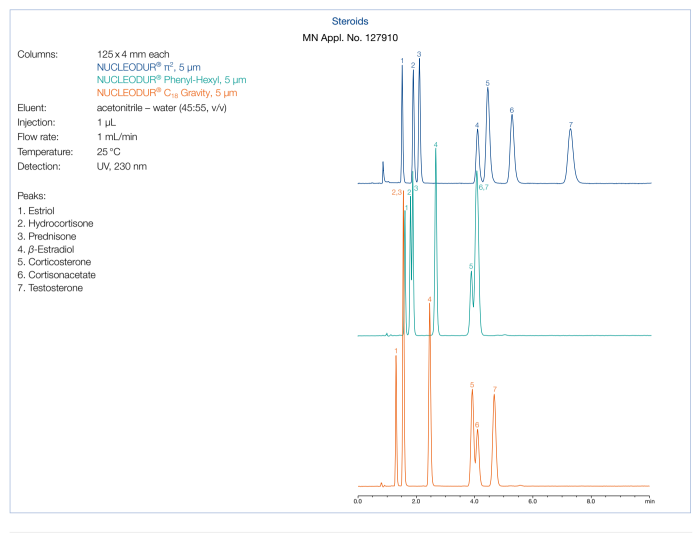
NUCLEODUR<sup>®</sup>  $\pi^2$  shows similar retention strength then C<sub>18</sub> modified phases and thereby displays a significantly stronger retention than phenyl phases. These interactions are even further enhanced in a methanol/water eluent.

NUCLEODUR<sup>®</sup>  $\pi^2$  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<sup>®</sup>  $\pi^2$ . NUCLEODUR<sup>®</sup>  $\pi^2$  is the stationary phase with the highest aromatic analyte selectivity.









Ordering informa	tion						
Eluent in column ace	etonitrile – wate	ər					
	ID	Length $\rightarrow$					
		50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® T	τ <sup>2</sup> , 5 μm part	ticle size 5 µm					
Analytical EC column	S						
	2 mm	760620.20	760621.20	760622.20	760623.20	760624.20	760625.20
	3 mm	760620.30	760621.30	760622.30	760623.30	760624.30	760625.30
	4 mm	760620.40	760621.40	760622.40	760623.40	760624.40	760625.40
	4.6 mm	760620.46	760621.46	760622.46	760623.46	760624.46	760625.46
EC guard columns*		4 x 2 mm: 7	761810.20	4 x 3 mm:	761810.30		
EC columns in packs	of 1, guard co	lumns in packs of 3.					
Guard column sy	/stems						
Guard columns for E	C columns wit	th ID	2 mm	3 mm	4 mm	4.6 mm	Guard column holder
* Column Protection	System (pack c	of) EC	2 4/2 (3)	4/3 (3)	4/3 (3)	4/3 (3)	718966



### NUCLEODUR® PFP hydrophobic pentafluorophenyl phase · USP L43

### 🔀 Key feature

- $\cdot$  Hydrophobic phase with alternative selectivity in comparison to classical  $C_{\rm 18}$  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

### 꾿 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.

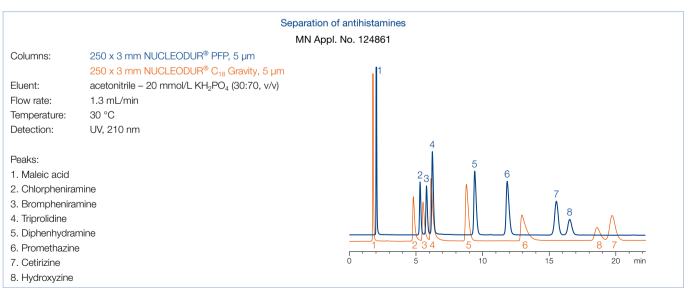
Thus NUCLEODUR<sup>®</sup> PFP offers an excellent selectivity especially for highly polar analytes like aromatic and unsaturated compounds, phenols or halogenated hydrocarbons.

While a typical C<sub>18</sub> phase just provides hydrophobic interactions between stationary phase and analyte NUCLEODUR® PFP offers four different retention mechanisms: polar interactions (H bonds), dipole-dipole,  $\pi$ - $\pi$ , 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.

<code>NUCLEODUR® PFP</code> offers a completely different retention behavior compared to alkyl modified silica and is often used for separations which provide insufficient results on traditional C<sub>18</sub> phases.

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







	Separation of phen	iol isomers
	MN Appl. No. 1	24531
Column:	125 x 4 mm NUCLEODUR® PFP, 5 µm	3 5
Eluent:	125 x 4 mm NUCLEODUR <sup>®</sup> C <sub>18</sub> HTec, 5 μm acetonitrile, 0.1 % formic acid – water, 0.1 % formic acid (35:65, v/v)	4 6
Flow rate:	1 mL/min	
Temperature:	35 °C	
Detection:	UV, 280 nm	
Peaks:		
1. o-Kresol	5. 2,5-Dimethylphenol 9. 3,4-Dichlorophenol	
2. m-Kresol	6. 2,6-Dichlorophenol 10. 2,4-Dibromophenol	
3. 3,4-Dimethyl	phenol 7. 2,3-Dichlorophenol 11. 3,5-Dibromophenol	
4. 3,5-Dimethyl	phenol 8. 2,4-Dichlorophenol	0 2 4 6 8 10 12 14 min

Ordering	information
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### Eluent in column acetonitrile - water

	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR <sup>®</sup> F	PFP, 1.8 μr	n particle size	1.8 µm · UHPLC					
Analytical EC column								
	2 mm	760431.20	760433.20	760435.20	760436.20		760438.20	
	3 mm	760431.30	760433.30	•••••	760436.30	•••••		
	4 mm	760431.40	760433.40		760436.40			
	4.6 mm	760431.46	760433.46		760436.46			
EC guard columns*	S*		4 x 2 mm:	761975.20	4 x 3 mm:	761975.30		
NUCLEODUR® F	PFP, 3 µm	particle size 3 µ	Im					
Analytical EC column	S							
	2 mm		760443.20		760446.20	760447.20	760448.20	760449.20
	3 mm		760443.30		760446.30	760447.30	760448.30	760449.30
	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 <sup>®</sup> F	PFP, 5 µm	particle size 5 µ	Im					
Analytical EC column	S							
	2 mm		760453.20		760456.20	760457.20	760458.20	760459.20
	3 mm	•	760453.30	•••••	760456.30	760457.30	760458.30	760459.30
	4 mm		760453.40		760456.40	760457.40	760458.40	760459.40
	4.6 mm		760453.46	760455.46	760456.46	760457.46	760458.46	760459.46
EC guard columns*			4 x 2 mm:	761977.20	4 x 3 mm:	761977.30		
Preparative VarioPrep	o columns							
	10 mm		762210.100			762211.100		762213.100
	21 mm		762210.210			762211.210		762213.210
L	32 mm							762213.320
	40 mm						762212.400	762213.400
VP guard columns **			10 x 8 mm:	762214.80	10 x 16 mm: 762214.160		15 x 32 mm	: 762216.320

### 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	

### NUCLEODUR<sup>®</sup> Sphinx RP bifunctional RP phase · USP L1 and L11

### Key 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

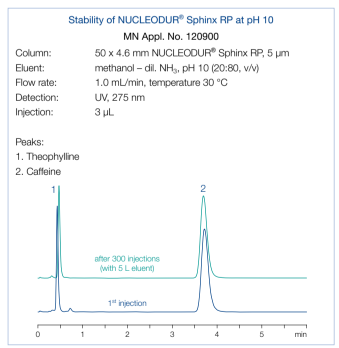
### Recommended application

• Quinolone antibiotics, sulfonamides, xanthines, substituted aromatics

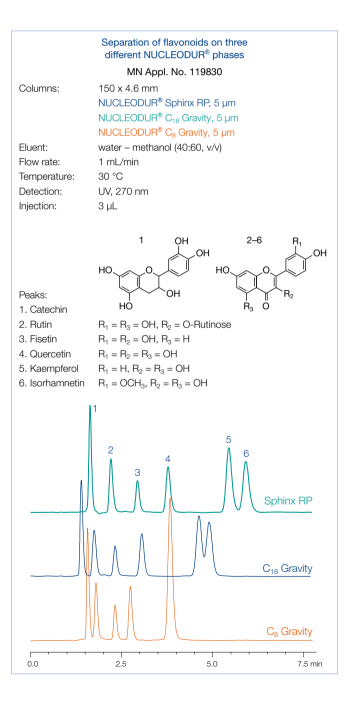
### Alternative RP selectivity

NUCLEODUR<sup>®</sup> 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  $\pi$ - $\pi$  interactions (aromatic ring system) expands the scope of selectivity in comparison with conventional reversed phase packings. NUCLEODUR<sup>®</sup> Sphinx RP is particularly suited for the separation of molecules containing aromatic and multiple bonds.

For the separation of polar compounds NUCLEODUR<sup>®</sup> Sphinx RP can be especially recommended and can also outperform many customary  $C_{18}$  phases. In addition, exhaustive endcapping steps minimize unwanted surface silanol activity and guarantee excellent peak shapes even for strong basic analytes.



Different from standard phenyl phases, NUCLEODUR<sup>®</sup> Sphinx RP is far more stable towards hydrolysis and is also suggested for LC/MS applications. Due to the additional intermolecular interactions NUCLEODUR<sup>®</sup> Sphinx RP is an interesting replenishment to the high density bonded phases NUCLEODUR<sup>®</sup> C<sub>8</sub>/C<sub>18</sub> Gravity and the polar endcapped NUCLEODUR<sup>®</sup> C<sub>18</sub> Pyramid.







### Ordering information

Eluent in column acetonitrile - water

	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR <sup>®</sup> S	Sphinx RP.	1.8 µm partic	cle size 1.8 µm · U	HPLC				
Analytical EC columr	-							
	2 mm	760821.20	760822.20	760825.20	760823.20		760824.20	
	3 mm	760821.30	760822.30		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 mm: 761920.30						
NUCLEODUR <sup>®</sup> S	Sphinx RP,	3 µm particle	size 3 µm					
Analytical EC columr	าร							
	2 mm		760806.20		760812.20	760807.20	760805.20	760808.20
	3 mm		760806.30		760812.30	760807.30	760805.30	760808.30
	4 mm		760806.40		760812.40	760807.40	760805.40	760808.40
	4.6 mm	•	760806.46	760813.46	760812.46	760807.46	760805.46	760808.46
EC guard columns*		•	4 x 2 mm:	761921.20	4 x 3 mm:	761921.30		•
NUCLEODUR <sup>®</sup> S	Sphinx RP,	5 µm particle	size 5 µm					
Analytical EC columr	าร							
	2 mm		760800.20		760809.20	760801.20	760802.20	760803.20
	3 mm	•	760800.30		760809.30	760801.30	760802.30	760803.30
	4 mm		760800.40		760809.40	760801.40	760802.40	760803.40
	4.6 mm		760800.46	760815.46	760809.46	760801.46	760802.46	760803.46
	••••	••••	4 x 2 mm <sup>.</sup>	761922.20	4 x 3 mm <sup>.</sup>	761922.30		
EC guard columns*			174 - 11010		1 / 0 /////			
•	o columns		1721111					
•	o columns 10 mm		762372.100			762375.100		762373.100
						762375.100 762375.210		
	10 mm		762372.100					762373.210
•	10 mm 21 mm		762372.100				762371.400	762373.210 762373.320
EC guard columns* Preparative VarioPrep	10 mm 21 mm 32 mm 40 mm		762372.100					762373.100 762373.210 762373.320 762373.400 762392.320

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	

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

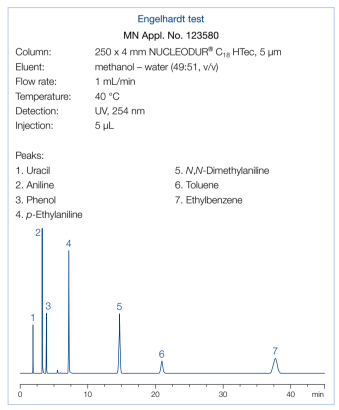
### 📩 Key feature

- 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

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<sup>®</sup> C<sub>18</sub> HTec scores in low bleed characteristics and is therefore highly suitable for LC/MS.



### 🔼 Technical data

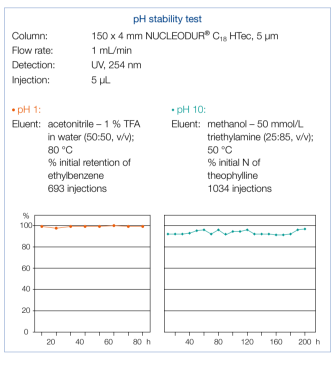
• High density octadecyl modification (C<sub>18</sub>); pore size 110 Å; particle sizes 1.8  $\mu$ m, 3  $\mu$ m, 5  $\mu$ m, 7  $\mu$ m and 10  $\mu$ m for analytical and preparative separations; carbon content 18 %, pH stability 1–11

### 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

### Stability and lifetime

Based on fully synthetic and extremely robust totally spherical NUCLEODUR<sup>®</sup> silica, NUCLEODUR<sup>®</sup> C<sub>18</sub> 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<sup>®</sup> C<sub>18</sub> HTec columns show a remarkably long lifetime in acidic (pH 1) as well as basic (pH 10) mobile phases.



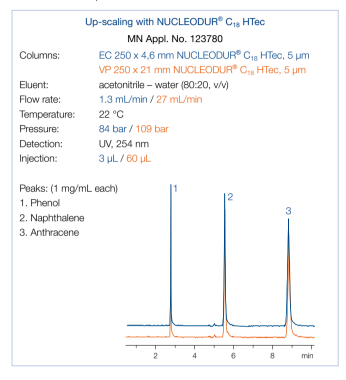
Due to innovative surface coating procedures NUCLEODUR<sup>®</sup> C<sub>18</sub> HTec offers excellent analytical separation properties and is the first choice for up-scaling to preparative column dimensions.





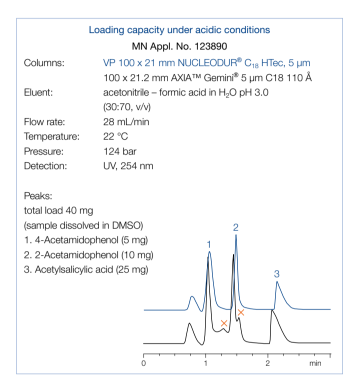
### Up-scaling

Due to highest quality standards in silica production and phase chemistry combined with optimized packing technology, NUCLEODUR<sup>®</sup> C<sub>18</sub> HTec allows exceptional transferability from analytical to preparative scale with respect to different particle sizes (e.g., 5, 7 or 10  $\mu$ m) 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<sup>®</sup>  $C_{18}$  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).



<u> </u>	
Ordering	information

Eluent in column acetonitrile - water

		L						
	ID	Length → 30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® C	C <sub>18</sub> HTec, <sup>-</sup>	1.8 µm particle	e size 1.8 µm · U⊦	IPLC				
Analytical EC column	s							
	2 mm	760301.20	760305.20	760304.20	760306.20		760308.20	
<u> </u>	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 <sup>®</sup> C	C <sub>18</sub> HTec, 3	3 µm particle s	ize 3 µm					
Analytical EC column								
	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		



### Ordering information

### Eluent in column acetonitrile - water

	ID	30 mm	50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
				7511111	100 11111	125 11111	130 mm	230 11111
NUCLEODUR <sup>®</sup> C		5 µm particles	size 5 µm					
Analytical EC column								
	2 mm	····•	760311.20		760313.20	760314.20	760315.20	760316.20
	3 mm		760311.30	<u>-</u>	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
EC guard columns*		4 x 2 mm:	761927.20	4 x 3 mm:	761927.30			
Preparative VarioPrep								
	10 mm		762551.100	·····	·····	762554.100	····•	762556.100
	21 mm		762551.210	·····	762553.210	762554.210		762556.210
P	32 mm	<u>.</u>	·····	·····	762553.320	<b>.</b>	762555.320	762556.320
	40 mm						762555.400	762556.400
	50 mm		·····		762553.500	····•	762555.500	762556.500
VP guard columns **		·····	10 x 8 mm:	·····	· · · · · · · · · · · · · · · · · · ·	762591.160	····•	·····
			15 00	760500 000	15 x 50 mm	762592.500		
-			15 x 32 mm:	102092.320	15 x 50 mm	102032.000		
		7 µm particle s		702092.020	13 x 30 mm	102392.000		
	columns	7 µm particle s	size 7 µm	102392.320	13 X 30 1111			
	columns 10 mm	7 μm particle s	size 7 μm 762561.100	102392.320		762564.100		
	o columns 10 mm 21 mm	7 µm particles	size 7 µm	102392.320	762563.210			762566.210
	10 columns 10 mm 21 mm 32 mm	7 μm particle s	size 7 μm 762561.100	102392.320		762564.100	762565.320	762566.210 762566.320
	columns           10 mm           21 mm           32 mm           40 mm	7 μm particle s	size 7 μm 762561.100	102392.320	762563.210 762563.320	762564.100	762565.400	762566.210 762566.320 762566.400
Preparative VarioPrep	10 columns 10 mm 21 mm 32 mm	7 μm particles	size 7 μm 762561.100 762561.210		762563.210 762563.320 762563.500	762564.100 762564.210		762566.210 762566.320 762566.400
Preparative VarioPrep	columns           10 mm           21 mm           32 mm           40 mm	7 µm particles	size 7 μm 762561.100 762561.210 10 x 8 mm:	762591.80	762563.210 762563.320 762563.500 10 x 16 mm	762564.100 762564.210 762591.160	762565.400	762566.210 762566.320 762566.400
Preparative VarioPrep	columns           10 mm           21 mm           32 mm           40 mm           50 mm		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.210 762563.320 762563.500 10 x 16 mm	762564.100 762564.210	762565.400	762566.100 762566.210 762566.320 762566.400 762566.500
Preparative VarioPrep	columns           10 mm           21 mm           32 mm           40 mm           50 mm		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.210 762563.320 762563.500 10 x 16 mm	762564.100 762564.210 762591.160	762565.400	762566.210 762566.320 762566.400
Preparative VarioPrep	2 columns 10 mm 21 mm 32 mm 40 mm 50 mm 20 18 HTec,		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.210 762563.320 762563.500 10 x 16 mm	762564.100 762564.210 762591.160	762565.400	762566.210 762566.320 762566.400
Preparative VarioPrep	2 columns 10 mm 21 mm 32 mm 40 mm 50 mm 20 18 HTec,		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm:	762591.80	762563.210 762563.320 762563.500 10 x 16 mm	762564.100 762564.210 762591.160	762565.400	762566.210 762566.320 762566.400 762566.500
Preparative VarioPrep	2 columns 10 mm 21 mm 32 mm 40 mm 50 mm 50 mm C <sub>18</sub> HTec, o columns		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm	762591.80	762563.210 762563.320 762563.500 10 x 16 mm	762564.100 762564.210 762591.160 762592.500	762565.400	762566.210 762566.320 762566.400 762566.500 762566.500
Preparative VarioPrep	2 columns 10 mm 21 mm 32 mm 40 mm 50 mm C <sub>18</sub> HTec, o columns 10 mm		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm 762571.100	762591.80	762563.210 762563.320 762563.500 10 x 16 mm 15 x 50 mm	762564.100 762564.210 762591.160 762592.500 762574.100	762565.400	762566.210 762566.320 762566.400 762566.500 762566.500 762576.100 762576.210
Preparative VarioPrep	2 columns 10 mm 21 mm 32 mm 40 mm 50 mm 50 mm 218 HTec, 0 columns 10 mm 21 mm		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm 762571.100	762591.80	762563.210 762563.320 762563.500 10 x 16 mm 15 x 50 mm 762573.210	762564.100 762564.210 762591.160 762592.500 762574.100	762565.400 762565.500	762566.210 762566.320 762566.400 762566.500 762566.500 762576.100 762576.210 762576.210
Preparative VarioPrep	10 mm           10 mm           21 mm           32 mm           40 mm           50 mm		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm 762571.100	762591.80	762563.210 762563.320 762563.500 10 x 16 mm 15 x 50 mm 762573.210	762564.100 762564.210 762591.160 762592.500 762574.100	762565.400 762565.500 762575.320	762566.210 762566.320 762566.400 762566.500 762566.500 762576.100 762576.210 762576.210 762576.320 762576.400
NUCLEODUR <sup>®</sup> C Preparative VarioPrep VP guard columns ** NUCLEODUR <sup>®</sup> C Preparative VarioPrep	columns           10 mm           21 mm           32 mm           40 mm           50 mm   C18 HTec, columns 10 mm 21 mm 32 mm 40 mm		size 7 μm 762561.100 762561.210 10 x 8 mm: 15 x 32 mm: size 10 μm 762571.100	762591.80 762592.320	762563.210 762563.320 762563.500 10 x 16 mm 15 x 50 mm 762573.210 762573.210 762573.320 762573.500	762564.100 762564.210 762591.160 762592.500 762574.100	762565.400 762565.500 762575.320 762575.320 762575.400	762566.210 762566.320 762566.400

### 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® C<sub>18</sub> HTec bulk material in 7 and 10 µm for self-packing of preparative columns see page 256.





### $NUCLEODUR^{\tiny{(B)}} C_{18} ec \cdot C_8 ec \cdot C_4 ec \text{ nonpolar phases for routine analysis} \cdot \text{USP L1 } (C_{18}) \cdot \text{L7 } (C_8) \cdot \text{L26 } (C_4) + (C_{18}) \cdot \text{L26 } ($

### 📩 Key feature

- Ideal and reliable standard RP phase for daily routine analysis and up-scaling for preparative HPLC
- Medium density Octadecyl (C<sub>18</sub>) and octyl (C<sub>8</sub>) with pore size of 110 Å with exhaustive endcapping for a wide range of applications
- $\cdot$  Octadecyl (C\_{18}) and butyl (C\_4) with pore size of 300 Å for the separation of biomolecules

### 🔼 Technical data

- Pore size 110 Å: particle sizes 3 μm and 5 μm, 7 μm, 10 μm, 12 μm, 16 μm, 20 μm, 30 μm
- 10  $\mu$ m, 12  $\mu$ m, 16  $\mu$ m, 20  $\mu$ m, 30  $\mu$ m and 50  $\mu$ m for preparative separations; carbon content 17.5 % for C<sub>18</sub>, 10.5 % for C<sub>8</sub>; 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

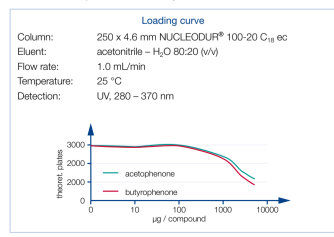
 300 Å: biomolecular macromolecules, like proteins and peptides

### 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<sub>18</sub> ec is available in 9 different particle sizes (3, 5, 7, 10, 12, 16, 20, 30 and 50  $\mu$ m) which cover the whole range from high speed analytical HPLC up to medium and low pressure prep LC. NUCLEODUR® C<sub>18</sub> 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<sup>®</sup> 100-20 C<sub>18</sub> ec column describes the correlation between the increase of column loading and the decrease of separation efficiency.



Separation of theophylline and caffeine at pH 10 30 x 3 mm NUCLEODUR® 100-5 C<sub>18</sub> ec Column: methanol - ag. NH<sub>3</sub> Eluent: (20:80, v/v), pH 10 Flow rate: 0.5 mL/min Temperature: 25 °C Detection: UV, 254 nm 1000 Ini. 750 Ini. 4 500 Inj. 3 250 Inj. Start

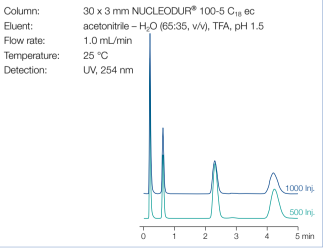
pH stability of NUCLEODUR® C18 ec

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

ő

2

3 min



181



### Chemical stability

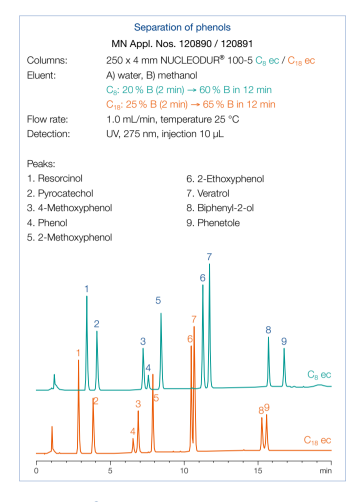
The utmost purity of the base silica and the exceptional silane bonding chemistry minimize the risk of dissolution, or hydrolysis at pH extremes.

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

### NUCLEODUR<sup>®</sup> octyl phases

In addition to NUCLEODUR<sup>®</sup> C<sub>18</sub> phases MACHEREY-NAGEL offers octyl modified NUCLEODUR<sup>®</sup> C<sub>8</sub> Gravity and NUCLEODUR<sup>®</sup> C<sub>8</sub> ec columns to expand the RP tool box. Based on the same spherical high purity silica the C<sub>8</sub> phases exhibit the same chemical and mechanical stability as the C<sub>18</sub> counterparts. Indeed NUCLEODUR<sup>®</sup> C<sub>8</sub> 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<sub>18</sub> phases). NUCLEODUR<sup>®</sup> C<sub>8</sub> ec and NUCLEODUR<sup>®</sup> C<sub>8</sub> 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<sup>®</sup>  $C_8$  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<sup>®</sup> phases for biochromatography

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

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

· Octyl phases (C<sub>8</sub>) show superior polar selectivity.

- $\cdot$  Octadecyl phases (C<sub>18</sub>) show superior hydrophobic selectivity.
- $\cdot$  Hydrophobic compounds show shorter retention times on C<sub>8</sub> phases.

Ordering informa		er					
	ID	Length → 50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR® 1	00-3 C <sub>18</sub> ec				120 1111		200 1111
Analytical EC column	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	





#### Ordering information

	ID	Length →					
		50 mm	75 mm	100 mm	125 mm	150 mm	250 mm
NUCLEODUR®	100-5 C <sub>18</sub> ec	coctadecyl phase, j	particle size 5 µm,	17.5 % C			
Analytical EC colum	ns						
	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: 7	761932.20	4 x 3 mm: 7	61932.30	
Preparative VarioPre	p 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
/P guard columns *	*		10 x 8 mm: 7	762090.80	10 x 16 mm:	762090.160	
-	••••••		15 x 32 mm: 7	· · · · · · · · · · · · · · · · · · ·	15 x 50 mm:	· · · · · · · · · · · · · · · · · · ·	
NUCLEODUR®	100-10 C₁₀ e	C octadecyl phase	, particle size 10 ur	n, 17.5 % C			
Preparative VarioPre							
	10 mm	762011.100			762302.100		762010.100
	21 mm	762011.210	•••••		762302.210		762010.210
	32 mm						762010.320
L	40 mm	•••••				762303.400	762010.400
	50 mm	•••••	·····	·····	·····	102000.400	762010.500
/P guard columns *	*		10 x 8 mm: 7	762000 80	10 x 16 mm:	762090 160	102010.000
Vi guara columna			15 x 32 mm: 1				
			13 × 32 11111.	02311.320	15 x 50 mm:	762311.500	
			10 × 02 11111.	02311.320	15 X 50 mm:	762311.500	
Ordering inform	ation		13 x 32 11111.	02311.320	15 x 50 mm:	762311.500	
-		ter	13 x 32 1111.	02311.320	15 X 50 mm:	762311.500	
-		ter Length →	10 x 02 mm.	02311.320	15 X 50 mm:	762311.500	
-	cetonitrile – wat		75 mm	100 mm	125 mm	150 mm	250 mm
Eluent in column ad	cetonitrile – wat ID	Length →	75 mm	100 mm			250 mm
Eluent in column ad	ID 100-3 C <sub>8</sub> ec	Length → 50 mm	75 mm	100 mm			250 mm
Eluent in column ad	ID 100-3 C <sub>8</sub> ec	Length → 50 mm	75 mm	100 mm			250 mm 760062.20
Eluent in column ad	cetonitrile – wat ID 100-3 C <sub>8</sub> ec ns	Length → 50 mm octyl phase, particl	75 mm	100 mm 6 C	125 mm		
Eluent in column ad	Letonitrile – wat ID 100-3 C <sub>8</sub> ec ns 2 mm	Length → 50 mm octyl phase, particl 760063.20	75 mm	100 mm 6 C 760059.20	<b>125 mm</b> 760060.20		760062.20
Eluent in column ad	tetonitrile – wat ID 100-3 C <sub>8</sub> ec ns 2 mm 3 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30	75 mm	100 mm 6 C 760059.20 760059.30	<b>125 mm</b> 760060.20 760060.30		760062.20 760062.30
Analytical EC colum	2 mm 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 %	100 mm 6 C 760059.20 760059.30 760059.40 760059.46	<b>125 mm</b> 760060.20 760060.30 760060.40	150 mm 760061.46	760062.20 760062.30 760062.40
Eluent in column ad NUCLEODUR® Analytical EC colum	cetonitrile – wat ID 100-3 C <sub>8</sub> 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 % 760064.46 4 x 2 mm: 1	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20	125 mm 760060.20 760060.30 760060.40 760060.46	150 mm 760061.46	760062.20 760062.30 760062.40
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR®	2 mm 2 mm 3 mm 4 mm 4.6 mm 100-5 C <sub>8</sub> ec	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm: 1	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20	125 mm 760060.20 760060.30 760060.40 760060.46	150 mm 760061.46	760062.20 760062.30 760062.40
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR®	2 mm 3 mm 4 mm 4.6 mm 100-5 C <sub>8</sub> 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 % 760064.46 4 x 2 mm: 1	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C	<b>125 mm</b> 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
Eluent in column ac NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR®	2 mm 3 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 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	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm: 1	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20	<b>125 mm</b> 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20	150 mm 760061.46	760062.20 760062.30 760062.40 760062.46 760703.20
Analytical EC columns*	2 mm 3 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 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	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm: 1	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30	125 mm 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30	150 mm 760061.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.20
Analytical EC columns*	2 mm 3 mm 4 mm 100-5 C <sub>8</sub> ec ns 2 mm 3 mm 4 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 2 mm 3 mm 4 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 % 760064.46 4 x 2 mm; e size 5 μm, 10.5 %	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40	125 mm 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40	150 mm 760061.46 61936.30	760062.20 760062.30 760062.40 760062.46 760703.20 760703.20 760703.30 760703.40
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum	2 mm 3 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 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	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm; 1 e size 5 μm, 10.5 % 760706.46	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46	125 mm 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.20 760701.30 760701.40 760701.46	150 mm 760061.46 61936.30 760702.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.20
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC columns EC guard columns*	2 mm 3 mm 4 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 2 mm 3 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 2 mm 3 mm 4 mm 4 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 % 760064.46 4 x 2 mm; e size 5 μm, 10.5 %	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46	125 mm 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40	150 mm 760061.46 61936.30 760702.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.20 760703.30 760703.40
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC columns EC guard columns*	2 mm 3 mm 4 mm 4.6 mm 100-5 C <sub>8</sub> ec ns 2 mm 3 mm 4.6 mm 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 760700.46	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm; 1 e size 5 μm, 10.5 % 760706.46	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46	125 mm 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	150 mm 760061.46 61936.30 760702.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum EC guard columns*	Cetonitrile – wat ID           100-3 C <sub>8</sub> ec ns           2 mm           3 mm           4 mm           4.6 mm           100-5 C <sub>8</sub> ec ns           2 mm           3 mm           4 mm           4.6 mm           9 columns           10 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.20 760700.40 760700.40 760700.40	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm; 1 e size 5 μm, 10.5 % 760706.46	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46	125 mm 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.40 760701.46 4 x 3 mm: 7 762061.100	150 mm 760061.46 61936.30 760702.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46 762062.100
Eluent in column ad NUCLEODUR® Analytical EC colum EC guard columns* NUCLEODUR® Analytical EC colum EC guard columns*	Cetonitrile – wat ID           100-3 C <sub>8</sub> ec ns           2 mm           3 mm           4 mm           4.6 mm           100-5 C <sub>8</sub> ec ns           2 mm           3 mm           4 mm           4.6 mm           9 columns           10 mm           21 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 % 760064.46 4 x 2 mm; 1 e size 5 μm, 10.5 % 760706.46	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46	125 mm 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	150 mm 760061.46 61936.30 760702.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46 762062.100 762062.210
Eluent in column ad NUCLEODUR® Analytical EC colum	Cetonitrile – wat ID           100-3 C <sub>8</sub> ec ns           2 mm           3 mm           4 mm           4.6 mm           100-5 C <sub>8</sub> ec ns           2 mm           3 mm           4 mm           4.6 mm           9 columns           10 mm	Length → 50 mm octyl phase, particl 760063.20 760063.30 760063.40 760063.46 octyl phase, particl 760700.20 760700.20 760700.40 760700.40 760700.40	75 mm e size 3 μm, 10.5 % 760064.46 4 x 2 mm; 1 e size 5 μm, 10.5 % 760706.46	100 mm 6 C 760059.20 760059.30 760059.40 760059.46 761936.20 6 C 760704.20 760704.30 760704.40 760704.46	125 mm 760060.20 760060.30 760060.40 760060.46 4 x 3 mm: 7 760701.20 760701.30 760701.40 760701.40 760701.46 4 x 3 mm: 7 762061.100	150 mm 760061.46 61936.30 760702.46	760062.20 760062.30 760062.40 760062.46 760703.20 760703.30 760703.40 760703.46 762062.100

EC and VarioPrep columns in packs of 1, guard columns see previous NUCLEODUR® phases.

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

NUCLEODUR® C<sub>18</sub> ec bulk material with 10–50 µm for self-packing of preparative columns see page 256.

The ordering information for C<sub>18</sub> and C<sub>4</sub> modified 300 Å NUCLEODUR<sup>®</sup> 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.



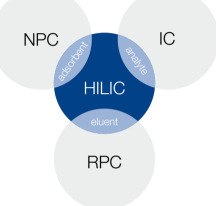


### NUCLEODUR® HILIC zwitterionic phase

### 📩 Key feature

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

### 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<sub>2</sub>, 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."

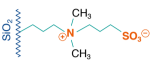
NUCLEODUR<sup>®</sup> HILIC is a special zwitterionic modified stationary phase based on ultra spherical NUCLEODUR<sup>®</sup> particles. The betaine character of the ammoniumsulfonic acid ligands results in total charge equalization and in an overall neutrally charged but highly polar surface

### 😕 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

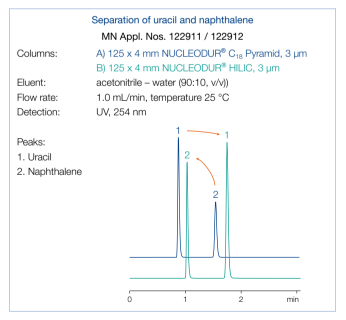
### Recommended application

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



### 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.



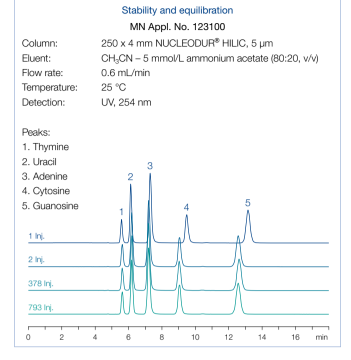


### Stability features

Due to an advanced and unique surface modification procedure (pat. pend.) NUCLEODUR<sup>®</sup> HILIC columns provide short equilibration times – after just 20 min equilibration already the 2nd injection shows stable and reproducible results.

Beyond this, NUCLEODUR<sup>®</sup> 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<sup>®</sup> HILIC is absolutely suitable for preparative and semi-preparative applications.

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



### Ordering information

Eluent in column ace	etonitrile – w ID	tater (80:20, v/v) Length → 30 mm	50 mm	75 mm		100 mm	125 mm	150 mm	250 mm
NUCLEODUR <sup>®</sup> H	III IC. 1.8					100 1111	125 1111	100 11111	230 mm
Analytical EC column				20					
	2 mm	760521.20	760523.2	20 760525	.20	760526.20		760528.2	0
	3 mm	760521.30	760523.3	30		760526.30	••••		
	4 mm	760521.40	760523.4	10		760526.40	•••••••••••••••••••••••••••••••••••••••		••••••
	4.6 mm	760521.46	760523.4	16		760526.46	•••••••••••••••••••••••••••••••••••••••	·····	••••••
EC guard columns*	•••••	•••	4 x 2	mm: 761960.20		4 x 3 mm: 7	61960.30		
NUCLEODUR <sup>®</sup> H	IILIC, 3 µr	n particle size 3	3 µm						
Analytical EC column									
,	2 mm		760532.2	20		760534.20	760531.20	760533.2	0 760530.20
	3 mm	••••	760532.3	30		760534.30	760531.30	760533.3	0 760530.30
	4 mm	•	760532.4	10		760534.40	760531.40	760533.4	0 760530.40
	4.6 mm	•	760532.4	16		760534.46	760531.46	6 760533.4	6 760530.46
EC guard columns*	•••••		4 x 2	mm: 761961.20		4 x 3 mm: 7	61961.30		•
NUCLEODUR® H	IILIC. 5 ur	n particle size 5	5 um						
Analytical EC column	-								
	2 mm		760552.2	20		760554.20	760551.20	760553.2	0 760550.20
	3 mm		760552.3	30		760554.30	760551.30	) 760553.3	0 760550.30
	4 mm		760552.4	10		760554.40	760551.40	) 760553.4	0 760550.40
	4.6 mm	••••	760552.4	16		760554.46	760551.46	6 760553.4	6 760550.46
EC guard columns*	•••••	••••	4 x 2	mm: 761962.20		4 x 3 mm: 7	61962.30		
Guard column sy	vstem								
Guard columns for E		with ID		2 mm	3 mm	4 mm		4.6 mm	Guard column holde
* Column Protection :	System (pac	k of)	EC	4/2 (3)	4/3 (3)	4/3 (3	)	4/3 (3)	718966

### NUCLEODUR<sup>®</sup> CN/CN-RP cyano-modified high purity silica phase · USP L10

### 📩 Key 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 C\_8 and C\_{18}

### Recommended application

Tricyclic antidepressants, steroids, organic acids

### Alternative bonded-phase functionality

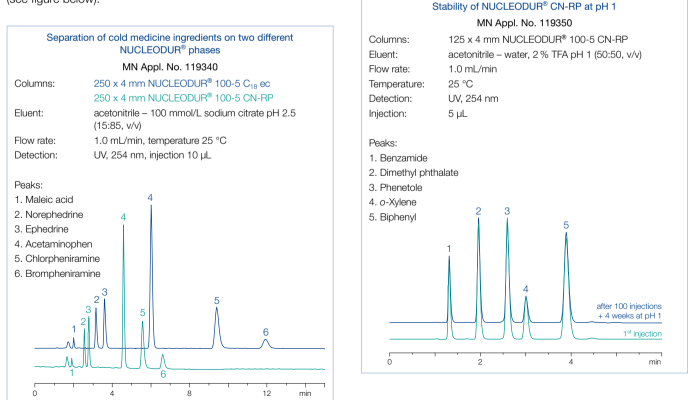
In reversed phase HPLC it is fairly common to start with C<sub>18</sub> or C<sub>8</sub> columns, if new methods have to be developed. However, superior polarity and selectivity properties often required for more sophisticated separations, are not always sufficiently provided by classical RP phases, 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<sup>®</sup> 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).

The polarity of NUCLEODUR<sup>®</sup> 100-5 CN-RP can be classified as intermediate based on multiple retention mechanisms such as dipole-dipole,  $\pi$ - $\pi$ , and also hydrophobic interactions [8]. Therefore, this phase shows a distinct selectivity for polar organic compounds as well as for molecules containing  $\pi$  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)

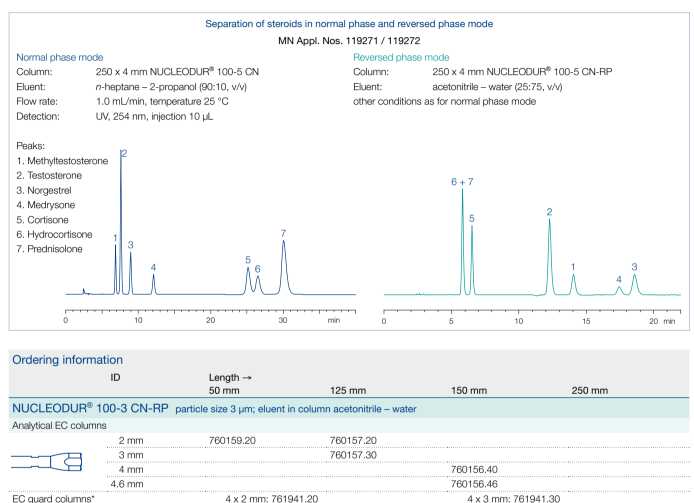






### Multi-mode columns

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.



EC guard columns\*

NUCLEODUR <sup>®</sup> 1	00-5 CN-RP particle	size 5 µm; e	luent in colur	nn acetonitrile – wat	er		
Analytical EC column	IS						
	4 mm			760153.40			760152.40
	4.6 mm			760153.46	760154.	46	760152.46
EC guard columns* 4 x 3 mm: 761944.30							
NUCLEODUR <sup>®</sup> 1	00-5 CN particle size	5 µm; eluen	t in column <i>n</i>	-heptane			
Analytical EC column	IS						
	4 mm		760151.40		760149.40		760150.40
	4.6 mm		760151.46		760149.46		760150.46
EC guard columns* 4 x 3 mm: 761943.30							
EC columns in packs	s of 1, guard columns in p	acks of 3.					
Guard column sy	/stem						
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

### NUCLEODUR<sup>®</sup> NH<sub>2</sub>/NH<sub>2</sub>-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

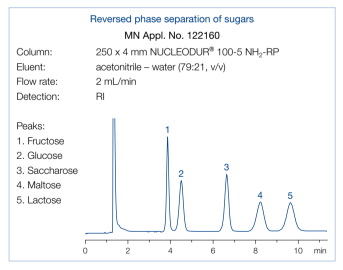
### Recommended application

- 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.

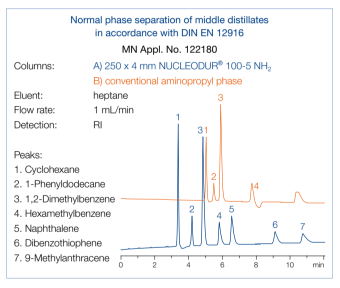
### Multi-mode columns

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<sup>®</sup> NH<sub>2</sub>, 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<sup>®</sup>  $NH_2$  is the separation of simple and complex sugars, sugar alcohols and other hydroxy compounds under RP conditions as well as hydrocarbons under NP conditions.

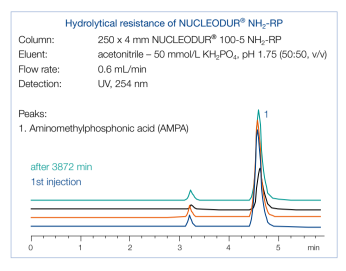


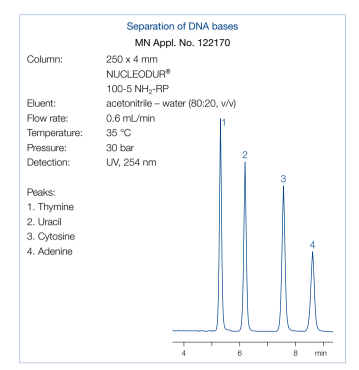
Due to the special method of surface modification NUCLEODUR<sup>®</sup> NH<sub>2</sub> features a pronounced stability at higher as well as at lower pH values. The following figure shows, that even after several days of exposure of the column material at pH 1.75 good separation efficiency and peak symmetry are maintained. The resulting high column life allows cost reduction due to lower column consumption.

This example shows the enhanced pH stability of NUCLEODUR<sup>®</sup> NH<sub>2</sub> 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*.









Based on superspherical NUCLEODUR<sup>®</sup> 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<sup>®</sup> NH<sub>2</sub> enables reliable analyses especially for routine work.

Ordering informa		L e ve exte							
	ID	Length → 100 mm		125 mm	150 mm		250 mm		
NUCLEODUR <sup>®</sup>	100-3 NH <sub>2</sub> -RP	particle size 3 µm; e	eluent in col	umn acetonitrile – wa	iter				
Analytical EC colum									
	2 mm	760740.20		760741.20					
	4.6 mm				760742.4	6	760739.46		
EC guard columns*		4 x 2 n	4 x 2 mm: 761951.20			4 x 3 mm: 761951.30			
NUCLEODUR <sup>®</sup>	100-5 NH <sub>2</sub> -RP	particle size 5 µm; e	eluent in col	umn acetonitrile – wa	iter				
Analytical EC colum	าร								
	2 mm			760730.20			760732.20		
	3 mm			760730.30			760732.30		
	4 mm			760730.40			760732.40		
	4.6 mm			760730.46	760731.4	6	760732.46		
EC guard columns*	4 x 2 n	4 x 2 mm: 761953.20			4 x 3 mm: 761953.30				
NUCLEODUR <sup>®</sup>	100-5 NH <sub>2</sub> partic	cle size 5 µm; eluen	t in column	<i>n</i> -heptane					
Analytical EC colum									
	4 mm			760720.40			760722.40		
	4.6 mm	•		760720.46	760721.4	6	760722.46		
EC guard columns*	iard columns*				4 x 3 mm: 761952.30				
EC columns in pack	s of 1, guard colum	ns in packs of 3.							
Guard column s	ystem								
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		



### NUCLEODUR<sup>®</sup> SiOH unmodified silica for normal phase · USP L3

### Key feature

- $\cdot$  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)</li>

### Recommended application

 Polar and midpolar compounds under normal phase conditions

Ordering informa	ation							
Eluent in column n-l	neptane							
	ID	Length →						
		50 mm		125 mm	150 mm		250 mm	
NUCLEODUR <sup>®</sup> 1	00-3 particle size 3	μm						
Analytical EC column	าร							
	4.6 mm	760170.46			760172.46	3	760173.46	
EC guard columns*					4 x 3 mm: 761966.30			
NUCLEODUR <sup>®</sup> 1	00-5 particle size 5	μm						
Analytical EC column	IS							
	4 mm						760007.40	
	4.6 mm	760023.46			760012.46	3	760007.46	
EC guard columns* 4 x 3 mm: 761967.30						30		
Preparative VarioPrep	o columns							
	10 mm	762077.100	-	762078.100			762007.100	
	21 mm	762077.210	-	762078.210			762007.210	
	40 mm				762075.40	00	762007.400	
VP guard columns *	VP guard columns * 10 x 8 mm: 762094.80			C	10 x 16 mm: 762094.160			
		15 x 32	15 x 32 mm: 762330.320					
EC and VarioPrep co	lumns in packs of 1, g	uard columns se	e below.					
Guard column sy	ystems							
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 ho		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.



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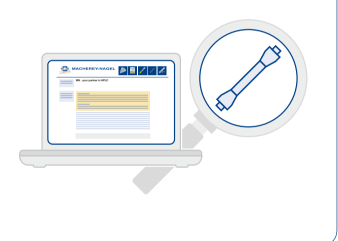
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