

Thermo Scientific Hypersil BDS Columns



Robust and rugged, delivering excellent reproducibility and peak symmetry



# Thermo Scientific Hypersil BDS Columns

When you are scaling the peaks in chromatography, every piece of equipment is critical. Since their introduction in 1989, Thermo Scientific Hypersil BDS columns have gained a reputation as one of the most robust, reproducible and reliable HPLC columns available. Based on highly base deactivated silica with endcapping, Hypersil™ BDS columns exhibit the following key features, separation after separation:

- Excellent reproducibility
- Reduced tailing
- Very robust and rugged with long column lifetimes
- Excellent peak symmetry for basic and acid compounds

This Technical Guide provides an overview of the Hypersil BDS product range and how quality and reproducibility of the phases and columns are monitored.

#### Base Deactivated Silica and Bonded Phases

The use of covalently-bonded silica stationary phases in HPLC allows the analysis of a broad range of analytes. Along with rapid equilibration times, and significantly improved mass transfer characteristics over liquid-liquid partition chromatography, this has resulted in the hugely successful advancement of HPLC as a modern day analytical technique. However, covalently-bonded silica stationary phases often have specific limitations.

Many chemical properties associated with derivatized silicas used in HPLC have a strong effect on analyte interactions. These properties are specific to either the derivatized ligand itself, or the remaining underivatized silanol groups on the silica surface. In particular, the number and acidity of these remaining silanol groups is of significance. It is the silanol groups that are responsible for the acid-base properties of the base silica, contributing to the overall polarity of the surface even when the surface method is derivatized. Their type and acidity play an important role in determining resolution and peak shape for various classes of compounds being analyzed.

Peak tailing and low efficiencies of both basic and acidic compounds can occur due to unwanted silanol interactions. The effect is most apparent with some of the earlier silicas developed for HPLC, in which silanol groups are quite acidic.

For these types of silicas, the observed effect on peak shape requires the mobile phase to include either a competing base such as triethylamine or a competing acid such as acetic acid. Both peak shape and column performance are improved dramatically when the appropriate competing agent is used. The effect of the additive is to compete with any silanol interactions that interfere with analyte retention and peak shape. Consequently, the additive must be present in fairly high concentrations, often as much as 1% volume fraction of the mobile phase. The use of additives in such a concentration can often have a deleterious effect on the column lifetime and also on reproducibility of the method.



These difficulties have provided the impetus for the development of an improved range of chromatographic silicas (base deactivated silicas) that allow the analysis of both basic and acidic compounds without the requirement for competing additives in the mobile phase. A proprietary treatment to the silica surface results in significant improvements to the homogeneity of the surface silanol population prior to derivatization. The result has been that the bonded silica surface no longer requires a competing acid or base in the mobile phase to achieve acceptable peak shapes for problematic analytes. This is illustrated in Figure 1.

Hypersil BDS packings were among the first base deactivated silica HPLC packings to offer the characteristics associated with these surface improvements, offering benefits such as:

- Reduced silanol interactions
- · Reduced peak tailing
- Reduced need for mobile phase additives
- Excellent peak symmetry
- Long column lifetimes
- Improved performance with basic, neutral and acidic compounds

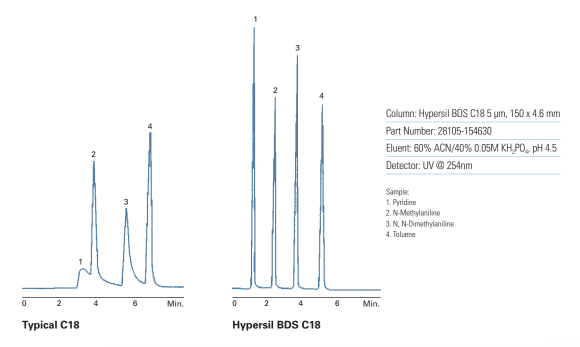
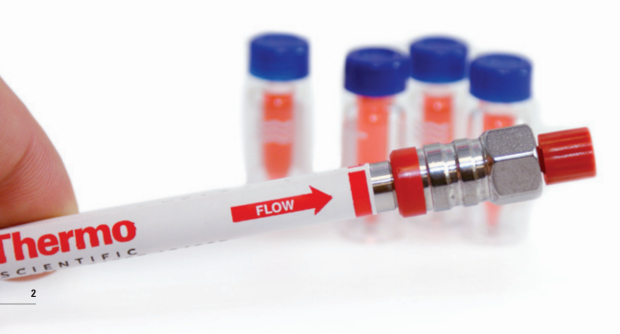


Figure 1: Hypersil BDS shows improved peak symmetry over traditional ODS for difficult analytes



#### The Base Deactivation Process

The popularity of columns packed with C18 derivatized silica is due to their wide breadth of application, encompassing non-polar and neutral, acidic, and basic analytes. The selectivity of a given C18 phase can depend on the type of silane used and the synthetic conditions, as both of these factors will affect the density of the bonded phase on the surface.

This density of the bonded phase is important since the greater the access of an analyte to the underlying silica support, the greater the opportunity for secondary interactions such as hydrogen bonding. There are approximately five silanol (Si-OH) groups per nm² of surface on the silica, corresponding to 8-9 mmol/m². It is stereochemically impossible to react more than ~50% of the silanol groups even with ligands as small as trimethylsilane (C1).

The surface composition of silica prior to derivatization is very important. As illustrated in Figure 2a, at most silica surfaces it is usual to have a variety of silanol groups: (1) lone silanols, (2) siloxanes, (3) geminal silanols and vicinal silanols. The presence of these silanols in a derivatized silica can result in unwanted silanol interactions with the analytes which can give rise to peak tailing and changes in retention and selectivity on a typical alkyl C18 packing, such as:

- Si -OH •••••NH<sub>2</sub> -R (Hydrogen bonding with base)
- Si OH••••••0=RCOH (Hydrogen bonding with acid)
- Si O- +NH<sub>3</sub>-R (Ion exchange with base)

Figure 2a

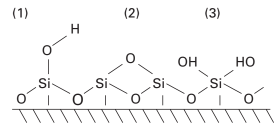
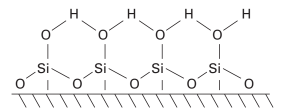


Figure 2: The base deactivation process produces a homogenous surface

#### Figure 2b





Using a proprietary production process, the surface of the Hypersil BDS silica is made much more homogeneous, so that all silanols are of the same type (vicinal silanols), as illustrated in Figure 2b (previous page). The resultant silica surface is more uniform and ready for surface derivatization. Special care is taken to ensure a high density of coverage followed by thorough end-capping in order to further reduce the possibility of any silanol interactions.

As a consequence of the base deactivation process, the silanols that are still present after surface derivatization become much more 'friendly' toward basic and acidic compounds, and the packing material becomes an excellent choice to develop highly reproducible methods. Silanols are less acidic and are less likely to be available for ion exchange interaction with ionized basic analytes, and are also less likely to hydrogen bond with polar analytes. With these reduced silanol interactions, Thermo Scientific Hypersil BDS columns are ideally suited for analysis of a wide range of analytes including both acids and bases, with peak shape and column performance significantly improved.

### Reproducibility

HPLC owes its success as an analytical technique to several factors. One of the most important has been the ability to transfer newly developed methods to other laboratories around the world. In this respect, column reproducibility has played a crucial role. Column performance parameters are key factors in determining reproducibility from column to column. A column with poor efficiency may lead to loss in resolution, while stationary phase differences may lead to a change in selectivity that can result in loss of resolution. Thermo Scientific strives continuously to provide HPLC columns of the highest standard with a strong focus on reproducibility.

Different brands of C18 media may differ from one another significantly. This is largely due to differences in the properties of the underlying silica. Differences in surface area and silanol population give rise to stationary phases that differ in carbon content, ligand type or silanol content.

Differences within a particular column brand may also occur simply due to the amorphous properties of the base silica itself. Strict control over the processes employed to manufacture both the stationary phase and columns are therefore of paramount importance. Our stringent quality control measures ensure that the required column-to-column reproducibility is achieved. In the following discussion, we describe some of the quality control measures that ensure the continued quality and reproducibility associated with Thermo Scientific Hypersil BDS columns.

#### **Batch Testing Procedure**

Hypersil BDS columns are manufactured to the highest standards, and are rigorously quality controlled. The fully documented ISO9001:2000 control procedures for both media and column production ensure that only the highest quality columns are released to end users. Derivatization only takes place once the BDS silica has passed almost thirty (30) physical and chromatographic test specifications. Once bonded, every production lot of the BDS C18 media is tested for its chromatographic properties and for carbon load. This testing is done both prior to and after end-capping has taken place.

The chromatographic test compares the selectivity, efficiency and asymmetry for the range of analytes against a standard column which is prepared from a blend of up to 50 previous batches of Hypersil BDS C18 packing. The test mixture employed contains compounds such as pyridine and dimethylaniline, which are known to be sensitive to the stationary phase silanol content and can cause peak tailing and varying selectivity on many older

HPLC phases. For each batch of BDS material, all selectivity parameters (k and alpha values) must be within 5% of those measured for the standard column, while efficiency parameters and asymmetry values must also meet strict specifications before it is made available for packing into columns. Figure 3 shows batch-to-batch reproducibility for the selectivity between toluene and NN Dimethyl aniline.

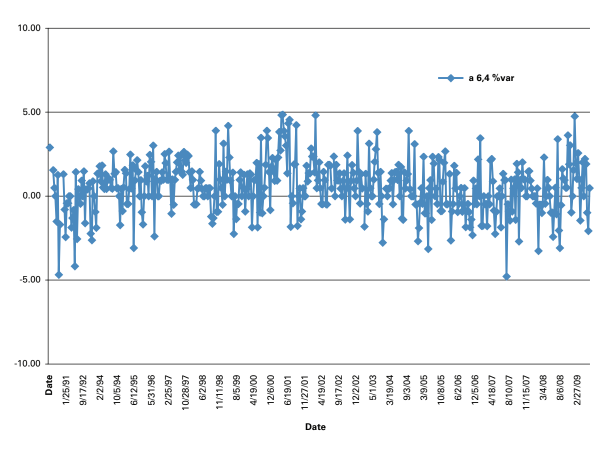


Figure 3: Percentage variation in selectivity (measured against a standard column) for batches of Hypersil BDS produced between 1989 and 2009





## **Column Testing Procedure**

Peak asymmetry and decreased column efficiency are usually observed when a column deteriorates, but may also occur if the column is poorly packed. Every Thermo Scientific Hypersil BDS column is individually tested prior to shipment to ensure the quality of the column. Figure 4 demonstrates column performance

in terms of efficiency for over 15,000 Hypersil BDS C18 (250 x 4.6 mm, 5  $\mu$ m) columns. The peak efficiency (measured for o-xylene) is consistently above 80,000 plates/m. Also noticeable is the trend towards higher efficiency which reflects our commitment to continually improve our column packing process.

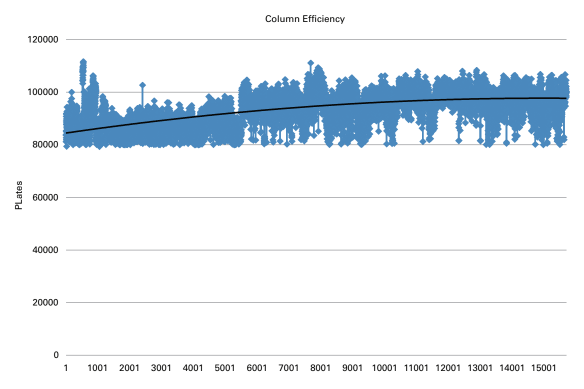


Figure 4: QC data illustrating efficiency for 15000 Hypersil BDS columns

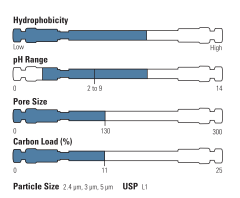


#### **Bonded Phases**

Hypersil BDS columns are available in four bonded phases. All Hypersil BDS columns, the base deactivation procedure and endcapping process minimize peak tailing, even for basic drugs. Each Hypersil BDS column comes with a Certificate of Authenticity.

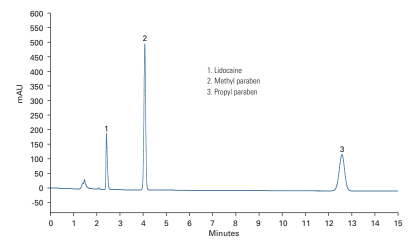
## Hypersil BDS C18 Columns

Hypersil BDS C18 media is an excellent reversed phase material for a variety of applications and is one of the most popular packing materials available. Hypersil BDS columns are a good choice for QA/QC labs as a robust general purpose column in applications where reproducibility and long column lifetimes are required. Hypersil BDS C18 columns are applicable to a wide range of analytes including acids, bases and neutrals as shown, and are popular in methods worldwide.



#### Ordering Information

Particle size Length (mm)		4.6 mm ID	4.0 mm ID	3.0 mm ID	2.1 mm ID	
2.4 µm	30	28102-034630	-	-	28102-032130	
	50	28102-054630	-	-	28102-052130	
	100	28102-104630	_	_	28102-102130	
	150	28102-154630	_	_	28102-152130	
3 µm	30	28103-034630	28103-034030	28103-033030	28103-032130	
	50	28103-054630	28103-054030	28103-053030	28103-052130	
	100	28103-104630	28103-104030	28103-103030	28103-102130	
	150	28103-154630	28103-154030	28103-153030	28103-152130	
5 µm	50	28105-054630	28105-054030	28105-053030	28105-052130	
	100	28105-104630	28105-104030	28105-103030	28105-102130	
	125	28105-124630	28105-124030	28105-123030	28105-122130	
	150	28105-154630	28105-154030	28105-153030	28105-152130	
	200	28105-204630	28105-204030	28105-203030	28105-202130	
	250	28105-254630	28105-254030	28105-253030	28105-252130	

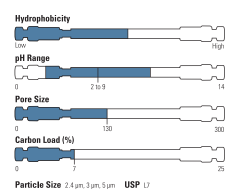


Column: Hypersil BDS C18 150 x 4.6 mm 5  $\mu$ m Mobile phase: 30% MeCN 70% 50 mM KH<sub>2</sub>PO<sub>4</sub> pH 3.5 Flow rate: 1.0 mL/min Detection: UV @ 254nm Inj vol: 20  $\mu$ l Temperature: 30°C Pressure: 92 bar

Figure 5: Separation of lidocaine and parabens

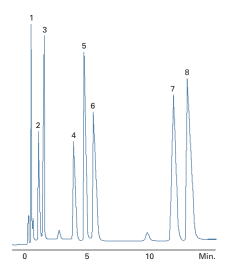
## Hypersil BDS C8 Columns

Hypersil BDS C8 columns offer the same high quality base-deactivated, fully endcapped phase as Hypersil BDS C18, with similar selectivity but slightly less retention. They are applicable to the analysis of acids, bases and neutrals including soft drink additives as shown.



#### **Ordering Information**

Particle size Length (mm)		4.6 mm ID	4.6 mm ID 4.0 mm ID		2.1 mm ID
2.4 μm	30	28202-034630	-	-	28202-032130
	50	28202-054630	_	_	28202-052130
	100	28202-104630	-	-	28202-102130
	150	28202-154630	-	-	28202-152130
3 µm	30	28203-034630	28203-034030	28203-033030	28203-032130
	50	28203-054630	28203-054030	28203-053030	28203-052130
	100	28203-104630	28203-104030	28203-103030	28203-102130
	150	28203-154630	28203-154030	28203-153030	28203-152130
5 µm	50	28205-054630	28205-054030	28205-053030	28205-052130
	100	28205-104630	28205-104030	28205-103030	28205-102130
	125	28205-124630	28205-124030	28205-123030	28205-122130
	150	28205-154630	28205-154030	28205-153030	28205-152130
	200	28205-204630	28205-204030	28205-203030	28205-202130
	250	28205-254630	28205-254030	28205-253030	28205-252130



Column: Hypersil BDS C8, 5 µm, 250 x 4.6 mm

Part Number: 28205-254630

Mobile Phase: A: 0.85% v/v  $\rm H_2SO_4$  in 17.5 mM  $\rm KH_2PO_4$  at pH 1.8

Flow Rate: 1.5 mL/min
Detection: UV at 254 nm

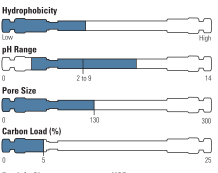
Sample:

- 1. Asorbic Acid
- 2. Quinine
- 3. Caffeine
- 4. Saccharin
- 5. Vanillin
- 6. Aspartame
- 7. Sorbic Acid 8. Benzoic Acid

Figure 6: Additives in soft drinks can be readily separated using a Hypersil BDS C8 column

## Hypersil BDS Phenyl Columns

Hypersil BDS Phenyl columns offer alternative selectivity to Hypersil BDS C18 and C8 columns. It is well established that phenyl phases offer clear advantages over alkyl chains when certain types of compounds must be resolved. The exceptional stability and unique selectivity of Hypersil BDS Phenyl can be used to accomplish a difficult separation.



Particle Size 2.4 µm, 3 µm, 5 µm USP L11

#### **Ordering Information**

Particle size Length (mm)		4.6 mm ID	4.6 mm ID 4.0 mm ID		2.1 mm ID	
2.4 μm	30	28902-034630	-	-	28902-032130	
	50	28902-054630	_	_	28902-052130	
	100	28902-104630	-	-	28902-102130	
	150	28902-154630	-	-	28902-152130	
3 µm	30	28903-034630	28903-034030	28903-033030	28903-032130	
	50	28903-054630	28903-054030	28903-053030	28903-052130	
	100	28903-104630	28903-104030	28903-103030	28903-102130	
	150	28903-154630	28903-154030	28903-153030	28903-152130	
5 µт	50	28905-054630	28905-054030	28905-053030	28905-052130	
	100	28905-104630	28905-104030	28905-103030	28905-102130	
	125	28905-124630	28905-124030	28905-123030	28905-122130	
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	200	28905-204630	28905-204030	28905-203030	28905-202130	
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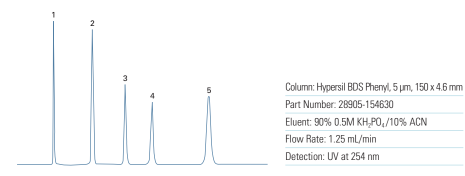


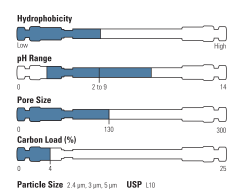
Figure 7: Analysis of procainamides using a Hypersil BDS Phenyl column

Sample:

- 1. Uracil
- 2. Procainamide
- N-Acetyl Procainamide
- 4. Caffeine
- 5. N-Propionyl Procainamide

## Hypersil BDS Cyano Columns

Hypersil BDS Cyano columns can be used in both reversed phase and normal phase chromatography. In reversed phase, they offer different selectivity compared to C18 or C8 phases. In normal phase, they are less retentive than silica columns.



#### **Ordering Information**

Particle size Length (mm)		4.6 mm ID	4.0 mm ID	3.0 mm ID	2.1 mm ID	
2.4 µm	30	28802-034630	-	-	28802-032130	
	50	28802-054630	_	_	28802-052130	
	100	28802-104630	_	_	28802-102130	
	150	28802-154630	_	_	28802-152130	
3 µm	30	28803-034630	28803-034030	28803-033030	28803-032130	
	50	28803-054630	28803-054030	28803-053030	28803-052130	
	100	28803-104630	28803-104030	28803-103030	28803-102130	
	150	28803-154630	28803-154030	28803-153030	28803-152130	
5 µm	50	28805-054630	28805-054030	28805-053030	28805-052130	
	100	28805-104630	28805-104030	28805-103030	28805-102130	
	125	28805-124630	28805-124030	28805-123030	28805-122130	
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	250	28805-254630	28805-254030	28805-253030	28805-252130	

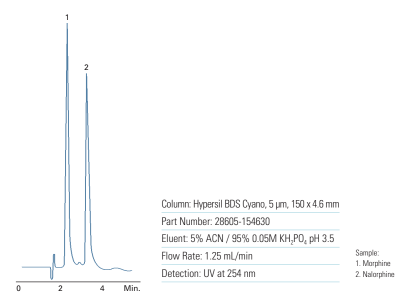


Figure 8: Polar compounds (morphine and nalorphine) are retained and separated using a Hypersil BDS Cyano column

### 2.4 µm Particle Size for Faster Separations

 $2.4~\mu m$  particles give higher efficiency than 3 or 5  $\mu m$  particles and this efficiency is delivered over a greater range of optimum linear velocity. This makes it possible to operate at higher flow rates without losing performance. Because shorter columns packed with  $2.4~\mu m$  particles give equivalent efficiency to longer columns packed with 5  $\mu m$  particles faster analysis and solvent savings for the chromatographer become a reality.

When transferring methods to columns packed with  $2.4 \mu m$  particles, the following three tips should be considered:

 To maintain an equivalent separation when transferring a method it is important to keep the reduced linear velocity constant between the original and new method.

- 2) 2.4 µm-based methods are most often transferred to smaller volume columns, so the same injection volume will take up a larger proportion of the new column, possibly leading to column omeprazole or band broadening. It is therefore important to scale down the injection volume to match the change in column volume.
- 3) Geometrical transfer of the gradient requires calculation of the number of column volumes of mobile phase in each segment (time interval) of the gradient in the original method to ensure that the new calculated gradient takes place over the same number of column volumes, for the new column.

Figure 9 illustrates method transfer to 2.4  $\mu$ m particles using omeprazole as an example. The separation using the column packed with 2.4  $\mu$ m particles, omeprazole elutes in 3.3 minutes, compared with 7.5 minutes for the column packed with 5  $\mu$ m particles and delivers 20% more efficiency.

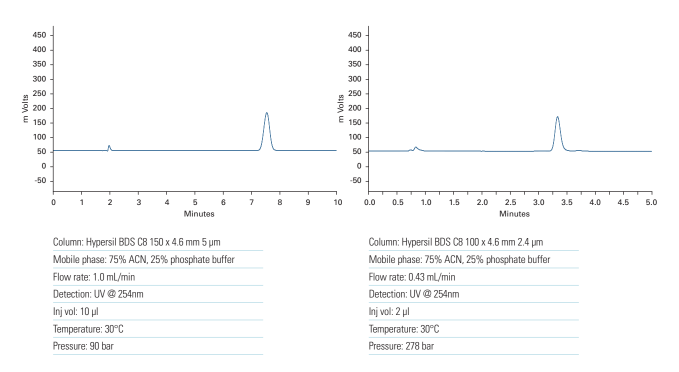
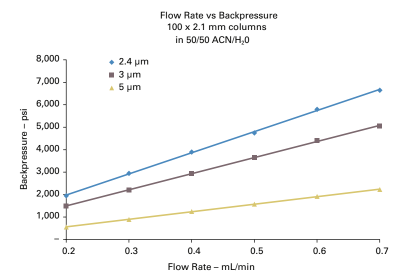


Figure 9: Columns packed with 2.4 µm particles give faster, more efficient chromatography than columns packed with 5 µm particles



One advantage of using 2.4  $\mu$ m particle size columns is that high speed, high efficiency separations are achievable using conventional HPLC systems, even for narrow columns. For example, the backpressure for a 100 x 2.1 mm ID column packed with 2.4  $\mu$ m particles is shown in Figure 10. The optimum flow rate for this column is 0.4 – 0.5 mL/min, which will give a backpressure within the limits for a conventional HPLC system.



We offer a convenient method transfer calculator at the Chromatography Resource Centre.

www.thermoscientific.com/columns



Figure 10: Column backpressure as a function of flow rate for 2.4, 3 and 5 μm particle packed columns

There are some system considerations to remember when using short columns packed with 2.4  $\mu$ m particles in order to maximise their performance. Firstly, to avoid dispersion which can lead to peak broadening, the system volume (connecting tubing ID and length, injection volume, UV detector flow cell volume)

should be minimized. Secondly, because peak widths are narrower with fast chromatography, the detector time constant and sampling rate need to be carefully selected. If fast gradients are being used, then a pump with a low dwell volume is desirable to transfer the gradient to the column quickly.

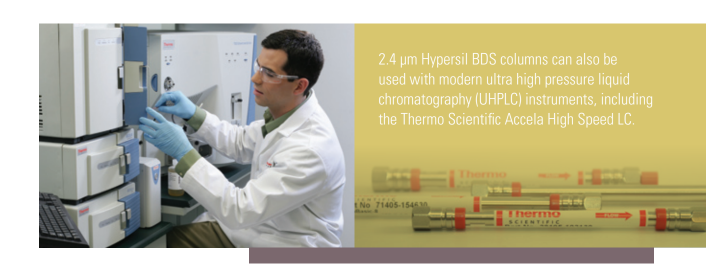
## **Guard Cartridges for Column Protection**

Drop-in guard cartridges and holders offer convenience, economy and effective protection for extending column lifetimes. The 10 mm design offers maximum protection with minimal increase in retention. Thermo Scientific Hypersil BDS drop-in guard cartridges are provided in packs of 4 each.



### **Ordering Information**

Phase	Particle size	Length (mm)	4.6 mm ID	4.0 mm ID	3.0 mm ID	2.1 mm ID
C18	2.4 µm	10	28102-014001	_	-	28102-012101
	3 µm	10	28103-014001	28103-014001	28103-013001	28103-012101
	5 μm	10	28105-014001	28105-014001	28105-013001	28105-012101
C8	2.4 µm	10	28202-014001	-	-	28202-012101
	3 µm	10	28203-014001	28203-014001	28203-013001	28203-012101
	5 μm	10	28205-014001	28205-014001	28205-013001	28205-012101
Phenyl	2.4 µm	10	28902-014001	-	-	28902-012101
	3 µm	10	28903-014001	28903-014001	28903-013001	28903-012101
	5 μm	10	28905-014001	28905-014001	28905-013001	28905-012101
Cyano	2.4 µm	10	28802-014001	-	-	28802-012101
	3 µm	10	28803-014001	28803-014001	28803-013001	28803-012101
	5 μm	10	28805-014001	28805-014001	28805-013001	28805-012101





#### **Chromatography Columns and Consumables**

As the world's sole manufacturer of Thermo Scientific Hypersil silica, we have set a very high standard in HPLC and continue to maintain it with innovative new products spanning across SPE, HPLC and GC. We offer one of the broadest selections of premier chromatographic phases and innovative hardware designs available, combined with superb technical support and customer service. Whether you use HyperSep for rapid sample preparation, Hypersil BDS for routine separations or are looking for something new for your most challenging methods like Thermo Scientific Hypersil GOLD or TRACE™ GC columns, we have the choices to meet your needs.

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