

## Sepax Technologies, Inc.

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### **Unix-C SEC-300 Column Manual**

#### Column Information

Utilizing proprietary surface technologies, Unix-C SEC-300 phase is made of uniform, hydrophilic, and neutral nanometer thick films chemically bonded to high purity and mechanically stabilized silica with the particle size of 1.8 µm. The combination of small particle size and large pore volume of Unix-C SEC-300 renders the highest separation efficiency and resolution of analytes. The well-controlled surface chemistry results in excellent lot-to-lot reproducibility. Our unique bonding chemistry, coupled with the maximized bonding density, allows Unix-C SEC-300 to provide high stability and negligible non-specific interactions. Typical applications for Unix-C SEC-300 columns include separation and analysis of biological molecules, hydrophobic, and water-soluble polymers.

## Column Stability and Performance

Unix-C SEC columns use full coverage bonded silica packing, which allows exceptionally high stability. They are compatible with most aqueous buffers, such as ammonium acetate, phosphate, tris, etc.

The neutral and hydrophilic Unix-C stationary phases have negligible nonspecific interactions with biological molecules, especially proteins. Combined with their high capacity, Unix-C SEC columns enable high efficiency and high recovery separations. A typical quality control chromatogram is shown in Figure 1 for a 4.6 x 300 mm Unix-C SEC column.

#### Column Characteristics

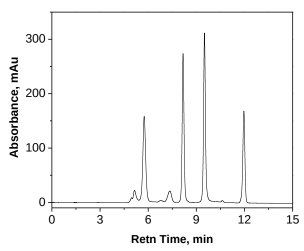
Silica: Spherical, high purity (<10 ppm metals)

Particle size:  $1.8 \mu m$ Pore size: 300 Å

Safety Precautions

The columns are normally operated under moderate pressure. Loose connections will cause leaking of buffers and injected samples, all of which should be considered hazardous. In the case of leaking, proper gloves should be worn while handling the columns. When opening the columns, proper protections should be used to avoid inhalation of the small silica particles.

Figure 1. Elution profiles of a protein mixture by Unix-C SEC-300 phase.



Column: Unix-C SEC-300, 4.6 x 300 mm Mobile phase: 150 mM sodium phosphate buffer, pH 7.0

Flow rate: 0.35 mL/min
Temperature: Room temperature
Detection: UV 214 nm
Injection:  $1 \mu L$ 

Sample: (1) Thyroglobulin aggregate; (2) Thyroglobulin (1.0 mg/mL), 670 kD; (3) BSA dimer, 132 kD; (4) BSA (1.0 mg/mL), 66 kD; (5) Ribonuclease A (1.0 mg/mL), 14 kD, and (6) Uracil (0.1 mg/mL), 120 D.

## Column Installation and Operation

The column should always be capped at both ends when it is not in use. When installing the column to the system, first remove the end caps. Unless a user has special purpose to reverse the flow direction, for example, removal of the inlet blockage, follow the flow direction as marked on the column. Column connections are an integral part of the chromatographic process. If ferrules are over tightened, not set properly, or are not specific for the fitting, leakage can occur. Set the ferrules for column installation to the HPLC system as follows:

- (a) Place the male nut and ferrule, in order, onto a 1/16" outer diameter piece of tubing. Be certain that the wider end of the ferrule is against the nut.
- (b) Press tubing firmly into the column end fitting. Slide the nut and ferrule forward, engage the threads, and finger-tighten the nut.
- (c) Repeat this coupling procedure for the other end of the column.

# Samples and Mobile Phases

To avoid clogging of the column, all samples and solvents should be filtered through 0.45  $\mu m$  or 0.2  $\mu m$  filter before use. Unix-C SEC columns are compatible with aqueous mobile phases or a mixture of organic solvent and water, such as methanol or

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acetonitrile and water. Always degas the mobile phase. A simple way for degassing is to sonicate it for 5 minutes under aspirator vacuum.

#### Column Care

**Shipping Solvent** New columns are shipped in water containing 0.02% (w/v) sodium azide (NaN3). During stocking and shipping, the resin bed may become dried out. To activate a new column, it is recommended to flush the column with 10-20 column volumes of the buffer starting at a low flow rate and gradually increasing until desired operating flow rate is met.

**Column Installation** Flush the column with the mobile phase while gradually increasing the flow rate from 0.1 mL/min to the operating flow rate allowing the column to equilibrate the pressure until the baseline is stable at each step. If the column back pressure and baseline fluctuate, this might be due to air bubbles trapped inside the column. Flush the column for longer time until the back pressure and base line are stable.

**Pressure** Even though the columns can operate at a pressure up to 4,500 psi, continuous use at high pressure may eventually damage the column. Since the pressure is generated by the flow rate, the maximum flow rate is limited by the back pressure. It is expected that the back pressure might gradually increase with its service. A sudden increase in back pressure suggests that the column inlet frit might be clogged. In this case it is recommended that the column be flushed with reverse flow in an appropriate solvent.

*Flow Rate* Standard operating flow rate is 0.1 - 0.35 mL/min.

 $\it pH$  For optimum performance and lifetime keep pH between 2.0 and 8.5.

**Temperature** The maximum operating temperature is 80 °C. The optimum operating temperature for the longest lifetime is 10 - 30 °C. Continuous use of the column at higher temperatures (>30 °C) can damage the column, especially under high pH (>8.0).

**Storage** When the column is not in use for an extended time, the column should be stored in water  $(H_2O)$  with 0.02% sodium azide or 10% methanol. Each column is shipped with two removable end plugs. To prevent drying of the column bed, seal both ends of the column with the end plugs provided.

**Cleaning** From time to time, some samples could get adsorbed onto the inlet frit or the packing material. When the adsorption accumulates to a certain level, it is usually indicated by an increase in back pressure and a broader peak. When this occurs, it is time to clean the column. The general procedure for column cleaning is as follows:

- 1. Disconnect the column from the detector.
- 2. Clean the column in the reverse flow direction.
- 3. Run the column at below 50% of the maximum recommended flow rate. Monitor the back pressure. If the pressure is much higher than the normal operating conditions, you need to lower the flow rate or change the washing buffer as the cleaning solutions may be of different viscosities.
- 4. Typically, 10-15 column volumes of cleaning solution are sufficient. Rinse well with 3-5 column volume of nanopure water between each solution.

Cleaning Solutions Low pH salt solutions help remove basic proteins. Organics are useful when removing hydrophobic proteins. Chaotropic agents help to remove strongly adsorbed materials (via hydrogen bonding). Only use chaotropic agents when neutral salts or organics have not improved resolution. Two cleaning solutions are recommended for general cleaning:

- 1. Concentrated neutral salt (e.g.,  $0.5\ M\ Na_2SO_4$ ) at low pH (e.g., pH 3.0)
- 2. Water soluble organic (MeOH, ACN, EtOH, 10 %-20 %) in aqueous buffer (e.g., 50 mM sodium phosphate, pH 7.0)

If both solutions fail to clean the column, use 6 M urea (filter before use).

- a. 2 cv 6 M urea at 0.35 mL/min (monitor pressure)
- b. 3 cv nanopure water at 0.35 mL/min
- c. 7 cv mobile phase at 0.35 mL/min

### **Column Protection**

In addition to filtering the sample and the mobile phase, the best way to protect the separation column is to install a guard column or a pre-column filter in front of it. In most cases a pre-column filter helps to remove the residual particulates that are in the sample, the mobile phase, or leached from the HPLC/UHPLC system, such as pump and injector seals. However, a guard column is highly recommended because it is more effective in trapping highly adsorptive sample components and residual particulates in the sample, the mobile phase or from the HPLC/UPLC system.

## Ordering Information:

Unix-C SEC-300 (1.8 µm 300 Å)

P/N	Length x ID (mm)
231300-4615	4.6 x 150
231300-4630	4.6 x 300

Unix-C Precolumn Filters and Frits

P/N	Description
102000-P346	Precolumn Filter with 0.5 um stainless steel frit
102001-P346	0.5 um stainless steel refill frits (for Precolumn Filter)