CNBr Pre-Activated Resins

Functionalized on a modern, high flow agarose base matrix for simplified ligand immobilization and customizable affinity chromatography purification solutions
Why Purolite®?

For over 35 years, Purolite has supplied specialty ion exchange resin technology to industries within complex regulatory environments, including biotechnology, pharmaceutical, food, fine chemical and electric power generation. Purolite is the only global company to focus 100% on resin technology.

Security of Supply

Ensuring reliable availability of products in case of emergency is vital to customers and of paramount importance to Purolite.

As a leading supplier of resin media to the world’s most regulated industries, Purolite has a real-world security-of-supply system in place to support your process requirements for business continuity in the instance of natural disaster or emergency.

Purolite has manufacturing facilities at 3 strategic global locations in the USA, Asia and Europe, and is currently building its 4th manufacturing plant in the UK. This facility will be the second largest agarose manufacturing plant globally, with a capacity of 100,000 L per annum.

Currently, approximately 90% of all biopharmaceuticals approved by the U.S. Food and Drug Administration utilise a single source of agarose resins from a single manufacturing site, presenting a security of supply risk to long-term clinical trial material production.

Purolite have addressed this industry-wide concern by providing the first proven and reliable alternative source of agarose resins, allowing customers to dual-source their products to mitigate their supply risks.

Regulatory Support

Purolite Life Sciences provides customers with regulatory support documentation for Praesto products used by our customers in GMP regulatory environments.

Comprehensive regulatory support files are available for each Praesto resin, and are provided under a confidential disclosure agreement.

The purpose of this Regulatory Support File (RSF) is to provide assistance with:

♦ Process development of clinical and commercial purification processes
♦ Manufacturing validation
♦ Quality control tests
♦ Standard Operating Procedure (SOP) for cleaning in place (CIP) and sanitization
♦ Application for various regulatory licenses or compliance
♦ Plant and document audits

Quality

Global ISO 9001:2008 standards ensure consistent operating practices across each of our plants. Compliance is monitored and maintained through a quality assurance and regulatory team who conduct internal audits to ensure operations meet the guidelines and protocols for equipment and procedures.
Additionally, our production team is given continual training on quality processes to ensure batch-to-batch consistency, and we host numerous customer audits each year to make sure that we are in compliance with user expectations.


**Raw Materials**

Our raw material suppliers are selected and qualified from leading manufacturers and are part of our global network of suppliers. Each key raw material has at least one alternative supplier and is managed through a globally coordinated inventory system to ensure security of supply.

Additionally, a quality control protocol is in place for testing new batches/lots of raw materials to confirm product specifications and lot-to-lot consistency.

Purolite Life Sciences also has long-term supply agreements in place for our Protein A ligands, which are sourced from Repligen Corporation.

Repligen provides dual-site supply for critical raw materials and has a long-standing history of successfully supplying a variety of Protein A ligands to the industry.
Pre-Activated Base Matrices

**Praesto® CNBr**

Pre-activated CNBr resin functionalized on a modern, high flow agarose base matrix for simplified ligand immobilization and fully customizable affinity chromatography purification solutions

**Overview**

To support in the development and manufacture of biopharmaceuticals, Purolite has developed a range of pre-activated agarose resins. These resins enable manufacturers to couple their own ligands to develop affinity chromatography solutions. NHS, Epoxy and CNBr pre-activated chemistries are available in three particle sizes - 45 µm, 65 µm and 90 µm.

**Praesto CNBr Resins**

**Praesto** CNBr resins have been designed to offer a simple solution for the immobilization of ligands onto an agarose chromatography matrix, which can be utilized to make customized affinity resins. This enables rapid scale-up from R&D proof of concept to larger scale bioprocess production columns.

The use of Cyanogen bromide (CNBr) pre-activated base matrices is a well-established, rapid and familiar technique for the coupling of bio-specific ligands to generate affinity chromatography purification media. This choice of pre-activation chemistry is well suited for research, diagnostic and academic applications. Coupling is via primary amino groups. **Praesto** CNBr resins have a cyanogen-active group which form an isourea linkage between ligand and resin. Many well-documented references (published over several decades) are publicly available.

**Key Performance Benefits**

- Very low levels of non-specific binding due to the highly hydrophilic properties of the agarose base matrix
- Rigid base matrix allows significantly (over 100%) higher flow velocities than other agarose resins, making it suitable for process scale operations
- Quick and straightforward ligand coupling
- CNBr agarose has been successfully used for over two decades
- Chemically stable due to multipoint attachment ligand chemistry coupling
- Modern range of resins maximizes facility productivity, improving process economics significantly

**Figure 1: Praesto CNBr Pre-Activated Resin Structure**

![Figure 1: Praesto CNBr Pre-Activated Resin Structure](image)
Matrix Characteristics

The *Praesto* CNBr range of pre-activated chromatography resins use a modern, highly cross linked-agarose matrix formulation. Due to the unique rigidity and open pore structure of the *Praesto* agarose base beads, the *Praesto* CNBr range is well suited for process-scale chromatography, allowing large columns to be operated. This is due to high cross linking, which enables processes to operate at very high flow rates compared to other commercially available resins.

Proteins and other molecules are covalently coupled directly to the pre-activated gel via primary amino groups.

Figure 2 shows the pressure flow comparison against Sepharose 4 Fast Flow (90 µm) and Sepharose 6 Fast Flow (90 µm). Even at process scale, with larger diameter columns and bed heights, the rigidity of *Praesto* allows at least 100% higher linear flow velocity. The ability to run at high flow rates increases productivity and improves facility throughput.

*Praesto* CNBr pre-activated resins are available in three particle sizes - 45 µm, 65 µm and 90 µm. Across the range of three bead sizes, porosity and ligand density is maintained. This enables the selection of an optimal particle size for a particular downstream process to maximize productivity, resolution, and pressure restraints.

Operation and Use

First, the *Praesto* CNBr resin requires swelling. 1 g of lyophilized powder typically provides 3.5-4 ml of final volume. Once swelled, the resin is then washed prior to ligand coupling. The coupling reaction is quick and spontaneous.

Instruction protocols are provided with the product and describe generic conditions, however, we recommend specific optimization for individual processes.
CIP, Shelf Life and Storage

Regular cleaning-in-place (CIP) is a key process step that regenerates the resin, extending lifetime and maintaining capacity through the removal of contaminants bound but not removed during a low elution pH. CIP should be optimized for each specific process, however in general the use of low and high pH solutions (e.g. 0.1 M sodium acetate containing 0.5 M NaCl, pH 4.5 and 0.1 M Tris HCl containing 0.5 M NaCl, pH 8.5) is suitable. Ethanol concentrations using several column volume washes between 30-70% can be used to remove strongly bound contaminants.

All Praesto CNBr resins are provided freeze-dried, and long term storage is recommended between 2-8°C, providing an expected shelf life of at least 24 months. Since 2014, a long-term stability study of the Praesto base matrix has been ongoing. The stability of the coupled matrix is directly dependant to the coupled ligand. CNBr chemistry has been successfully used for over 20 years.

This bond is stable between pH 2-11 short term and pH 3-11 long term.

The Praesto® Range

The Praesto range offers a selection of modern, high-flow Affinity and Ion Exchange agarose resins, delivering exceptional results from Protein A to high-resolution polishing steps. The range also includes a full selection of Praesto Pure base matrices, and pre-activated resins in a variety of source chemistries.

All Praesto products provide an advanced, high-flow, highly cross-linked agarose base matrix. The entire range benefits from excellent pressure/flow characteristics and stability for optimal recovery of active proteins.

Discover Praesto at: www.purolite.com/life-sciences
Ordering Information

To place your order simply contact us via email or telephone using the information on the back page of this brochure and quote your order number from the table below. If you wish to discuss your purification challenges with a specialist, we have dedicated experts on-hand across the globe to provide knowledgeable, same-day technical assistance.

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Protein Coupling to Pre-Activated *Praesto* Resins

**Ligand Coupling Methodology**

- Magnetic stirrer bars should be avoided as agarose resin is susceptible to damage from grinding. Damaged agarose can result in inability to couple the ligand to the resin and poor performance of the coupled resin in the application use.

- Direct heating of the solution should be avoided.

- Coupling conditions between pH 7-9 can be used.

- A standard buffer which can be used is 0.1 M NaHCO3, 0.5 M NaCl (pH 8.3).

- After coupling the ligand to the resin any remaining CNBr groups need to be deactivated from the resin, this is achieved by washing with a Tris buffer.

**Coupling Procedure**

Dissolved the ligand to be coupled in a small amount of coupling buffer.

Wash *Praesto* CNBr on a filter with 10 equivalent volumes of ice cold 1 mM HCl and then wash with coupling buffer.

Re-suspend the gel in coupling buffer and stir. Once the slurry is stirring efficiently add the dissolved ligand. Stir for 2 – 4 hour at ambient temperature or for at least 16 hours at 4°C.

The slurry composition should ideally consist of a 0.5-1:1 ratio of buffer to preactivated resin – i.e. 50 ml of gel in a total slurry volume of 75 – 100 ml. If the slurry is to dilute then reaction times are increased and incomplete coupling can occur.

Once the reaction is complete wash the coupled media on a filter with 5 equivalent volumes of coupling buffer, transfer the dewatered gel back to the reaction set up and add an equal volume of 1 M ethanolamine or 0.1 M Tris-HCl and leave to stand for 2 hours.

Wash the coupled resin with at least 3 equivalent volumes of acetate buffer (pH 4) followed by 3 equivalent volumes of Tris-HCl buffer (pH 8) and then 3 equivalent volumes of water. For long term storage wash the resin with 20% Ethanol solution and store the resin in 20% Ethanol solution.
Packing and Column Evaluation of Immobilized Resins

Column Packing

Packing Tricorn columns

• The following instructions are for packing a Tricorn 10/300 (GE Life Sciences) column with a 30 cm bed height.
• For more details about packing Tricorn columns, please see the GE Life Sciences instructions: Tricorn Empty High Performance Columns (28-4094-88).

Materials and Equipment

• Praesto Pure90, Praesto Pure65 or Praesto Pure45
• Tricorn 10/100 packing equipment
• Tricorn 10/300 column
• Plastic beaker
• Plastic syringe
• Measuring cylinder
• 0.5M NaCl solution
• A Chromatography system, such as ÄKTA™ system, or a stand-alone pump such as Pump P-900, depending on the flow rate required, can be used for packing.

Packing Procedure

1. Wash the sample with 5 times with 50 ml of 0.5 M NaCl solution to remove the 20% ethanol storage solution.
2. Decant off remaining NaCl wash solution and add 0.5 M NaCl solution to obtain a 70% slurry concentration.
3. Calculate the required slurry volume for a 30 cm packed bed.
   a. Determining the slurry volume for column packing.
   b. Determine the desired packed bed height.
   c. Calculate the column volume (Cv) of a packed column by the following equation:
      i. Cross-sectional area of the column (CSA)×bed height (Bh)
      ii. Multiply the column volume by a compression factor (C.F) (Cv × C.F)
      (C,F = 1.12 to 1.15 dependent on particle size. 45 µm = 1.12, 65 µm = 1.15 and 90 µm = 1.15)
      iii. Divide by the slurry concentration (normally between 50% to 70%).
   d. Example calculation
      Column: Tricorn 10/300
      Desired bed height: 30 cm
      Slurry concentration: 70%
      Compression Factor (90 µm): 1.15
      (CSA × Bh × C,F) / Slurry Concentration
      ((0.5)² π) x 30 x 1.15)/0.7 = 38.7 ml
      Required slurry volume for a 30 cm packed bed = 38.7 ml.
4. Unpack a Tricorn 10/300 column, assemble and connect Tricorn 10/100 packing equipment as per the manufacturer’s instructions (GE Life Sciences).

* ÄKTA and Tricorn are trademarks of GE Healthcare
5. Stir column media gently with a plastic spatula (DO NOT use a magnetic stirrer bar to ensure homogeneity) and pour down a plastic spatula into the top of the packing column until the column and packing column are completely full. Leaving an inverted meniscus at the top of the packing column.

6. Insert connector, with filter attached, at a 45° angle to prevent air bubbles forming at the top of the column and screw the top cap of the packing column.

7. Using an ÄKTA™ system, start a flow rate of 0.5 ml/min of 0.5 M NaCl packing solution through position 1 of the column valve. Once a flow is established, connect 0.5 mm tubing from column position 1A to the top of the packing column.

8. Remove the stop plug from the bottom of the column and replace with 0.5 mm tubing running into a waste container.

9. Adjust the flow rate to 1.25 ml/min and run until the resin has settled, then increase the flow rate to * ml/min and run for ** minutes to pack the resin. (Praesto Pure45 does not require a settling step, please skip to step 10)

10. Stop the flow and mark the point at which the resin has settled. If necessary, remove excess medium by re-suspending the top of the packed bed and remove with a Pasteur pipette or spatula.

11. Insert column adaptor into the top of the column at a 45° angle and screw plunger down to the marked point and reconnect tubing to the top of the column from position 1A of the column valve.

12. Connect tubing from the bottom of the column to position 1B of the column valve.

13. Pack for a further 20 minutes at *** ml/min. At the end of 20 minutes mark the point at which the resin has settled.

14. Detach the tubing connected to the column and place a stop plug in the bottom of the column. Remove the lock on the top of the adaptor and screw the plunger down to the point marked in step 13.

15. Reconnect the tubing as described in step 7.

<table>
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<tr>
<th>Product</th>
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<th>Time**</th>
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<tr>
<td>Praesto CNBr90</td>
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<td>6 ml/min</td>
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**Column Efficiency Testing**

The column efficiency should be tested immediately after packing and at regular intervals during use to monitor any deterioration.

The preferred method for determining the efficiency of a packed column is through the use of the height equivalent to a theoretical plate (HETP) and the asymmetry factor (AS).

The HETP and AS values are determined by applying a sample such as 1 – 3% acetone in demineralised water to the packed column.

A sample of 0.4 to 0.8 M NaCl in demineralized water can also be used.

A sample volume of approximately 1% of the column volume and a flow velocity of between 30 to 50 cm/h will give the optimal results.
Calculating HETP and $A_s$

Below is the calculation by which HETP and $A_s$ are determined. This is done using the UV curve (or if using a NaCl sample, the conductivity curve is used).

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

$L$ = bed height (cm)  

$N$ = number of theoretical plates  

$V_r$ = volume eluted from the start of the sample application to the peak maximum.

$W_h$ = The width of the recorded peak at half of the peak height.

$V_r$ and $W_h$ have the same units.

The reduced plate height is calculated by the following equation:

$$h = \frac{HETP}{d_{50v}}$$

$d_{50v}$ = mean particle size (cm)

The reduced plate is often taken into consideration when evaluating column packing efficiency. As a guide a value of $< 4$ well packed can indicate a well packed column. A value $< 3$ is considered a very good result.

The peak corresponding to the acetone or NaCl sample should be symmetrical with an asymmetry factor as close to 1 as possible.

An acceptable is $0.8 < A_s < 2.0$

$$A_s = \frac{b}{a}$$

$a$ = ascending part of the peak width at 10% of peak height.  

$b$ = descending part of the peak width at 10% of peak height.

A change in the shape of the peak is usually the first indication of bed deterioration as a result of excessive use.

Figure 1. Shows an example UV chromatogram of a 1 – 3% acetone sample during a column efficiency test.

The calculated plate number will vary according to the test conditions and it should only be used as a reference value. It is important that test conditions and equipment are kept constant so that results are comparable. Changes of solute, solvent, eluent, sample volume, flow velocity, temperature will all affect the results.