



## Epoxy-Activated Beads

### Multi purpose activated support

Cat. 1004-5/10/25

ver. 1.1

### Introduction

The **Epoxy-Activated Beads** is a preactivated support that contains high density of epoxy groups used for immobilization of various ligands. The preactivated support can be used to immobilize proteins, carbohydrates and various ligands via stable linkage to amine, SH and OH groups.

### Epoxy-Activated Beads characteristics

Matrix: Sepharose™ CL-4B

Activate group: Oxiran.

Active group density: 10–25 μmole/ml

Mean bead size: 45 -165 μm

Bead structure: Highly cross-linked spherical agarose, 6%

Max back pressure: 0.3 MPa, 3 bar

Max. flow rates: 4 ml/min/cm<sup>2</sup>

Recommended flow rate: 1-2 ml/min/cm<sup>2</sup>

Stability of the matrix: pH 2-11.

Storage: 20% ethanol as preservatives

### Protocol: Immobilization of ligand to Epoxy-Activated Beads

#### A. Buffers needed

In general coupling should be performed in carbonate, borate or phosphate buffers. Sodium hydroxide may be used for making solutions of high pH. Avoid using Tris or free-amine containing buffers during conjugation, as these groups may couple to the epoxy groups. Organic solvents may be needed to dissolve the ligand. Dimethylformamide and dioxane may be used to up to 50% of the final volume. The same concentration of organic solvents should be included in the coupling buffer.

Buffers required for conjugation of ligand to 10ml of **Epoxy-Activated Beads**.

1. Double distilled water (100 ml).

2. Coupling buffer (50 ml): Sodium bicarbonate or carbonate buffer pH-8-10. Neutral conjugation buffer may also be used, although basic coupling buffer is preferred: Use ddH<sub>2</sub>O, or phosphate buffer such as PBS.
3. Blocking solution (100 ml): 1 M ethanolamine pH 8.0.
4. Wash buffer (200 ml): PBS pH 7.4.

#### B. Recommended protocol

1. Gently mix the beads to form homogenous mix and remove by pipette the desired amount of beads. Wash beads 3 times with double distilled. Washing could be done either using a suitable filter funnel or by centrifugations cycles of 1 minute long each one, done at approximately 600-1000xg.
2. Dissolve the ligand to be immobilized in Coupling buffer. Organic solvent may be used. For large ligands prepare solution of 2-10 mg/ml use and for small ligands prepare solution of at least 200  $\mu$ moles ligand per ml Coupling buffer.
3. Add two volumes of ligand solution to one volume of washed **Epoxy-Activated Beads** in polypropylene tube and mix gently. Sample 100ul upper supernatant (without beads) as time zero reference sample and store in the refrigerator.
4. Mix slowly overnight at 4°C to 45°C (stability of the ligand limits the maximum temperature) preferably with the use of a rocker. Do not use magnetic stirrer for mixing.
5. Sample 100ul upper supernatant (without beads) for conjugation efficiency determination and wash away excess ligand using Coupling buffer in order to remove unbound biomolecules.
6. Add approximately two gel volumes of Blocking solution. Let it stand for at least 4 hours or overnight at 4°C.
7. Wash gel three times with Wash buffer 2( minutes each time with 5 volumes) at room temperature, in order to remove unbound ethanolamine.

#### C. Storage

Store gel refrigerated in any desired buffer added with 0.1% azide w/v until use.