



**BESTCHROM**

博 格 隆

**Chelating Bestarose FF  
Metal chelate  
chromatography resin  
Instruction for use**



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## 1. Introduction

Chelating Bestarose FF is an immobilized metal affinity chromatography resin, which is made by covalently crosslinking ligand Iminodiacetic acid (IDA) into Bestarose FF matrix. The target protein is based on the separation of side chain histidine, cysteine and tryptophan with the transition metal ions (Cu<sup>2+</sup>, Co<sup>2+</sup>, Ni<sup>2+</sup>, Zn<sup>2+</sup> etc.) immobilized on the resin.

The ligand of Chelating Bestarose FF resin can provide 3 coordination sites to chelate with metal ions while provide three ionic bond binding sites to purify the target protein with high affinity. The same type of IMAC Bestarose FF resin provides 4 coordination sites to chelate with metal ions, and 2 ionic bond binding sites to purify the target protein, that is to say, Chelating Bestarose FF resin has the same ligand density and same metal ion conditions. IMAC Bestarose FF has a stronger affinity. All samples that cannot be adsorbed in IMAC Bestarose FF resin can be combined with Chelating Bestarose FF. However, because IMAC Bestarose FF resin has one more metal ion chelation site, it has stronger binding strength to metal ions, and can also be compatible with DTT β-mercaptoethanol. Thus, when purifying recombinant histidine-tagged protein, the resin should be selected according to the characteristics of the protein.

## 2. Technical characteristics

Appearance	White slurry, can be layered
Matrix	Cross-linked agarose, 6%
Particle size <sup>+</sup>	45-165μm
Chelating ability	Cu <sup>2+</sup> : ~34μmol/mL resin
Working pH	4~8.5
Chemical stability <sup>++</sup>	Stable in common aqueous buffers:1M NaOH <sup>+++</sup> ,6M GuHCl,8M Urea.
Physical stability	Changes in volume due to changes in pH or ionic strength are negligible
Pressure flow velocity	>300cm/h (0.1MPa BXK50, H=15cm, 25℃)
Max. pressure	0.3MPa
pH stability <sup>++++</sup>	3-13(working) 2-14(CIP)
Temperature tolerance	Working temperature:2~40℃, Can't freeze, Can tolerate 121℃ high pressure sterilization (20min)
Storage <sup>++++</sup>	2~30℃, 20% ethanol or 2% benzyl alcohol
Recommended flow velocity	60~200cm/h

+Particle size is normally distributed, and particles within this range account for more than 90% of the total.

++No chelated metal ion

+++1M NaOH only be used for cleaning.

++++ CIP refers to pH stability when metal ions are removed.

+++++ 2% benzyl alcohol is only used for international transport or special requirements from customer

### 3. Method of chromatographic

#### 3.1 Column packing

**Note: It is best to equilibrate the resin slurry to room temperature before column packing.**

- According the column volume to calculate the amount of resin.

Resin volume=column volume×1.15(Compression factor=1.15)

According to the volume of the settlement resin required, the suspended slurry of the resin required is calculated by the follow:

Required resin slurry<sup>1</sup> volume = Settlement resin volume ÷ Resin slurry<sup>1</sup> concentration.The original concentration of resin slurry<sup>1</sup> is shown in the follow table.

Pack size	Resin slurry <sup>1</sup> concentration (%)
25mL、100mL、500mL、1L、5L、10L	80
20L、40L	75

**1: It refers to the original packaging resin slurry sold by Bestchrom.**

**Note:For non-original packaging, customer can calculate the required volume according to the actual concentration of resin slurry.**

- Washing the resin:Thoroughly shake the resin and weigh certain volume of resin calculated by the method mentioned above. Pour it into a funnel, drain the liquid, and wash with about 3mL distilled water/mL resin for 3 times. Use a glass stick or stirrer to stir each time when adding distilled water, which helps to wash the shipping solvent away.
- Prepare the packing slurry: Transfer the washed resin from the funnel into a beaker or other appropriate container, add distilled water to obtain a 50%~75% slurry, stir well and set aside for use.
- Take a cleaned B XK column (B XK series columns with diameters ranging from 1cm to 30cm can satisfy different scale chromatography applications). Take B XK16/20 for example, purge the bubbles trapped at the end-piece net by draining some distilled water through the column outlet. Leave about 1cm water at the bottom of the column and close the bottom outlet. Adjust the column so that it is perpendicular to the ground.
- Slowly pour the slurry into the column at one time (use a packing reservoir if necessary). Do not bring any air bubbles into the column.

**Packing reservoir: Empty glasstube with same diameter as the B XK column.**

- Fill the remainder of the column with packing solution. Connect the packing reservoir to the chromatography system, open the flow velocity, drain the bubbles in the hose, close the flow

velocity, and tighten the top cover of the packing reservoir.

- ◇ After pouring, stir well again with Stirrer, and then wash the resin particles on the inner wall of the column from top to bottom with the packing solution, and let the resin settle naturally until there is about 1cm of clarifying solution on the suspension. Mount the adapter and connect the adapter to the chromatography system or peristaltic pump. Lower the adapter to descend to contact with the clarifying solution and tighten the sealing ring after it is fully immersed in the clarifying solution. With the outlet of the top piece is opened, slowly move the adapter down until all bubbles are drained.

**Note: This operation is only applicable to BXK 100 and above columns. Flushing the inner wall reduces the resin particles sticking between the seal ring and the column wall, avoiding the risk of leakage.**

- When the bed height is 10cm, the flow velocity can be set to 75cm/h. Open the bottom plug, start the pump and run the setting flow velocity until the bed is stabilized, mark the bed height.
- Remove the packing reservoir (if any), Install the adaptor lower the adaptor to about 0.5cm above the resin surface, set the flow velocity at 260cm/h, and continue to press the column using the above flow velocity until the bed is completely consolidated, mark the consolidated bed height.
- Stop the pump, open top plug, close the bottom plug, loosen the O-ring seal slightly, press the adaptor to about 0.3cm below the marked position, tighten the O-ring seal, close adaptor stop plug, and complete the column packing.

### 3.2 Evaluation of Packing

- The packing quality of chromatographic column can be confirmed by column efficiency measurement and evaluation. The tests are required after the column packing, during the column working life and when the separation and purification performance weakens. The method usually relies on the height equivalent to a theoretical plate(HETP) and the asymmetry factor(As).
- Acetone or NaCl solution can be used as sample for the testing. Sample solution and mobile phase can be prepared according to the following table.

	Acetone method	NaCl method
Sample	1.0% (v/v)acetone in water	0.8M NaCl in water
Sample volume	1.0%CV	1.0%CV
Mobile phase	Water	0.4M NaCl in water
Flow velocity	30cm/h	30cm/h
Monitor	UV280 nm	Conductivity

- Method for measuring HETP and As:  
Use UV curve or the conductivity curve to calculate the height equivalent of theoretical plate (HETP), number of theoretical plates(N) and the asymmetry (As):

$$HETP=L/N$$

$$N=5.54(V_R/W_h)^2$$

Note:  $V_R$  = retention volume

$W_h$  = half-peak width

$L$  = column height

$N$  = the number of theoretical plates

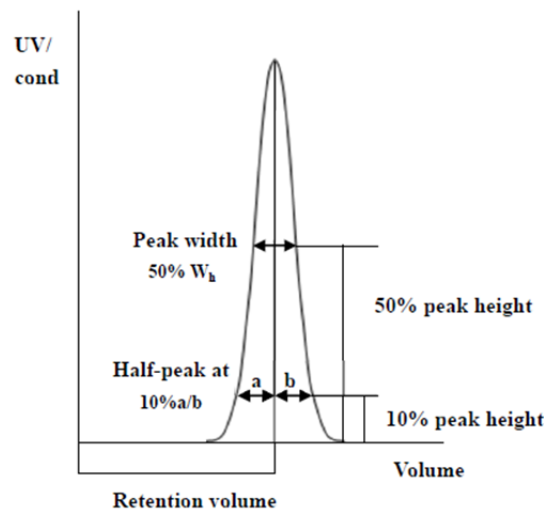
(The units of  $V_R$  and  $W_h$  should be the same)

$$As=b/a$$

Note:

$a$ = 1st half peak width at 10% of peak height

$b$ = 2nd half peak width at 10% of peak height



- Evaluation the column packing

As a guideline, if the value of HETP is less than 3 times the average particle size( $d_{50}$ ) of the resin and the  $As$  is between 0.8~1.8, the column is very efficient. The unsatisfactory results should be analyzed and the column should be repacked.

### 3.3 Chromatographic method

- Chelated metal ion:

- Rinse 2CV with purified water,

- Pass 0.5CV 0.2M metal ion solution through the chromatography column;

**The general metal ion environment is neutral (pH 7~8). Zinc ion selects  $pH \leq 5.5$  to avoid the solubility of high pH salt. Iron ion selects  $pH \approx 3$  to avoid the formation of insoluble matter.**

- Use 5CV of purified water to remove unbound metal ions;

- The chromatography column was cleaned with at least 5 times column volume acidic buffer solution (20mM NaAc, 1M NaCl, pH4.0) until the effluent pH was 4.0, and the ions that were not tightly bound were washed to avoid leakage during chromatography.

- Equilibrate the column with equilibration buffer for later use.

- Buffer solution: Phosphate buffer is preferred, with a pH range of neutral (7~8), and EDTA and citrate are avoided. Common additive reagents and concentrations that do not affect and affect metal chelation chromatography are listed in Table 1 and Table 2 respectively.

In order to reduce the non-specific binding of the host protein to the resin, low concentrations of imidazole (20~40 mM) are usually added to the equilibration buffer and the sample.

NaCl of 0.15 ~ 0.5M must be added to the buffer solution to eliminate ion exchange.

Table 1 Additives that do not affect protein binding to immobilized metal ion affinity resin

Additives	Concentration	Additives	Concentration
Phosphate、Tris、Borate、 HEPES	20-100mmol/L	Non-ionic detergent	2%
NaCl	2mol/L	Triton X-100	2%
KCl	1mol/L	Tween-20	2%
Guanidine hydrochloride	6mol/L	Octyl glucoside	2%
Urea	8mol/L	Dodecyl maltoside	2%
glycerin	50%	C <sub>12</sub> E <sub>8</sub> ,C <sub>10</sub> E <sub>6</sub>	2%
Isopropanol	60%	PMSF (Protease inhibitor)	
Ethanol	30%	Pepsin inhibitor (Pepsin inhibitor)	1mmol/L
Amphoteric detergent (CHAPS)	1%	Leupeptin (Protease inhibitor)	1μmol/L
1% Benzamidine (Protease inhibitor)	1mmol/L		0.5μg/mL

Table 2 Additives that may disrupt protein binding to the immobilized metal ion affinity resin

Additives	Concentration	Additives	Concentration
2-mercaptoethanol	—	Histidine	Can be used instead of imidazole
Strong reducing agents (DTT and DTE)	—	Glycine	—
Chelating agents (EDTA and EGTA)	0.1mmol/L, take Ni <sup>2+</sup> from the resin	Glutamine	—
Ionic detergent (cholate, SDS)	—	Arginine	—
Sodium azide	3mmol/L	Ammonium chloride	—
Citrate	Can tolerate low concentrations		

- Sample preparation: In order to prevent blocking of the column, the sample needs to be filtered by microporous membrane of 0.45μm before loading, the pH and conductivity of the sample are adjusted to be consistent with the equilibration buffer, The loading volume is determined according to the substance content in the sample and the binding load of Chelating Bestarose FF.
- Rinse: Rinse with equilibration buffer until the UV absorption value drops to an appropriate value.
- Sampling: sample the prepared samples according to the set conditions.

**In order to reduce the impact of metal ion shedding on the protein, the test is to rinse 5CV with purified water before loading the sample, Then wash 5CV with elution buffer, and finally equilibrate the column with equilibration buffer before loading.**

- Elution: Substances bound to the chromatography column may be eluted in one of three ways, depending on the situation.

Competitive elution: linearly or progressively increase the concentration of substances with affinity to metal ions, such as 0~2M  $\text{NH}_4\text{Cl}$ , 0~0.5M imidazole, 0~0.5M histidine. Gradient elution is best performed at a constant pH of the equilibrium buffer.

The pH of the buffer can be lowered for elution: with the decrease of pH, the weak binding protein and the strong binding protein are eluted successively. When the pH of buffer drops below 4, the metal ions dissociate from the resin to achieve elution. (If the target protein is sensitive to low pH, it is recommended to add 1/10 volume of 1M Tris-HCl to the elution collection, pH9.0 for neutralization)

EGTA or EDTA solution with a chelating agent of 0.05M can dissociate metal ions from the resin to achieve elution. This method can also be used for elution of denatured or precipitated proteins. This method is not recommended. The metal ions in the eluted products can be removed by a desalting column.

- Regeneration: Impurity residue and shedding of metal ions will affect the column's chromatographic performance and reduce the load. Metal ions need to be re-chelated after every one to five cycles, depending on production needs.
  - Strip metal ion buffer (50mM PB, 0.5M NaCl, 0.1-0.2M EDTA, pH 7.0) in 2-5CV;  $\text{Fe}^{3+}$  tends to form insoluble in neutral solution, so it is recommended to use 50mM EDTA overnight to remove metal ions.
  - The residual EDTA was removed by 2~3CV of 0.5M NaCl passing through the column.
  - Use 0.5CV 0.2M metal ion solution chromatography column;
  - Remove unbound metal ions with 5CV purified water;
  - The chromatography column was cleaned with 5CV tric elution buffer;
  - Balance the chromatography column with a balancing buffer and set aside.

#### 4. Cleaning-in-place(CIP)

With the increasing use of chromatography resin, the accumulation of contaminants on the chromatography column is also increasing. Cleaning-in-place can prevent the accumulation of contaminants and maintain a stable working state. Determine the frequency of CIP according to the degree of contamination of the resin (if the contamination is serious, CIP should be carried out after each use to ensure repeatability of the results).

- First remove metal ions;
- To remove the protein adsorbed by ion exchange: wash the column with 2M NaCl solution of 2-3 times the bed volume, and then wash the column with distilled water of 3 times the bed volume;
- Precipitated or denatured material: can be removed with 1M NaOH for 0.5-1h;



- Hydrophobic binding substance: 2CV 70% ethanol or 30% isopropanol to wash the column, immediately with at least 5CV of filter-sterilized equilibration buffer, reverse washing.

## 5. Sterilization

Since the 20% ethanol or 2% benzyl alcohol preservation solution does not have sterilization and depyrogenation, it is recommended that Chelating Bestarose FF resin can be treated with 70% ethanol for more than 12 hours before or during use, or the resin after removing metal ions can be treated with 1M NaOH 0.5 -1h to reduce the risk of microbial contamination.

## 6. Storage

Chelating Bestarose FF is supplied in 20% ethanol or 2% benzyl alcohol. It should be stored in 20% ethanol and sealed at 2-30°C after use, in order to prevent ethanol volatilization and microbial growth, it is recommended to replace the storage solution every 3 months.

## 7. Disposal and Recycling

Chelating Bestarose FF is difficult to degrade in nature, incineration is recommended to protect the environment.

## 8. Order information

Product	Code No.	Pack size
Chelating Bestarose FF	AA206305	25mL
	AA206307	100mL
	AA0302	500mL
	AA0303	1L
	AA0304	5L
	AA0305	10L
	AA206315	20L

Prepacked columns	Code No.	Pack size
EzFast Chelating FF	EA206301	1×1mL
	EA206303	1×5mL
	EA206351	5×1mL
	EA206353	5×5mL
EzScreen Chelating FF	EA03025	1×4.9mL
	EA03035	5×4.9mL
EzLoad 16/10 Chelating FF	EA206304	1 pcs
EzLoad 26/10 Chelating FF	EA206306	1 pcs
EzLoad 10/20 Chelating FF	EA053	1 pcs