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(FOR RESEARCH USE ONLY. DO NOT USE IT IN CLINICAL DIAGNOSIS !)

Butyl Focurose HPL

Catalog No: E-CM-HI11

This manual must be read attentively and completely before using this product.

May you have any problems, please contact our Technical Service Center for help.

Phone: 240-252-7368(USA) 240-252-7376(USA) Email: <u>techsupport@elabscience.com</u> Website: <u>www.elabscience.com</u>

Please kindly provide us the lot number (on the outside of the box) of the kit for more efficient service.

Please read this manual carefully before use to ensure the performance and successful operation. If you have any questions, please contact our Technical Support.

Product introduction

Butyl Focurose HPL-coupled hydrophobic ligands can interact with some hydrophobic groups on the surface of proteins or antibodies under high ionic strength conditions (high ionic strength will increase the interaction between ligands and hydrophobic groups), thus to achieve the purpose of separation and purification. Butyl Focurose HPL is mainly used for the separation and purification of macromolecular substances (antibodies, viruses, etc.).

Advantages

- 1. Rapid, easy to use (one-step purification).
- 2. Compared with reversed-phase chromatography, the ligand concentration in the hydrophobic interaction chromatography media is low and the elution conditions are mild, which helps to maintain the biological activity of biomolecules.
- 3. Wide application. It can be used in moderate purification and fine purification alone, and can be repeatedly used in combination ion exchange chromatography media.
- 4. Large aperture, high capacity, and high resolution.

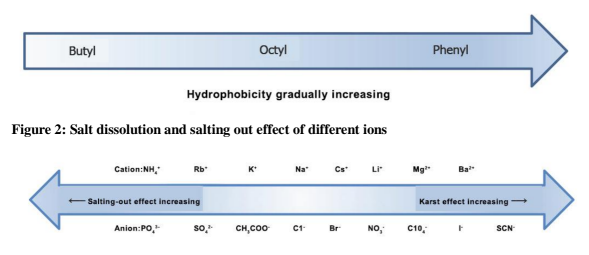
Matrix	Highly rigid agarose
Particle size range	45-165 μm
Average particle size	75 μm
Ligand density	40 µmol/mL (media)
pH stability	2-14 (short-term)
	3-13 (long-term)
Chemical stability	All of the commonly used buffers, 1M Acetic
	acid, 1M NaOH, 8M urea, 6M guanidine
	hydrochloride, 30% Isopropanol, 70% Ethanol
Maximum flow rate	300 cm/h
Storage buffer	20% Ethanol
Storage temperature	$4 \sim 30^{\circ}$ C ($4 \sim 8^{\circ}$ C is preferred)

Table 1: Performance index

Influence	Function mechanism	Suggestion
Ligand structure	The binding ability between different ligands and proteins is different.	Pre-experiment is recommended to screen suitable media, which can be referred to in Figure 1.
Ligand density	The higher the concentration of ligand, the stronger the binding ability will be.	Pre-experiment is recommended to screen the optimum ligand concentration.
Sample properties	The hydrophobicity of protein depends on the distribution of hydrophobic groups on its surface.	/
Salt concentration	The higher the salt concentration, the stronger the binding between the ligand and the protein will be, but the excessive high concentration of salt may result in protein precipitation.	Check the solubility and stability of protein under different salt concentrations.
Salt type	Different kinds of salts can result in different binding effects.	(NH ₄) ₂ SO ₄ and NaCl are preferred, other selections can refer to Figure 2.
Temperature	The higher the temperature, the stronger the protein hydrophobic will be.	Temperature must be maintained to be the same. Room temperature is recommended.
рН	Excessive high or low pH may affect the solubility and stability of protein, and pH can affect the binding effect.	The recommended pH range is 5.0- 8.5 on the premise of ensuring the solubility and stability of protein.

Table 2: Factors affecting hydrophobic chromatography

Figure 1: Hydrophobic properties comparison of ligands with different concentrations



Operation (take 1 mL column and 5 mL column for example)

1. Wash (water)

Wash the media with $5\sim10$ CV (column volume) of purified water with a flow rate of 0.5 mL/min (HT 1 mL) or 2.0 mL/min (HT 5 mL).

Note: This operation is used to remove the 20% ethanol in media.

2. Equilibration

Balance the media with $5 \sim 10$ CV of equilibrium liquid with a flow rate of 0.5 mL/min (HT 1 mL) or 2.0 mL/min (HT 5 mL) until the baseline turns stable and then set to zero.

Note: This procedure is used to balance the media. Ensure that the pH and component in media are in accordance with sample.

3. Sample application

Apply the sample with flow rate of 0.5 mL/min (1 mL) or 1.0 mL/min (5 mL) after centrifugation and filtration (0.45 μ m). Wash with the equilibrium liquid until the baseline tends to zero.

Note: Make sure that the ionic strength and pH of sample solution must be the same with the equilibrium liquid.

4. Elution (Choose the elution mode according to the equipment condition)

Linear gradient elution (with chromatographic system): Elute with 20 CV of 0%-100% eluent at a flow rate of 0.5 mL/min (1 mL) or 2.0 mL/min (5 mL) and collect the eluted solution.

Stepwise elution (with peristaltic pump): Stepwise elute by gradually decreasing the salt concentration with 20 CV of eluent at a flow rate of 0.5 mL/min (1 mL) or 2.0 mL/min (5 mL) and collect the eluted solution.

Note: Linear gradient elution is strongly recommended. A series of eluent* with different salt concentration should be prepared for stepwise elution.

*Eluent I : 0.05M PB, 1.50M (NH₄)₂SO₄, pH 7.0.

Eluent II : 0.05M PB, 1.25M (NH₄)₂SO₄, pH 7.0.

Eluent III: 0.05M PB, 1.00M (NH₄)₂SO₄, pH 7.0.

Eluent IV: 0.05M PB, 0.75M (NH₄)₂SO₄, pH 7.0.

Eluent V : 0.05M PB, 0.50M (NH₄)₂SO₄, pH 7.0.

Eluent VI: 0.05M PB, 0.25M (NH₄)₂SO₄, pH 7.0.

Eluent VII: 0.05M PB, pH 7.0.

5. Wash with water

Wash the media with $5\sim 10$ CV of purified water with a flow rate of 0.5 mL/min (1 mL) or 2.0 mL/min (5 mL).

Note: This procedure is used to remove the eluent in media.

6. Storage

Wash the media with 5~10 CV of 20% ethanol with a flow rate of 0.5 mL/min (1 mL) or 2.0 mL/min (5 mL) and the store the media.

Note: 20% ethanol can prevent the growth of microorganism. Media preserved with 20% ethanol can be stored at $4 \sim 30^{\circ}$ C ($4 \sim 8^{\circ}$ C is better).

7. Preparation of buffer

Equilibrium liquid: 0.05M PB, 1.70M (NH₄)₂SO₄, adjust the pH to 7.0. Store the prepared equilibrium liquid at room temperature.

Note: Some proteins may precipitate under high salt concentration. It is recommended to determine the protein solubility and stability under different salt concentration. pH should be ≤ 8.0 when using $(NH_4)_2SO_4$

Eluent: 0.05M PB, adjust the pH to 7.0. Store the prepared eluent at room temperature.

Storage buffer: 20% ethanol, store at room temperature.

Cleaning

The excellent performance of media (e.g. loading ability, mobility, column efficiency, etc.) can be recovered after cleaning the strong coupling substance (e.g. some strong coupling protein, denatured protein, lipids, etc.).

It is recommended to wash the media after used for each 5 times. The exact washing frequency should be adjusted according to the cleanliness of the purified sample.

1. Wash the media with 5~10 CV of purified water.

Note: This procedure is used to remove the 20% ethanol in media.

2. Wash the media with 5~10 CV of 1M NaOH and stand for 1 hour, then wash the media to neutral with purified water.

Note: This procedure is used to remove the precipitates or denatured substances accumulated in the media.

3. Wash the media with 5~10 CV of 70% ethanol or 30% isopropyl alcohol and stand for 1 hour, then wash the media with purified water immediately.

Note: This procedure is used to remove the strong hydrophobic binding substances.

4. Store the media after washed with 5~10 CV of 20% ethanol.

Note: 20% ethanol can prevent the growth of microorganism. Media preserved with 20% ethanol can be stored at $4 \sim 30^{\circ}$ C ($4 \sim 8^{\circ}$ C is preferred).

Problem **Possible cause** Suggestion Overloading of sample Decrease the sample volume. volume. Speed of sample loading is too Reduce the flow speed of sample The target compound fast. loading. does not combine with Protein or lipids accumulate in Wash the media timely and effectively. the media or the the media. combining amount is Increase the salt concentration of The salt concentration of low when purifying equilibrium liquid, or change the equilibrium liquid is low or the equilibrium liquid, or choose another hydrophobicity of target is type of media with stronger weak. hydrophobicity. The target compound does not Confirm whether the target combine combine with the media or the with media or not. combining amount is low. Decrease the flow rate and prolong the Insufficient elution time. No target compound retention time of eluent. Insufficient elution volume. Increase the elution volume. was collected or only a small amount of target Decrease the salt concentration of compound was equilibrium liquid, or change the salt collected The target and media combined type, or choose another type of media too strong. with weaker hydrophobicity, or add additives into the eluent (a small amount of detergent or low concentration organic reagent). Sample has not been Samples must be centrifuged or pretreated. filtered before loading. Dilute the sample properly with equilibrium liquid to decrease the High viscosity of sample. viscosity. Increase the washing volume until the Low purity of target Insufficient washing. baseline smooth and keep consistence compound with equilibrium liquid. Impurity protein or lipids Wash the media timely and effective. accumulate in the media. Poor elution condition, fast elution speed and abrupt Adjust the elution condition. elution gradient.

Trouble shootings

	Bad loading effect of column	Reload or re-purchase.	
	resin.		
	The top of separation column	Reload the column or reduce the	
	has a large volume of sample.	volume of the sample.	
	Wrong type of media.	Select the suitable hydrophobic media.	
	There is microbial grow in the	Correctly store the media after used.	
	media.		
Decrease of loading volume	Speed of sample loading is too	Reduce the flow speed of sample	
	fast.	loading.	
	Protein or lipids accumulate in the media.	Wash the media timely.	
	Ligand oxidized or dropped off	Wash the media timely or reload with	
	due to excessive use.	new media	
The chromatographic	The media was loaded too		
peak rises slowly	tight.	Reload the column.	
The chromatographic	The media was loaded too	Reload the column.	
peak trails	loose.		
The column bed cracks	Leakage occurred or a large	Check whether there is leaded or	
	volume of bubbles was	Check whether there is leakage or bubble, reload the column.	
or being dry	introduced.	bubble, leload the column.	
	Protein or lipids accumulate in	Wash the media or filter membrane	
	the media.	timely.	
		Adjust the content of equilibrium	
Flow of the column is	Protein precipitates in the	liquid and wash buffer to maintain the	
	media.	stability of target compound and	
exceedingly slow		combining efficiency of media.	
	There is microbial grow in the media.	Filter and degas all the reagents.	
		Samples must be centrifuged or	
		filtered before loading.	