

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

# **IMAC Focurose 6FF**

Catalog No: E-CM-AF25

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.

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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**

IMAC Focurose 6FF can be used for separation and purification with the interactions between chelated metal ion and certain amino acids (mainly include histidine, cysteine, tryptophan) on the side chain of protein, and it is suitable for separation and purification of His-tagged protein and biological molecules which interact with metal ion.

## **Advantages**

- 1. Quick and easy (one-step purification).
- 2. Wide application, simple-operation. Suitable for gravity column and prepacked column.
- 3. Multiple selection. Various metal ions (e.g. Cu<sup>2+</sup>, Zn<sup>2+</sup>, Fe<sup>2+</sup>, Co<sup>2+</sup>, Ca<sup>2+</sup>, etc.) can be used.
- 4. Lower Ni<sup>2+</sup> abscission probability and wider compatibility with reagents (Table 2) when compared with IDA Focurose 6FF.

Note: Avoid of using phosphate buffer when coupled with Ca<sup>2+</sup> (precipitate will occur).

## Performance index

Matrix	Highly cross-linked 6% agarose	
Particle size range	45-165 μm	
Average particle size	90 μm	
Binding capacity	40 mg (His-tag protein)/mL (media)	
II , 1'1', \$	3-12 (long-term)	
pH stability*	2-14 (short term)	
Chemical stability	0.01M HCl, 0.01M NaOH (1 week)	
	1M NaOH, 70% Ethanol (12 hours)	
	2% SDS (1 hour)	
	30% Isopropyl alcohol (30 minutes)	
Flow rate	600 cm/h	
Pressure	≤ 0.3 MPa	
Storage buffer	20% Ethanol	
Storage temperature	4~30°C (4~8°C is preferred)	

<sup>\*:</sup> The stability of media when unchelated with metal ions.

**Table 2: Compatibility with common reagents** 

ompatibility with common r	eagents
	0.05M sodium phosphate, pH 7.4
Buffer	0.1M Tris-HCl, pH 7.4
	0.1M Tris-acetate, pH 7.4
	0.1M HEPES, pH 7.4
	0.1M MOPS, pH 7.4
	0.1M sodium acetate, pH 4
Denaturant	8M Urea
	6M Gua-HCl
	2% Triton X-100
	2% Tween 20
Detergent	2% NP-40
	2% Cholate
	1% CHAPS
	0.005M DTE
	0.005M DTT
Reductant*	0.02M β-mercaptoethanol
	0.005M TCEP
	0.01M reduced glutathione
	0.5M Imidazole
	20% Ethanol
	50% Glycerol
Other additives	0.1M Na <sub>2</sub> SO <sub>4</sub>
	1.5M NaCl
	0.001M EDTA**
	0.06M Citrate

<sup>\*</sup> During the operation with IMAC Focurose 6FF, it is allowed to add low concentration of reductant, but it must be avoided of being immersed or stored for a long time with solution with reductant.

<sup>\*\*</sup> During the operation with IMAC Focurose 6FF, it is allowed to add nominal concentration of metal ions chelating agent (e.g. 0.001M EDTA) into samples of small volume, but it must be avoided of adding or loading a large volume of samples containing chelating agent into the purified solution.

#### Chelation method

Different metal ions can be selected for chelation, and different metal ion solutions  $(NiSO_4/CuSO_4/CoSO_4/ZnCl_2/CaCl_2/FeCl_2)$  can be selected for different samples.

1. Wash with 5~10 CV (column volume) of purified water.

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

2. Wash with 5~10 CV of 0.1M metal ion solution, and then wash the media with 5~10 CV of purified water after standing for 0.5 hours.

Note: This operation is used to couple and clean metal ions.

3. Wash with 5~10 CV of 20% ethanol and the store the media.

Note: 20% ethanol can prevent the growth of microorganism. Media preserved with 20% ethanol can be stored at  $4\sim30^{\circ}$ C ( $4\sim8^{\circ}$ C is preferred).

## 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 (1 mL) or 2.0 mL/min (5 mL).

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

#### 2. Equilibration

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

Note: This procedure is used to balance the media. Make sure that the pH and component in media are in accordance with equilibrium liquid.

## 3. Sample application

Apply the sample with flow rate of 0.2 mL/min (1 mL) or 1.0 mL/min (5 mL) after centrifugation and filtration (0.45  $\mu$ m). Wash with the equilibrium liquid until the baseline tends to zero.

Note: The binding capacity of proteins varies with the type of lysates, the properties of target proteins, flow rate, temperature, and pH. Low flow rates often increase the binding efficiency of samples.

#### 4. Wash

Wash out the impurity substance with  $5\sim10$  CV of washing solution with a flow rate of 0.5 mL/min (1 mL) or 2.0 mL/min (5 mL) and collect the washed solution.

Note: The washing solution is used to wash out some non-specific adsorbed impurity proteins.

## 5. Elution

Elute with  $5\sim10$  CV of eluent with a flow rate of 0.5 mL/min (1 mL) or 2.0 mL/min (5 mL) and collect the eluted solution.

Note: Low flow rate may increase the concentration of target protein in the eluted solution.

#### 6. Wash with water

Wash the media with  $5\sim10$  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.

## 7. Storage

Wash the media with  $5\sim10$  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  $4\sim30^{\circ}$ C ( $4\sim8^{\circ}$ C is preferred).

## 8. Preparation of buffer

**Equilibrium liquid:** 0.02M PB, 0.5M NaCl, adjust the pH to 7.4. Store the prepared equilibrium liquid at room temperature.

Note: NaCl in equilibrium liquid is used to inhibit the ion exchange of media.

**Washing buffer:** 0.02M PB, 0.5M NaCl, 0.02-0.04M Iminazole, adjust the pH to 7.4. Store the prepared equilibrium liquid at room temperature.

**Note:** Add 0.02-0.04M Iminazole to the washing buffer (yield preferred) or to the equilibrium liquid directly (purity preferred) according to the final application demands.

**Eluent:** 0.02M PB, 0.5M NaCl, 0.5M Iminazole, adjust the pH to 7.4. Store the prepared eluent at room temperature.

**Note:** 0.05-0.25M Iminazole in eluent is enough to elute the target protein in general condition.

# 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 eluent (wash after using) or 20% ethanol (wash before using).
- 2. Wash the media with 5~10 CV of 0.02M Tris-HCl, 0.1M EDTA (pH 8.0), then wash the media with 5~10 CV of purified water immediately.
  - Note: This procedure is used to remove Ni<sup>2+</sup>.
- 3. Wash the media with 5~10 CV of 1.0M NaOH. Wash the media until the pH to neutral with purified water after standing for 1 hour.
  - Note: This procedure is used to remove the precipitates or lipids accumulated in the media.
- 4. Wash the media with 5~10 CV of 0.1M NiSO<sub>4</sub>, then wash the media with 5~10 CV of purified water after standing for 0.5 hour.
  - Note: This procedure is used to couple Ni<sup>2+</sup>.
- 5. 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  $4\sim30^{\circ}$ C ( $4\sim8^{\circ}$ C is preferred).

**Trouble shootings** 

Problem	Possible cause	Suggestion
	Overloading of sample volume.	Decrease the sample volume.
	Speed of sample leading is too fast	Reduce the flow speed of sample
	Speed of sample loading is too fast.	loading.
	Protein or lipids accumulate in the	Wash the media timely and
	media.	effectively.
The target compound	The expression condition is too	An empty vector is suggested to
does not combine with	severe that the His-tag was	set as control of expression and
the media or the	enwrapped and cannot combine with	purification to check whether the
combining amount is	the media.	expression condition is suitable.
low when purifying	The initial sample does not contain	Verify through the gene sequence
	His-tagged protein.	or His-tag antibody.
		The target protein has not been
	The target protein occurs in the flowthrough.	expressed successfully, or the pH
		and components in sample and
		equilibrium liquid were incorrect
	The target compound does not	Confirm whether the target
	combine with the media or the combining amount is low.	combine with media or not.
	Unsuitable elution condition.	Increase the iminazole
		concentration in eluent.
No target compound	Insufficient elution time.	Decrease the flow rate and prolong
was collected or only a		the retention time of eluent.
small amount of target compound was	Insufficient elution volume.	Increase the elution volume
	Target protein was washed out	Decrease the iminazole
collected	during the washing.	concentration in washing buffer.
	The target compound accumulates in the elution buffer.	Check the stability of target
		compound in the elution buffer
		(salt concentration, pH, etc.) Add
		some additives into the eluent,
		such as 0.2%Triton X-100 or 0.5%
		Tween 20.
Low purity of target compound	Sample has not been pretreated.	Samples must be centrifuged or
		filtered before loading.
	High viscosity of sample.	Dilute the sample properly with
		equilibrium liquid to decrease the
		viscosity.
	Insufficient washing.	Increase the washing volume until

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		the baseline smooth and keep
		consistence with equilibrium
		liquid.
	Impurity protein or lipids accumulate	Wash the media timely and
	in the media.	effective.
	Impurity substances have strong-	Purify with other type of media
		(e.g. ion–sieve or molecular-
	affinity with Ni <sup>2+</sup> .	sieve).
		Determine the stability of target
	The target compound degrades	compound and add protease
	The target compound degrades	inhibitors.
	Bad loading effect of column resin.	Reload or re-purchase.
	Bad loading effect of column resin.	Add appropriate additive to reduce
	Non-specificity absorption of	the non-specific absorption, such
	impurity materials.	as 0.5% Triton X-100, 1.0% Tween
	impurity materials.	
	TT	20 or 50% glycerol.
	The top of separation column has a	Reload the column or reduce the
	large volume of sample.	volume of the sample
	There is microbial grow in the	Correctly store the media after
	media.	used
	Speed of sample loading is too fast.	Reduce the flow speed of sample
		loading.
	Protein or lipids accumulate in the	Wash the media timely.
	media.	, , usia use siscosu usino-j,
Decrease of loading	Ligand dropped off due to excessive use.	Re-couple with new media.
volume		
	The expression condition is too	An empty vector is suggested to
	severe that the His-tag was	set as control of expression and
	enwrapped and cannot combine with	purification to check whether the
	the media.	expression condition is suitable.
The chromatographic	The media was local-day disha	Daland the column
peak rises slowly	The media was loaded too tight.	Reload the column.
The chromatographic		B
peak trails	The media was loaded too loose.	Reload the column.
The column bed cracks	Leakage occurred or a large volume	Check whether there is leakage or
or being dry	of bubbles was introduced.	bubble, reload the column.
	Protein or lipids accumulate in the	Wash the media or filter
Flow of the column is exceedingly slow	media.	membrane timely.
	Protein precipitates in the media.	Adjust the content of equilibrium
	1 Totalii precipitates in the media.	1 Lajast the content of equilibrium

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		liquid and wash buffer to maintain
		the stability of target compound
		and combining efficiency of
		media.
	771	Filter and degas all the reagents.
	There is microbial grow in the	Samples must be centrifuged or
media.	media.	filtered before loading.