Instructions 28-9413-30 AD

HiPrep IMAC FF 16/10

Introduction

HiPrepTM IMAC FF 16/10 is a ready to use column, prepacked with uncharged IMAC SepharoseTM 6 Fast Flow. This column is ideal for preparative purification of histidine-tagged recombinant proteins and untagged, naturally occurring proteins. HiPrep IMAC FF 16/10 provides fast, simple and easy separations in a convenient format, and IMAC Sepharose 6 Fast Flow is ideal for scaling up.

Column data

 Medium
 IMAC Sepharose 6 Fast Flow

 Bead structure
 Highly cross-linked 6% agarose

Mean particle size 90 µm

Dynamic binding capacity* Approx. 40 mg (histidine) ₆-tagged protein/ml medium (Ni²--charged). Untagged protein: Approx. 25 mg/ml

medium (Cu^{2+} charged), or approx. 15 mg/ml medium ($7n^{2+}$ or Ni^{2+} charged)

Metal ion capacity Approx. 15 µmol Ni²+/ml medium

 Bed volume
 20 ml

 Bed diameter × height
 16 × 100 mm

 Column hardware
 Polypropylene

 Recommended flow rate^{† †}
 1–10 ml/min (30–300 cm/h)

 Maximum flow rate^{† †}
 10 ml/min (300 cm/h)

Maximum pressure over the packed bed during

operation, Δp[‡]

HiPrep column hardware pressure limit‡

Compatibility during use Chemical stability

(for medium without metal ion)

pH stability

(for medium without metal ion)
Storage

0.15 MPa, 1.5 bar, 22 psi 0.5 MPa, 5 bar, 73 psi

See Table in "Buffers and compatibility" section. 1 M NaOH, 70% acetic acid. Tested for 12 h.

2% SDS. Tested for 1 h. 30% 2-propanol. Tested for 30 min.

short term (at least 2 h): 2–14 long term (≤ one week): 3–12 4°C to 30°C in 20% ethanol

* Dynamic binding capacity conditions

Samples: Histidinel_s-tagged proteins: Capacity data were obtained for a protein (Mr, 28 000) bound from an *E. coli* extract, and a pure protein (Mr, 43 000;

(m, 28 000) bound from the Econ extract, and a part protein (m, 48 000 applied at 1 mg/ml in binding buffer; capacity at 10% breakthrough).

<u>Untagged protein:</u> Capacities determined at 10% breakthrough for human apo-transferrin applied at 1 mg/ml in binding buffer.

Column volume: 0.25 or 1 ml

Flow rate: 0.25 or 1 ml/min, respectively

Binding buffer: 20 mM sodium phosphate, 500 mM NaCl, 5 mM imidazole,

(1 mM for untagged protein) pH 7.4

Elution buffer: 20 mM sodium phosphate, 500 mM NaCl, 500 mM imidazole,

(50 mM for untagged protein) pH 7.4

Note: Dynamic binding capacity is metal-ion- and protein-dependent.

- † Water at room temperature. Flow rate is determined by $v \bullet \eta \le 10$ ml/min where v = flow rate and $\eta = v$ is cosity.
- # Many chromatography systems are equipped with pressure gauges to measure the pressure at a particular point in the system, usually just after the pumps. The pressure measured here is the sum of the precolumn pressure, the pressure drop over the medium bed, and the post-column pressure. This pressure is always higher than the pressure drop over the bed alone. We recommend keeping the pressure drop over the bed below 1.5 bar. Setting the upper limit of your pressure gauge to 1.5 bar will ensure the pump shuts down before the medium is overpressured. If necessary, post-column pressure of up to 3.5 bar can be added to the limit without exceeding the column hardware limit. To determine post-column pressure, proceed as follows:

To avoid breaking the column, the post-column pressure must not exceed 3.5 bar. $\label{eq:column}$

- 1. Connect a piece of tubing in place of the column.
- Run the pump at the maximum flow you intend to use for chromatography. Use a buffer with the same viscosity as you intend to use for chromatography. Note the back pressure as total pressure.
- 3. Disconnect the tubing and run at the same flow rate used in step 2. Note this back pressure as precolumn pressure.
- 4. Calculate the post-column pressure as total pressure minus precolumn pressure. If the post-column pressure is higher than 3.5 bar, take steps to reduce it (shorten tubing, clear clogged tubing, or change flow restrictors) and perform steps 1–4 again until the post-column pressure is below 3.5 bar. When the post-column pressure is satisfactory, add the post-column pressure to 1.5 bar and set this as the upper pressure limit on the chromatography system.





First time use

- 1. Charge with metal ions (see below).
- 2. Set an appropriate pressure limit.
- 3. Equilibrate the column with 100 ml binding buffer.

HiPrep IMAC FF 16/10 can be used directly on ÄKTA™ design systems without the need for extra connectors.

Try these conditions first

Binding buffer for histidine-tagged proteins*: 20 mM sodium phosphate, 500 mM NaCl, 20-40 mM imidazole, pH 7.4 (The imidazole concentration for optimal purity/yield is proteindependent, see "Optimization)".

Elution buffer for histidine-tagged proteins*:

f: 20 mM sodium phosphate, 500 mM NaCl, 500 mM imidazole, pH 7.4

Flow rate: 1-10 ml/min (30-300 cm/h)

* For untagged target proteins, the imidazole concentrations that should be used are usually lower than the above, both for binding (sometimes no imidazole used) and elution.

Note: Especially for untagged target proteins, low-pH elution is an alternative to competitive elution with imidazole, for example a linear gradient from pH 7.4 to pH 4.

De-gas and filter all solutions through a 0.45-µm filter to increase column lifetime. High purity imidazole gives very low or no absorbance at 280 nm.



Buffers and compatibility

IMAC Sepharose 6 Fast Flow charged with Ni²⁺ is compatible with:

Reducing agents 5 mM DTE (See Blank run below) 5 mM DTT

20 mM ß-mercaptoethanol

5 mM TCEP (Tris[2-carboxyethyl] phosphine)

10 mM reduced glutathione

Denaturing agents 8 M urea* 6 M Gua-HCl*

Detergents 2% Triton™ X-100 (nonionic)

2% Tween™ 20 (nonionic) 2% NP-40 (nonionic) 2% cholate (anionic) 1% CHAPS (zwitterionic) 500 mM imidazole 20% ethanol

20% ethanol 50% glycerol 100 mM Na₂SO₄ 1.5 M NaCl

1 mM EDTA[†] 60 mM citrate[†]

Buffer substances 50 mM sodium phosphate, pH 7.

50 mM sodium phosphate, pH 7.4 100 mM Tris-HCl, pH 7.4 100 mM Tris-acetate, pH 7.4 100 mM HEPES, pH 7.4 100 mM MOPS, pH 7.4 100 mM sodium acetate, pH 4*

* Tested for one week at 40°C

Other additives

The strong chelator EDTA has been used successfully in some cases at 1 mM. Generally, chelating agents should be used with caution (and only in the sample, not the buffer). Any metal-ion stripping may be counteracted by adding a small excess of MgCl $_2$ before centrifugation/filtration of the sample. Note that stripping effects may vary with the applied sample volume.

Optional: Blank run

Note: Perform a blank run without reducing agents before applying buffers/ samples containing reducing agents. Likewise, a blank run is recommended for critical purifications where metal ion leakage during purification must be minimized

Use binding buffer and elution buffer without reducing agents.

- If the column has been stored in 20% ethanol after metal ion charging, wash it with 100 ml distilled water.
- Wash with 100 ml the buffer that has been chosen for protein elution, for example., imidazole elution buffer or low-pH elution buffer
- 3. Equilibrate with 100–200 ml binding buffer. Imidazole equilibration can be monitored by absorbance, for example. at 220 nm.

Charging with metal ions



- Charge the water-washed column by loading 10 ml 0.1 M metal-ion solution in distilled water. For example chlorides and sulfates can be used. For choice of metal ion, see "Optimization".
- Wash with 100 ml distilled water and 100 ml binding buffer (washing with binding buffer – to adjust pH – should be done even if the metal-charged column is only to be stored in 20% ethanoll
- In some cases, a blank run may be needed for optimal performance, see "Blank run".

Note: The column does not have to be stripped and recharged between each purification, if the same protein is going to be purified; it may be sufficient to strip and recharge it after approximately five purifications, depending on the sample properties, sample volumes, metal ion, etc.



Metal ion stripping

Stripping buffer: 20 mM sodium phosphate, 500 mM NaCl, 50 mM EDTA, pH 7.4

- .. Wash with at least 100–200 ml stripping buffer
- 2. Wash with at least 100–200 ml binding buffer
- . Wash with at least 100–200 ml distilled water
- Clean the column, see "Cleaning-in-Place (CIP)" and/or recharge with metal ions.



Avoid

Chelating agents in buffers, for example EDTA, EGTA, and citrate. Unfiltered solutions.

Sample preparation

Centrifuge at 10 000 \times g (or higher) for 10 min and/or filter the sample through 0.45- μ m filter. If possible, dilute the sample in binding buffer. The sample should contain imidazole at the same concentration as in the binding buffer.



Delivery/storage

The column is supplied in 20% ethanol.

If the column is to be stored for a longer period, clean it according to the procedure described under "Cleaning-in-Place (CIP)". Then equilibrate with at least 50 ml 20% ethanol.

Note: HiPrep columns cannot be opened or refilled.



Optimization

Perform your first run according to "First time use" and "Try these conditions first". If the results are unsatisfactory, consider the following:

Action	Effect
Increase the imidazole concentration in the sample and binding buffer.	Decreases the amount of contaminants binding to the medium.
Increase the imidazole concentration in the binding/wash buffer.	Washes out contaminants bound to the medium more effectively.

Elute with a stepwise or linear imidazole gradient to determine the optimal imidazole concentrations to use for binding and washing; add imidazole to the sample to the same concentration as in the binding buffer. Wash before elution with binding buffer containing the highest possible concentration of imidazole that does not cause elution of the target protein.

Note: There is an optimal imidazole concentration at binding and wash that will balance high purity and high yield. This optimal concentration is different for different histidine-tagged proteins.

Note: The possibility of reusing HiPrep IMAC FF 16/10 depends on the properties of the samples and on the metal ion used. Beware of cross-contamination if using the same column for purification of more than one target protein.

 Ni^{2+} is usually the first-choice metal ion for purifying most histidine-tagged proteins. For purification of untagged proteins, Cu^{2+} ions have frequently been used. When the binding characteristics of an untagged target protein are not known, it is advisable to test other metal ions (for example, Zn^{2+} , Ni^{2+} , Co^{2+}) to establish the most suitable metal ion to use. In some instances, a weak binding to a metal ion can be exploited to achieve selective elution (higher purity) of a target protein. In some special applications, Fe^{3+} and Co^{2+} have also been used.

Cleaning-in-Place (CIP)

Decreased binding capacity and/or increased back pressure may be due to an accumulation of debris or of precipitated, denatured, or non-specifically bound proteins. These problems can be solved using the procedures described below. For difficult cases, use reversed flow direction.

Note: Before cleaning, strip off the metal ions by using the recommended procedure. Stripping, without any additional CIP procedures, may sometimes give a satisfactory cleaning effect.

- Removal of ionically bound substances:
 Wash with approximately 20 ml 1.5 M NaCl. Then wash the column with approximately 200 ml distilled water.
- Removal of precipitated and/or hydrophobically-bound substances and lipoproteins: Wash with 1 M NaOH, contact time usually 1–2 h (longer time may be required to inactivate endotoxins); then wash with approximately 200 ml binding buffer, followed by 100–200 ml distilled water.
- Removal of hydrophobically-bound proteins, lipoproteins, and lipids: Wash with 100-200 ml 30% isopropanol for at least 15-20 min; then wash with approximately 200 ml distilled water. Alternatively, wash with 40 ml detergent in a basic or acidic solution. Use, for example, 0.1-0.5% nonionic detergent in 0.1 M acetic acid, contact time 1-2 h. After treatment, always remove residual detergent by washing with at least 100 ml 70% ethanol. Then wash with approximately 200 ml distilled water.

Note: HiPrep columns cannot be opened or refilled.

Troubleshooting

The following tips serve as a guide.

Note: Proteins generally unfold when using high concentrations of urea or Gua-HCl (as described below). Refolding on-column (or after elution) is proteindependent. Samples containing urea can be analyzed directly by SDS-PAGE whereas samples containing Gua-HCl must be buffer-exchanged to a buffer with urea before SDS-PAGE. Solid urea or Gua-HCl can be added to the sample to minimize dilution.

Symptom Remedy

Column has clogged

Sample is too viscous

Protein is difficult to dissolve or precipitates during

Cell debris in the sample may clog the column. Clean the column according to "Cleaning-in-Place". It is important to filter and/or centrifuge the sample before loading, see "Sample preparation".

If the lysate is very viscous due to the presence of a high concentration of host nucleic acid, continue sonication until the viscosity is reduced, and/or add DNase I to 5 μ g/ml, Mg²-to 1 mM and incubate on ice for 10–15 min. Alternatively, draw the lysate through a syringe needle several times.

The following additives may help: 2% Triton X-100, 2% Tween 20, 2% NP-40, 2% cholate, 1% CHAPS, 1.5 M NaCl, 50% glycerol, 20 mM β-mercaptoethanol, 1-3 mM DTT or DTE (up to 5 mM possible but is protein-dependent), 5 mM TCEP, 10 mM reduced glutathione, 8 M urea, or 6 M Gua-HCl. Mix gently for 30 minutes to aid solubilization of the tagged protein (inclusion bodies may require longer mixing). Note that Triton X-100 and NP-40 (but not Tween) have a high absorbance at 280 nm. Furthermore, detergents cannot be easily removed by buffer exchange.

No histidine-tagged protein in the purified fractions

SDS-PAGE of samples collected

bacterial lysate indicates that

most histidine-tagged protein is

located in the centrifugation

The eluted protein is not pure

(multiple bands on SDS

polyacrylamide gel)

pellet

during preparation of the

Elution conditions are too mild (histidine-tagged protein still bound): Elute with an increasing imidazole or decreasing pH gradient to determine the optimal elution conditions.

Protein has precipitated on the column: For the next experiment, decrease the amount of sample, or decrease protein concentration by eluting with a linear imidazole gradient instead of steps. Elute under denaturing (unfolding) conditions (use 4–8 M urea or 4–6 M Gua-HCI).

Nonspecific, hydrophobic or other interaction:
Add a nonionic detergent to the elution buffer (e.g. 0.2%
Triton X-100) or change the NaCl concentration.

Concentration of imidazole in the binding buffer is too high: The protein is found in the flowthrough. Decrease the imidazole concentration in the binding buffer.

Target protein may not be histidine-tagged as expected: Verify DNA sequence of the gene. Analyze samples taken before and after induction of expression with, e.g., anti-His antibodies in Western blotting.

Histidine-tag may be insufficiently exposed: The protein is found in the flowthrough; perform purification of unfolded protein in urea or Gua-HCl as for inclusion bodies.

Buffer/sample composition is incorrect: The protein is found in the flowthrough; check pH and composition of sample and binding buffer. Ensure that the concentration of chelating or strong reducing agents, as well as imidazole in the sample, is not too high.

Sonication may be insufficient: Check cell disruption by microscopic examination or monitor by measuring the release of nucleic acids at 260 nm. Adding lysozyme (up to 0.1 volume of a 10 mg/ml lysozyme solution in 25 mM Tris-HO, pH 8.0) prior to sonication may improve results. Avoid overheating and foaming as this may denature the tagged protein. Over-sonication can also lead to copurification of host proteins with the tagged protein.

Protein may be insoluble (inclusion bodies): The protein can usually be solubilized (and unfolded) from inclusion bodies using common denaturants such as 4–6 M Gua-HCl, 4–8 M urea, or strong detergents. Prepare buffers containing 20 mM sodium phosphate, 8 M urea or 6 M Gua-HCl and suitable imidazole concentrations, pH 7.4–7.6. Buffers with urea should also include 500 mM NaCl. Use these buffers for sample preparation, as binding buffer and as elution buffer. For sample preparation and binding buffer, use 20 mM imidazole or the concentration selected during the optimization trials (including urea or Gua-HCl).

Partial degradation of tagged protein by proteases:
Add protease inhibitors (use EDTA with caution,

see "Compatibility").
Contaminants have high affinity for the metal ion used:

See "Optimization" instructions. A shallow imidazole gradient (20 column volumes or more), may separate proteins with similar binding strengths. If optimized conditions do not remove contaminants, further purification by ion exchange chromotography (using HiTrap™ Q HP or HiTrap SP HP) and/or gel filtration (using Superdex™ Peptide, Superdex 75 or Superdex 200) may be necessary.

Contaminants are associated with tagged proteins:

Add detergent and/or reducing agents before sonicating cells, or shortly afterwards if foaming is a problem. Increase detergent levels (for example up to 2% Triton X-100 or 2% Tween 20), change the NaCl concentration, or add glycerol (up to 50%) to the wash buffer to disrupt nonspecific interactions.

Change metal ion: The metal ion used may not be the most suitable, see Optimization.

Intended use

Product

The HiPrep IMAC FF 16/10 is intended for research use only, and shall not be used in any clinical or *in vitro* procedures for diagnostic purposes.

No. per pack

Code No.

Ordering information

HiPrep IMAC FF 16/10	1 x 20 ml	28-9365-52
Companion products		
HiTrap IMAC FF	5 x 1 ml	17-0921-02
HiTrap IMAC FF	5 × 5 ml	17-0921-02
IMAC Sepharose 6 Fast Flow	25 ml	17-0921-04
IMAC Sepharose 6 Fast Flow	100 ml	17-0921-07
HisTrap™ FF crude	5 x 1 ml	11-0004-58
HisTrap FF crude	100 x 1 ml*	11-0004-59
HisTrap FF crude	5 x 5 ml	17-5286-01
HisTrap FF crude	100 x 5 ml*	17-5286-02
HisTrap FF	5 × 1 ml	17-5319-01
HisTrap FF	100 × 1 ml*	17-5319-02
HisTrap FF	5 × 5 ml	17-5255-01
HisTrap FF	100 × 5 ml*	17-5255-02
HisPrep™ FF 16/10	1 x 20 ml	28-9365-51
Ni Sepharose 6 Fast Flow	5 ml	17-5318-06
Ni Sepharose 6 Fast Flow	25 ml	17-5318-01
Ni Sepharose 6 Fast Flow	100 ml	17-5318-02
Ni Sepharose 6 Fast Flow	500 ml	17-5318-03
HiPrep 26/10 Desalting	1 x 53 ml	17-5087-01
HiPrep 26/10 Desalting	4 x 53 ml	17-5087-02
HiTrap Desalting	5 × 5 ml	17-1408-01
HiTrap Desalting	100 × 5 ml*	11-0003-29

 $^{^{\}star}\,$ Pack size available by special order.

Accessories

To connect columns with 1/16" connectors to FPLC™ System:				
Union M6 female/1/16" male	5	18-3858-01		
HiTrap/HiPrep 1/16" male connector for ÄKTA design	8	28-4010-81		

Related printed literature

he Recombinant Protein Purification Handbook, Principles and Methods	18-1142-75
ffinity Chromatography Handbook, Principles and methods	18-1022-29
ffinity Chromatography Columns and Media, Selection guide	18-1121-86
Ii Sepharose and IMAC Sepharose, Selection guide	28-4070-92
repacked chromatography columns for ÄKTA design,	
Selection guide	28-9317-78

Further information

For more information, please visit

www.gelifesciences.com/protein-purification www.gelifesciences.com/purification_techsupport

Refer also to the handbooks above, contact our technical support team, or your local representative.

www.gelifesciences.com/protein-purification www.gelifesciences.com/purification_techsupport

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Purification and preparation of fusion proteins and affinity peptides comprising at least two adjacent histidine residues may require a license under US patents 5,284,933 and 5,310,663, including corresponding foreign patents (assignee: Hoffmann-La Roche, Inc).

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