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Protein purification strep-tag on gravity column

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1 Works for me dx.doi.org/10.17504/protocols.io.7qahmse

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ABSTRACT

Protein purification of a protein with a strep-tag on a gravity column.

GUIDELINES

After lysis, sample should be kept on gel as much as possible

MATERIALS

NAME	CATALOG #	VENDOR
MilliQ water		
Roche Complete Protease Inhibitor EDTA-Free tablets	5056489001	Sigma Aldrich

MATERIALS TEXT

50 % Strep-Tactinsepharose solution

BEFORE STARTING

Have performed Protein expression using E. coli strain BL21DE3 by Robert Hoofman

buffers

- Buffer W:
 - 100 mM Tris-HCl (pH 8.0)
 - 150 mM NaCl
 - 1 mM EDTA

Buffer E:






- 100 mM Tris-HCl (pH 8.0)
- 150 mM NaCl
- 1 mM EDTA
- 2.5 mM biotin

Protein extraction

- Place post-induction culture from Protein expression using E. coli strain BL21DE3 on ice



- Centrifuge culture at 5000xg for 00:10:00 at 4 °C

1m

- 4 The pellet is resuspended in  **1 ml** buffer W per  **100 ml** cell culture containing one crushed cOmplete mini tablet.
- 5 The cells are sonificated (VS70 T rod, 25% 1 sec on 2 sec off for  **00:05:00** . On ice water). For small amounts use the MS72 rod.
From this point be very sure to keep the cells on ice as much as possible.
- 6 The cell extract is centrifuged for  **00:45:00** at 30 000 g.
- 7 The supernatant is collected and filtered first with 0.45 um filter and then with 0,22 um filter. If there are small sample volumes: add  **3 ml** buffer W.

4m

Protein purification

- 8 Fill centrifuge Column with 200-300 µL Strep- Tactinsepharose (so 400-600 µL 50% suspension). This is your CV.
- 9 Equilibrate with 3 Column bed Volume (CV) of Buffer W.
- 10 Load 10 CV of cell free extract on the column
- 11 Wash the resin 5 times by 1 CV of buffer W
Collect this in Eppendorf tubes per 1 CV. Apply  **2 µl** of the first washing fraction and  **8 µl** of each subsequent fraction to an analytical SDS-PAGE.
- 12 Apply 0.5 CV of Buffer E on column 6 times to elute the protein of interest
8 µl samples of each fraction can be used for SDS-PAGE analysis. Most of the purified Strep-tag® II fusion protein usually elutes in the 2nd to 5th fraction.
- 13 Determine protein concentration flow through on nanodrop with the molecular weight and extinction coefficient
- 14 Run on an appropriate SDS-page gel



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