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## Protein purification strep-tag on gravity column

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1 Works for me

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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 Hooftman

# 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

1m

Centrifuge culture at 5000xg for **© 00:10:00** at § 4 °C

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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.
	Trom the point severy care to keep the color of fee as mash as possible.
6	The cell extract is centrifuged for <b>© 00:45:00</b> 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 $3 \text{ ml}$ buffer W.
Protein purification	
8	Fill centrifuge Column with 200-300 $\mu$ L Strep- Tactinsepharose (so 400-600 $\mu$ L 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 $2 \mu$ of the first washing fraction and $8 \mu$ 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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