





ChroSpin - ProteinG 👄

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dx.doi.org/10.17504/protocols.io.pyfdptn



ABSTRACT

Purification Guide for the Isolation of Antibodies with ChroSpin Columns by DALEX Biotech.

Easy and guick small scale antibody purification from various sources and species.

Each ChroSpin column has a binding capacity of > 2 mg (tested with human polyclonal Ig, binding varies between species and clones).

The proprietary resin does not shrink or swell in aqueous buffers.

High pressure stability.

pH stability short term 2 - 8, long term 3 - 8.

Excellent thermal stability up to 15 minutes at 80 °C in aqueous buffers at neutral pH.

Can be dried for long term storage (80 °C for > 2 h).

EXTERNAL LINK

https://dalex-biotech.com/

PROTOCOL STATUS

Working

Official product protocol by DALEX Biotech.

GUIDELINES

For optimal binding and purity, the pH of the sample should be 7-8 and should contain 150-300 mM NaCl. An easy way to achieve this is by adding 1/11 volume 10x PBS to your sample.

Purification works best with an antibody concentration of 2-3 mg/ml in your sample.

It is advisable that all fractions are collected (Sample, flow through, wash, and eluate) in separate tubes for analysis, e.g. SDS-PAGE.

MATERIALS TEXT

Materials provided in the kit:

ChroSpin columns

Wash buffer

Flution buffer

Neutralization buffer

Sanitization solution

Materials not provided in the kit:

Tween-20

10x PBS

Deionized water

20 % ethanol

The buffers in the kit include sodium azide (CAS No. 26628-22-8) as a preservative.

For safety information on this chemical(s) check http://www.dguv.de/ifa/gestis-database

BEFORE STARTING

Make sure your sample is free of particulate matter. You can remove particles by centrifugation or filtration (0.45 μm).

What do you want to do?

1 Do you want to purify antibodies or sanitize your column? Please choose below.

step case

Purification

no description provided

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Equilibration

2 Remove the bottom cap, add 500 μl wash buffer to the column, close the lid, invert sharply 3 - 5 times, and centrifuge 30 - 60 seconds at 1000 g.



If you work with a used column, remove the storage solution by centrifugation first.

Load and Wash

3 Add up to 500 µl sample to the top of the column and incubate the column with end-over-end mixing or occasional inversion for 3 minutes.



NOTE

For fast binding antibodies the incubation time can be decreased to 30 seconds. For slow binding antibodies the incubation time can be increased to 10 minutes. Longer incubation time is possible, however, this will also increase background.

For optimal binding and purity, the pH of the sample should be 7 - 8 and should contain 150 - 300 mM NaCl. An easy way to achieve this is by adding 1/11 volume 10x PBS to your sample.

Purification works best with an antibody concentration of 2 - 3 mg in your sample.

4 Centrifuge the column for 30 - 60 seconds at 1000 g and place the column into a clean receiver tube.

NOTE

It is advisable to keep the flow-through and wash fractions for later analysis, e.g. SDS-PAGE.

5 Add 500 µl wash buffer to the column, close the lid, and invert 5 times. Centrifuge the column for 30 - 60 seconds at 1000 g and empty the receiver tube.

NOTE

In case of unspecific hydrophobic and/or ionic interactions include up to 1 % Tween-20 and/or up to 0.5 M NaCl in the wash

L L	buffer.
6 Re	epeat the previous step.
	ENOTE For increased purity, repeat the washing step up to 5 times.
Elution	
7 Pla	ace the column into a clean receiver tube, add 200 μl elution buffer to the column, close the lid, and invert three times. Centrifuge the olumn for 30 - 60 seconds at 1000 g.
0	eave receiver tube in place. Add 200 μl elution buffer to the column, close the lid and centrifuge for 30 - 60 seconds at 1000 g. epeat once more.
9 Ad	dd two drops of neutralization solution to the eluate and mix gently.
Cleaning	and Storage
50	ash the column by adding 500 µl elution buffer and centrifuge at 1000 g for 30 - 60 seconds. Repeat this step with 500 µl wash buffer and 00 µl deionized water. Then, close the bottom of the column, add 500 µl 20 % ethanol or wash buffer (contains 0.05 % (w/v) sodium azide). ose the lid and store at room temperature or at 4 - 8 °C.
Dr	ternative for long-term storage: y the open (top and bottom) column in an oven at 80 °C for at least 2 hours or over night. Make sure the bottom stopper is completely dry, o. Put on the column's outlet, close the lid and store the column closed at room temperature.
Sanitizati	ion
	step case —
Sa	anitization
Af	ter five purification cycles or after a detectable decrease in capacity a sanitization of the column is recommended.
_	iter purification: wash the column with 500 μl elution buffer, centrifuge the column for 30 - 60 seconds at 1000 g, and repeat with 500 μl ash buffer.
If t	the column was stored before: remove the bottom cap and centrifuge the column for 30 - 60 seconds at 1000 g. Then, add 500 µl of wash after and centrifuge the column for 30 - 60 seconds at 1000 g.
3 Cld	ose the bottom of the column, add 500 μl sanitization solution. Close the lid and incubate for one hour at RT.

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