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created by Lizbé Koekemoer

Parallel rapid expression and purification of proteins for crystallography (PREPX): 48x 100 mL cultures V.2

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ASAP Discovery



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ABSTRACT

This protocol details the parallel rapid expression and purification of 24x proteins for crystallography (PREPX) at 100 mL culture scale. Recombinant proteins are expressed in *Escherichia coli* using the autoinduction method and then purified in parallel using a IMAC, desalt, tag cleavage, reverse IMAC and gel filtration work flow.

ATTACHMENTS

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GUIDELINES

Method overview

Standard workflow is expression via autoinduction followed by purification using IMAC/PD-10/revIMAC and serial gel filtration

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Protocol status: Working We use this protocol and it's working

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PROTOCOL integer ID: 94883

Keywords: parallel protein purification, Recombinant protein, Escherichia coli

MATERIALS

 His Gravitrap columns (Cytiva) supplier item 11003399 - for capture and purification of histidine tagged proteins, maximum binding capacity of 40 mg of tagged protein per column

His GraviTrap | Cytiva (cytivalifesciences.com)

- PD-10 buffer reservoir (Cytiva) supplier item 18321603 increases the His Gravitrap and PD-10 desalting column load volumes to 40mL
 Buffer Reservoir | Cytiva (cytivalifesciences.com)
- PD-10 column spin adapters (Cytiva) supplier item 28923245 used to prevent wobbling of columns in nalgene racks
 Spin Adapters for Gravity Columns | Cytiva (cytivalifesciences.com)
- PD-10 desalting columns (Cytiva) supplier item 17085101 used for rapid buffer exchange

PD-10 desalting columns packed with Sephadex G-25 resin | Cytiva (cytivalifesciences.com)

- Nalgene™ Unwire™ Test Tube Racks: Resmer™ Manufacturing

 Technology, for 30mm tubes, white **Thermo Fisher Catalog #5970-0030**
- AIM Terrific Broth Base including Trace elements Formedium Catalog #AIMTB0210
- W Ultra Yield 2.5L Flask, Sterile Generon Catalog #931136-B

Optional but useful

BENCHMIXER™ XL MULTI-TUBE VORTEXER Benchmark
Scientific Catalog #BV1010

Materials (1 L cultures) for Expression:

- 24 well tissue culture plates with LB-agar+antibiotics
- ☐ 5 L of autoclaved autoinduction TB + 20 g/L glycerol + antibiotics
- 🗸 5 mL of 10 % Antifoam 204 (Sigma) in ethanol
- 48x <u>A</u> 250 mL Corning baffled flasks (fitted with loose foil cover**)



Materials for Purification:

5 L of Base Buffer

	А	В
Г	HEPES	10 mM
Г	Glycerol	5%
Г	NaCl	500 mM
	TCEP, pH 7.5	0.5 mM

- △ 100 mL of [M] 3 Molarity (M) imidazole pH 7.5.
- △ 100 mL of 10 % Triton X-100 in water.
- \perp 50 mg/mL Lysozyme solution (100 x).
- △ 1 mg/mL homemade benzonase (1000x). Maybe susbtituted for 10 mg/mL of commercial DNase I
- 48 His GraviTrap columns to be purified (Cytiva) fitted with LabMate extender (Cytiva) and PD-10 spin adapter (Cytiva) in 24 place Nalgene rack.
- 48 PD-10 desalting column to be purified fitted with LabMate extender and PD-10 spin adapter in 24 place Nalgene rack.
- 48 x 50 mL centrifuge tubes per litre of culture to be purified in a 24 place Nalgene rack.

SAFETY WARNINGS

Triton x-100 is currently restricted for use in the EU and cannot be used without an exemption certificate REACH Annex XIV (Jan 2021). It can be readily subsituted with IGEPAL CA-630 (which is likely to be subject to the same restrictions in the near future). Alternatives that also maybe used are Tergitol 15-S-9 or Tween-20 or octyl glucoside.



12h 20m

Either transform BL21 (DE3) with the appropriate plasmid OR streak from glycerol stock onto 24 well agar 4h plate and incubate (*) Overnight ₿ 37°C *.

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Note

- * Freshly transformed or re-streaked cells always give better yields than growing overnights directly from frozen glycerol stocks.
- Grow 4 1.5 mL Overnight in suitable container (2 mL 96-well plate, 15 ml faclon) of each clone in superbroth + 1 % glucose + the appropriate antibiotics.
 - 3 Use 🚨 1 mL to inoculate 🚨 100 mL AIM-TB (+ Antibiotics + Antifoam 204) in a baffled flask.
- 4 Grow 250 rpm, 37°C, 04:00:00 shaking using loose foil cover**.

4h



AERATION IS ESSENTIAL!.

Note

- **An upturned 50 mL plastic beaker with a 1.5 mL microcentrifuge tube taped to the side of the flask to act as a spacer can also be used.
- **A breathable membrane such as an AirOtop enhanced flask seal may also be used.
- 5 Grow 40-48 h 3 250 rpm, 18°C shaking.



6 Harvest at 3 4000 x g, 4°C, 00:20:00 in 50 ml falcon tubes and discard supernatant.

20m



Harvest remainder of culture in same tube at 4000 x g, 4°C, 00:20:00, discard supernatant and freez pellet in falcon tube at -80 °C.



Final wet cell weight is typically 5.5 g of culture

Cell lysis

3h 30m

8 Thaw pellets.

9 Add 🗸 20 mL Base Buffer to each tube containing pellet ([M] 10 millimolar (mM) HEPES,



[M] 500 millimolar (mM) NaCl, 5 % Glycerol, [M] 0.5 millimolar (mM) TCEP, (μ) 7.5) + Δ 0.5 mg/mL Lysozyme, Δ 1 μg/ml Benzonase or Δ 10 μg/ml DNase I, [M] 20 millimolar (mM) imidazole.

Vortex to dissolve pellet and leave 00:30:00 Room temperature

30m

Add 4 mL 10 % Triton X-100*** to each tube and bring volume to 40 mL using Base Buffer

Note

***Triton x-100 is currently restricted for use in the EU and cannot be used without an exemption certificate REACH Annex XIV (Jan 2021). It can be readily substituted with IGEPAL CA-630 (which is likely to be subject to the same restrictions in the near future). Alternatives that also maybe used are Tergitol 15-S-9 or Tween-20 or octyl glucoside.

Freeze -80 °C 1-2 h or overnight if preferred.







Thaw in Room temperature water bath 01:00:00 and mix.

1h



1h



Note

Higher speed centrifugation can also be performed if desired, e.g. 20,000 g but transfer to suitable centrifuge tubes will be necessary.

Purification

3h 30m

Apply supernatant from a single tube to 1 mL His GraviTrap column (Cytiva) fitted with LabMate extender.

Note

Imidazole concentration can be increase to 40 mM in most cases, but may affect yield.

Wash ☐ 10 mL Base Buffer + [M] 20 millimolar (mM) Imidazole**.

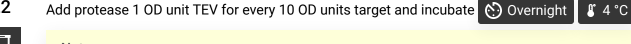


Note

**10 mL of a 40 mM or 70 mM imidazole wash can also be done, but this is very target dependent and may lead to significant reduction in final yield BUT can also increase purity substantially, worth trying if your purity is poor.

- Slot His GraviTrap column into PD10 column (Cytiva) fitted with LabMate extender (pre-equilibrated in Base Buffer + [M] 20 millimolar (mM) Imidazole).
- Elute protein with 4 2.5 mL of Base Buffer + [M] 500 millimolar (mM) Imidazole directly onto PD10 column.

- 19 Remove His GraviTrap column.
- 20 Place PD10 into 50 mL falcoln tube and add A 3.5 mL Base Buffer + [M] 20 millimolar (mM) **Imidazole** and collect.
- 21 Measure A280.
- 22 Add protease 1 OD unit TEV for every 10 OD units target and incubate 🚫 Overnight





Note

Some targets exhibit significant affinity for IMAC columns even after TEV cleavage try increasing the imidazole concentration to 40 or 70 mM or use an MBP-TEV construct so that the protease can be removed using an amylose column rather than reverse IMAC.

- 23 Run back over His GraviTrap column equilibrated in Base Buffer + [M] 20 millimolar (mM) Imidazole.
- 24 Wash column 2.5 mL [M] 20 millimolar (mM) Imidazole.
- 25 Check purity of A 6 mL pool.

- 26 Concentrate to A 1 mL ish.
- Transfer to 1.6 mL glass autosampler vial ensure at least 1.1 mL in vial!.
- Run through serial gel filtration system injecting 4 1 mL
- Take peak fraction(s) only (1-2 mL) and concentrated to 10-20 mg/mL if possible.

Column regeneration: PD-10

Wash PD-10 columns with ☐ 50 mL - ☐ 100 mL of Milli-Q water.

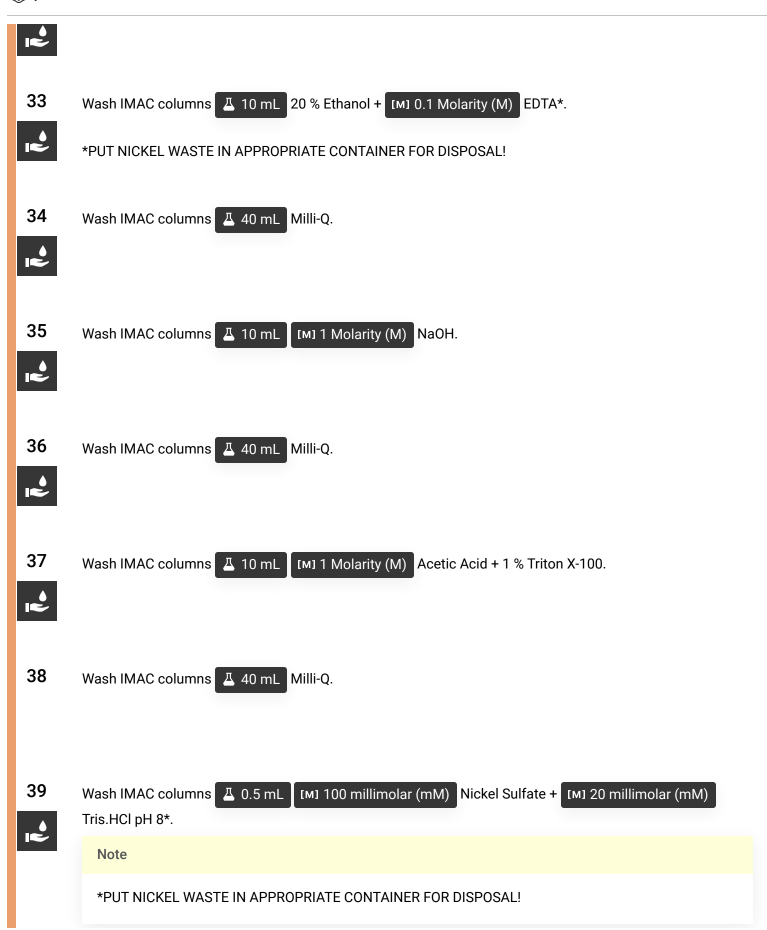


30

31 Store all columns in water at [4 ° C]. For long term storage use 20 % Ethanol

Column regeneration: His GraviTrap

Wash IMAC columns 40 mL Milli-Q.





Wash IMAC colums 40 mL Milli-Q.



Store all columns in water at \$\mathbb{E} 4 \cdot \mathbb{C}\$. For long term storage use 20 % Ethanol