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Purification of BCL2L13-GFP

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We use this protocol and it's working

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Abstract

This protocol details the purification of BCL2L13-GFP.

Materials

Lysis buffer:

	A	B
	Tris-HCl	50 mM
	pH	7.4
	NaCl	300 mM
	Triton X-100	1%
	glycerol	5%
	MgCl ₂	2 mM
	DTT	1 mM
	β-mercaptoethanol	2mM
	cOmplete EDTA-free protease inhibitors (Roche)	
	CIP protease inhibitor (Sigma)	
	DNase (Sigma)	

Wash buffer:

	Tris-HCl	50 mM
	pH	7.4
	NaCl	300 mM
	DTT	1 mM

High salt wash buffer:

	Tris-HCl	50 mM
	pH	7.4
	NaCl	700 mM
	DTT	1 mM

SEC buffer:

	Tris-HCl	25 mM
	pH	7.4
	NaCl	300 mM
	DTT	1 mM

- pET-DUET1 vector (available on Addgene)  pETDuet-1 TIM9,10 addgene Catalog #170280

- BCL2L13 W275A/I278A (Δ LIR1)(available on Addgene)
- BCL2L13 Y213A/I216A/W275A/I278A (Δ LIR1+2) (available on Addgene)
- BCL2L13 I224A/L227A/W275A/I278A (Δ LIR1+3) (available on Addgene)
- BCL2L13 W275A/I278A/I307A/V310A (Δ LIR1+4) (available on Addgene)
- BCL2L13 I224A/L227A/W275A/I278A/I307A/V310A (Δ LIR1+3+4) (available on Addgene)
- Rosetta pLysS cells (Novagen Cat# 70956-4)

 Rosetta™(DE3)pLysS Competent Cells - Novagen **Merck Catalog #70956-4**

- SORVAL RC6+ centrifuge with an F21S-8×50Y rotor (Thermo Scientific)
- Glutathione Sepharose 4B beads (GE Healthcare)
- 10 kDa cut-off Amicon filter (Merck Millipore)

 Amicon® Ultra Centrifugal Filter, 10 kDa MWCO **Merck MilliporeSigma (Sigma-Aldrich) Catalog #UFC801008**

- GFP-tagged BCL2L13-GFP (available from Addgene),
- BCL2L13(W275A/I278A)-GFP (Δ LIR1) (available from Addgene),
- BCL2L13(Y213A/I216A/W275A/I278A)-GFP (Δ LIR1+2) (available from Addgene),
- BCL2L13(I224A/L227A/W275A/I278A)-GFP (Δ LIR1+3)(available from Addgene),
- BCL2L13(W275A/I278A/I307A/V310A)-GFP (Δ LIR1+4) (available from Addgene),
- BCL2L13(I224A/L227A/W275A/I278A/I307A/V310A)-GFP (Δ LIR1+3+4)(available from Addgene)

Purification - BCL2L13-GFP

20h 46m

1 To purify GFP-tagged

- BCL2L13-GFP (available from Addgene),
- BCL2L13(W276A/I279A)-GFP (Δ LIR1) (available from Addgene),
- BCL2L13(Y213A/I216A/W276A/I279A)-GFP (Δ LIR1+2) (available from Addgene),
- BCL2L13(I224A/L227A/W276A/I279A)-GFP (Δ LIR1+3)(available from Addgene),
- BCL2L13(W276A/I279A/I307A/V310A)-GFP (Δ LIR1+4) (available from Addgene),
- BCL2L13(I224A/L227A/W276A/I279A/I307A/V310A)-GFP (Δ LIR1+3+4)(available from Addgene),

fuse the cytosol-exposed domain of BCL2L13 (1-463aa) to a C-terminal GFP-tag through cloning into a pET-DUET1 vector (available on Addgene).

2 Introduce the point mutants by in vitro mutagenesis to generate

- BCL2L13 W276A/I279A (Δ LIR1)(available on Addgene),
- BCL2L13 Y213A/I216A/W276A/I279A (Δ LIR1+2) (available on Addgene),
- BCL2L13 I224A/L227A/W276A/I279A (Δ LIR1+3) (available on Addgene),
- BCL2L13 W276A/I279A/I307A/V310A (Δ LIR1+4) (available on Addgene),
- BCL2L13 I224A/L227A/W276A/I279A/I307A/V310A (Δ LIR1+3+4) (available on Addgene).

3 After the transformation of the pET-DUET1 vector encoding BCL2L13-GFP wild-type or mutants in E. coli Rosetta pLysS cells (Novagen Cat# 70956-4), grow the cells in 2x Tryptone Yeast extract (TY) medium at 37 °C until an OD₆₀₀ of 0.4 and then continued at 18 °C .



4 Once the cells reaches an OD₆₀₀ of 0.8, induce the protein expression with 100 micromolar (μ M) isopropyl β -D-1-thiogalactopyranoside (IPTG) for 16:00:00 at 18 °C . Collect the cells by centrifugation and resuspend in lysis buffer.

16h



Lysis buffer:

A	B
Tris-HCl pH 7.4	50 mM
NaCl	300 mM

A	B
Triton X-100	1%
Glycerol	5%
MgCl ₂	2 mM
DTT	1 mM
β-mercaptoethanol	2mM
cOmplete EDTA-free protease inhibitors (Roche)	
CIP protease inhibitor (Sigma)	
DNase (Sigma)	

5 Sonicate the cell lysates twice for 30 s and clear by centrifugation at  18000 rpm, 4°C, 00:45:00 in a SORVAL RC6+ centrifuge with an F21S-8×50Y rotor (Thermo Scientific).

45m



5.1 Sonicate the cell lysates for  00:00:30 (1/2).

30s

5.2 Sonicate the cell lysates for  00:00:30 (2/2).

30s

6 Collect the supernatant and incubate with pre-equilibrated Glutathione Sepharose 4B beads (GE Healthcare) for  02:00:00 at  4 °C with gentle shaking to bind BCL2L13-GFP.

2h



7 Centrifuge the samples to pellet the beads and remove the unbound lysate.



8 Wash the beads twice with wash buffer, once with high salt wash buffer, and two more times with wash buffer.



Wash buffer:

A	B
Tris-HCl pH 7.4	50 mM
NaCl	300 mM



A	B
DTT	1 mM

High salt wash buffer:

A	B
Tris-HCl pH 7.4	50 mM
NaCl	700 mM
DTT	1 mM

- 9 To cleave off the GST-tag  Overnight , elute the GFP-tagged cargo receptor from the GSH beads by the addition of TEV protease.



- 10 To collect the supernatant, collect the beads by centrifugation.



- 11 Wash the beads twice with  4 mL of wash buffer, and collect the supernatant.



- 12 Pool the supernatant fractions, filter through a 0.45 μ m syringe filter, concentrate with 50 kDa cut-off Amicon filter (Merck Millipore), and load onto a pre-equilibrated Superdex 200 Increase 10/300 GL column (Cytiva).

- 13 Elute the proteins with SEC buffer.

SEC buffer:

A	B
Tris-HCl pH 7.4	25 mM
NaCl	300 mM
DTT	1 mM

- 14 Analyze fractions by SDS-PAGE and Coomassie staining. Pool fractions containing purified BCL2L13-GFP.



- 15 After concentrating the purified protein, aliquote the protein and snap-frozen in liquid nitrogen.



Note

Store the proteins at  -80 °C .