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# Calf-Intestinal Alkaline Phosphatase Treatment in situ-Killinger 2024

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ASAP Collaborative Rese...

Killinger



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We use this protocol and it's

working

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#### **Abstract**

Alpha-synuclein phosphorylated at serine 129 (PSER129) occurs in two pools, non-aggregated (physiological) and aggregated (disease). This protocol allows for the selective dephosphorylation of non-aggregated PSER129 and enhances the specificity and sensitivity immunodetection of aggregated PSER129. Thus, this protocol can be used to differentiate physiological from aggregated PSER129.

#### **Materials**

Dilution media:

	A	В
	Nacl	150 mM
Г	Tris-HCl, pH 7.4	50 mM
	Triton-X100	0.5%

#### CIAP buffer:

A	В
Nacl	100 mM
Tris- Hcl	50 mM
Mgcl2, pH 7.9	10 mM

■ Alkaline Phosphatase Calf Intestinal (HC), 1,000u Promega Catalog #M2825



## Day 1

1d 1h 10m

Wash free-floating tissue (3 x 10 minutes) in dilution media.



Dilution media:

A	В	
Nacl	150 mM	
Tris-HCl pH 7.4	50 mM	
Triton-X100	0.5%	

1.1 Wash free-floating tissue 00:10:00 in dilution media (1/3).

10m

1.2 Wash free-floating tissue 00:10:00 in dilution media (2/3).

10m

1.3 Wash free-floating tissue 00:10:00 in dilution media (3/3).

10m

2 Incubate the samples with 1% Triton X-100 in DM for 00:10:00.

10m

3 Wash in DM (5) 00:10:00 .

10m

4 Wash the tissues in CIAP buffer (2x10 minutes).



CIAP buffer:

A	В
Nacl	100 mM
Tris- Hcl	50 mM
Mgcl2, pH 7.9	10 mM

Autoclave and store | | Room temperature |.



4.1 Wash the tissues in CIAP buffer 00:10:00 (1/2). 10m 4.2 Wash the tissues in CIAP buffer 00:10:00 (2/2). 10m 5 Incubate the tissues with CIAP at a dilution of 1:333 for 24:00:00 at 37 °C on a shaker. 1d ■ CIAP concentration per bottle: 20 u/µl (Promega, Cat.# M2825). ■ In 🚨 500 µL CIAP buffer, add 🚨 1.5 µL CIAP (30 units). Day 2 3h 20m 6 Wash in DM (2 x 10 minutes). 6.1 Wash in DM (1/2). 10m 6.2 Wash in DM (2/2). 10m 7 Heat water bath to \$\mathbb{\m 1h 30m 8 Place the dish containing sodium citrate buffer in the water bath and heat it for 00:10:00. 10m ■ Sodium Citrate Buffer, pH 6.0 (1L): 
☐ 2.94 g Sodium citrate-Trisodium salt (Dihydrate) in △ 1000 mL DI water. pH 6.0. Add △ 0.5 mL Tween-20. Mix well. 9 Wash the tissues in sodium citrate buffer 00:10:00. 10m



- 10 Incubate the tissues in the heated sodium citrate buffer for 00:30:00. 30m 11 Cool the dish containing tissues to Room temperature (at least 00:20:00). 20m 12 Wash in DM for 10 min x 2 times. 12.1 Wash in DM for 00:10:00 (1/2). 10m 12.2 Wash in DM for 00:10:00 (2/2). 10m
  - 13 Tissues are now ready for downstream assays.