

Naveni pY PD1, AP

KIT INSTRUCTIONS

GENERAL GUIDELINES:

- Do not mix NaveniBright reagents with other Naveni[™] product lines.
- Reaction volume depends on the sample size.
- Use best practices when pipetting to minimize reagent consumption.
- Centrifuge vials before pipetting.
- Thoroughly defrost all buffer mixtures at room temperature, vortex well and spin down before use.
- Vortex and spin down all enzymes (1 and 2) before use.
- Keep enzymes on ice or a frozen cold block.
- Wait to add enzymes until immediately before adding to the sample.
- Remove excess washing buffer from samples before adding reagent.
- Do not allow slides/samples to dry.
- Preheat the humidity chamber before each step.
- Incubation times or temperatures other than those specified may give erroneous results.
- As with any product derived from biological sources, proper handling procedures should be used.
- Wear appropriate Personal Protective Equipment to avoid contact with eyes and skin.
- Chromogens may be carcinogenic and should be handled with care.
- Unused solutions should be disposed of according to local regulations.

REQUIRED BUT NOT SUPPLIED

- VectaMount[®] Express Mounting Medium (H-5700-60) from Vector Laboratories.
- Isopropanol 99,5 %.
- Endogenous alkaline phosphatase quenching solution.
- TBS and TBS-T Tris-buffered saline and Tris-buffered saline supplemented with 0,05% Tween, respectively.

IMPORTANT:

Appropriate precautions should be taken to avoid antibody cross-contamination. Crosscontamination is the primary source of unspecific background due to the high sensitivity of the assay.

Avoid bulk washing methods when multiple antibodies are used. Wash any technical controls separately.



KIT COMPONENTS:

Box 1.1:

Material	Article Number	Amount	Storage*
NaveniBright Blocking Buffer (1x)	NB.1.100.01	4000 μl	
NaveniBright Supplement 1	NB.1.100.03	500 μl	+4 to +8°C
NaveniBright Antibody Diluent (1x)	NB.1.100.02	8000 μl	
NaveniBright Supplement 2	NB.1.100.04	1000 μl	DO NOT
Probe Diluent (1x)	NF.1.100.03	4000 μl	FREEZE
NaveniBright Probe Anti-M (40x)	NB.1.100.06	100 µl	
NaveniBright Probe Anti-R (40x)	NB.1.100.07	100 µl	

Box 1.2:

Material	Article Number	Amount	Storage*
NaveniBright AP/HRP diluent	NB.1.100.08	8000 μl	
NaveniBright AP Reagent	NB.1.100.10	100 µl	+4 to +8°C
NaveniBright AP Substrate Diluent	NB.1.100.09	8000 μl	DO NOT
NaveniBright AP Substrate 1	NB.1.100.11	140 µl	FREEZE
NaveniBright AP Substrate 2	NB.1.100.12	100 µl	

Bag 1.3:

Material	Article Number	Amount	Storage*
Nuclear Stain	NB.1.100.16	6000 μl	+4 to +8°C DO NOT FREEZE

Box 2:

Material	Article Number	Amount	Storage*
Navenibody PD1 (200x)	PD1.2.01	20 µl	
Navenibody pTyr M (200x)	PD1.2.02	20 µl	
NaveniBright Buffer 1 (5x)	NB.2.100.17	800 µl	at
NaveniBright Enzyme 1 (40x)	NF.2.100.11	100 µl	-25 to -15°C
NaveniBright Buffer 2 (5x)	NB.2.100.18	800 µl	
NaveniBright Enzyme 2 (40x)	NF.2.100.15	100 µl	

* When stored as directed, the product is stable at least for 3 months after receipt



1. Sample preparation	1.1	After antigen retrieval, add enough alkaline phosphatase blocking solution (for quenching) (not provided) to cover each sample. Incubate for 10 min at room temperature, or according to manufacturer's user guide.
	1.2	Wash slides for 2x5 min in 1x TBS-T**.
2. Blocking	2.1	Prepare blocking solution by adding 5 μ l of Supplement 1 to every 40 μ l of Blocking Buffer (1x).
	2.2	Add the prepared blocking solution to the entire sample area (approximately $40 \ \mu l$ for each 1 cm ² area).
	2.3	Incubate for 60 min at +37 °C in a preheated humidity chamber.
3. Navenibody incubation	3.1	Prepare Navenibody solution by adding 5 μ l of Supplement 2 to every 40 μ l of Antibody Diluent (1x).
	3.2	Use the prepared Navenibody solution to dilute Navenibody PD1 and Navenibody pTyr M to 1x (dilute 1:200 each).
	3.3	Decant the blocking solution from slides and wash for 2x3 min in 1x TBS-T.
	3.4	Add enough of Navenibodies to cover the sample area.
	3.5	Incubate overnight at +4 °C in a humidity chamber.
	26	Decent the antihedy colution and wach clides for 2vE min in 1v TPS T in a

3.6 Decant the antibody solution and wash slides for 3x5 min in 1x TBS-T in a staining jar under gentle agitation.

Kit component		Blocking Solution	Navenibody solution	
Blocking Buffer		40 μl	-	
Supplement 1		5 μl	-	
Antibody diluent		-	40 µl	
Supplement 2		-	5 μl	
	Total:	45 μl	45 μl	
4. Probe incubation	eac 4.2 Add 4.3 Incu 4.4 Dec	epare the probes by diluting Probe anti-M and Probe anti-R (dilute 1:40 ch) in Probe Diluent (1x). Id enough of the probes to cover the sample area. Subate for 60 min at +37 °C in a preheated humidity chamber. Second the solution and wash slides for 3x5 min in 1x TBS-T in a staining jar der gentle agitation.		
5. Reaction 1	5.2 Prej gen 5.3 Add 5.4 Incu	ute Buffer 1 1:5 in distilled water. Vortex and spin down. epare Reaction 1 by adding <mark>Enzyme 1</mark> (dilute 1:40) to the diluted buffer. Mix otly by pipetting and spin down. d enough Reaction 1 to cover the sample area. ubate for 30 min at 37 °C in a preheated humidity chamber. ish slides for 2x3 min in 1x TBS-T in a staining jar under gentle agitation.		
6. Reaction 2	6.2 Prej gen 6.3 Add	te Buffer 2 1:5 in distilled water. Vort pare Reaction 2 by adding <mark>Enzyme 2</mark> (tly by pipetting and spin down. enough Reaction 2 to cover the sam bate for 90 min at +37 °C in a prehea	dilute 1:40) to the diluted buffer. Mix ple area.	

	0	Incubate for 30 min at room temperature with slow agitation.		
8. Substrate development	under gentle agitation. 8.2 Prepare the substrate solution by mixing 62,5x) and AP Substrate Reagent 2 (dilut	 Decant the solution and wash slides for 2x2 min in 1x TBS in a staining jar under gentle agitation. Prepare the substrate solution by mixing AP Substrate Reagent 1 (dilute 62,5x) and AP Substrate Reagent 2 (dilute 80x) in AP Substrate Diluent. * See calculation example for minimal volume: 		
	Kit Component	Substrate Solution		
	AP Substrate Diluent	80 µl		
	AP Substrate 1	1,3 µl		
	AP Substrate 2	1,0 µl		
	Total:	82,3 μl		
9. Nuclei staining	 8.3 Decant wash buffer from the slides. 8.4 Add enough substrate solution to cover t 8.5 Incubate the slides at room temperature 8.6 Decant of the substrate solution from the deionized water under gentle agitation. 9.1 Decant wash buffer from the slides. 9.2 Add enough bushese states to see the set of the substrate slides. 	for 15 to 25 min. ** slides and wash slides for 2x2 min in		
	9.2 Add enough Nuclear stain to cover the sa	-		
	9.3 Incubate for 2 to 10 seconds at room tem	•		
	9.4 Rinse the slides under running tap water	(not delonized water).		
10. Dehydration and mounting ¤	 10.2 Rapid dehydrate slides with 2x1 min was 10.3 Blot excess isopropanol from slides a Mounting Medium (H-5700-60). 	 Rapid dehydrate slides with 2x1 min wash in isopropanol. Blot excess isopropanol from slides and apply VectaMount[®] Express Mounting Medium (H-5700-60). 		
	min.10.5 Analyze using a brightfield microscope, fluorescence imaging, use a filter set for	Texas Red.		
	10.6 After imaging, store the slides at room te	emperature. Signal is stable for years.		

7.1 Decant the solution and wash slides for 2x5 min in 1x TBS, followed by 1x10

min in 0,1x TBS in a staining jar under gentle agitation.

7.4 Add enough AP solution to cover the sample area.

7.3 Decant wash buffer from the slides.

7.2 Dilute the NaveniBright AP reagent 1:300 in AP/ HRP diluent.

⁺ For alternative substrates, prepare according to manufacturer's user guide.

** Substrate incubation time should be optimized for each assay.

*** Excessive nuclear staining may obscure developed signals.

7. AP Incubation

* Slides must be mounted with VectaMount[®] Express Mounting Medium (H-5700-60) from Vector Laboratories. Usage of other dehydration methods or mounting medium may lead to reduced signal intensity.