

NPT.HER2

Naveni pTyr Her2

General guidelines

- Reactions volume depends on the sample size.
- Use best practices when pipetting to minimize reagents consumption.
- Centrifuge vials before pipetting.
- The final working concentration of different detergents for permeabilization (ex. Tween-20, Triton X100, digitonin) should be explored to find the optimal conditions to best preserve the cell structure and protein of interest.
- Select the Buffer C vial with the appropriate detection fluorophore for your microscope filter set.
- Available fluorophores are Texas Red, Atto 647, and Atto 488.
 The standard kit is equipped with Texas Red; if other fluorophore is needed, contact Navinci Diagnostics at contact@navinci.se
- Thoroughly defrost all buffer mixtures at room temperature, vortex well before use.
- Vortex and spin down all enzymes (A, B, and C) before use.
- Keep enzymes on ice or a frozen cold block.
- Wait to add enzymes until immediately before adding to the sample.
- Buffer C is a light-sensitive reagent. Always keep it protected from light.
- 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.
- Naveni assays might be combined with traditional immunofluorescence techniques, providing that all primary antibodies are from different species and the fluorescence detection systems use different fluorophores.
- 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.
- Unused solutions should be disposed of according to local regulations.

Kit components

Box 1:

Storage: +4 to +8°C.



Material	Art.no	Amount
Blocking Buffer (1x)	NF.1.100.01	4000 µl
Navenibody Diluent (1x)	NPT.1.100.01	4000 µl

Box 2:

Storage: -25 to -15°C. Protect from light.



Material	Art.no	Amount
Her2 Navenibody (40x)	NPT.2.18	100 μΙ
pTyr R Navenibody (40x)	NPT.2.21	100 μΙ
Buffer A (5x)	NF.2.100.08	800 µl
Enzyme A (40x)	NF.2.100.09	100 μΙ
Buffer B (5x)	NF.2.100.10	800 µl
Enzyme B (40x)	NF.2.100.11	100 μΙ
Buffer C (5x), Texas Red	NF.2.100.12	800 μΙ
Enzyme C (40x)	NF.2.100.15	100 μΙ

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



Important:

Cross-contamination is the main source of unspecific background due to the high sensitivity of the assay.



For more information, or to place an order, visit www.navinci.se/products
Email: contact@navinci.se

Instructions for use

1. Permeabilization(not provided):

 $\bf 1.1$ Only for fresh frozen cell slides: Permeabilize cells with 0,05% Triton X-100 in PBS for 5 min at room temperature.

1.2 Wash slides for 2x2 min with 1x PBS.

2. Blocking

2.1 Add Blocking Buffer (1x) to the entire sample area (approximately $40~\mu l$ for each 1cm2 area).

2.2 Incubate for 30 min at +37 °C in a pre-heated humidity chamber.

3. Navenibody incubation

3.1 Prepare Navenibodies by diluting Her2 Navenibody (40x) and pTyr R Navenibody (40x) in Navenibody Diluent (1x) (dilute 1:40 each).

3.2 Add enough of the Navenibodies to cover the sample area.

3.3 Incubate for 60 min at +37 °C in a pre-heated humidity chamber.

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

**TBS-T (Tris-buffered saline supplemented with 0.05% Tween 20)

4. Reaction A

- **4.1** Start preparing Reaction A by diluting Buffer A (5x) 1:5 in water. Vortex and spin down.
- **4.2** Add Enzyme A (dilute 1:40). Mix gently by pipetting and spin down.
- 4.3 Add enough Reaction A to cover the sample area.
- **4.4** Incubate for 60 min at +37 °C in a pre-heated humidity chamber.
- **4.5** Decant the solution and wash slides for 2x3 min with 1x TBS-T in a staining jar under gentle agitation.

5. Reaction B

- **5.1** Start preparing Reaction B by diluting Buffer B (5x) 1:5 in water. Vortex and spin down.
- ${\bf 5.2}$ Add ${\bf Enzyme~B}$ (dilute 1:40). Mix gently by pipetting and spin down.
- **5.3** Add enough Reaction B to cover the sample area.
- **5.4** Incubate for 30 min at 37 °C in a pre-heated humidity chamber.
- **5.5** Wash slides for 2x3 min with 1x TBS-T in a staining jar under gentle agitation.

6. Reaction C: Protect from light!

- **6.1** Select the Buffer C vial with the appropriate detection fluorophore for your microscope filter set. Do not use more than one Buffer C vial.
- **6.2** Start preparing Reaction C by diluting Buffer C (5x) 1:5 in water. Vortex and spin down.
- **6.3** Add Enzyme C (dilute 1:40). Mix gently by pipetting and spin down.
- **6.4** Add enough Reaction C to cover the sample area.
- **6.5** Incubate for 90 min at +37 °C in a pre-heated humidity chamber.
- **6.6** Decant the solution and wash slides for 2 min with 1x TBS in a staining jar under gentle agitation.

7. Nuclei staining (not provided): Protect from light!

- **7.1** Start preparing a Nuclei staining solution according to the manufacturer's instruction. Vortex and spin down.
- 7.2 Decant wash buffer from the slides.
- 7.3 Add enough Nuclei staining solution to cover the sample area.
- **7.4** Incubate according to the manufacturer's instruction.
- **7.5** Decant the solution and wash slides for 2x 10 min with 1x TBS in a staining jar under gentle agitation.
- **7.6** Wash slides for 15 min with 0.1x TBS in a staining jar under gentle agitation.

8. Mounting (not provided)

- 8.1 Decant excess wash buffer from the slides.
- **8.2** Mount the slides with a coverslip using a Fluoroshield anti-fade mounting medium.
- **8.3** Image your slides in fluorescence or confocal microscope, using 20x objective or higher and filter set for DAPI and a corresponding fluorophore (FITC for Atto 488, Texas Red, or Cy5 for Atto 647, respectively).

