# Proti-Ace<sup>™</sup> 2Kit



User Guide HR2-432 (pg 1)

## **Application**

*In situ* proteolysis and proteolytic screening of protein samples for crystallization and structure determination

#### **Features**

- 6 proteases
- Stable, optimized, freeze dried protease formulation
- Enhanced stability
- Proti-Ace Dilution Buffer
- Optimized protocol for in situ proteolysis or proteolytic screening

## **Kit Contents**

Proti-Ace 2 Reagent 1\* (Qty 3):

1 mg/ml Proteinase K,
10 mM TRIS hydrochloride pH 7.5,
1 mM Calcium acetate hydrate

# Proti-Ace 2 Reagent 2\* (Qty 3):

1 mg/ml Clostripain (Endoproteinase-Arg-C),
10 mM HEPES pH 7.5,
2 mM TCEP hydrochloride,
2 mM L-Cysteine,
1 mM Calcium chloride dihydrate

## Proti-Ace 2 Reagent 3\* (Qty 3):

1 mg/ml Pepsin,1 mM Hydrochloric acid

## Proti-Ace 2 Reagent 4\* (Qty 3):

1 mg/ml Thermolysin,
10 mM Calcium acetate hydrate pH 8.0,
50 mM Sodium chloride

## Proti-Ace 2 Reagent 5\* (Qty 3):

1 mg/ml Bromelain,
5 mM HEPES pH 7.5,
2 mM L-Cysteine,
1 mM Ethylenediaminetetraacetic acid disodium salt dihydrate

# Proti-Ace 2 Reagent 6\* (Qty 3):

1 mg/ml Actinase,
5 mM HEPES pH 7.5,
10 mM Sodium chloride

# Proti-Ace Dilution Buffer (Qty 6):

• 1.6 ml of 10 mM HEPES pH 7.5, 500 mM Sodium chloride

#### Discussion

A proteolytic fragment or domain of a protein may crystallize more readily or form better diffracting crystals than the intact protein. Proteases can be used to generate small, active fragments or domains of the target protein for crystallization. The fragment or domain can be used directly for crystallization experiments. Or the proteolytic sample analyzed by gel electrophoresis and/or mass spectrometry for mass and sequence for subsequent cloning, expression, purification and crystallization. Using proteolysis to enhance sample crystallization, the current overall success rate for yielding a deposited crystal structure is currently better than 12%.

# Instructions for Proteolytic Screening

Proteolytic screening is a procedure involving limited proteolysis of the sample versus a portfolio of proteases, followed by denaturing gel electrophoresis (SDS-PAGE) and/or mass spectrometry (MS) to identify regions of a gene corresponding to the protease resistant domain/fragment for subsequent cloning, expression, purification and crystallization.

- 1. Select all six, a subset or a single protease from the Proti-Ace 2 kit for Proteolytic Screening.
- 2. Add 100  $\mu$ l of deionized water to each of the selected Proti-Ace 2 enzymes to create a 1 mg/ml Protease Stock solution.
- 3. Into empty micro centrifuge tubes, create a 1:100 dilution (0.01 mg/ml) of each 1 mg/ml Protease Stock from Step 2 by adding 5  $\mu$ l of the 1 mg/ml Protease Stock plus 495  $\mu$ l of Proti-Ace Dilution Buffer (10 mM HEPES pH 7.5, 500 mM Sodium chloride).
- 4. Pipette 10  $\mu$ l of the 1:100 protease stock into 10  $\mu$ l aliquots of protein (10 mg/ml) for each protease to be screened.
- 5. Incubate at 37°C for 60 minutes.
- 6. Stop the reaction by adding SDS-PAGE sample buffer for SDS-PAGE analysis or a final concentration of 10% v/v trichloroacetic acid for MS analysis. Refer to your SDS-PAGE and MS protocols for the appropriate volume and concentration of SDS-PAGE or MS sample buffer for quenching.
- 7. Analyze the digests by SDS-PAGE and/or MS. Identify the small active fragment (SAF) or protease resistant domains. Clone the corresponding region of the gene. Express, purify and crystallize this gene product. Alternatively, scale up the proteolysis and purify the digest to produce a pure homogeneous sample of the SAF or domain for crystallization.

In the event of insufficient digestion, repeat steps 1-3 using a higher protease concentration such as 1:10 dilution of each Protease Stock (5  $\mu l$  of 1 mg/ml Protease Stock plus 45  $\mu l$  of Proti-Ace Dilution Buffer). Also consider longer incubation times, up to 24 hours.

In the event of over digestion, repeat steps 1-3 using a lower protease concentration such as 1:1,000 dilution of each Protease Stock (10  $\mu$ l of the 1:100 Protease Stock plus 90  $\mu$ l of Proti-Ace Dilution Buffer). Also consider shorter incubation times and/or lower incubation temperature (4 to 25°C).

<sup>\*</sup> When 100  $\mu$ l of deionized water is added to each Proti-Ace 2 freeze dried reagent

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## Instructions for In Situ Proteolysis

*In situ* proteolysis is a procedure where trace amounts of protease are included with the sample to be crystallized and mixed with crystallization reagents for screening or optimization experiments.<sup>2-4</sup>

- 1. Select the desired protease(s) from the Proti-Ace 2 kit to be used for *in situ* proteolysis.
- 2. Add 100  $\mu$ l of deionized water to each of the selected Proti-Ace 2 enzymes to create a 1 mg/ml Protease Stock solution.
- 3. Add the protease to the protein crystallization sample. Add 10  $\mu$ l of the 1 mg/ml Protease Stock solution to 90  $\mu$ l of 10 mg/ml protein to create a 1:100 w/w dilution.
- 4. Set the crystallization experiment using the protease:sample mixture.

# Optimization of In Situ Proteolysis for Crystallization

- a. Vary the protease:sample ratio. Typical protease:sample ratios are 1:100, 1:1,000 and 1:10,000.
- b. Alter the incubation time. Typical incubation times are between 0 and 24 hours
- c. Alter the incubation temperature. Typical incubation temperatures are between 4 and 37°C.
- d. For protein concentrations other than 10 mg/ml one can either use the preferred sample concentration with the protease:sample dilutions described in steps 1-4 or one can dilute the Proti-Ace 2 enzymes to a perfect 1:100 and/or 1:1,000 ratio based on the actual protein concentration. For example, if the protein concentration is 20 mg/ml one can add 50  $\mu l$  of deionized water in step 2 to create a 2 mg/ml Protease Stock solution and then proceed with steps 3 and 4 to screen 1:100 protease:sample.

#### Storage of the Proti-Ace 2 Kit

The unique freeze dried formulation of the Proti-Ace 2 kit offers a much improved protease stability compared to liquid protease formulations. Recommended storage: Room temperature up to 30 days,  $4^{\circ}\text{C}$  up to 12 months, -20°C up to 24 months. Once the proteases are made into solution the recommended storage is: Room temperature to  $4^{\circ}\text{C}$  up to 24 hours, -20°C up to 12 months.

#### References

- 1. Allan D'Arcy, personal communication, 1989-2009
- 2. In situ proteolysis for protein crystallization and structure determination. Dong, A et al. Nature Methods 4, 1019 1021 (2007)
- 3. In Situ Proteolysis to Generate Crystals for Structure Determination: An Update. Amy Wernimont, Aled Edwards. PLoS ONE 4(4): e5094. doi:10.1371/journal.pone.0005094
- 4. The use of in situ proteolysis in the crystallization of murine CstF-77. Tong et al. Acta Cryst. (2007). F63, 135-138
- 5. A brief history of protein crystal growth. McPherson, A. Journal of Crystal Growth, vol. 110, issue 1-2, pp. 1-10, 1991

- 6. Preparation and analysis of protein crystals. McPherson, A. John Wiley & Sons, Inc. 1982. ISBN 089464355X
- 7. A crystallizable form of the Streptococcus gordonii surface antigen SspB C-domain obtained by limited proteolysis. Forsgren et al. Acta Cryst. (2009). F65, 712–714
- 8. Preliminary X-ray analysis of a human VH fragment at 1.8 angstrom resolution. Gaur, Kupper, Fischer & Hoffman. Acta Cryst. (2004). D60, 965-967
- 9. Replication Protein A Characterization and Crystallization of the DNA Binding Domain. Pfuetzner et al. The Journal of Biological Chemistry, Vol. 272, No. 1, Issue of January 3, pp. 430–434, 1997.

#### **Related Products**

**HR2-432-01** Proti-Ace 2 Reagent 1: 1 mg/ml Proteinase K, 10 mM TRIS hydrochloride pH 7.5, 1 mM Calcium acetate hydrate\*

HR2-432-02 Proti-Ace 2 Reagent 2: 1 mg/ml Clostripain (Endoproteinase-Arg-C), 10 mM HEPES pH 7.5, 2 mM TCEP hydrochloride, 2 mM L-Cysteine, 1 mM Calcium chloride dihydrate\*

HR2-432-03 Proti-Ace 2 Reagent 3: 1 mg/ml Pepsin, 1 mM Hydrochloric acid\*

**HR2-432-04** Proti-Ace 2 Reagent 4: 1 mg/ml Thermolysin, 10 mM Calcium acetate hydrate pH 8.0, 50 mM Sodium chloride\*

**HR2-432-05** Proti-Ace 2 Reagent 5: 1 mg/ml Bromelain, 5 mM HEPES pH 7.5, 2 mM L-Cysteine, 1 mM Ethylenediaminetetraacetic acid disodium salt dihydrate\*

**HR2-432-06** Proti-Ace 2 Reagent 6: 1 mg/ml Actinase, 5 mM HEPES pH 7.5, 10 mM Sodium chloride\*

**HR2-429-07** Proti-Ace Dilution Buffer: (10 mM HEPES pH 7.5, 500 mM Sodium chloride), 1.6 ml

\* When 100  $\mu$ l deionized water is added to the supplied freeze dried Proti-Ace 2 reagent.

#### **Technical Support**

Please e-mail (tech@hrmail.com), fax (1-949-425-1611), or telephone (1-949-425-1321 option 2) your request to Hampton Research. Fax and e-mail Technical Support are available 24 hours a day. Telephone technical support is available 8:00 a.m. to 4:30 p.m. USA Pacific Standard Time.

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