

Piranha™ Targeted Protein Degradation System

Cat # PTPD500A-1, PTPD510A-1, PTPD520A-1, PTPD600A-1, PTPD513VA-1, PTPD527VA-1

User Manual

Storage: Please see individual components

Version 2 8/2/2019 A limited-use label license covers this product. By use of this product, you accept the terms and conditions outlined in the License and Warranty Statement contained in this user manual.

Contents

Product DescriptionProduct Description	2
List of Components	2
Storage	3
General Information	3
Protocol for Piranha™ System	3
Example Data and Applications	6
References	
Keterences	٠ ک
Technical Support	8
Licensing and Warranty Statement	8

Product Description

Genetic approaches to modulate protein activity in cells have played a pivotal role in our understanding of genes and gene networks, and have helped scientists understand the basis for many physiologically important events, both in normal and disease states. Two well-characterized approaches – CRISPR/Cas9 (genomic level) and RNAi (transcriptional level) have been extensively used to interrogate protein function indirectly in both *in vivo* and *in vitro* models. However, these approaches, while powerful, modulates protein activity indirectly, and relies on downstream events to exert their effects on the protein encoded by the genes. As a result, true phenotypes may be masked or undesirable/unexpected phenotypes (i.e. off-target effects) may result due to indirect modulation of the protein target itself. Researchers relying on these genetic approaches to understand functionality need to ask themselves the following question – Is this a real effect?

For the first time, researchers are now in the position to answer this question using a powerful, yet simple system to directly interrogate protein levels in the target cells of interest. Taking advantage of the underlying TRIM21-based protein degradation system¹ present in most eukaryotic cells, SBI is proud to offer PiranhaTM Targeted Protein Degradation System – the first validated, commercial system devoted to directly target proteins for degradation based on the TRIM21 pathway.

Using your own antibody plus SBI's Piranha[™] mRNA and a quick electroporation into target cells, targeted proteins of interest can be rapidly degraded in under 1 hr* after electroporation, and phenotypes observed shortly after. Specificity is governed by the antibody used, so previously untouchable post-translational modifications such as phosphoproteins can now be specifically targeted. In addition, cells traditionally refractory to genetic manipulation (such as non-dividing primary cells) can be safely studied, giving researchers another powerful tool to study protein function and resulting phenotypic outcomes.

*Based on published data by Clift et al. 2018. Degradation rate of target protein and resulting phenotypic effects will depend on particular protein(s) being targeted and influenced by protein turnover and any compensatory mechanisms. An initial timecourse experiment to find optimal conditions to assay for protein knockdown and phenotypes is highly recommended.

List of Components

Pi<u>ranha™ mRNA</u>

Cat #	ltem	Volume/Qty	Storage Temperature
PTPD500A-1	Piranha [™] Electroporation- ready mRNA (10 μg)	10 μL	-80°C
PTPD510A-1	Piranha [™] Electroporation- ready RFP-tagged mRNA (10 μg)	10 μL	-80°C
PTPD520A-1	Piranha [™] Electroporation- ready GFP-tagged mRNA (10 μg)	10 μL	-80°C

Piranha[™] Stable Cell Line

Cat #	ltem	Volume/Qty	Storage Temperature
PTPD600A-1	Piranha™ HEK293T GFP/Puromycin Stable Cell Line (>1x10^6 cell/vial)	1 vial	-80°C for up to 1 month Liquid Nitrogen (gas phase) for up to 12months

Piranha[™] Pre-Packaged Virus

Cat #	Item	Volume/Qty	Storage Temperature
PTPD513VA-1	pCDH-CMV-Piranha [™] -EF1- GFP-T2A-Puro virus (>1x10^6 IFUs)	2 x 25 μl	-80°C
PTPD527VA-1	pCDH-EF1a-Piranha [™] -T2A- Puro virus (>1x10^6 IFUs)	2 x 25 μl	-80°C

Storage

All listed components are shipped on dry Ice and should be stored at recommended temperatures as stated above. Properly stored components are stable for 12 months from the date received.

General Information

The Piranha system is designed to be modular in nature, offering maximal flexibility to the researcher. You can choose between mRNA, stable cell line, or pre-packaged lentivirus particles. For example, mRNA and antibody against protein of interest can be co-electroporated into cells for transient knockdown. Additionally, you can have a platform cell line that stably expresses Piranha protein, and you can electroporate different antibodies into the cell line in parallel to assess protein knockdown across multiple targets. Finally, you can develop your own platform cell line with the pre-packaged lentivirus containing Piranha cDNA.

Materials required but not provided:

- 1. Electroporator and accessories (e.g. ThermoFisher Neon Transfection System, Cat #MPK5000)
- 2. Concentrated antibody targeting your protein of interest (>0.5 µg/ul is recommended for best performance)

Protocol for Piranha[™] System

Note: The protocol below assumes the use of a Neon Transfection System and accessories (buffers, etc.) Use of other electroporation systems will require similar or equivalent components to be used for cell preparation.

A. Piranha mRNA and stable cell line

- 1. Cultivate the required number of cells (see below).
- 2. 24 to 48 hours prior to electroporation, transfer $1-5 \times 10^6$ cells (depending on growth rate) into a new 10 cm dish with fresh growth medium such that the cells are 70–90% confluent on the day of the experiment. Be sure to use low passage number, actively-dividing cells (if applicable).
- 3. Pre-warm an aliquot of culture medium containing serum, PBS (without Ca²⁺ and Mg²⁺), and Trypsin/EDTA solution to 37°C.
- 4. Aspirate the media from cells and rinse the cells using PBS (without Ca²⁺ and Mg²⁺).
- 5. Trypsinize the cells as normal.
- 8. Take an aliquot of trypsinized cell suspension and count cells to determine the cell density.
- 9. Transfer the cells to a 1.5 ml microcentrifuge tube or a 15 ml conical tube and centrifuge the cells at $100-400 \times g$ for 5 minutes at room temperature. Make sure that you have enough cells to accommodate $^{\sim}8 \times 10^{5}$ cells per electroporation reaction.
- 10. Aspirate supernatant, and wash cells with PBS (without Ca^{2+} and Mg^{2+}) by centrifugation at 100–400 × g for 5 minutes at room temperature.
- 11. Aspirate the PBS and resuspend the cell pellet in Resuspension Buffer R at a final density of 8×10^7 cells/ml. Gently pipette the cells to obtain a single cell suspension.

Note: Avoid storing the cell suspension for more than 15–30 minutes at room temperature, which reduces cell viability and transfection efficiency.

12. Prepare 6-well plates by filling the wells with 2 ml of culture medium containing serum and supplements without antibiotics and pre-incubate plates in a humidified 37°C/5% CO2 incubator.

Electroporation protocol:

- 1. Set up a Neon Tube (ThermoFisher) with 3 ml of Electrolytic Buffer E into the Neon Pipette Station (ThermoFisher).
- 2. Mix 10 μ l from the cell suspension (8 × 10 7 /ml) with 0.5 μ l of Piranha mRNA (1-1.5 μ g/ μ l) and 1 μ l of antibody (0.5-1 μ g/ μ l, or PBS as negative control). The mixture is taken up into a 10 μ l Neon Pipette Tip (ThermoFisher) and electroporated using the following settings: 1400V, 20ms, 2 pulses.
- 3. Immediately transfer the electroporated cells into the pre-incubated plate.
- 4. Be sure to change the Neon Pipette Tip after using it twice and Neon Tube after 10 usages.

^{*}Electroporation should be conducted and optimized according to your electroporator model and cell type of interest.

B. Piranha Pre-Packaged Lentivirus

For efficient delivery of pre-packaged lentiviral particles containing Piranha cDNA into your target cells, SBI recommends the following protocol for transduction.

Recommended reagent: TransDux MAX Lentivirus Enhancer Reagent (SBI Cat #LV860A-1)

Transduction of Target Cells (24-well transduction)

Day 1

1. Plate 50,000 cells per well in a 24 well plate in culture medium.

Day 2

- 1. Cells should be between 50 to 70% confluent.
- 2. Aspirate medium from cells.
- 3. Combine TransDux™ and TransDux MAX Enhancer with culture medium to a final concentration of 1x. [Example: Add 2.5 μL of TransDux™ and 100 μL of MAX Enhancer to 400 μL culture medium and then transfer to each well].
- 4. Add virus to each well at different MOIs or different volumes, depending on experimental aims.
- 5. Incubate at 37°C for 72hrs.

Day 5

Look at the cells for reporter expression if the specific construct has a reporter like GFP and/or begin Puromycin antibiotic selection to establish stable cell line expressing Piranha protein. For Puromycin selection, it is recommended to use a kill curve assay for the cell line of interest to determine minimal dose of Puromycin needed for selective pressure. Once **the** stable cell has been established, and cells are properly growing under selection, please refer to "Protocol for PiranhaTM mRNA and Stable Cell Line" for **additional protocol details.**

Example Data and Applications

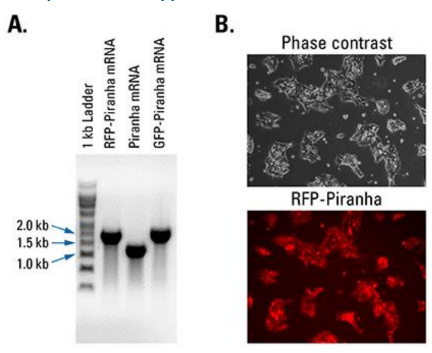


Figure 1. A) QC of Piranha mRNA (untagged, RFP- and GFP-tagged) by denaturing RNA gel electrophoresis, showing expected size of mRNA products B) HEK293T cells electroporated with RFP-tagged Piranha mRNA, showing robust delivery of the mRNA into cells via electroporation.

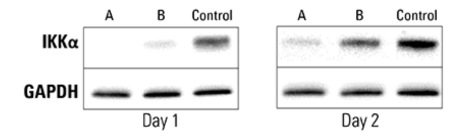


Figure 2. Western blot results for knockdown of IKKα protein in HEK293T cell line stably expressing Piranha protein and validated antibody recognizing IKKα protein electroporated into cell line. Two different electroporation conditions (using NeonTM Transfection System) were tested and IKKα protein levels were assayed 1 and 2 days after electroporation. Within 1 day, complete ablation of endogenous IKKα protein is seen in one condition (A =1400v 2p) and near ablation seen in another condition (B = 1150v 2p) relative to control lane, demonstrating the efficiency of protein depletion by the Piranha system.

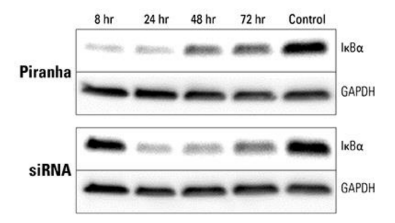


Figure 3. Western blot results demonstrating efficiency and speed of knockdown of IkB α protein using Piranha system vs traditional siRNA targeting. HeLa cell lines were co-electroporated with Piranha mRNA + validated antibody recognizing IkB α and IkB α protein levels were assayed at the listed timepoints after electroporation. Ablation of IkB α protein is seen as soon as 8 hrs (compared to 24 hrs with siRNA).

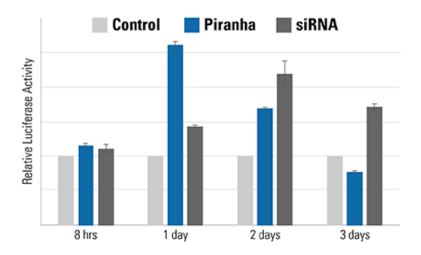


Figure 4. NFκB luciferase reporter assay in MDA-MB-231 breast cancer cells demonstrating faster response rate for Piranha system vs siRNA targeting IκB α protein. IκB α sequesters NFκB in the cytoplasm, and degradation of IκBa triggers translocation of NFκB to the nucleus to activate transcription. Luciferase levels increase significantly one day after electroporation for the Piranha system vs siRNA, demonstrating the speed of phenotypic response after protein degradation.

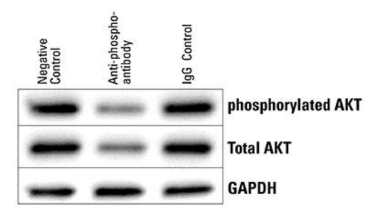


Figure 5. Western blot results demonstrating knockdown of phospho-specific protein (pAKT S473) with the Piranha system. HEK293T stable cell line expressing Piranha protein was electroporated with buffer only (negative control), pAKT S473 antibody, or non-specific IgG control. Total and phospho AKT levels were assessed, and within 5 hrs post-electroporation, knockdown of phospho-AKT is seen with decreased total AKT levels (consistent with majority of AKT being the phosphorylated form in HEK293T cells).

References

1. Clift D *et al.* A Method for the Acute and Rapid Degradation of Endogenous Proteins. Cell 2017 Dec 14; 171(7): 1692-1706.e18. doi: 10.1016/j.cell.2017.10.033. Epub 2017 Nov 16.

Technical Support

For more information about SBI products and to download manuals in PDF format, please visit our web site: http://www.systembio.com

For additional information or technical assistance, please call or email us at:

System Biosciences (SBI) 2438 Embarcadero Way Palo Alto, CA 94303

Phone: (650) 968-2200 Toll-Free: (888) 266-5066 Fax (650) 968-2277

E-mail:

General Information: info@systembio.com
Technical Support: tech@systembio.com
Ordering Information: orders@systembio.com

Licensing and Warranty Statement

Limited Use License

Use of the PiranhaTM Targeted Protein Degradation System (*i.e.*, the "Product") is subject to the following terms and conditions. If the terms and conditions are not acceptable, return all components of the Product to System Biosciences (SBI) within 7 calendar days. Purchase and use of any part of the Product constitutes acceptance of the above terms.

The purchaser of the Product is granted a limited license to use the Product under the following terms and conditions:

- The Product shall be used by the purchaser for internal research purposes only. The Product is expressly not designed, intended, or warranted for use in humans or for therapeutic or diagnostic use.
- The Product may not be resold, modified for resale, or used to manufacture commercial products without prior written consent of SBI.
- This Product should be used in accordance with the NIH guidelines developed for recombinant DNA and genetic research.

SBI has pending patent applications related to the Product. For information concerning licenses for commercial use, contact SBI

Purchase of the product does not grant any rights or license for use other than those explicitly listed in this Licensing and Warranty Statement. Use of the Product for any use other than described expressly herein may be covered by patents or subject to rights other than those mentioned. SBI disclaims any and all responsibility for injury or damage which may be caused by the failure of the buyer or any other person to use the Product in accordance with the terms and conditions outlined herein.

Limited Warranty

SBI warrants that the Product meets the specifications described in this manual. If it is proven to the satisfaction of SBI that the Product fails to meet these specifications, SBI will replace the Product or provide the purchaser with a refund. This limited warranty shall not extend to anyone other than the original purchaser of the Product. Notice of nonconforming products must be made to SBI within 30 days of receipt of the Product.

SBI's liability is expressly limited to replacement of Product or a refund limited to the actual purchase price. SBI's liability does not extend to any damages arising from use or improper use of the Product, or losses associated with the use of additional materials or reagents. This limited warranty is the sole and exclusive warranty. SBI does not provide any other warranties of any kind, expressed or implied, including the merchantability or fitness of the Product for a particular purpose.

SBI is committed to providing our customers with high-quality products. If you should have any questions or concerns about any SBI products, please contact us at (888) 266-5066.

© 2018 System Biosciences (SBI), All Rights Reserved

This page intentionally left blank.

This page intentionally left blank.



System Biosciences (SBI) 2438 Embarcadero Way Palo Alto, CA 94303

 Phone:
 (650) 968-2200

 Toll-Free:
 (888) 266-5066

 Fax:
 (650) 968-2277

E-mail:

General Information: info@systembio.com
Technical Support: tech@systembio.com
Orders@systembio.com

orders@systembio.com