pHIP(Var7)-P1AP suppresses tumor cell proliferation in MDA-MB-231 triple-negative breast cancer by targeting protease activated receptor 1

MingMing Yu1 · YueHua Chen2 · ZhenGuang Wang1 · XiaoDong Ding3

Received: 10 December 2019 / Accepted: 31 January 2020 / Published online: 7 February 2020
© The Author(s) 2020

Abstract
Purpose Protease-activated receptor 1 (PAR1) is a signaling protein ubiquitously present on the surface of tumor cells, and its homologous protein fragment, PAR1-activating peptide (P1AP), can inhibit protein signal transduction of PAR1/G in tumor cells. pHI (Low) insertion peptide (pHIP) can target the acidic tumor microenvironment (TME) and can be used as an excellent carrier to deliver P1AP to tumor cells for therapeutic purposes.

Methods PAR1 expression on the surface of MDA-MB-231 cells and human MCF10A mammary epithelial cells was observed. The binding between fluorescent-labeled pHIP(Var7)-P1AP and MDA-MB-231 cells under different pH values was analyzed. The effect of pHIP(Var7)-P1AP on the proliferation of MDA-MB-231 cells was analyzed under the conditions of pH 7.4 and 6.0.

Results PAR1 was highly expressed on the surface of MDA-MB-231 cells. In an acidic environment (pH 6.0 and 5.0), fluorescent-labeled pHIP(Var7)-P1AP and MDA-MB-231 cells had a high binding ability, and the binding ability increased with the decrease in pH. In an acidic environment (pH 6.0), pHIP(Var7)-P1AP significantly inhibited MDA-MB-231 cell proliferation. With 0.5 μg, 1 μg, 2 μg, 4 μg, and 8 μg of pHIP(Var7)-P1AP, the cell proliferation inhibition rates were 3.39%, 5.27%, 14.29%, 22.14%, and 35.69%, respectively.

Conclusion PAR1 was highly expressed on the surface of MDA-MB-231 cells. pH (low) insertion peptides (pHIP) · Protease-activated receptor 1 (PAR1) · Triple-negative breast cancer (TNBC) · Cell proliferation

Keywords pH (low) insertion peptides (pHIP) · Protease-activated receptor 1 (PAR1) · Triple-negative breast cancer (TNBC) · Cell proliferation

Abbreviations
PAR1 Protease activated receptor 1
pHIP PH (low) insertion peptide
TME Tumor microenvironment
TNBC Triple-negative breast cancer

Introduction
Triple-negative breast cancer (TNBC) refers to breast cancer tumor cells that do not express estrogen receptor (ER), progesterone receptor (PR), and human epidermal growth factor receptor 2 (Her-2). It accounts for 15–20% of breast cancer cases. TNBC has some clinical features, such as strong invasiveness, fast progression, and poor prognosis. The endocrine therapies and targeted therapies commonly used for the treatment of breast cancer are ineffective for TNBC. Therefore, the development of new therapeutic drugs targeting TNBC needs to be addressed in an urgent manner to improve the treatment strategies [1].

Protease-activated receptor 1 (PAR1) participates a variety of tumor invasion and metastasis processes and is a potential target for tumor treatment [2]. Pepducin PZ-128 is a palmitoylated peptide composed of a lipid and a PAR1 homologous protein fragment, PAR1-activating peptide...
(P1AP, KKSRALF). It has been shown that PZ-128 can inhibit PAR1/G protein signal transduction in tumor cells by targeting the third intracellular loop of PAR1 [2]. pH (Low) insertion peptide (pHLIP) family members are a novel type of vector that can target the acidic tumor microenvironment (TME) [3, 4]. The molecular mechanism of targeting is based on pHLIP insertion into the tumor cell membrane in a TME with low pH and the formation of a pH-dependent transmembrane α-helices [5]. pHLIP inserts its C-terminus across the cell membrane into tumor cells, where therapeutic molecules can be delivered.

In this study, the C-terminus of pHLIP(Var7) and the N-terminus of P1AP were linked by a disulfide bond, and pHLIP(Var7)-P1AP was obtained. pHLIP(Var7) inserted P1AP, which was linked to the C-terminus, into MDA-MB-231 cells after cleavage of the disulfide bond. The PAR1/G protein signal transduction was inhibited by targeting the intracellular loop of PAR1, achieving the purpose of treating MDA-MB-231 TNBC.

Materials and methods

Materials

PAR1 rabbit monoclonal antibody was purchased from Affinity Biosciences. Cy3-conjugated goat anti-rabbit IgG was purchased from CWBIO. 4′,6-Diamidino-2-phenylindole (DAPI) was purchased from KeyGen Biotech. 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) was purchased from Sigma-Aldrich.

Design, synthesis, and purification of (FITC)pHLIP(Var7)-P1AP

pHLIP(Var7)Cys and CysP1AP sequences were prepared by solid phase polypeptide synthesis. (pHLIP(Var7)Cys: AEEQNPWARYLEWLFPTETLLLELC; CysP1AP: CKKSRALF), pHLIP(Var7)Cys and CysP1AP were linked by disulfide bond to obtain pHLIP(Var7)-P1AP. The N-terminus of pHLIP(Var7)-P1AP was labeled with fluorescein isothiocyanate (FITC), and the C-terminus was modified by amidation to eventually obtain (FITC)pHLIP(Var7)-P1AP: (FITC)AEEQNPWARYLEWLFPTETLLLELC-CKKSRALF-NH₂. Peptides were purified via reversed-phase high-performance liquid chromatography (RP-HPLC) (Gemini-NX 10 µm, C18, 100 A, 4.6×250 mm; flow rate 1.0 mL/min; phase A: 0.1% trifluoroacetic acid in 100% acetonitrile; phase B: 0.1% trifluoroacetic acid in 100% water; gradient, 30 min from 28/72 A/B to 100/0 A/B). The purity of the peptide was determined by RP-HPLC as provided, and its identity was confirmed via mass spectrometry.

Cell culture and PAR1 expression on the cell surface

The human breast cancer cell line MDA-MB-231 and normal human breast epithelial cell line MCF-10A were purchased from the Cell Research Institute of the Chinese Academy of Sciences. MDA-MB-231 cells were cultured in L-15 medium containing 10% fetal bovine serum (FSB) at 37 °C without CO₂. MCF-10A cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM)/F12 (1:1) + horse serum (5%) + insulin (10 µg/mL) + epidermal growth factor (20 ng/mL) + cholera toxin (100 ng/mL) + hydrocortisone (0.5 µg/mL) at 37 °C with 5% CO₂.

PAR1 expression on the surface of MDA-MB-231 and MCF-10A cells

The cultured cells were washed with phosphate-buffered saline (PBS) three times, fixed in 4% paraformaldehyde for 15 min, and washed with PBS three times. The cells were permeabilized with 0.5% Triton X-100 (prepared with PBS) for 20 min at room temperature. The cells were washed with PBS three times, and 5% bovine serum albumin (BSA) was added; then, the cells were incubated for 30 min at 37 °C. The blocking buffer was aspirated by pipette. Adequate amounts of diluted PAR1 rabbit monoclonal antibody (1:200) was added to the culture dish, and the cells were incubated at 37 °C for 3 h. The cells were washed with PBS three times, diluted fluorescence Cy3-conjugated goat anti-rabbit IgG (1:200) was added dropwise, and the cells were incubated at 37 °C for 30 min. The cells were washed with PBS three times. DAPI was added dropwise, and the cells were incubated in the dark for 5 min to stain the cell nuclei. Excess DAPI was washed away with PBS. The cells in the culture dish were blocked with 50% glycerol, and images were obtained under a fluorescence microscope (OLYMPUS).

Analysis of (FITC)pHLIP(Var7)-P1AP binding to MDA-MB-231 cells

MDA-MB-231 cells were inoculated into 24-well plates (5.0×10⁴ cells/well). After the cells attached to the plates, the culture medium was removed, and phosphate buffer (25 mmol/L, pH 7.4) was used to wash the cells. The cells were divided into three groups, and L-15 medium (pH of 7.4, 6.0, or 5.0) was added to the wells, and the cells were incubated overnight at 37 °C without CO₂. A total of 30 µL of 100 nM (FITC)pHLIP(Var7)-P1AP was added to each well, and the plate was placed in a shaking mixer in an incubator for 2 h. All cells were washed five times with...
the corresponding pH (7.4, 6.0, 5.0)-specific phosphate buffer (25 mmol/L) to remove any unbound probes; then, fluorescence imaging was performed.

**Antiproliferation assay**

MDA-MB-231 cells were inoculated into 24-well plates (5.0 × 10⁴ cells/well). After the cells attached to the plates, the culture medium was removed, and phosphate buffer (25 mmol/L, pH 7.4) was used to wash the cells. L-15 medium with a pH of 7.4 or 6.0 was added to the wells, and the cells were incubated overnight at 37 °C without CO₂. Different amounts of pHLIP(Var7)-P1AP (0, 0.5 μg, 1 μg, 2 μg, 4 μg, and 8 μg) were added into the wells, and then, the plate was placed in an incubator for 72 h. All cells were washed five times with the corresponding pH (7.4 or 6.0)-specific phosphate buffer (25 mmol/L) to remove any unbound probes. Cell viability was determined by colorimetric MTT assay. A total of 20 μL of 5 mg/mL MTT solution was added to the treated cells, and the cells were incubated at 37 °C for 4 h. The obtained crystals were dissolved in 150 μL of dimethyl sulfoxide (DMSO), and the absorbance was measured at 490 nm using a microplate reader to calculate the cell proliferation inhibition rate.

**Statistical analysis**

SPSS 24.0.0.0 (IBM) statistical software was used for data processing. Variables are expressed as the mean (M) ± standard deviation (SD). Using one-way ANOVA to compare variables, p values less than 0.05 were considered statistically significant.

**Results**

**Synthesis and purification of (FITC)pHLIP(Var7)-P1AP**

(FITC)pHLIP(Var7)-P1AP was successfully synthesized. pHLIP(Var7)Cys and CysP1AP were prepared by solid phase polypeptide synthesis. pHLIP(Var7)-P1AP was obtained by the formation of a disulfide bond between the two peptides through oxidation. After labeling the N-terminus of pHLIP(Var7)-P1AP with FITC, (FITC)pHLIP(Var7)-P1AP was obtained.

HPLC analysis of the product indicated the formation of several compounds, i.e., a major product (96.8777%, retention time of 11.193 min) accompanied by at least two smaller peaks with retention times of 11.033 min and 11.508 min. The mass spectrometry (ESI–MS) analysis of (FITC)pHLIP(Var7)-P1AP showed three mass peaks with m/z of 753.8 ([M+6H]⁶⁺), 904.3 ([M+5H]⁵⁺), 1130.2 ([M+4H]⁴⁺). The experimentally observed molecular weight (4516.8, 4516.5) correlated well with the theoretical molecular weight (4517.17).

**PAR1 expression on the surface of MDA-MB-231 and MCF-10A cells**

PAR1 was highly expressed on MDA-MB-231 cells, and no obvious PAR1 expression was found on MCF-10A cells (Fig. 1).

![Fig. 1 PAR1 expression was positive in MDA-MB-231 cells, and PAR1 expression was negative in MCF-10A cells (400×)](image-url)
Analysis of (FITC)pHLIP(Var7)-P1AP binding to MDA-MB-231 cells

For the acidic pH values (pH 5.0 and 6.0), the probe and MDA-MB-231 cells had a high binding ability; the binding ability at pH 5.0 was higher than that at pH 6.0. At pH 7.4, the probe and MDA-MB-231 cells had only slight binding (Fig. 2).

Antiproliferation assay

Compared with pH 7.4, pHLIP(Var7)-P1AP significantly inhibited MDA-MB-231 cell proliferation at pH 6.0 ($p < 0.05$) (Fig. 3). At pH 6.0 with probe doses of 0.5 μg, 1 μg, 2 μg, 4 μg, and 8 μg, the cell proliferation inhibition rates for MDA-MB-231 cells were (3.39 ± 0.7)%, (5.27 ± 1.1)%, (14.29 ± 0.1)%, (22.14 ± 1.2)% and, (35.69 ± 1.2)%, respectively.

Discussion

TNBC occurs at a young age and is prone to early metastasis. Due to the lack of corresponding receptors, conventional endocrine therapies and targeted therapies are not effective, leading to a research hotspot in the field of breast cancer [1]. PAR1 is a G protein-coupled receptor (GPCR), exists on the surface of a variety of tumor cells and is a potential target for...
tumor treatments [2]. This study verified that a large amount of PAR1 exists on the surface of TNBC MDA-MB-231 cells, while there is almost no PAR1 expression on the surface of normal human MCF-10A breast cells; therefore, PAR1 is expected to become a potential target for the diagnosis and treatment of MDA-MB-231 TNBC.

The PAR1 inhibitor pepducin PZ-128 (palmitic acid-P1AP) is a class of lipid peptide that can enter cells and inhibit PAR1/G protein signal transduction [2]. PZ-128 is in phase II clinical trials. Currently, it is under development to target the PAR1 on the platelet surface to inhibit platelet function, which is essential for the effective treatment of patients with acute coronary syndrome (ACS) and patients undergoing percutaneous coronary intervention (PCI) [6]. Other studies confirmed that PZ-128 can inhibit the growth and metastasis of breast cancer, lung cancer, and ovarian cancer cells [7]. The exact mechanisms of PZ-128 with respect to GPCR remain unclear, but some researchers have proposed that peptides can directly interact with their homologous receptors, allowing the receptors to have an active or non-active conformation [8]. The effective part of PZ-128 that can inhibit the signal transduction of PAR1/G protein is the P1AP heptapeptide sequence, and the contained palmitic acid helps PZ-128 enter cells. PZ-128 is not able to specifically target tumor cells. If P1AP can be delivered into tumor cells using a tumor-specific carrier instead of palmitic acid, probe entry into normal cells can be minimized and background and adverse effects can be decreased in vivo.

The TME is acidic [9, 10]. The intracellular pH of almost all solid tumors is neutral to alkaline, while the extracellular pH is acidic and can be as low as 6.0 [3, 4, 11]. The mechanisms of an acidic TME include hypoxia-induced anaerobic glycolysis, aerobic glycolysis (Warburg effect), increased CO₂ production due to uncontrolled cell growth and increased ion pump activity on the cell membrane [9, 11]. An acidic TME is stable and is unaffected by the clonal selection of tumors. Therefore, it is also considered a promising marker for tumor-targeted detection [3, 5]. Research has shown that pHLIP family peptides can target TMEs. This peptide is derived from the bacterial rhodopsin C-helix, originally known as BRC peptide [12]. pHLIP can sense the pH near the cell membrane and insert into the cell membrane spontaneously to form a helical structure when the pH is near the cell membrane and insert into the cell membrane spontaneously to form a helical structure when the pH is near the cell membrane. The pHLIP variants studied, four sequences have high tumor-targeting properties, namely, wild type (WT), variant 3 (Var3), variant 7 (Var7), and ATRAM [22]. One study [19] linked pHLIP (WT) to the P1AP peptide by a non-cleavable linker (chloroacetyl chloride) to obtain pHLIP(WT)-P1AP, which can inhibit the growth of human breast cancer cells (MDA-MB-231 and MCF7 cells) that highly express PAR1 receptor. We made some changes based on a previous study [19], as follows. 1. The vector used in this study was generated with the Var7 sequence of pHLIP. pHLIP(Var7) is the shortest and most polar peptide sequence in the pHLIP family (targeting the tumor), and it has the advantages of easy synthesis, fast blood clearance, etc. pHLIP(Var7) was first applied in a study targeting GPCR. 2. A disulfide bond, instead of chloroacetyl chloride, was used to link pHLIP(Var7) to the P1AP peptide. Disulfide bonds can be lysed in cells; thus, P1AP can be delivered to cells. This method simplifies the synthesis of pHLIP(Var7)-P1AP and is conducive to widespread application.

The cell binding assay in this study showed that (FITC) pHLIP(Var7)-P1AP and MDA-MB-231 cells had high binding ability in an acidic environment (pH 5.0 and pH 6.0); however, at pH 7.4, there was almost no binding, suggesting that pHLIP(Var7) can target acidic tissues, which is consistent with previous studies regarding pHLIP. The antiproliferation assay showed that when the pH was 6.0, pHLIP(Var7)-P1AP significantly inhibited MDA-MB-231 cell growth, suggesting that under acidic conditions, pHLIP(Var7) can effectively deliver the P1AP peptide into MDA-MB-231 cells and that the disulfide bond was cleaved inside the cell and, thus, the P1AP peptide was released into the cells. P1AP played a cytotoxic role by inhibiting PAR1/G protein signal transduction.

Based on previous studies, this study designed and synthesized a novel therapeutic molecule, pHLIP(Var7)-P1AP, that can inhibit the growth of MDA-MB-231 cells. This study confirmed that at the cellular level, pHLIP(Var7)-P1AP can target MDA-MB-231 cells and can effectively inhibit the growth of MDA-MB-231 cells; however, it remains unclear whether MDA-MB-231 solid tumors can be inhibited. This study did not conduct a therapeutic study on animals due to certain restrictions, but future studies will be conducted when conditions permit. We also hope that other researchers can carry out further studies.

**Conclusion**

For TNBC, effective treatments are lacking, the prognosis is poor, and the development of new targets has important significance for future treatment strategies. PAR1 is highly expressed on the surface of MDA-MB-231 cells; therefore, PAR1 could be considered a target for the treatment of MDA-MB-231 TNBC. pHLIP(Var7)-P1AP can effectively target MDA-MB-231 cells in an acidic environment and inhibit the growth of MDA-MB-231 cells by inhibiting the
signal transduction of PAR1/G protein. pHLIP(Var7)-P1AP is expected to become a new valuable drug for the treatment of TNBC.

Acknowledgements I give special thanks to my wife Hua for her companionship and enthusiastic support.

Compliance with ethical standards

Conflict of interest The authors declare that they have no competing interests.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. Foulkes WD, Smith IE, Reis-Filho JS (2010) Triple-negative breast cancer. N Engl J Med 363(20):1938–1948
2. Yang E, Boire A, Agarwal A, Nguyen N, O’Callaghan K, Tu P et al (2009) Blockade of PAR1 signaling with cell-penetrating pepducins inhibits Akt survival pathways in breast cancer cells and suppresses tumor survival and metastasis. Cancer Res 69(15):6223–6231
3. Pathak AP, Gimi B, Glunde K, Ackerstaff E, Artemov D, Bhujwalla ZM (2004) Molecular and functional imaging of cancer: advances in MRI and MRS. Methods Enzymol 386:3–60
4. Penet MF, Glunde K, Jacobs MA, Pathak AP, Bhujwalla ZM (2008) Molecular and functional MRI of the tumor microenvironment. J Nucl Med 49(5):687–690
5. Vävere AL, Biddlecombe GB, Spees WM, Garbow JR, Wijesinghe D, Andreev OA et al (2009) A novel technology for the imaging of acidic prostate tumors by positron emission tomography. Cancer Res 69(10):4510–4516
6. Zhang P, Gruber A, Kasuda S, O’Callaghan K, Cox DH et al (2012) Suppression of arterial thrombosis without affecting hemostatic parameters with a cell-penetrating PAR1 pepducin. Circulation 126(1):83–91
7. Covic L, Kuliopulos A (2018) Protease-activated receptor 1 as therapeutic target in breast, lung, and ovarian cancer: pepducin approach. Int J Mol Sci 19(8):2237
8. Dimond P, Carlson K, Bouvier M, Gerard C, Xu L, Covic L et al (2011) G protein-coupled receptor modulation with pepducins: moving closer to the clinic. Ann N Y Acad Sci 1226:34–49
9. Fukumura D, Jain RK (2007) Tumor microenvironment abnormalities: causes, consequences, and strategies to normalize. J Cell Biochem 101(4):937–949
10. Izumi H, Torigoe T, Ishiguchi H, Uramoto H, Yoshida Y, Tanabe M et al (2003) Cellular pH regulators: potentially promising molecular targets for cancer chemotherapy. Cancer Treat Rev 29(6):541–549
11. Cairns R, Papandreou I, Denko N (2006) Overcoming physiologic barriers to cancer treatment by molecularly targeting the tumor microenvironment. Mol Cancer Res 4(2):61–70
12. Hunt JF, Earnest TN, Bousché O, Kalghatgi K, Reilly K, Horváth C et al (1997) A biophysical study of integral membrane protein folding. Biochemistry 36(49):15156–15176
13. Reshetnyak YK, Segala M, Andreev OA, Engelman DM (2007) A monomeric membrane peptide that lives in three worlds: in solution, attached to, and inserted across lipid bilayers. Biophys J 93(7):2363–2372
14. Wijesinghe D, Engelman DM, Andreev OA, Reshetnyak YK (2011) Tuning a polar molecule for selective cytoplasmic delivery by a pH (low) insertion peptide. Biochemistry 50(47):10215–10222
15. Moshnikova A, Moshnikova V, Andreev OA, Reshetnyak YK (2013) Antiproliferative effect of pHLIP-amantin. Biochemistry 52(7):1171–1178
16. Karabazzhak AG, An M, Yao L, Langenbacher R, Moshnikova A, Adochite RC et al (2014) pHLIP-FIRE, a cell-insertion-triggered fluorescent probe for imaging tumors demonstrates targeted cargo delivery in vivo. ACS Chem Biol 9(11):2545–2553
17. Cheng CJ, Bahal R, Babar IA, Pincus Z, Barrera F, Liu C et al (2015) MicroRNA silencing for cancer therapy targeted to the tumour microenvironment. Nature 518(7537):107–110
18. Song Q, Chuan X, Chen B, He B, Zhang H, Dai W et al (2016) A smart tumor targeting peptide-drug conjugate, pHLIP-SS-DOX: synthesis and cellular uptake on MCF-7 and MCF-7/Adr cells. Drug Deliv 23(5):1734–1746
19. Burns KE, Thevenin D (2015) Down-regulation of PAR1 activity with a pHLIP-based allosteric antagonist induces cancer cell death. Biochem J 472(3):287–295
20. Burns KE, Robinson MK, Thevenin D (2015) Inhibition of cancer cell proliferation and breast tumor targeting of pHLIP-monomethyl auristatin E conjugates. Mol Pharm 12(4):1250–1258
21. Burns KE, McCleerey TP, Thevenin D (2016) pH-selective cytotoxicity of pHLIP-antimicrobial peptide conjugates. Sci Rep 6:28465
22. Wyatt LC, Moshnikova A, Crawford T, Engelman DM, Andreev OA, Reshetnyak YK (2018) Peptidites of pHLIP family for targeted intracellular and extracellular delivery of cargo molecules to tumors. Proc Natl Acad Sci USA 115(12):E2811–E2818

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.