Peptide-based tumor inhibitor encoding mitochondrial p14ARF is highly efficacious to diverse tumors

Ken Saito,¹ Hidekazu Iioka,¹ Chie Kojima,² Mikako Ogawa³ and Eisaku Kondo¹

¹Division of Molecular and Cellular Pathology, Niigata University Graduate School of Medical and Dental Sciences, Niigata; ²Department of Applied Chemistry, Graduate School of Engineering, Osaka Prefecture University, Osaka; ³Laboratory of Bioanalysis and Molecular Imaging, Graduate School of Pharmaceutical Sciences, Hokkaido University, Sapporo, Japan

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Correspondence
Eisaku Kondo, Division of Molecular and Cellular Pathology, Niigata University Graduate School of Medical and Dental Sciences, 1-757 Asahimachi-dori, Chuo-ku, Niigata 951-8510, Japan.
Tel: +81-25-227-2102; Fax: +81-25-227-0761; E-mail: ekondo@med.niigata-u.ac.jp

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p14ARF is one of the major tumor suppressors conventionally identified both as the mdm2-binding molecule restoring p53 function in the nucleus, and as a nucleophosmin-binding partner inside the nucleolous to stabilize ribosomal RNA. However, its recently reported mitochondrial localization has pointed to novel properties as a tumor suppressor. At the same time, functional peptides are gaining much attention in nanomedicine for their in vitro utility as non-invasive biologics. We previously reported the p14ARF-specific peptide that restored the sensitivity to gefitinib on the gefitinib-resistant lung cancer cells. Based on the information of this prototype peptide, here we generated the more powerful anti-tumor peptide “9-Cat8-p14 MIS,” which comprises the minimal inhibitory sequence of the mitochondrial targeting p14ARF protein in combination with the proteolytic cleavage site for cathepsin B, which is activated in various tumor cells, fused with the nine-polyarginine-domain for cell penetration, and demonstrated its novel action of regulating mitochondrial function in accordance with localization of endogenous p14ARF. The p14 MIS peptide showed a potent tumor inhibitor in vitro and in vivo against not only lung cancer cells but also tumor cells of diverse lineages, via modulating mitochondrial membrane potential, with minimal cytotoxicity to non-neoplastic cells and tissues. Hence, this mitochondrially targeted p14 peptide agent provides a novel basis for non-invasive peptide-based antitumor therapeutics.
Cyclosporin A (Sigma-Aldrich, Tokyo, Japan). Fisher Scientific, Waltham, MA, USA), ROS detection agent, mitochondrial marker), CellLight Lysosomes-RFP (lysosomal marker), (Molecular Probes), CellLight Mitochondria-RFP (mitochondrial marker), Lyso Tracker Red Lysosomal Probe (Molecular Probes), Hoechst 33342 dye (Molecular Probes), Cleaved PARP detection (Thermo Fisher Scientific). Dyes/indicators were: Alexa Fluor 555-conjugated goat anti-rabbit IgG (H+L) antibody (Jackson ImmunoResearch, West Grove, PA, USA), Alexa Fluor 488-conjugated rabbit anti-mouse IgG (H+L) antibody for RFP detection (Jackson ImmunoResearch, West Grove, PA, USA), Alexa Fluor 555-conjugated goat anti-rabbit IgG (H+L) antibody for cleaved PARP detection (Thermo Fisher Scientific), Dyes/indicators and related reagents were: Hoechst 33342 dye (Molecular Probes, Eugene, OR, USA), Mito Tracker Red CMXRos (Molecular Probes), Lyso Tracker Red Lysosomal Probe (Molecular Probes), CellLight Mitochondria-RFP (mitochondrial marker), CellLight Lysosomes-RFP (lysosomal marker), mitochondrial membrane potential indicator, JC-1 (Thermo-Fisher Scientific, Waltham, MA, USA), ROS detection agent, CellROX Deep Red reagent (ThermoFisher Scientific) and Cyclosporin A (Sigma-Aldrich, Tokyo, Japan).

Peptide synthesis. All peptides were synthesized at the laboratory of the division of life science, Sigma-Aldrich Japan by standard Fmoc chemistry on an ABI 433A Peptide Synthesizer (Applied Biosystems, Foster City, USA). FITC-labeled peptides were purified to HCl form by reverse-phase HPLC to >95% purity for both in vitro and in vivo applications. Peptide identities were confirmed by mass spectrometry.

Immunohistochemistry and immunofluorescence. Antigen retrieval of paraffin-embedded sections, immunohistochemistry and signal development were performed as described previously.(15) The primary antibodies for p14ARF and Ki-67 were diluted at 1:500 in CanGet signal solution (Toyobo, Tokyo, Japan). Secondary antibodies were: Alexa Fluor 488-conjugated goat anti-rabbit IgG (H+L) antibody for RFP detection (Jackson ImmunoResearch, West Grove, PA, USA), Alexa Fluor 555-conjugated goat anti-rabbit IgG (H+L) antibody for cleaved PARP detection (Thermo Fisher Scientific), Dyes/indicators and related reagents were: Hoechst 33342 dye (Molecular Probes, Eugene, OR, USA), Mito Tracker Red CMXRos (Molecular Probes), Lyso Tracker Red Lysosomal Probe (Molecular Probes), CellLight Mitochondria-RFP (mitochondrial marker), CellLight Lysosomes-RFP (lysosomal marker), mitochondrial membrane potential indicator, JC-1 (Thermo-Fisher Scientific, Waltham, MA, USA), ROS detection agent, CellROX Deep Red reagent (ThermoFisher Scientific) and Cyclosporin A (Sigma-Aldrich, Tokyo, Japan).

Statistical analysis. Statistical differences were analyzed by paired Student’s t-test (MS Excel), and a value of P < 0.05 was regarded as statistically significant.

Results

Mitochondrial expression and its loss of p14 on various lineages of cancer cells. Because the endogenous expression of...
p14 in mitochondria in previous studies was restricted to a few tumor lines, we examined whether it is frequently observed in tumor cells of diverse origins. While loss of p14 expression was observed in PK8, BxPC3 (both pancreatic adenocarcinoma), MCF7 (breast cancer), LoVo (colon adenocarcinoma) and A549 (lung adenocarcinoma), mitochondrial p14 expression was detected in HeLa (uterine cervical carcinoma), PC-9 (lung adenocarcinoma), T47D (breast cancer), HT-29 (colon adenocarcinoma) and HAK-1B (hepatocellular carcinoma) cells with or without its nucleolar expression, which was corroborated by colocalization with HSP60 by immunofluorescence (Fig. 1a). To precisely confirm the mitochondrial expression (not only by immunofluorescence using the antibodies or Mito-tracker as in former studies) we employed immunoelectron microscopy. It revealed that p14 is localized not to the outer membrane but the inside (likely at cristae) of mitochondria (Fig. 1b). In contrast to tumor cells, p14 is consistently expressed in mitochondria in all lineages of normal (non-

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**Fig. 1.** Intracellular localization of endogenous p14ARF in tumor cells of diverse origins and non-neoplastic cells. (a) Double immunofluorescence using both anti-p14ARF (1:500 dilution) and anti-HSP60 (1:2000 dil.) antibodies for various lineages of human tumor cells. (b) Immunoelectron microscopy for HeLa cells using rabbit anti-p14 polyAb (1:100 dil.) and 10 nm gold particle-conjugated goat anti-rabbit IgG polyAb. (c) Double immunofluorescence for non-neoplastic cells of normal origins expressing p14 and HSP60. (d) Expression of endogenous p14ARF both in the tumor cells and normal lineage cells by RT-PCR. β-actin was amplified as an endogenous control to p14 mRNA.
neoplastic) cells examined, including MMNK-1 (cholangiocyte), NHDF (dermal fibroblast), NuLi-1 (bronchial epithelium) and HPNE (pancreatic duct epithelium), but not TIME (vascular endothelium) (Fig. 1c). The mRNA expression in each cell line including tumor cells and non-neoplastic cells corresponded to that of p14 protein, respectively (Fig. 1d).

**Design and mitochondrial targeting of the p14 MIS peptide.** We previously defined the primary functional region, within the entire amino acid sequence encoded by p14	extsuperscript{ARF}, as "APAAVALVLMLLSQRLLGQQP" from the amino acid position of 38th (alanin) to position 65 (glycine); it functions as a growth inhibitory peptide when combined with the cell-penetrating polyarginine (nine d-Arginines; r9) domain, against gefitinib-resistant lung adenocarcinoma cells. To bolster the antitumor action of this prototype, we first inserted the cathepsin B-cleavable motif "GFLG" between the cell-penetrating domain and the p14 amino acid sequence in order to release the functional p14 sequence and to maximally exert its inhibitory effect following intracellular incorporation. This design was influenced by the fact that cathepsin B, one of the

![Graph](attachment:graph.png)
lysosomal cysteine proteases of the papain family, is highly activated in tumor cells.\(^{(17,18)}\) We next attempted to shorten the sequence of the prototype peptide to the minimal functional amino acid sequence within the p14 derived segment. Because “AVAL,” at the 41st through 44th amino acid positions of the p14 protein, is known to be the mitochondrial localization motif,\(^{(11)}\) we constructed the truncated peptides with or without this motif, and compared the efficiency of

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**Fig. 3.** In vitro challenge by the p14 MIS peptide (r9-CatB-p14 MIS) against various tumor cell lineages and normal cells of several different origins. (a) Growth suppression of tumor lines (1 × 10^5 cells) by 10 µM of the peptide 48 h after peptide introduction. Pancreas adenocarcinoma lines (PK-8, BxPC-3), uterine squamous cell carcinoma (HeLa), lung adenocarcinoma (PC-9, A549), breast adenocarcinoma (MCF-7, T47D), colon adenocarcinoma (LoVo, HT-29) and hepatocellular carcinoma (HAK-1B). The result of non-neoplastic cells, MMNK-1, NHDF, NuLi-1, TIME and HPNE, as described in the Materials and Methods, are shown. Means and SD of triplicates are shown. (b) Microscopic cellular image of growth alteration of the 10-µM peptide-treated tumor cells corresponding to the result shown in (a). (c) Magnitude of growth suppression by each inhibitory peptide (10 µM) with different designs (protease-cleavable spacer sequence and p14 sequence) on four malignant tumor lines of distinct origins. MTT assays were performed 48 h after peptide introduction.
growth suppression among the four peptides; namely, prototype r9-p14 38-65 with the original “GPG” spacer, r9-CatB-p14 38-65 with the “GFLG” spacer, truncated r9-CatB-p14 41-57 retaining “AVAL,” and the most truncated form, r9-CatB-p14 MIS (see Results and Fig. S1a). To evaluate the growth suppression by these four p14 peptides, the lung adenocarcinoma lines, both gefitinib-sensitive PC-9 and gefitinib-resistant RPC-9 clones, were employed for the treatment because they were already demonstrated to be very responsive to the original form, the r9-p14 38-65 peptide, in our previous study.\(^{(15)}\)

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**Fig. 4.** r9-CatB-p14 MIS peptide functions to downregulate mitochondrial membrane potential (Δψm) of the tumor cells based on JC-1 dye staining. (a) Native proliferative status of Δψm detected by JC-1 in tumor and non-neoplastic cells. (b) Significant reduction of Δψm on tumor cells triggered by introducing 10 μM of r9-CatB-p14 MIS. (c) Quantitative analysis of fluorescence reduction in JC-1 staining in response to 10 μM of p14 MIS peptide. Means and SD of triplicates are shown. (d) Generation of ROS in the peptide-sensitive tumor cells. (e) RT-PCR evinced expression of the mitochondrial function-related gene, ATPAF1 and UCP-2, in tumor lines and non-neoplastic cells employed in the peptide study.
In the MTT assay, the most potent inhibition was obtained with r9-CatB-p14 MIS, the shortest form (Fig. S1b). It showed 80% inhibition of PC-9 and 85% inhibition of RPC-9 at 10-µM concentration, which raised the inhibition 1.9 to 2.5-fold compared to the original r9-p14 38-65 peptide (Figs 2a, S1b). Taking all these results together, the most efficient amino acid sequence of the peptide was decided in rrrrrrrr-GFLG-LVLMLLRQRLG (r; D-Arg), named “r9-GFLG-p14 MIS,” as shown in Figures 2(b) and S1a. Indeed, when r9-CatB-p14 MIS was introduced to tumor cells, PC-9 and BxPC3, it was more efficiently targeted to mitochondria inside tumor cells. This was demonstrated by intracellular localization of the fluorescent signal derived from the FITC-labeled peptides 24 h after their introduction, as shown in Figure 2(c). Specifically, only a small proportion of the N-term FITC labeled original form “r9-p14 38-65” having Gly-Pro-Gly as a spacer, a non-cleavable form by cathepsin B, was localized to mitochondria as well as N-term FITC-labeled r9-CatB-p14 MIS (Fig. 2c, upper panel), while most of the rest of the r9-p14 38-65 was predominantly localized not to mitochondrias, but to lysosomes (Fig. 2c, lower panel). Thus, the C-terminal FITC-labeled r9-CatB-p14 MIS was prominently localized to mitochondria after tumor cell uptake. The result indicated that r14 MIS sequence is targeted to mitochondria after its cellular entry, and the cathepsin B-cleavable form “r9-p14 38-65” is inefficiently targeted to mitochondria. Consequently, r9-CatB-p14 MIS was revealed to be the most efficient form of the p14-derived antitumor peptide that facilitates mitochondrial localization for potent inhibition of tumor cell growth (Fig. S2).

Challenge of the r9-CatB-MIS antitumor peptide against tumor cells of diverse origin in vitro. Upon finding that the peptide is properly recruited to the mitochondria after cellular entry, the anti-proliferative effect on tumor cells of several lineages was examined. Insofar as the p14ARF anti-proliferative effect on tumor cells of several lineages was properly recruited to the mitochondria after cellular entry, the peptide was mainly due to the loss of mitochondrial localization of the cleaved p14 MIS sequence by isolating the r9 domain. Consequently, r9-CatB-p14 MIS was revealed to be the most efficient form of the p14-derived antitumor peptide that facilitates mitochondrial localization for potent inhibition of tumor cell growth (Fig. S2).

Sequence analysis and amplification of the peptide sensitivity in the resistant tumor cells. Because the amino acid sequence “LVLMLLRQRLG” encoded between the 45th to 56th position of the p14ARF protein has a three leucine (“blue lettered” “LVMLLLRSQRLG” encoded between the 45th to 56th position) stretch in a canonical position at a four amino acid interval, we examined whether this leucine positioning is critical for its antitumor function, by replacement with alanine (A) or glycine (G), where these amino acids may contribute to emulating the helical structure (Fig. 5a). These derivatives, AAA and L50G, did not significantly affect peptide-resistant tumor cells, HAK-1B, but BxPC3 cells showed patterns of hampered growth and form consistent with the original inhibitory function indicating the unecessity of forming helix by these leucines (Fig. 5a). Because Cyclosporin A (CsA), a representative immunosuppressant, is recognized to increase ΔΨm, we treated the peptide-resistant HAK-1B cells with 1 µM of CsA prior to introduction of the r9-CatB-p14 MIS, and evaluated the effect of co-treatment. Treatment of HAK-1B with CsA prominently increased ΔΨm in HAK-1B, while ΔΨm was not significantly altered in the BxPC3 that originally showed high ΔΨm (Fig. 5c). CsA treatment restored growth suppression by the r9-CatB-p14 MIS in the peptide-resistant HAK-1B cells, from 25% inhibition up to 60% inhibition, an efficacy on a par with the peptide-sensitive BxPC3 cells (Fig. 5d). Substantial change
in ΔΨm was quantitatively shown between the cells cotreated by CsA with the peptide and the cells treated by CsA alone (Fig. 5e). By contrast, downregulation of ΔΨm and growth suppression by the same treatment were minimized in several lineages of non-neoplastic cells except in the case of NuLi-1 (Fig. S4a–c). Mitochondrial dysfunction by the combinatorial
Fig. 6. Tumor suppression in vivo by the p14 MIS peptide. (a) The therapeutic protocol for tumor-bearing mice with repeated intravenous peptide administration. GFP-expressing BxPC-3 cells were allowed to grow intraperitoneally for a week before peptide treatment. (b) In vivo distribution of metastatic tumors in the mouse abdominal cavity with or without the peptide treatment; upper row, an example of mice treated with control peptide (r9 alone); lower row, mouse treated with r9-CatB-p14 MIS. (c) GFP imaging of the intraperitoneal metastatic tumors collected from each mouse model treated with r9 alone or with r9-CatB-p14 MIS; the total weight of collected tumors per mouse from each group. (d) Histological examination of the tumor tissue sections from each peptide-treated mouse model. HE stain and immunohistochemistry using Ki-67 antibody.
treatment of CsA and the r9-CatB-p14 MIS yielded generation of reactive oxygen species (Fig. 5f).

**Tumor suppression in vivo by transduction of r9-CatB-p14 MIS.** Based on the *in vitro* functional assay of r9-CatB-p14 MIS, the mouse tumor model xenografted with stable GFP-expressing BxPC3 cells was prepared and subjected to an administration regimen for *in vivo* challenge by the peptide (Fig. 6a).

Before moving to the therapeutic experiment, we first confirmed that r9-CatB-p14 MIS was delivered to the tumor tissue by detecting a fluorescent signal of the FITC-labeled peptide *in vivo* (Fig. S5). Mice were kept without therapeutic treatment for a week after intraperitoneal implantation of $2 \times 10^6$ of the tumor cells, allowing the growth of tumor cells *in vivo*. One week later, each mouse was intravenously administrated 300 μg of the r9-CatB-p14 MIS peptide per day, 6 days in total, then mice were autopsied on the day next to the last treatment. As shown in the fluorescence image of the peritoneal cavity, multiple GFP-positive disseminated tumor foci were observed (Fig. 6b, arrow head) in the peptide-treated mouse as well as in the control peptide consisting of the nine poly-d-arginine sequence (r9) alone (Fig. 6b). However, each tumor focus was significantly smaller in size, and the total weight of the intraperitoneally grown tumors with the peptide treatment was only one-quarter (i.e., 75% inhibition of tumor growth) that of the control peptide-treated mouse (Fig. 6c). Macroscopically visualized nodules were histologically confirmed as the adenocarcinoma lesions originating from implanted human BxPC3 cells. Consistent with the tumor growth suppression by the peptide, mitotically active cancer cells were rarely observed in the tumor tissue treated by r9-CatB-p14 MIS peptide (Ki-67-positivity was <1% in total number of cells), whereas prominent mitotic activity was detected in approximately 50% of Ki-67-positive tumor cells in the control tumor tissue treated with cell-penetrating domain, r9, alone (Fig. 6d).

**Discussion**

The tumor suppressor gene, p14ARF, functions to halt cell division and trigger apoptosis in response to various oncogenic stresses in normal cells. Reflecting its indispensable role in prevention of tumor development, loss of the function through mutation or epigenetic silencing is observed in most forms of human cancer. As a canonical pathway, the p14ARF protein is recognized to activate p53 tumor suppressor by antagonizing the mitochondrial death factor p53INP (2). Other than MDM2, are exposing its novel p53-independent tumor suppressor activities. In particular, mitochondrial p14 protein have been recently gained much attention that suggests novel function of p14 in contrast to conventional p14 function in nucleus.\(^\text{(11,12,23,24)}\)

In this study, we first examined endogenous p14 expression in a spectrum of tumor cell lineages, some of which showed loss or marked downregulation of p14 expression; we then investigated the mitochondrial expression of p14 with or without its conventional nuclear/nucleolar expression in p14-positive tumor cells (Fig. 1a,d). Employing immunoelectron microscopy, we further corroborated the intramitochondrial localization of p14ARF as direct evidence of p14 localization at the mitochondrial inner membrane and at the matrix (Fig. 1b).

We previously identified the functional core sequence within the entire p14ARF ORF, which enables targeting of the p14 gene product to mitochondria to trigger apoptosis in tumor cells. From the previous findings, we next sought a more efficient and practical mitochondrial targeting system against tumor cells using the potent antitumor peptide “r9-CatB-p14 MIS.” This tumor suppressor peptide encodes the minimal inhibitory amino acid sequence “LVLMLRSQRLG” derived from the 45th to 50th amino acid position of p14ARF, which successfully targeted itself to mitochondria with the cathepsin B-cleavable motif “GFLG” as a spacer, and strongly suppressed the hyperactivity of tumor mitochondria, indicated by downregulation of ΔΨm. Previously, “AVAL” was reported to be the hypothetical mitochondrial localization motif.\(^\text{(11)}\) In this study, the r9-CatB-p14 MIS lacking “AVAL” significantly showed mitochondrial localization. Thus, the MIS sequence seems to be sufficient and these four amino acids are revealed to be dispensable for mitochondrial migration.

The magnitude of growth inhibition by the p14 MIS peptide was initially expected to be dependent on the loss of expression of mitochondrial p14 in tumor cells; that is, the peptide was expected to be highly effective in tumors lacking endogenous p14 expression. However, the potent growth inhibition by p14 MIS peptide was obtained not only in p14-negative tumor lines (e.g., PK8, BxPC3 and MCF-7) but also in p14-expressing lines, such as PC-9 (Figs 1d, s3a), which provokes questions on the precise molecular mechanisms of tumor-selective growth suppression by the candidate peptide. Recognizing that a point of action for the p14 MIS peptide is the regulation of the mitochondrial membrane depolarization (downreguration of ΔΨm), we screened for prospective molecules relating possibly to mitochondrial activation, and retrieved “ATP6F1.” ATP synthase mitochondrial F1 complex assembly factor 1,\(^\text{(21)}\) which critically correlated with sensitivity of tumor cells to r9-CatB-p14 MIS peptide (Fig. 4e). Although the p14 MIS peptide amino acid sequence predicts a functional α-helical structure in view of a triple leucine run at a five residue interval, substitution of these leucines with alanine (A) or glycine (G)
did not affect the original function of the p14 peptide, as variants AAA and L50G also showed comparable growth suppression to the original peptide (WT) (Fig. 5a,b). This suggests that the peptide does not functionally require the predicted helix arrangement for its intracellular function, contrary to usual expectations.

The investigated anti-tumor peptide, r9-CatB-p14MIS, as shown in the scheme summarizing the mode of its action (Fig. 7), offers two unique advantages in that the peptide automatically transits to mitochondria after intracellular entry and functions to downregulate the hyper-activated ΔΨm in neoplastic cells, inducing growth arrest and/or triggering apoptosis in diverse tumor cell lineages without appreciably affecting non-neoplastic cells. Growth inhibition by the p14 MIS was consistently observed, particularly in cancer cells with a high ΔΨm (up to 70% inhibition at 10 μM peptide), probably due to selective sensitivity for reduction of ΔΨm. p14 MIS response of tumor cells with native low ΔΨm potential was weaker than that of the tumor cells with high ΔΨm; however, cotreatment with cyclosporine A and p14 MIS significantly augmented the inhibitory effect in these low responders (e.g. there was a threefold enhancement [from 20% to 60%] in HAK-1B hepatoma cells [Fig. 5c–f]). We then sought in vivo tumor suppression using the r9-CatB-p14MIS peptide on tumor-bearing mice. First, the FITC-labeled peptide by intravenous administration showed successful accumulation in the multiple pancreatic adenocarcinoma lesions in the peritoneal cavity (Fig. S5).

The antitumor effect after six repeated administrations based on the therapeutic protocol with daily peptide injection (300 μg/i.v./day) resulted in approximately 75% inhibition of tumor growth compared with that of r9 alone-treated group (Fig. 6b,c). Consistent with macroscopic tumor regression, histology of the peptide-treated tumor evinced a marked decrease in number of Ki-67 antigen-positive cells compared to the control r9-treated tumors (Fig. 6d).

Recent advances in nanotechnology have yielded important clues toward countering malignant tumors, and functional oligopeptides appear to offer strong potential as effective biologics, wherein their dwindling scales (approximately 3–5 kDa in molecular weight) are consistent with the miniaturization efforts of nanobiology. In this study, we attempted to develop a novel antitumor peptide, r9-CatB-p14 MIS, and explore its application for peptide-based tumor therapeutics. The peptide showed dramatic growth suppression against intraperitoneally disseminated tumors by downregulating mitochondrial ΔΨm, while being intrinsically non-invasive to normal tissues in vivo and suggesting a safe physiologic compatibility for clinical applications, on the strength of its derivation from the p14 ARF tumor suppressor protein sequence.

Accumulated molecular biological studies indicate that the mitochondrion is a crucial organelle in determining cell fate, and is, therefore, a burgeoning focus of interest in cancer therapeutics.25 While our future studies will direct increasing attention to the underlying molecular interaction mechanisms, we believe that our present report as such will offer critical insights for treating malignancies using peptide-based molecular agents targeting tumor cell mitochondria, and add significantly to frontline biomedical strategies against cancer.

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Disclosure Statement

The authors have no conflict of interest to declare.

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Supporting Information

Additional Supporting Information may be found online in the supporting information tab for this article:

Fig. S1. Amino acid sequence of designed p14 peptides and their effect on growth suppression of the tumor cells.

Fig. S2. Colocalization of r9-CatB-p14 MIS with the lysosomal or mitochondrial markers in tumor cells after its intracellular incorporation.

Fig. S3. Growth suppression and induction of apoptosis in the p14 peptide-treated tumor cells.

Fig. S4. Cyclosporin A (CsA) potentiates mitochondrial membrane potential.

Fig. S5. In vivo uptake of the r9-CatB-p14 MIS peptide via intravenous administration.