TEM1-targeting PEGylated PLGA shikonin nanoformulation for immunomodulation and eradication of ovarian cancer

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Introduction

As a leading cause of death worldwide, cancer continues to be the global burden. Various types of malignancies, in particular solid tumors, continue to increase with the growth and aging of the world’s population.1 In solid tumors, the resistance to cytotoxic agents cell death and the reprogramming of metabolic pathways are the main hallmarks of cancer cells, leading to chemotherapy inefficacy.2 Despite a rigorous chemotherapy course, many patients might inevitably induce marked toxicity mainly due to non-specific accumulation of anticancer drugs in the normal cells/tissue and drug resistance.3,4 The occurrence of such side effects may lead to the failure of treatment modalities. To avoid non-specific inadvertent toxic impacts of the conventional chemotherapy modalities, advanced multifunctional nanomedicines with active and/or passive targeting potential have successfully been used.5-7 Smart targeted nanoparticles (NPs)/ nanosystems (NSs) have been reported to accumulate highly in the tumor microenvironment (TME) via both the passive

Abstract

Introduction: Tumor endothelial marker 1 (TEM1) is expressed by tumor vascular endothelial cells in various cancers.

Methods: Here, we developed poly(lactic-co-glycolic acid) (PLGA) nanoparticles (NPs) PEGylated with polyethylene glycol (PEG) and functionalized with anti-TEM1 antibody fragment (78Fc) and loaded them with necroptosis-inducing agent shikonin (SHK) (78Fc-PLGA-SHK NPs).

Results: The nanoformulation showed a smooth spherical shape (~120 nm; the ζ potential of –30 mV) with high drug entrapment and bioconjugation efficiencies (~92% and ~90%, respectively) and a sustained-release profile in serum. Having significant toxicity in vitro (e.g., MS1 and TC1 cells), the nanoformulation dramatically increased the cytotoxicity in the TC1 murine lung carcinoma subcutaneous and intravenous/metastatic models as aggressive tumor models. The injection of the 78Fc-PLGA-SHK NPs to the MS1-xenograft mice resulted in significantly higher accumulation and effects in the TEM1-positive tumor targets, while they were excreted via urine track without retaining in the liver/spleen. In the TC1 subcutaneous model, C57/B16 mice treated with the 78Fc-PLGA-SHK NPs revealed a significant therapeutic effect. The mice, which were tumor-free after receiving the nanoformulation, were re-challenged with the TC1 cells to investigate the immune response. These animals became tumor-free a week after the injection of TC1 cells.

Conclusion: Based on these findings, we propose the 78Fc-PLGA-SHK NPs as a highly effective immunostimulating nanomedicine against the TEM1-expressing cells for targeted therapy of solid tumors including ovarian cancer.

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targeting by the enhanced permeation and retention (EPR) effect and the active targeting via interaction with unique cancer antigens/molecular markers, resulting in maximal therapeutic effects in cancerous cells with minimal side effects.8,14 Of anticancer NSs, biodegradable polymeric NPs have widely been studied in terms of their potential in the safe delivery of loaded anticancer agents to the target cells while they elicit negligible side effects.8 Among biodegradable polymers, poly(lactic-co-glycolic acid) (PLGA) is one of the most widely investigated biocompatible polymers,15,17 and has been considered to serve as delivery systems.8,18,19

For active targeting of cancer cells, NSs need to be decorated with a desired homing agent specific to cognate cancer molecular marker. In addition to epithelial cells, tumor microvasculature endothelial cells might display markers that differ from that of the normal stromal cells.20,21 The oncomarkers expressed are called tumor vascular markers (TVMs), which might be exploited as targets for antibody-based tumor diagnosis and therapy.22-25 In large part because of the relative stability of TVM-expressing cells within the tumor neovasculature that play a key role in the maintenance and progression of the tumor.26 Further, the leaky capillary of tumor neovasculature facilitates the circulation of Ab-conjugated nanomedicines, resulting in their easy access to TVMs. Among the TVMs, tumor endothelial marker 1 (TEM1), known as CD248/endoelisin, is a very promising candidate for the targeted therapy of tumor microvasculature.27,28 The functional expression of the TEM1 has been shown to associate with tumor microvascular thrombus (TVM), including pericytes and myofibroblasts,21,25,29 and TMV endothelial cells in various human cancers.20,30-32 However, it is not expressed by other adult normal cells/tissue.29,33 TEM1 plays a central role in the progression of the tumor via various mechanisms, including the induction of proliferation and angiogenesis,34 as well as migration and metastasis. Such impacts occur through the interaction of TEM1 with matrix proteins such as fibronectin, collagen type I and IV,35 and Mac-2 BP/90K.32 For instance, TEM1-deficient mice transplanted with the tumor in the abdominal sites were shown to present a striking reduction in the growth, invasiveness, and metastasis of the tumor.36 All these findings support the concept that the TEM1-mediated targeted therapy might offer a great tool for the diagnostic and/or therapy of solid tumors such as ovarian cancer.

It should be noted that the battle with cancer can substantially be improved through a combination of anticancer cytotoxicity and immunosurveillance, while very few anticancer agents display such properties. Of various products assessed for anticancer and immunomodulatory effects, shikokin (SHK) appears to provide unique pharmacological effects such as wound-healing potential together with antitumor, anti-inflammatory, antioxidant, antithrombotic, and antimicrobial effects.37,38 Given that the SHK functions on different molecular targets and exhibits a broad range of activities, it is considered a highly interesting substance.39-43 Further, as a highly lipophilic naphthaquinone pigment, SHK inhibits (i) the activity of pyruvate kinase M2,44,45 (ii) upregulates the activity of p53 and downregulates the functional expression of cyclin-dependent protein kinase 4,46 (iii) upregulates BCL2-associated X protein and downregulates B-cell lymphoma 2,37 and (iv) inhibits the pathways of extracellular signal-regulated kinases and protein kinase B.47 SHK was shown to incite significant cellular toxicity in vitro and in vivo,48 which is not specific to the cancer cells only. Like any other cytotoxic agent, to be effective on the target cells, SHK molecules must be delivered efficiently and specifically to the cancer cells that can be attained employing nanoscale targeted drug delivery systems (DDSs). In this current work, we developed the SHK-loaded, anti-TEM1-armed PLGA NPs as described previously,26 and investigated their biological effects both in vitro and in vivo. Furthermore, we studied the cell death mechanism(s) induced by SHK and looked into the immunostimulatory properties of this innovative nanoformulation in vivo.

Materials and Methods

Acid terminated PLGA (lactide:glycolide 50:50, Mw 38000-54000), ethyl acetate (EtAc), Pluronic® F68 (PF68), N-hydroxysulosuccinimide sodium salt (NHS), 4-morpholineethanesulfonic acid (MES hydrate), D-(-)-trehalose dehydrate, phalloidin–tetramethylrhodamine isothiocyanate, goat serum, cardiogreen, bovine serum albumin (BSA), 0-(2-aminoethyl)polyethyleneglycol 3000 (PEG -OH), N-(3-dimethylaminopropyl)-N’-ethyl carbodiimide hydrochloride (EDC), sulforhodamine B (SRB), and tissue culture dishes with vents CELLSTAR were purchased from Sigma Aldrich Co. (St. Louis, MO, USA). Shikonin, crystal violet, JC-1, RIP3 antibody (H-43), RIP antibody (H-207), HPV16 E6 (N-17) P, and HPV16 E7 (C-20) P were purchased from Santa Cruz Biotechnology Inc. (Santa Cruz, CA, USA). BD Bioscience Transwell™; Annexin V: PE apoptosis detection kit I, anti-IFN-γ Rat IgG1 κ Mouse Ab, 3 mL BD™ slip-tip disposable syringe, rat IgG1 κ anti-mouse IFN-γ biotinylated Ab, streptavidin–alkaline phosphatase, ham IgG1 κ anti-mouse CD3e Ab clone 145-2C11, Syr Ham IgG2 λ1 anti-mouse CD28 clone, and anti-CD31 rat anti-mouse Ab were obtained from BD Biosciences (San Jose, CA, USA). Fisherbrand Cover Glasses Circles (No. 1 to 0.13 to 0.17 mm thick; Size: 12 mm), and 1-Step NBT/BCIP Solution Pierce™ were purchased from Thermo Fisher Scientific (Waltham, MA, USA). The XTT cell proliferation kit II, and the in-situ cell death detection kit AP were purchased from Roche Applied Science (Penzberg, Germany). Necrostatin-1 and 2,7-dichlorodihydrofluorescein diacetate were obtained from Cayman Chemical Co. (Ann Arbor, MI, USA). NucBlue® fixed Cell ReadyProbes® reagent, Alexa Fluor® 594 goat anti-hamster IgG (H+L),
and PloLong diamond antifade mountant with DAPI were purchased from Molecular Probes Biotechnology Co. (Eugene, OR, USA). Wheat germ agglutinin, Alexa Fluor® 488 conjugate, and ACK lysing buffer were obtained from Life Technologies/Invitrogen Corporation (Carlsbad, CA, USA). DNase, RNase, protease-free ddH₂O and Falcon® 70µm cell strainer white sterile individually packaged 50/ Case were purchased from Mediatech Inc. (Manassas, VA, USA). Annexin V-FITC early apoptosis detection kit was obtained from Cell Signaling Technology (Danvers, MA, USA). The recombinant human IL-2 was obtained from Peprotech (Rocky Hill, NJ, USA). Acridine orange staining solution was purchased from ImmunoChemistry Technologies LLC (Bloomington, MN, USA). TNF-α (Q178) polyclonal antibody was purchased from Bioworld Antibodies Inc. (Woburn, MA, USA). Formvar/Carbon 400 Mesh, Cu, 50/bx was purchased from Electron Microscopy Sciences (Hatfield, PA, USA). 5-Bromo-4-chloro-3-indolyl-β-D-galactosidase was purchased from PanReac AppliChem (Darmstadt, Germany). MultiScreen-IP Filter Plate was purchased from EMD Millipore (Billerica, MA, USA). DAPI fluoromount G was purchased from SouthernBiotech (Birmingham, AL, USA). Rapamycin was purchased from InvivoGen (San Diego, CA, USA). 78Fc scFv_4H9 (2.435 µg/µL; 40 µL) was provided by Dr. Chunsheng Li (University of Pennsylvania). The cell lines were obtained from American Type Culture Collection (Manassas, VA, USA) and MS1 cells were engineered in-house to express the HuTem1 marker.

Nanoparticle formulation and physicochemical evaluation

The preparation and evaluation of NPs were performed employing a single emulsion-solvent evaporation technique, as described previously. Briefly, SHK (15 mg/mL) was dissolved in 30 µL DMSO. About 100 mg of PLGA was dissolved in a 3 mL oil phase (chloroform: ethyl acetate 2:1). Then, the SHK solution was slowly added to the oil phase. About, 3% of surfactants (i.e., 1:1 ratio of Pluronic F68 and polyvinyl alcohol) were dissolved in 10 mL double distilled water. To this end, the oil phase was added and mixed with the water phase (10 mL) over an ice bath. The mixture was then sonicated at 40 W output for 4 minutes (2x2 minutes) employing a probe sonicator (VCX 130; Sonics and Materials, Inc, Newtown, CT, USA). Afterward, to remove the oil phase, the mixture was evaporated under a reduced pressure of 150 mbar, employing a rotary evaporator (30 rpm) (Hei-VAP Advantage ML/G5B; Heidolph North America, Grove Village, IL, USA). Further, the same procedure was employed for the formulation of the drug-free PLGA NPs. The influence of variables on the NP formulations was further evaluated. For the lyophilization of the NPs, they were resuspended in 1 mL trehalose (250 mM) as a cryoprotectant supplemented with 0.5% Tween 80, which was applied to prevent the lyophilization-induced aggregation and keep it safe for later use.

Transmission electron microscopy (TEM) analysis

To morphologically characterize NPs, TEM analysis was conducted. Briefly, 10 µL of NPs sample was suspended 1 mL distilled water, and then, one drop of the suspension was placed over a carbon 400 mesh TEM grid and allowed to dry. The NPs on the mesh were visualized on a Tecnai 12 microscope at 120 KeV at indicated magnifications (FEI, Hillsboro, OR, USA), equipped using Gatan’s US1000 CCD camera (Gatan, Inc., Pleasanton, CA, USA).

Particle size and zeta potential analyses

For the analysis of the size and zeta potential of NPs, each sample (20 µL) was diluted to 1 mL of distilled water. A Zetasizer, Nano-ZS (Malvern Instruments, Malvern, UK) was recruited to measure the size and zeta potential of the NPs.

Drug content (DC%), drug entrapment (DE%), drug loading (DL%), and yielding (YE%) efficiencies

The amount of SHK in NPs was analyzed using the ultraviolet (UV) spectrophotometric method, where opted amounts of SHK-loaded NPs were dissolved in acetonitrile (1 mg/mL). To complete the release of the entrapped drug, the samples were kept on a shaker at 37°C for 72 hours. Then, to extract the drug, the samples were removed by centrifugation at 10000 × g, 4°C for 5 minutes. After collecting the supernatant, a UV spectrophotometer (Genesys 6; Thermo Fisher Scientific) was employed to measure the absorbance at a wavelength of 520 nm. The DC%, DE%, DL%, and yielding efficiency (YE%) values were calculated for all the formulations (Table S1).

Release kinetics analysis

To explore the kinetics profile, the release of SHK from PLGA NPs were evaluated at various pHs (pH 4.4, 5.4, 6.4, and 7.4) utilizing phosphate-buffered saline (PBS) release buffer (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na2HPO4, 1.47 mM KH2PO4) supplemented with fetal bovine serum (FBS). All the experiments were performed at 37°C. In practice, NPs (25 mg) were distributed in 5 mL of PBS buffer, and the samples were divided into equal aliquots (1 mL each). These tubes were kept on a shaker at 37°C and 150 rpm. At designated time intervals (i.e., 1, 2, 4, 8, 12, 24, 48, and 72 hours), the samples were centrifuged at 10000 × g at 4°C for 5 minutes. Then, the extracted supernatant was read exploiting the Genesys 6 UV spectrophotometer, at a wavelength of 520 nm (SHK) for PBS and 600 nm for FBS samples. Finally, various kinetics models (Table S2) were recruited to analyze the data, as described previously.

PEGylation and 78Fc conjugation

A one-step approach was used for the PEGylation and Ab conjugation of NPs (Fig. 1A), as described previously.
Briefly, 10 mg of acid-terminated PLGA NPs was re-suspended in 2-4 mL MES (pH 6). It was then activated with 250 µL EDC (1 mg/mL) and 250 µL NHS (1 mg/mL) at room temperature under stirring at 35 rpm for 2 hours. The samples were washed (×2) with 50 mM sucrose by centrifugation at 10,000×g (4°C) for 5 minutes. The activated NPs were re-suspended in PBS and incubated with 250 µL NH2-PEG3000-OH (1 mg/mL) and 100 µL 78Fc at a molar ratio of 10:1 at room temperature under stirring at 35 rpm for 2 hours. The NPs were washed (×2) using 50 mM sucrose (pH 8) by centrifugation at 10,000×g (4°C) for 5 minutes, to remove the unconjugated molecules, and the supernatant was analyzed to determine the conjugation efficiency through UV spectrophotometry.

**Determination of Ab conjugation efficiency**

To analyze the conjugation efficiency, we capitalized on the Bradford method using Coomassie Blue dye. Briefly, Coomassie plus reagent (300 µL) was introduced to the Ab-conjugated NPs (i.e., 10 mg dispersed NPs in 200 µL PBS pH 7.4) and incubated for 10–15 minutes. Then, a microplate reader (EL800; BioTek, Winooksi, VT, USA) was used to read the absorbance at 595 nm, and the readings were compared to a standard plot of BSA solution at different concentrations (ranging from 10 to 1000 µg/mL).

**Cell culture**

Cell lines including MS1, MS1HuTem1, TC1, and T cells were cultured in well-plates using RPMI 1640 (GlutaMAX™) supplemented with 10% fetal bovine serum, 100 units/mL penicillin G, and 100 µg/mL streptomycin. T cells were cultured in 24-well plates at a seeding density of 5.0×10⁴ cells/well, which were exposed to 20 U/mL of IL-2 to help their proliferation before the assays. The cultured cells were kept in a humidified chamber at 37°C for 12 hours. They were then washed with PBS (×2). The cells were fixed by the addition of formaldehyde (3.7% in PBS) (1 mL in the chamber, 1 mL in the lower chamber) at room temperature for 2 minutes. Formaldehyde was discarded and chambers were washed x2 with PBS. Cells were permeabilized by the addition of 100% ethanol at room temperature for 20 minutes. Ethanol was discarded and chambers were washed x2 with PBS. Giemsa stain was added (1 mL in the chamber, 1 mL in the lower chamber) at room temperature for 15 minutes. Giemsa was discarded and chambers were washed x2 with PBS. The non-migrated cells were scraped off with cotton swabs, while the migrated cells were counted under a light microscope.

For cell invasion assay, the chambers were detached using 0.25% Trypsin-EDTA solution, washed with DPBS, and re-suspended in serum-free cell culture media containing 0.1% BSA (1.0 × 10⁶ cells/mL). About 50 µL of Matrigel was added to a 24-well transwell insert and solidified in a 37°C incubator for 15-30 minutes, to form a thin gel layer. About 50 µL of cell suspension was added on top of the Matrigel coating and incubated at 37°C and 5% CO₂ for 10 minutes, to allow the cells to settle down. Using a pipette, 600 µL of full media was added to the membrane without damaging it. The transwell insert was washed (×2) with 1.0% acetic acid solution, the samples were added with 200 µL of 10 mM Tris base solution (pH 10.5) and kept under shaking for 30 minutes. Finally, the absorbance was read by the EL800 microplate reader at 570 nm, and the viability of cells was calculated.

**Cell migration and cell invasion assay**

TC1 and MS1 cells were treated with SHK (0.5, 1.5, 2.5, 5 µM), doxorubicin (DOX) 100 nM as the positive control, or full media as the negative control for 6 h before the assays. For migration assay, transwell migration chambers were placed in 24-well plates. The serum-free medium (100 µL) was added into the chamber, followed by the addition of 200 µL pretreated cells in suspension in serum-free medium (2.5×10⁶ cells/mL). In the lower chamber, 750 µL of full media were added. The cells were incubated at 37°C for 12 h. They were then washed with PBS (×2). The cells were fixed by the addition of formaldehyde (3.7% in PBS) (1 mL in the chamber, 1 mL in the lower chamber) at room temperature for 2 minutes. Formaldehyde was discarded and chambers were washed x2 with PBS. Cells were permeabilized by the addition of 100% ethanol at room temperature for 20 minutes. Ethanol was discarded and chambers were washed x2 with PBS. Giemsa stain was added (1 mL in the chamber, 1 mL in the lower chamber) at room temperature for 15 minutes. Giemsa was discarded and chambers were washed x2 with PBS. The non-migrated cells were scraped off with cotton swabs, while the migrated cells were counted under a light microscope.

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**Internalization of NPs**

To analyze the internalization of 78Fc-armed NPs by TEM1-positive/-negative MS1 cells, the flow cytometry technique was employed.²⁸ To this end, the cells were treated with free SHK or SHK-loaded NPs for 2 hours, and then, the plates were kept on the ice, and the internalization analysis was performed utilizing BD Canto II flow cytometer using the PE filter.

**Cytotoxicity evaluation**

The cytotoxic impacts of free SHK and SHK-loaded PLGA NPs were analyzed using sulforhodamine B (SRB) assay, as described previously.²⁸ The cells were cultivated at a seeding density of 3.0×10⁴ cells/cm² in 96-well plates. About 24 hours post-cultivation, they were treated with different concentrations of SHK or equivalent SHK-loaded NPs. The treatments were then removed after 2 hours of initial exposure, and the cells were incubated for 24 hours. After fixing the cells with 10% trichloroacetic acid at 4°C for 60 minutes, they were washed (×4) with distilled water. Then, the samples were stained with 0.057% w/v in 1.0% v/v acetic acid solution at room temperature for 30 minutes. Once washed (×4) with 1.0% acetic acid solution, the samples were added with 200 µL of 10 mM Tris base solution (pH 10.5) and kept under shaking for 30 minutes. Finally, the absorbance was read by the EL800 microplate reader at 570 nm, and the viability of cells was calculated.
was dipped into 70% ethanol for 10 minutes, to allow cell fixation. Afterward, a cotton-tipped applicator was utilized to eliminate the remaining ethanol from the top of the membrane. Then, the transwell membrane was kept to dry for 15 minutes. The crystal violet solution (0.2%) was used to stain the cells. Then, the transwell insert was gently washed several times with distilled water to remove the crystal violet excess and the transwell membrane was allowed to air-dry. Finally, an inverted microscope was employed to count the cells and get an average sum of the cells migrated through the membrane towards the chemo-attractant and attached to the membrane.

Colony formation assay
The cells were cultivated at a seeding density of 2.5 × 10^4 / flask in 25-T flasks for 4 hours to allow them to attach but not to multiply. Then, the cells were received the following treatments for 5 hours full media (i.e., the negative control), doxorubicin 100 nm (i.e., the positive control), SHK 0.3, 1.5, 2.5, and 5 µM. About 5 hours after initial treatment addition, samples were harvested. The number of cells in the resulting cell suspension was counted and diluted in sterile tubes so that approximately 5.0 × 10^5 cells could be pipetted into the 6-well plates. Samples were pipetted in 6-well plates in triplicates and placed in an incubator. They were kept until the cells in the control dishes have formed sufficiently large clones (8 days). Once clones were formed samples were washed twice and 2 mL of fixation/staining solution (6.0% glutaraldehyde and 0.5% crystal violet) were added to each well at room temperature for 30 minutes. Then, the glutaraldehyde crystal violet mixture was removed and carefully rinsed with tap water. The samples were air-dried at RT. The plating efficiency (PE) and surviving factor (SF) values were calculated as follows.

PE = (number of colonies formed/ number of cells seeded) × 100%

SF = number of colonies formed after treatment/ number of cells seeded × PE

Fluorescence microscopy
Cells at a seeding density of 1.0×10^4 were grown on a coverslip for 24 hours. The next day, the cells were treated for 2 hours with SHK 2.5, 5, and 10 µM. The treatments were removed from the samples washed twice and fixed with 2% Paraformaldehyde with Triton X. Then, the nuclei were stained with DAPI and the cellular membrane Wheat Germ Agglutinin, Alexa Fluor® 488 Conjugate according to the manufacturer’s instruction. Samples were mounted with PloLong Diamond Antifade Mountant and images were taken with a Leica TCS SP8 confocal microscope (Leica Microsystems GmbH, Wetzlar, Germany).

β-Galactosidase (13-Gal) staining
To detect senescent cells, samples were stained for β-galactosidase. The cells were grown in 6 well plates 2.0 × 10^4 cells/well. The next day, the cells were treated with the full media and doxorubicin 100 nM (as the negative and positive controls, respectively), SHK 2.5 µM, 5 µM, and 10 µM for 4 hours. After 4 hours treatment, the samples were washed with PBS and fixed with 2% paraformaldehyde at room temperature for 5 minutes. Then, they were washed and incubated with the fresh senescence-associated 3-Gal (SA-,3-Gal) stain solution 1 mg/mL at 37°C for 12 hours. After staining, cells were washed, air-dried, and images were taken using an inverted wide-field microscope.

JC1 staining
The cells were cultured in 6-well plates 5.0 × 10^4 cells/ml in a CO₂ incubator overnight at 37°C. The next day, the cells were treated with full media (negative control), camptothecin 20 µM (positive control), SHK 0.3, 1.5, 2.5, and 5 µM. For the flow cytometry analysis, 5 hours after treatments, the samples were washed twice and 100 µL/ mL JC-1 staining solution was added to each well. Samples were incubated in a CO₂ incubator at 37°C for 15 minutes. Samples were washed, harvested, resuspended in PBS, and analyzed promptly. In healthy cells with functional mitochondria, the red JC-1 molecules can be accumulated and detected in the FL2 channel. However, the apoptotic cells or the unhealthy ones with damaged mitochondria contain mainly green JC-1 monomers and are detectable in the FL-1 channel. For the fluorescence microscopy analysis, 5 hours after treatment, samples were washed twice and 100 µL/mL JC-1 staining solution was added to each sample, and samples were incubated in a CO₂ incubator at 37°C for 20 minutes. The cells were imaged directly in the culture medium since phenol red does not interfere with fluorescence staining. The accumulated JC-1 molecules in the healthy cells were detected using the Texas Red filter of the fluorescence microscope, while the JC1 monomers in the unhealthy/apoptotic cells were detected using the FITC filter of the fluorescence microscope.

Reactive oxygen species (ROS) formation evaluation
For ROS formation analysis, 2’,7’-dichlorofluorescein diacetate (DCFDA) staining was performed. TC1 cells were grown in 6-well plates to reach 70% confluency; then treated with SHK (2.5, and 5.0 µM) and H₂O₂ (100 mM) as the positive control, and incubated for 3 h before staining with DCFDA. Before experiments, 20 mM DCFDA stock was prepared in absolute ethanol and kept at 4°C until use. For flow cytometry, the cells were trypsinized and re-suspended at the density of 2×10^5 cells/mL. Then, 1 µL of 20 mM DCFDA was added to each sample (1 mL/sample). The samples were incubated at 37°C in dark for 15 minutes, then treated with PBS as a control. The samples were washed (×2), then re-suspended in PBS, and analyzed through a flow cytometer equipped with a 488
The samples were immediately read with PBS and images were captured through a fluorescence microscope (FITC filter).

Acridine orange staining
The cells were cultured in 6 well plates with a density of 5.0 ×10^3 cells/mL in a CO₂ incubator overnight at 37°C for 24 hours. On the next day, the cells were received the following treatments, full media as the negative control, rapamycin 500 nm as the autophagy inducer (positive control), SHK with different concentrations (0.3 μM, 1.5 μM, 2.5 μM, and 5 μM) for 5 hours. Flow cytometry was performed to assess the generation of acidic vesicular organelles (AVOs). The cells were stained with acridine orange solution (1 μg/mL) for 15 minutes. They were then detached from the plate with trypsin-EDTA and resuspended in a phenol red-free growth medium (1.0×10^3 cells/mL). The samples were immediately read through a flow cytometer (BD Canto II) at excitation 488 nm, and red fluorescence >600 nm versus dot plot of green fluorescence at 530 nm. For imaging of the autophagic vacuoles, 5 hours post-treatment, samples were washed twice and then incubated with acridine orange solution (1 μg/mL) in serum-free medium at 37 °C for 15 minutes. The acridine orange was removed, samples were washed (×2), and the cell images were acquired employing an inverted fluorescence microscope. For the analyses, the cytoplasm and nucleus of the cells were stained in bright green (detected with the FITC filter), whereas the acidic autophagic vacuoles were stained in bright red (detected with the Texas Red filter).

Flow cytometry analysis of apoptosis necrosis
To identify any transition from the apoptotic condition to the necrotic stage, a flow cytometry assay was applied utilizing phospholipid-binding annexin V protein that can show high affinity to phosphatidylserine in the presence of Ca²⁺, which is the major sign of early apoptosis. To this end, the cultured cells in a six-well plate were treated with different concentrations of SHK alone or SHK NPs. About 12 hours post-treatment, the cells were delicately trypsinized and centrifuged at 200 × g for 5 minutes. Afterward, the cells (around 1.0 × 10⁵ cells) were washed with 5.0 mL ice-cold PBS by centrifugation at 200 × g for 5 minutes. Then, they were resuspended in 500 μL of binding buffer (1x), which was supplemented with 5 μL of the apoptosis detection reagent (annexin V-EnzoGold) and 5 μL of the necrosis detection reagent, and incubated at room temperature for 15 minutes. To detect apoptosis and necrosis, the flow cytometry analyses were carried out using the FL2 and FL3 channels, respectively. The positive control cells (i.e., using UV irradiation for 15 minutes) and the untreated control cells were used for the comparison.

In vivo biodistribution analysis
To investigate the targeting efficacy of the 78Fc-armed, SHK-loaded NPs, we used a model developed previously. PE Gylated PLGA NPs loaded with ICG or free ICG (0.5 nmoles ICG in 1x DPBS) were injected (100 μL/injection) intravenously in female athymic nu/nu mice (strain C 88 Nude -/-, 18g, 4-weeks, Charles River Laboratories) (n=4/group). Infrared detection images were taken before injection and immediately after injection, 50 minutes, 2 hours, 6 hours, 12 hours, 24 hours, 48 hours, and 72 hours after injection. About 72 hours after injection, no signal was detected, mice were sacrificed and organs (brain, heart, lungs, thymus, kidneys, spleen, liver, oviduct, ovaries) were harvested and an infrared image of the organs was taken to detect an infrared signal. Images were taken using the Li-COR Pearl Impulse instrumentation provided by the University of Pennsylvania Small Animal Imaging Facility (SAIF) Optical/Bioluminescence Core, supported by NIH grant CA016520.

Targeting efficacy in the mouse model
Human TEM1 highly- expressing MS1 and MS1 cell lines, both expressing Luciferase at the same level were provided by Dr. Chunsheng Li. Tumor vascular model subcutaneous grafts expressing hTEM1 were established by the injection of 2.0 × 10⁷ MS1-hTEM1 or MS1 in each flank of nude female mice. Before tumors were implanted, the female athymic nu/nu mice (strain C 88 Nude -/-, 18g, 4-6 weeks, Charles River Laboratories) were acclimatized at the SAIF vivarium for 1 week, after which the mice were supplied with food and water ad libitum. Cellular growth was monitored by luminescence detection on a daily basis. Two weeks after the injection of cells, hemangiomas were established in both flanks, and treatments were injected intravenously into the retro-orbital complexes. The treatments were as follows: saline-control group; 5mg/kg free SHK; PLGA PEGylated empty NPs, PLGA PEGylated SHK-loaded NPs content of SHK/injection 5 mg/kg; 78Fc-armed PLGA PEGylated SHK-loaded NPs, content of SHK/injection 5 mg/kg; 100 μL/injection; n=7 mice/group. Treatments were injected intravenously on days 1, 3, 5, 7, and 9 after the hemangiomas were established. Small animal imaging detecting cellular luminesce and SHK fluorescence were being taken every other day using Perkin Elmer IVIS Spectrum instrumentation provided by the University of Pennsylvania SAIF Optical/Bioluminescence Core, supported by NIH grant CA016520. On day 14, the animals were sacrificed and hemangiomas were collected. Mouse weight was monitored daily for any indication of weight loss and/or behavioral change.

Therapeutic efficacy in the mouse model
The subcutaneous tumors with the overexpression of endogenous murine TEM1 were developed through the
injection of 4.0×10⁴ TC1 cells into the right flank of female nude mice (4-6 weeks old). Before the implantation of tumors, female athymic nu/nu mice (strain O88 Nude-/-, 18 g, Charles River Laboratories) were acclimatized at the SAIF vivarium for 1 week. Mice were carefully supplied with food and water ad libitum. The injected TC1 cells expressed firefly luciferase, enabling BLI imaging. On day 7, after cell inoculation TC1 tumors were established. The animals were intravenously injected via the retro-orbital complexes with various treatments, including saline-control group; 5 mg/kg free SHK; PLGA PEGylated empty NPs; PLGA PEGylated SHK-loaded NPs content of SHK/injection 5 mg/kg; 78Fc-armed PLGA PEGylated SHK-loaded NPs, the content of SHK/injection 5 mg/kg; 100 μL/injection; n=7 mice/group. The treatments were administered on days 1, 3, 5, 7, and 9 after tumors were established, while, to detect luminescence and SHK fluorescence, images were taken every other day. Animal weight was monitored daily. On the day 14th, the animals were sacrificed and tumors were collected.

**Therapeutic efficacy in the metastatic mouse model**

The metastatic tumors in the lung were developed by intravenous injection of TC1 cells (1.0×10⁵ luciferase-expressing TC1 cells) via the tail vein. These mice grow BLI visible lung tumors within 7 days after the cell inoculation, which otherwise succumbed in 3-4 weeks if no treatment is provided. Before tumors were implanted, female athymic nu/nu mice (strain O88 Nude-/-, 18 g, 4-6 weeks old, Charles River Laboratories) were acclimatized at the SAIF vivarium for 1 week. The animals were supplied with food and water ad libitum. On day 3 after TC1 cell injection, treatments were injected intravenously into the retro-orbital complexes, as follows: saline-control group; 5mg/kg free SHK; PLGA PEGylated empty NPs; PLGA PEGylated SHK-loaded NPs content of SHK/injection 5 mg/kg; 78Fc-armed PLGA PEGylated SHK-loaded NPs, the content of SHK/injection 5 mg/kg; 100 μL/injection; n=7 mice/group. The treatments were administered on days 1, 3, 5, 7, and 9 after tumors were established, while, to detect luminescence and SHK fluorescence, images were taken every other day. Animal weight was monitored daily. On day 16, the animals were sacrificed and the lungs were collected.

**Therapeutic efficacy in the immunocompetent mice model**

The subcutaneous tumors with the overexpression of endogenous murine TEM1 were established by the injection of 1.0×10⁴ TC1 cells into the right flank of female 8-week-old C57BL/6 mice. Mice were acclimatized at the SAIF vivarium for 1 week before tumors were implanted. Seven days after cell inoculation tumors were established. Treatments were injected intravenously into the retro-orbital complexes, as follows: saline-control group; 5mg/kg free SHK; PLGA PEGylated empty NPs; PLGA PEGylated SHK-loaded NPs content of SHK/injection 5mg/kg; 78Fc-armed PLGA PEGylated SHK-loaded NPs, content of SHK/injection 5 mg/kg; 100 μL/injection; n=7 mice/group. The treatments were administered on days 1, 3, 5, 7, and 9 after tumors were established, while, to detect luminescence and SHK fluorescence, images were taken every other day. Animal weight was monitored daily. On day 14th, animals were sacrificed and tumors and spleens were collected. The experiment was repeated three times. For the TC1 re-challenge, the animals treated with 78Fc-armed NPs and were tumor-free on the last day of the experiment were collected and monitored. After a month, about 1.0×10⁴ TC1 cells were injected into the right flank of these mice, and the tumor growth was monitored by luciferase detection every other day.

**Therapeutic efficacy in the re-challenged animals**

To further investigate if the immune system was indeed active after the therapy, instead of sacrificing the tumor-free mice, we collected and observe them for tumor growth. After 3 weeks, we attempted to re-challenge them by injecting TC1 cells subcutaneously. The tumor growth was monitored by luciferase detection, while no further treatment was given to the animals.

**TUNEL assay**

To detect the apoptotic dead cells in situ, the TUNEL assay was employed using the Cell/Death Detection Kit (Roche Applied Science; catalog no. 11684809910) according to the manufacturer’s guidelines. Briefly, frozen sections with about 6 μm thickness from the OCT-embedded tissue were allowed to air-dry under the hood for 30 minutes. The samples were then fixed with 2% paraformaldehyde in PBS at room temperature for 30 minutes. Each sample was treated with 100 μL of 3% H₂O₂-methanol solution at room temperature for 20 minutes and washed (×3) with PBS. To block the nonspecific binding sites, each sample was further treated with a 10% normal goat serum blocking solution at room temperature for 1 hour. The samples were analyzed through a Nikon Ti fluorescence microscope and a Leica TCS SP8 confocal microscope.

**Immunofluorescence assay**

Tumor samples were sliced using a cryostat, and tissue slides were frozen. Frozen tissue slides were allowed to air-dry for 30 minutes. The tissue slides were fixed in 4% paraformaldehyde in 1x warm PBS (pH 7.4) at room temperature for 15 minutes. Slides were washed (×3) with 1x PBS. Samples were incubated in 3% H₂O₂ in methanol at room temperature for 20 minutes, followed by washing (×3) in PBS. Normal goat serum (10%) was used for blocking (1 hour incubation at room temperature). Blocking solution was aspirated and a mixture of primary antibodies (Abs) was added (150 μL final volume of anti-CD31 dilution 1:200 (suggested 1:300), anti-ICAM 1:500 dilution, and anti-VCAM 1:700 dilution, in 5%
Goat serum). Samples were incubated with primary Abs at 4°C overnight. The following day, the slides were washed (×3) with PBS. Then, a mixture of the secondary Abs was added (5 μg/mL final concentration in 5% goat serum) and samples were incubated at room temperature in dark for 1 h. Slides were washed (×3) in PBS. Finally, the stained tissue was mounted with PloLong Diamond Antifade Mountant with DAPI.

**IFN-γ enzyme-linked immunosorbent spot assay (ELISpot)**
To carry out the ELISpot analysis, 96-well MAIP plates were coated with a 2.5 μg/mL solution of rat anti-mouse IFN-γ overnight. Bulk splenocytes were collected and plated at 1.0 × 10^6 cells/well in triplicate. The samples were then incubated with 2 μg/mL peptides at 37°C for 20 hours. Afterward, the samples were washed with PBS and 0.05% Tween 20. The samples were then exposed to anti-mouse biotin-conjugated anti–IFN-γ antibody and incubated at 4°C overnight. The following day, streptavidin-alkaline phosphatase conjugate was introduced for 30 minutes. The samples were developed through the addition of nitro blue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate. Finally, the spots were analyzed employing an automated ELISpot reader (Autoimmun Diagnostika GmbH, Strassberg, Germany).

**Study approval**
Before conducting any in vivo studies, all animal studies protocols were approved by the IACUC and University Laboratory Resources at the University of Pennsylvania. Mice were treated following the Animal Care and Use Committee guidelines of the University of Pennsylvania.

**Results**

**Development and characterization of shikonin nanoparticles**
SHK-loaded PLGA NPs were prepared, PEGylated, and functionalized with anti-TEM1 78Fc scFv (Fig. S1), as described previously. One-step PEGylation and 78Fc conjugation resulted in the production of NPs with a high drug content and a conjugation efficiency of ~90% (Table S1). The unmodified and 78Fc-armed functionalized PLGA NPs displayed smooth spherical morphology with an average size of around 130 nm and zeta potential of approximately –35 mV (Fig. S1). After the PEGylation and conjugation of 78Fc, DC%, DE%, DL%, and YE% of SHK-loaded NPs were somewhat reduced (Table S1). Among various kinetic models used to fit the SHK release profile, the SHK release profile was found to obey Wagner’s log-probability (Table S2). The pattern of drug release exhibited that 20% of the drug molecules could be released within the first 2 hours. Nevertheless, about 80% of the drug was found to be released after 48 h. The release of SHK from the NPs appeared to be somewhat pH-dependent with the highest release at pH 7.4 (Table S2).

**Internalization and cytotoxicity of SHK NPs**
The cellular uptakes of the free SHK, SHK-loaded PEGylated PLGA NPs, and 78Fc-armed SHK-loaded PEGylated PLGA NPs in the mammalian cells (MS1HuTem1, MS1, and TC1 cell lines) are shown in Fig. 1. Confirming our previous findings, the engineered NPs enhanced the entry of SHK into the cells, while the functionalization of NPs with 78Fc could further enhance the internalization of NPs in the TEM1-expressing cells such as MS1HuTem1 or TC1 cells, but not in the TEM1-negative MS1 cells (Fig. 1A, B and C).

Based on the cytotoxicity analysis, the SHK nanoformulation was found to display higher cytotoxicity in the MS1, MS1HuTem1, and TC1 cell lines (Fig. 1D, E, and F, respectively). The empty NPs did not induce any cytotoxic effects, indicating the safety of nanoformulation itself. As shown in Fig. 1, the TEM1-targeting NPs showed significantly higher cytotoxicity in the TEM1-overexpressing cells, implying the high capacity of the 78Fc-armed SHK-loaded PEGylated PLGA NPs in the active targeting of the TEM1-expressing cells in vitro. On the other hand, while the free SHK showed a higher impact on the T lymphocytes, the SHK nanoformulation resulted in significantly lower toxicity (Fig. 1G), indicating that the cytotoxic effect of the nanoformulation is largely dependent on the active targeting mechanism and internalization of NPs into the target cells.

**Inhibition of cell migration and invasion**
As the most relevant approach for the establishment of de novo strategies for the effective diagnosis and therapy of cancer, we looked at the SHK effects on the migration and invasiveness of the tumor cells (Fig. 2). To mimic the migration of cancer cells in vivo, we studied the effects of SHK on the cell movement through the extracellular matrices and found SHK to induce substantial inhibitory effects on the invasive migration as a fundamental function underlying the cellular processes, including metastasis and invasion of cancer cells (Fig. 2). SHK at a concentration of 5 μM (i.e., IC_{50}) was able to effectively inhibit both the cell migration and invasion (Fig. 2B and C, respectively).

**SHK effects on the formation of colony and heterochromatin foci and senescence**
The colony formation analysis, for studying the anchorage-independent growth as a hallmark of carcinogenesis, revealed that SHK was able to inhibit the formation of colonies significantly as compared to DOX used as the positive control (Fig. 3A and B). The cytotoxicity of SHK was further investigated to find the possible mechanisms behind such cytotoxicity. The fluorescence microscopy analysis resulted in somewhat morphological changes in the nucleus of the cells upon treatment with SHK (Fig. 3C). Clear DNA-stained dense foci were observed, which were readily distinguishable...
from the chromatin in the normal cells resembling the senescence-associated heterochromatin foci (SAHF) in the SHK-treated cells (Fig. 3C), indicating of possible induction of the senescence and dilapidation by SHK. The senescence analysis through β-galactosidase staining revealed that only a few SHK-treated cells displayed increased β-galactosidase metabolism compared to the positive senescence inducer DOX-treated control cells (Fig. 3D). In the treated MS1 cells with 100 nM DOX, the whole cellular population showed increased β-galactosidase metabolism (Fig. 3E). These results indicate that SHK is not able to induce senescence.

Induction of mitochondrial depolarization and autophagosomes by SHK

SHK, at a concentration above 2.5 μM, was found to induce some important morphological changes, including mitochondrial depolarization in the treated TC1 cells (Fig. 4A). The effect was found to be concentration-dependent (Fig. 4B). A strong signal of SHK was detected in the cytoplasm with an area of condensed chromatin in the nuclei of cells treated with SHK (Fig. 4C).

Upon analysis by using fluorogenic dye, DCFDA was used to measure the hydroxyl/peroxyl and other reactive oxygen species (ROS) activities within the cell, in which the

Fig. 1. The internalization and cytotoxicity of shikonin (SHK)-loaded nanoparticles (NPs) in different cell lines. A) Flow cytometry analysis of SHK and SHK-loaded NPs uptake (P1: the untreated control cells, P2: the cells treated with SHK in TEM1-positive, MS1HuTem1 endothelial cells, and TEM1-negative, MS1 endothelial cells, as well as TC1 cells. B) The uptake of NPs by MS1 and MS1HuTem1 cells. C) The uptake of NPs by TC1 cells. D) Treated cells with 2.5µM SHK. E) Treated cells with 5 µM SHK. F) TC1 cells treated with 2.5µM, 5µM, and 10µM SHK. G) T cells treated with 5µM SHK. The cultured cells were exposed to treatments for 2 h then the treatments were removed and cells were subjected to SRB cytotoxicity assay 24h after the addition of the treatment.
Fig. 2. The effects of shikonin (SHK) on the tumor cells migration and invasion in different cell lines. A) The inhibitory effects of SHK on the migration and invasion abilities of the MS1 and TC1 cells cultivated in the Matrigel-coated transwells. B) The quantified inhibitory effects of SHK on the migration of cells. C) The quantified inhibitory effects on the invasion of cells.

Fig. 3. The SHK biological impacts. A) The formation colony in the treated MS1 and TC1 cells 8 days after the treatment. Eight days after the treatment, the cells were fixed and stained with Crystal violet. Threshold 50 cells and higher were considered a single colony. B) The quantification of the colony formation in the MS1 and TC1 cells. C) The fluorescence microscopy image of DAPI staining of MS1 cells treated with SHK 2.5 μM. The DNA foci formation after SHK treatment is pointed with yellow arrows. D) β-galactosidase staining of MS1 cells treated with 2.5 μM SHK. E) β-galactosidase staining of MS1 cells treated with 100 nM DOX. Data are means ± SD of four independent experiments. *P < 0.05.
formation of ROSs was observed in the SHK-treated cells as compared to the untreated negative and H$_2$O$_2$-treated positive cells (Fig. 4D). The SHK-induced mitochondrial depolarization appeared to elicit the formation of ROSs in a dose-dependent manner (Fig. 4E and 4F).

Based on the acridine orange staining analysis, the AVOs were formed in the cells treated with SHK, in large part because of the induction of autophagosomes elicited by ROS in the SHK-treated MS1 and TC1 cells (Fig. 4G and 4H, respectively). The autophagosomes were formed in the cells treated with lower concentrations of SHK (2.5 µM) but not in the cells treated with the toxic concentrations of SHK (Fig. 4J), indicating that the lower concentrations of SHK could function as an autophagy inducer. The breakdown of the mitochondrial transmembrane potential might be attributed to the mitochondrial permeability transition pores opening, which might result in the release of cytochrome c into the cytosol. Such a phenomenon, in turn, can prompt the activation of other downstream events in the apoptotic cascade. The induction of mitochondrial depolarization by SHK was further validated with the JC1 staining, which revealed that SHK could cause a shift in the mitochondrial membrane potential in a dose-dependent manner, resulting in enhanced mitochondrial depolarization (Fig. 5).

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**Fig. 4.** Induction of mitochondrial depolarization, reactive oxygen species (ROSs), and the acidic vacuolar organelles (AVO) in the cells treated with shikonin (SHK). A) Fluorescence microscopy images of the TC1 cells treated with SHK (1.5 µM, 5 µM, and 10 µM) for 5 h and stained with JC1. The depolarized and normal mitochondria are shown as green and red, respectively. B) Quantification of green/red fluorescence ratio of SHK-treated TC1 cells. C) Confocal microscopy image of the TC1 cells treated SHK (5 µM for 3 h). The cells were fixed and stained with Wheat germ agglutinin (membrane, green) and DAPI (nucleus, blue), and SHK (red). SHK localized in the cytoplasm. D) Flow cytometry results after 3 h treatment. E) Light and fluorescent microscopy images of DCFDA stained cells after 3h treatment. F) Quantification of ROS-positive cells treated with SHK. G and H) Flow cytometry analysis of fluorescence ratio in the TC1 cells and the MS1 cells, respectively. The cells were analyzed after 5h treatment with various concentrations of SHK. J) Fluorescence microscopy images of the untreated (left) and the SHK (2.5 mM)-treated (right) cells. *P < 0.05.
Matthaiou et al. (2022) performed flow cytometry analysis to detect mitochondrial depolarization in TC1 and MS1 cells. They found that mitochondrial depolarization increased with higher SHK concentrations. Specifically, in the SHK 5µM treated cells, approximately 50% of the cell population showed depolarized mitochondria. Additionally, they used an apoptosis/necrosis assay by annexin-V FITC/PI to study the effect of SHK 5µM or staurosporine (an apoptosis inducer) 1µM on cell lines. The results showed an increase in the percentage of apoptotic/necrotic cells in both MS1 and TC1 cell lines after treatment. *P < 0.05.
Apoptosis and necrosis effects of NPs in vitro
The apoptosis and necrosis analyses, performed using the annexin-V FITC/PI in MS1 and TC1 cell lines after treatment with SHK 5μM or staurosporine (apoptosis inducer) 1 μM at 37°C for 2 hours, three populations of cells observed in both cell lines (Fig. 6A). These cells included (i) the viable cells with no apoptotic or necrotic signs (i.e., annexin V-cyanine 3 and NDR negative); (ii) the apoptotic cells (i.e., annexin V-cyanine 3 positive and NDR negative); and (iii) the cells underwent the late-stage apoptosis and early necrosis (i.e, annexin V-cyanine 3 and NDR positive). The ratio of the apoptotic to the necrotic cells in the SHK-treated MS1 and TC1 cells was 1:1, while the ratio of the apoptotic to the necrotic cells treated with staurosporine was 2:1 (Fig. 6B and C). These findings suggest that SHK can induce both cell death mechanisms.

In vivo biodistribution and targeting efficacy in a mouse model
The biodistribution analysis of the engineered NPs loaded with indocyanine green (ICG) injected intravenously via the retro-orbital route revealed no signal of ICG in the mice after 24 hours. Once the organs were harvested, there was no high detection of ICG in any of the organs observed, suggesting that the NPs did not maintain by the liver after 72 hours. However, there was somewhat retention of the ICG in the liver in the control group (free ICG treated) (data not shown). The active targeting efficacy of the 78Fc-armed SHK-loaded NPs was confirmed using a mice model subcutaneously injected with the MS1 and the MS1HuTem1 cells respectively at the right and left sides (Fig. 7A). The luciferase was detected on the first day of the detection and the last day of the experiment (Fig. 7A). On the first day of the experiment, the two sides appeared to have the same number of cells; however, after treatment with the 78Fc-armed NPs, the MS1HuTem1 cells were significantly eliminated (Fig. 7B), indicating the high in vivo targeting efficacy of the 78Fc-armed SHK-loaded NPs, as well as the efficacy of SHK as a cytotoxic agent. The luminescence quantification confirmed the results.

Fig. 7. In vivo targeting efficacy of anti-TEM1 78Fc-armed SHK-loaded PLGA NPs. Mice were bearing MS1 cells subcutaneous at their right side and MS1HuTem1 cells at their left side. A) Representative image of luminescence detection of the MS1 and MS1-huTem1 cell lines. B) Quantification of cellular luciferase expression. C) Luminescence detection on the first (top) and the last day (bottom) of the experiment. D) Quantification of the MS1HuTem1/MS1 luminescence detection ratio of control and anti-TEM1 Fc armed SHK-loaded PLGA NPs treated groups. E) Detection of SHK fluorescence 24h after the first injection (left) and on the last day of the experiment (right). SHK high fluorescent signal is detected in the MS1-huTem1 hemangioma, even 5 days after treatments, while the MS1-huTem1 hemangioma disappeared. F) Representative image of detection of fluorescence of SHK serial dilutions in PBS (top) human serum (middle) and FBS (bottom). SHK: Shikonin, PLGA: poly(lactic-co-glycolic acid, NPs: nanoparticles. *P< 0.05.
(Fig. 7B). The MS1 and MS1HuTem1 cells were developed to have the same growth rate in vivo and to express the same levels of luciferase (Fig. 7C and 7D). Mice weight was monitored daily, and neither any fluctuation of animal weight nor any behavioral change was observed, implying that this nano-therapy did not impose severe toxicity. As shown in Fig. 7E, the SHK high fluorescent signal was detected in the MS1HuTem1 hemangioma even 5 days after treatment, while we couldn’t detect any luciferase signal (live cells) in that area the same day. Furthermore, we noticed that when SHK was diluted in the serum, it could emit at 620 nm, indicating SHK as a compound easy to detect by in vivo imaging (Fig. 7F).

**Therapeutic efficacy of NPs in the mouse and metastatic mouse models**

The 78Fc-armed SHK-loaded NPs therapeutic efficacy analysis in the TC1 subcutaneously injected nu/nu mice model showed the luciferase expression in the 78Fc-armed SHK-loaded NPs treated group in comparison with the PBS-treated as the control on the first and the last day of the experiment (Fig. 8A and B). The quantity of the luciferase expression in the treated group with the 78Fc-armed SHK-loaded PLGA NPs was significantly lower than that of the PBS treated group (Fig. 8C). The tumor weight and size in the treated group with the 78Fc-armed SHK-loaded PLGA NPs were significantly reduced.
as compared to the control group (Fig. 8D and E).

The 78Fc-armed SHK-loaded NPs’ therapeutic efficacy in the TC1 lung metastasis model analysis revealed that the engineered NPs impose great efficacy in the treated animals (Fig. 8F and 8G). During the last day of the experiment, the tumor nodules at the lungs of the animals treated with the 78Fc-armed SHK-loaded PLGA NPs were significantly decreased or even diminished as compared to the control group (Fig. 8F and 8G). The quantity of luminescence revalidated such findings (Fig. 8F). The weight of the animals treated with the 78Fc-armed SHK-loaded PLGA NPs remained stable throughout the experiment as compared to the control group that showed weight loss and some behavioral changes (Fig. 8H).

**Therapeutic efficacy of NPs in the immunocompetent mice**

The SHK immunomodulatory capacity was investigated in the C57BL6 mice subcutaneously injected with the TC1 cells with results showing a lowered/diminished luciferase expression in the treated animals with the 78Fc-armed SHK-loaded PLGA NPs on day 14 (Fig. 9A and 9B). The quantity of luminescence validated the results (Fig. 9C and D). The animals treated with the engineered NPs showed good weight stability as compared to the PBS-treated group (Fig. 9E and 9F). On the last day of the experiment, approximately 72% of the mice population treated with the 78Fc-armed SHK-loaded PLGA NPs were tumor-free, while the rest 28% of the mice population had
extremely small tumors as compared to the control group or the PEGylated NPs group (Fig. 9G and 9H), indicating the therapeutic effects of the targeted nanoformulation in the immunocompetent C57BL6 mice.

**Therapeutic efficacy of NPs in re-challenged animals**

A few days after the TC1 cells re-challenge, the cellular growth started decreasing and by day 14 the TC1 cells were eliminated (Fig. 10A). The analysis of the tumor site where TC1 cells were injected on the last day of the experiment validated the results (Fig. 10B), which were further validated by the quantity of luminescence (Fig. 10C). These results indicate that the immune system of the C57BL6 mice remained to be functional after the treatment with the 78Fc-armed SHK-loaded PLGA NPs, enabling animals to recognize and eliminate the tumor cells.

**Induction of apoptosis and necrosis in vivo**

The results of the TUNEL assay showed that the slides of frozen tissue samples treated with the 78Fc-armed SHK-loaded PEGylated PLGA NPs were stained positive for TUNEL reagent as compared to the PBS treated group as the control. This finding confirms the induction of apoptosis *in vitro* by the DNA fragmentation resultant from apoptotic signaling cascades (Fig. 11).

Further, based on the immunofluorescence and TUNEL analyses, the induction of apoptosis *in vivo* in the slides of frozen tissue samples treated with 78Fc-armed SHK-loaded NPs was detected as compared to the PBS treated group as the negative control. It was found that SHK treated cells could exhibit apoptotic: necrotic cell death at approximately 1:1 ratio (Fig. 12).

Further, the IFN-γ enzyme-linked immunosorbent spot assay analysis was conducted to study whether the murine immune system was able to recognize and eliminate the TC1 cells. The ELISpot analysis, which was performed on the splenocytes of the 78Fc-armed SHK-loaded NPs treated tumor-free mice, showed a weak response to the E6/E7 peptides which was not statistically significant as compared to the positive control.

**Discussion**

Despite the constant advances in the field of oncology,
cancer remains to be one of the leading causes of death worldwide. After a conventional chemotherapy course, cancer patients are prone to relapse and/or develop drug resistance. Although chemotherapy modalities have been accepted as an effective treatment strategy for most types of cancers, such an intervention is often linked with some undesired side effects. The major reason for the occurrence of adverse reactions is the cytotoxic nature of most of the anticancer chemotherapeutic agents and their nonspecific distribution to healthy tissues/organs. Many studies have attempted to develop targeted DDS and achieve specific/selective delivery of chemotherapeutics into the TME. Given that the tumor neovasculature exhibits a fenestrated endothelial feature, in combination with the poor lymphatic drainage at the TME, nanomedicines can accumulate passively within the TME through the EPR effect. The nanoscale delivery system can further be decorated with a targeting moiety to serve as a nanocarrier with active targeting potential. Different nanosized DDSs have been employed for the delivery of cytotoxic agents. Unlike non-biodegradable polymer-/lipid-based DDSs that are considered as toxic entities, most of the biodegradable polymers like PLGA happen to be a much safer drug carrier for the delivery of cytotoxic chemotherapeutics. In the present study, we developed an advanced nanoformulation platform as reported previously for the targeted delivery of SHK – a known natural secondary metabolite with cytostatic, cytotoxic, and immunomodulatory impacts in a concentration-dependent manner. The SHK was shown to activate both receptors and the mitochondrial-mediated apoptosis effectively and circumvent the cancer drug resistance via the initiation of necroptotic cell death. To limit the SHK cytotoxicity solely within the TME, we formulated the SHK-loaded PEGylated PLGA NPs armed with a TEM1-specific antibody fragment, 78Fc scFv (Fig. S1). Based on the composition (polymer and surfactant type and concentration) and formulation method, PLGA NPs were formulated with good loading efficiency (Table S1) and a size range of 100-140 nm with a smooth surface (Fig. S1). The release of SHK from PLGA NPs in both PBS and the human serum was investigated at different pHs to disclose the release profile within the vesicular machinery of the target cells (e.g., endosomal compartments) and the TME, where the pH is acidic. The release rate was found to be slower at the lower pH, indicating a sustained-release profile in an acidic condition (Table S2). Such behavior can lead to the accumulation of drug molecules at the TME, where the pH is somewhat acidic. It has been reported that NPs of this size can actively accumulate at the tumor site in approximately 2 hours, during which period the systemic release of SHK is speculated to be around 20%. However, the systemic release of a small amount of SHK might be in favor of the cancer therapy goals via stimulating the immune system. Once the NPs reached the TME through size-dependent passive targeting mechanism, they can attach to the TEM1-expressing cells through the targeting agent and actively accumulate within the TME where the remaining ~80% of SHK load can be released. Given that the cytotoxic effects of SHK are non-specific, the main purpose of the current study was to deliver SHK actively specifically to the target cancerous cells and investigate its impacts on these cells. The PLGA NPs were PEGylated to become stealth and

![TUNEL assay on frozen tissue samples. A) The DNAse treated positive control (left), the 78Fc treated group sample (middle), the PBS-treated group (right). Images were taken using confocal microscope magnification 10x, images insert magnification is 100x. B) Tissue detected fluorescence intensity quantified using Fiji software.](image-url)
long-circulation particles, and conjugated with anti-TEM1 78Fc scFv recruiting the EDC/NHS chemistry for the active targeting purpose (Fig. S1), which could actively target the TEM1-positive, MS1HuTem1 endothelial cells (Fig. 1). Our findings showed that the 78Fc-armed SHK-loaded PEGylated PLGA NPs could considerably bind to the TEM1-overexpressing cells and induce markedly high toxicity in the target MS1HuTem1 cells (Fig. 1). Our previous results on the effects of SHK led us to seek the mechanisms underlying the toxic impacts of SHK. Thus, we first looked at the cytotoxicity of SHK alone in the treated cells. We witnessed the inhibitory effects of SHK, at a concentration of 5μM (i.e., IC\textsubscript{50}), on the migration and invasion abilities of MS1 and TC1 cells cultivated in the Matrigel-coated transwells (Fig. 2). Besides, the malignant transformation analyses revealed that SHK could significantly inhibit the formation of colonies of the MS1 and TC1 cells comparable to that of DOX (Fig. 3). It can be assumed that the targeted delivery of SHK could impose its inhibitory impacts solely on the target cancer cells. SHK was also found to elicit some morphological changes in the cellular nuclei (Fig. 4), perhaps through chromatin remodeling. Even if the SHK-induced heterochromatin foci resembled the SAHF, SHK did not induce profound senescence in the treated cells (Fig. 4). The morphological analysis showed that SHK at a concentration around IC\textsubscript{50} can induce substantial cytotoxicity possibly through induction of ROSs (Fig. 4). Based on our findings, high amounts of SHK were able to accumulate in the cytoplasm and induce ROSs, which led us to assume that the mitochondria could be a possible biological target of SHK. The JC-1 staining experiments for studying the effect of SHK on the mitochondria showed a concentration impact of SHK resulting in depolarization of mitochondria (Figs. 4 and 5). It should be noted that SHK was able to induce ROS in a dose-dependent manner (Fig. 4). We also witnessed that the induction of ROSs by SHK induced somewhat mitochondrial degradation by autophagy and further mitochondrial depolarization and degradation. Moreover, in recent years, a growing piece of evidence discusses the oxidative stress function(s) as the converging point of autophagy, in which ROS is counted as one of the major intracellular signal transducers that might sustain the autophagy. Thus, we investigated the effect of SHK...
treatment in the formation of autophagosomes and found that it could induce the formation of autophagic vacuoles only in low (non-toxic) concentrations (Fig. 4). Further, SHK was able to induce both necrosis and apoptosis at a ratio 1:1 in the treated cellular population (Fig. 6). We ponder that the mitochondrial depolarization by SHK can result in apoptosis, which is deemed to be one of the SHK mechanisms of action (Figs. 5 and 6). This finding is in agreement with other researchers’ findings. Our results are in clear consensus with previous reports reporting on the potential of SHK in the induction of both necrosis and apoptosis (necroptosis) in the cancer cells.

To investigate the targeting efficacy of the 78Fc-armed SHK-loaded PEGylated PLGA NPs in vivo, we used an animal model developed previously. The biodistribution study showed that PLGA NPs were cleared from the mice after 72h. At the targeting efficacy mouse model employed, after treatment with the 78Fc-armed SHK-loaded PEGylated PLGA NPs, the MS1HuTem1 cells were eliminated (Fig. 7). These results demonstrate the high in vivo targeting efficacy of the 78Fc-armed SHK-loaded NPs, as well as the therapeutic efficacy of SHK as a cytotoxic agent. Furthermore, we noticed that a high fluorescent signal of SHK was detected in the MS1-huTem1 hemangioma, even 5 days after the treatment (Fig. 7), while we could not detect any luciferase signal (live cells) in the same area on the last day. Such observations suggest that the 78Fc-armed SHK-loaded PEGylated PLGA NPs not only bind to their target actively but also retain within the tumor site for a few days. At the therapeutic efficacy mouse model employed in the current study, the tumors treated with the 78Fc-armed SHK-loaded PEGylated PLGA NPs were significantly reduced as compared to the PBS-treated control group (Fig. 7). In the metastatic model, we studied the 78Fc-armed SHK-loaded PEGylated PLGA NPs that imposed significant therapeutic efficacy. The tumor nodules at the lungs of the 78Fc-armed SHK-loaded PEGylated PLGA NPs treated groups were significantly decreased or even eliminated (Fig. 8). The therapeutic efficacy mouse model was further studied using immunocompetent C57BL6 mice. In this model, we noticed that the 78Fc-armed SHK-loaded PEGylated PLGA NPs treatment had an even better therapeutic outcome compared to the nu/nu strain (Fig. 9). Among other hypotheses to explain these results, we assumed that the better therapeutic outcome could be attributed to the synergy of the immune system with the engineered targeted nanoformulation. The results of the tumor re-challenge study indicated that the targeted nanoformulation of SHK could induce an immune response towards the TC1 cells (Fig. 10). However, additional investigations need to be accomplished for a better understanding of such modulation/stimulation of the immune system by SHK before its translation into clinical applications.

Previous studies have emphasized that various nanoscaled formulations loaded with anticancer agents can become the game-changer in the field of cancer therapy soon. Given that the SHK was found to cause necroptosis in vitro, we decided to investigate the mechanism of death induced by SHK in vivo. Thus, frozen tissue samples from tumors treated with the 78Fc-armed SHK-loaded PEGylated PLGA NPs were stained with TUNEL reagent, anti-RIPK1, and anti-RIPK3 Abs. The slides from the 78Fc-armed SHK-loaded PEGylated PLGA NPs treated groups were stained positive for both TUNEL and the RIP kinases (Figs. 11 and 12). These data indicate that SHK nano delivery can induce both apoptosis and necroptosis in vivo.

**Conclusion**

SHK can bind to the mitochondria, which can cause mitochondrial depolarization leading to apoptosis. Further, SHK can induce the production of intracellular ROS and the appearance of autophagosomes in low concentrations. Moreover, SHK shows a great capability to induce apoptosis and necroptosis in the cell lines/models examined in the current study. Furthermore, SHK was able to inhibit cell migration and invasion in vitro. To confine the biological impacts of SHK on cancerous cells, we developed the 78Fc-armed SHK-loaded PEGylated PLGA NPs which showed extremely high targeting potential and therapeutic efficacy in all the in vivo models employed in the current study. Finally, the results of the tumor re-challenge experiment suggest that the targeted nanoformulation of SHK imposes immunostimulatory effects in vivo. Upon our findings, the engineered biodegradable PEGylated and anti-TEM1 78Fc scFv armed SHK-loaded PLGA NPs are proposed as a novel multifunctional nanomedicine for the targeted therapy of solid tumors with overexpression of TEM1.

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Ethical statement
All animal studies were approved by the IACUC and University Laboratory Animal Resources at the University of Pennsylvania. The authors declare that the raw data required to reproduce these reported results and the processed data required to reproduce the reported findings are available upon request.

Competing interests
YO and JB act as the Editor-in-Chief of the journal, and LEK is a member of the editorial board of the journal. The peer-review process of this work was carried out in a double-blind manner as part of the journal editorial policy.

Authors’ contribution
GC, CL, and YO designed the study. EIM conducted all the experiments. JB, YG, RS, and LEK assisted in data interpretation and analyses. EIM drafted the manuscript. EIM, JB, and YO revised the manuscript. GC, RS, and YO supervised the study.

Supplementary Materials
Supplementary file 1 contains Figure S1 and Tables S1-S2.

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TEM1-targeting PEGylated PLGA shikonin nanoformulation for ovarian cancer

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