PKHB1, a thrombospondin-1 peptide mimic, induces anti-tumor effect through immunogenic cell death induction in breast cancer cells

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ABSTRACT
Breast cancer is the most commonly diagnosed cancer and the leading cause of cancer death in women worldwide. Recent advances in the field of immuno-oncology demonstrate the beneficial immunostimulatory effects of the induction of immunogenic cell death (ICD). ICD increases tumor infiltration by T cells and is associated with improved prognosis in patients affected by triple negative breast cancer (TNBC) with residual disease. The aim of this study was to evaluate the antitumoral effect of PKHB1, a thrombospondin-1 peptide mimic, against breast cancer cells, and the immunogenicity of the cell death induced by PKHB1 in vitro, ex vivo, and in vivo. Our results showed that PKHB1 induces mitochondrial alterations, ROS production, intracellular Ca2+ accumulation, as well calcium-dependent cell death in breast cancer cells, including triple negative subtypes. PKHB1 has antitumor effect in vivo leading to a reduction of tumor volume and weight and promotes intratumoral CD8 + T cell infiltration. Furthermore, in vitro, PKHB1 induces calreticulin (CALR), HSP70, and HSP90 exposure and release of ATP and HMGB1. Additionally, the killed cells obtained after treatment with PKHB1 (PKHB1-KC) induced dendritic cell maturation, and T cell antitumor responses, ex vivo. Moreover, PKHB1-KC in vivo were able to induce an antitumor response against breast cancer cells in a prophylactic application, whereas in a therapeutic setting, PKHB1-KC induced tumor regression; both applications induced a long-term antitumor response. Altogether our data shows that PKHB1, a thrombospondin-1 peptide mimic, has in vivo antitumor effect and induce immune system activation through immunogenic cell death induction in breast cancer cells.

Background
Breast cancer is the most frequent type of cancer among women; its innate and acquired treatment resistance to current therapies is the principal problem to treat it, causing the greatest number of cancer-related deaths. 1 While systemic therapies have increased the survival rates of breast cancer patients, the dramatic variations in response rates of patients with distinct clinicopathologic parameters, 2 as well as innate or acquired resistance to current therapies, 3 make relevant the search for new effective treatments for the different molecular subtypes of breast cancer, in particular those associated with poor prognosis.

Recently, several clinical studies have demonstrated the beneficial immunostimulatory effects of inducing immunogenic cell death (ICD), 4,5 recognized as a critical determinant for the efficiency of cancer therapies. Indeed, this peculiar type of cell death is capable of stimulating a long-term antitumor immune response against dead cancer cell antigens. 6,7 Additionally, inducing ICD increases tumor tissue infiltration by T cells, which plays an essential role in mediating a positive response to chemotherapy and is associated with improving clinical outcomes in all subtypes of breast cancer. 5,8

We recently designed PKHB1 through a structure–activity relationship studies around the C-terminal Binding Domain (CBD) of thrombospondin-1 (TSP-1). PKHB1 is a peptide mimic stable in the serum of mice and human and able to induce a cell death involving CD47 activation in different cancer cells, especially in hematological malignancies. 9–12 The ability of PKHB1 to induce cell death was also observed in leukemic cells from patients with aggressive and chemo-resistant phenotypes, without affecting non-tumoral cells from humans or mice. 9,11,13 Additionally, PKHB1 induces ICD in T cell acute lymphoblastic leukemia. 11,14 If this peptide is currently under

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development in pre-clinical studies addressing CLL (x-Pharma), little is known about the cell death capacity, mechanism, immunogenicity, and the antitumor effect of PKHB1 in solid cancers with different molecular characteristics and poor prognosis such as breast cancer (including the triple negative subtype).

Therefore, the aim of this study was to evaluate the antitumor potential of PKHB1 in breast cancer cells (in vitro and in vivo) including triple negative subtypes and to determine whether it induces antitumor immune system activation through ICD induction (ex vivo and in vivo).

Methods

Peptide synthesis

PKHB1 and 4NGG peptides were synthesized manually, using Fmoc-protected amino acids and standard solid phase peptide synthesis (SPPS) (supplemental material and methods), as described previously.6

Cell culture

MCF-7, MDA-MB-231, and 4T1 cell lines were obtained from the ATCC. MCF-7 and MDA-MB-231 cell lines were maintained in DMEM-F12 medium ( Gibco by Life Technologies, Grand Island, NY, USA), while 4T1 cell line was maintained in RPMI-1640 medium, both were supplemented with 10% of fetal bovine serum, 2 mM L-glutamine, 100 U/mL penicillin-streptomycin ( Gibco by Life Technologies, Grand Island, NY, USA), and incubated at 37°C in a controlled humidified atmosphere with 5% CO₂. Cell count was performed following the ATCC’s standard protocols.

Cell death induction and inhibition analysis

5 × 10⁴ cells were plated in 24 wells dishes and left untreated or treated for 2 h with 100 μM, 200 μM, 300 μM, or 400 μM of PKHB1 (KRFYVVMWKK), or 300 μM of 4NGG (KRFYGGMWKK). Annexin-V-allophycocyanin (Ann-V-APC 0.1 μg/ml; BD Pharmingen, San Jose CA, USA), and propidium iodide (PI, 0.5 μg/ml Sigma-Aldrich) were used to assess phosphatidylserine exposure, cell death, and cell viability quantification, respectively, in a BD AccuriC6 flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA) (total population 10,000 cells). Data was analyzed using FlowJo software (LLC, Ashland, OR, USA).

The calcium chelator BAPTA (5 mM), the pan-caspase inhibitor Z-VAD-FMK (Z-VAD, 50 μM), the antioxidant N-Acetyl Cysteine (NAC, 5 mM), the necroptotic inhibitor Necrostatin-1 (Nec-1, 50 μM), the phospholipase C (PLC) inhibitor U73122 (1.25 μM) and the ER receptor inhibitors dantrolene (50 μM) and 2-aminoethoxydiphenyl borate (2-APB, 40 μM) were incubated 30 minutes with the indicated agent, before treatment with PKHB1 (CC₅₀), epirubicin (42.5 μM for MCF-7 and MDA-MB-231 and 5 μM for 4T1 for 24 h), or H₂O₂ (25 μM for all cell lines for 24 h) when indicated.

Intracellular Ca²⁺ levels assay

5 × 10⁴ cells/well in 24 wells dishes (Life Science) were left untreated or pre-incubated with 2.5 mM BAPTA, and then treated for 2 h with PKHB1 (CC₅₀) or left untreated in medium. Then, cells were detached, washed with RINGER buffer without Ca²⁺, and resuspended in 200 μL of the same RINGER buffer with 0.001 μg/mL of Fluo-4 AM (Life Technologies) and 0.001 μg/mL of Pluronic F-127 (Life Technologies), incubated 37°C for 30 min. Next, cells were washed with RINGER buffer w/o Ca²⁺ and assessed by BD Accuri C6 flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA) (total population 10,000 cells), and data was analyzed using FlowJo software (LLC, Ashland, OR, USA).

In vivo model

This study was approved by The Animal Research and Welfare Ethics Committee (CEIBA), of the College of Biological Sciences, number: CEIBA-2018-003. All experiments were conducted according to Mexican regulation NOM-062-ZOO-1999. Female BALB/c mice (6- to 8-week-old; 22 ± 2 g weight) were maintained in controlled environmental conditions (25°C and 12 h light/dark cycle) and supplied with rodent food (LabDiet, St. Louis, MO, USA) and water ad libitum, and they were monitored daily for health status. Mice were randomly assigned to different groups for all studies. All experiments were designed in accordance with the ARRIVE guidelines for animal care and protection (supplemental material).15

Tumor establishment

5 × 10⁵ live 4T1 cells in 100 μL of PBS were injected subcutaneously in the left hind. Tumor volume and mice weight were measured three times per week using a caliper (Digimatic Caliper Mitutoyo Corporation, Japan) and a digital scale (American Weigh Scale-600-BLK, USA), respectively. Tumor volume was determined with the formula: tumor volume (mm³) = (Length × width²)/2. When tumor reached 70–120 mm², 3 days after inoculation with tumor cells, mice were treated daily with 400 μg of PKHB1 in 200 μL of sterile water by intraperitoneal injection, control mice were treated with 200 μL of sterile water. Sixteen days after inoculation with tumor cells, mice were anesthetized with ketamine (i.p. 80 mg/kg body weight) and xylazine (i.p. 10 mg/kg body weight) and were euthanized by cervical dislocation. Tumors from Control or PKHB1-treated mice, were obtained and fixed in 3.7% neutral formalin, embedded in paraffin, sectioned (5 μm thickness) and stained with H&E (MERCK). Histopathological analyses were done by an external veterinarian pathologist (National professional certificate 2,593,012).

T cells evaluation

Sixteen days after tumor inoculation, mice treated (n = 6) or untreated (n = 6) were anesthetized and sacrificed as described above. Blood was obtained by cardiac puncture and isolation of the peripheral blood mononuclear cells (PBMCs) was
performed by density gradient centrifugation using Ficoll-
Hypaque-1119 (Sigma-Aldrich, St Louis, MO, USA). The
spleen, lymph node, and tumor were harvested and filtered
through a cell strainer (70 μM) with PBS (PBMCs were obtained
from the spleen as described above), then 1 × 10^6 cells/mL were
plated and the percent of CD3+, CD4+ and CD8+ T cells was
observed by flow cytometry with the Mouse T lymphocyte subset antibody cocktail CD3 (clone 145–2C11), CD4 (clone RM4-
5), and CD8 (clone 53–6.7) (from BD Bioscience) following the
manufacturer’s instructions.

**Myeloid-derived suppressor cells (MDSCs) and Tregs evaluation**

For MDSCs assessment, PBMCs were obtained from the
blood of mice as described above. Cells were labeled with a
cocktail of CD11b-PE (clone M1/70), Gr-1-APC (clone
RB6/8C5), and Ly-6 G-FITC (clone 1A8) using the Mouse
MDSC Flow kit (from Biolegend) following the manufac-
turer’s instructions.

For Tregs evaluation, PBMCs were obtained from the blood
of mice as described above. Cells were labeled using a True
Nuclear One Step Staining Mouse Treg Flow kit (FOXP3-
AlexaFluor488, CD25-PE, CD4-PerCP; Biolegend) following
the manufacturer’s instructions.

Cells were assessed in a BD Accuri C6 flow cytometer (BD
Biosciences, Franklin Lakes, NJ, USA), and data was analyzed
using FlowJo software (LLC, Ashland, OR, USA).

**Calreticulin, HSP70, and HSP90 exposure**

5 × 10^4 cells/well were plated in 24-well plates and treated
with PKHB1 (CC50) for 2 h or epirubicin (42.5 μM for
MCF-7 and MDA-MB-231 and 5 μM for 4T1 for 24 h).
Then, cells were detached, washed, and incubated for 1 h at
room temperature (RT) with 2 μg/mL of anti-Calreticulin
(FMC-75, Enzo Life Science), 0.8 μg/mL anti-HSP70 (F-3,
Santa Cruz Biotechnology), and 0.8 μg/mL anti-HSP90
(F-8, Santa Cruz Biotechnology) in FACS buffer; cells
were washed and incubated for 30 min in darkness at RT
with goat anti-mouse IgG (Alexa Fluor 488) (H + L, Life
Technologies) (1:1500) in FACS buffer; cells were then
washed and incubated in the dark for 10 min at RT with 7-AAD
(Life Technologies) (1:1000) in FACS buffer. The surface
exposure of CALR, HSP70 and HSP90 was determined
by flow cytometry in non-permeabilized (7-AAD-
negative) cells.

**Immunofluorescence microscopy**

2.5 × 10^5 cells/well in 6-well dishes were left untreated
(Control) or treated with PKHB1 (CC50) and incubated for
2 h. Then, cells were washed with PBS and stained with
Calreticulin-PE antibody (FMC-75, 2 μg/ml) and Hoechst
33,342 (0.5 μg/ml) (Thermo Scientific Pierce, Rockford, IL,
USA), incubated for 1 h in FACS buffer at RT, washed twice,
maintained in PBS, and assessed by confocal microscopy
(Olympus X70; Olympus, Tokyo, Japan).

**ATP and High-mobility group box 1 release assay**

2.5 × 10^5 cells/well in 6-well dishes were left untreated
(Control) or treated with PKHB1 (CC50) for 2 h. Supernatants were recovered, centrifuged at 1600 rpm/10 min-
utes and used to assess extracellular ATP by a luciferase assay
(ENLITEN kit, Promega, Madison, WI, USA), or HMGBl
using the HMGB1 ELISA kit for MDA-MB-231, MCF-7 and
4T1 cells (BioAssay ELISA kit human or mouse, respectively;
US Biological Life Science Salem, MA, USA) following the
manufacturer’s instructions. Bioluminescence was assessed in
a microplate reader (Synergy HT, Software Gen5; BioTek,
Winooski, VT, USA) at 560 nm, and absorbance was assessed
at 450 nm.

**T cell isolation**

Mice were anesthetized and sacrificed as described above, and
blood was obtained by cardiac puncture. PBMCs isolation was
performed as described above. Murine CD3+ cells were iso-
lated from total PBMCs by positive selection using magnetic-
activated cell sorting (MACS) microbead technology with anti-
CD3ε-biotin and anti-biotin microbeads (Miltenyi Biotec;
>98% purity and >98% viability), as stated by manufacturer’s
instructions.

**Differentiation of bone marrow-derived dendritic cells
(BMDCs)**

After sacrifice of anesthetized mice (n = 6), bone marrow was
removed from the femur and tibia by flushing into RPMI-1640.
Eluted cells were cultured for 5 days with 20 ng/mL of IL-4 and
GM-CSF (R&D Systems, Minneapolis, MN, USA) until
approximately 70% of the cells were CD11c+.

**Evaluation of DCs maturation**

CD11c+, MHC-II, CD80, and CD86 were evaluated by flow
cytometry with the fluorescent label-conjugated antibodies,
antiCD11c-Alexa-fluor 488 (N418, R&D Systems), anti-MHC
Class II-PE (REA813, Miltenyi Biotec), anti-CD80-FITC (16–
10A1, R&D Systems), and antiCD86-APC (GL1) from BD
Biosciences (San Jose, CA, USA). In brief, 1 × 10^6 DCs
/mL were stained in 100 μL of FACS buffer with the indicated
antibodies at RT for 30 minutes and then washed twice
with PBS, centrifuged at 1600 rpm/10 min, resuspended in
100 μL of FACS buffer and assessed by Flow Cytometer as
described previously. For MHC-II and CD80 evaluation by
flow cytometry, CD11c was added with MHC-II or CD86, we
then gated CD11c+ cells and we next assessed MHC-II or
CD86 MFI.

**PKHB1-KC and EPI-KC preparation**

4T1 cells (1.5 × 10^6 cells/mL per mice) were plated and, after
adherence, cells were then treated with 400 μM of PKHB1 for
2 h or 10 μM of EPI for 24 h to obtain 80–90% of killed cells
(KC). After treatments, cells were obtained, centrifuged at
1600 rpm/10 min and resuspended in 100 μL of serum-free
medium/mice. Cell death was confirmed using Trypan blue staining and flow cytometry. Finally, the PKHB1-KC or EPI-KC were inoculated by subcutaneous injection in the right flank.

**Freeze and thaw-killed cells preparation**

4T1 cells (3 × 10^6 cells/mL per mice) were first frozen at −80°C for 15 min, then thawed 10 min at 37°C in a water bath. The freeze–thaw (F-T) cycles were repeated three times in rapid succession. After the final thaw, killed cells were resuspended in PBS.

**DCs’ co-culture with PKHB1-KC, EPI-KC, or FT-KC**

DCs were resuspended in fresh medium (1 × 10^6 cells/mL), left untreated (control) or incubated with 3 × 10^6 4T1 killed cells/mL obtained after treatment with PKHB1, EPI, or FT, to give a range of 1:3 (DCs to killed cells); co-culture was left for 24 h. Then the supernatant was obtained, and the well was washed twice with PBS before the next co-culture.

**DCs + T lymphocytes co-culture**

Control DCs or DCs previously co-cultured with PKHB1-KC, EPI-KC, or FT-KC were maintained in fresh medium at 1 × 10^6 cells/mL. Then, allogeneic BALB/c mCD3+ cells were added at 3 × 10^6 cells/mL to give a range of 1:3 (DCs to CD3+ cells), co-culture was left for 96 h. Then, lymphocytes were collected (in the supernatant), washed with PBS, and resuspended in fresh medium at 5 × 10^6 cells/mL for their use in the next co-culture.

**T-Lymphocytes + 4T1 cells co-culture**

1 × 10^5 cells/mL viable 4T1 cells were plated. Then, allogeneic BALB/c mCD3+ cells were added to each well at 5 × 10^5 cells/mL, unprimed (previously co-cultured with control DCs) or primed (previously co-cultured with DCs-PKHB1-KC, EPI-KC, or FT-KC) to give a range of 1:5 (tumor to effector). Co-culture was left for 24 h.

**Cytokine release assay**

Supernatants from the indicated co-cultures were obtained for the assessment of IL-2, IL-4, IL5, and TNFa (BD CBA Mouse Th1/Th2 Cytokine Kit, San Jose, CA, USA) by flow cytometry following manufacturer’s instructions. IFNy was assessed using an ELISA kit (Sigma-Aldrich) and the Synergy HTTM (BioTek Instruments, Inc., Winooski, VT, USA) plate reader at 570 nm wavelength, following manufacturer’s instructions.

**Calcein assay**

4T1 cells (1 × 10^5 cells/mL) were stained with 0.1 mL/mL Calcein-AM from BD Biosciences (San Jose, CA) in FACS buffer at 37°C and 5% CO₂ for 30 min, washed twice with PBS. Thus, primed or unprimed T cells were added in a 1:5 (tumor to effector) ratio. Co-culture was incubated at 37°C and 5% CO₂ for 24 h. Finally, calcein positive or negative 4T1 cells were assessed in a BD AccuryC6 flow cytometer (BD Biosciences) (total population 10,000 cells). Data was then analyzed using FlowJo software.

**Prophylactic vaccination**

Vaccination was carried out as follows: PKHB1-KC (n = 10) or EPI-KC (n = 10) were obtained as previously described, and then inoculated s.c. in 100 µL of serum-free medium into the right hind leg (day −7), 7 days later, viable (5 × 10^5) 4T1 cells were inoculated into the left hind leg (day 0). Tumor volume and weight were measured as described above.

**PKHB1-KC and EPI-KC treatment**

Tumor was established by subcutaneous injection of 5 × 10^5 4T1 cells in 100 µL of PBS, in the left hind. Tumor volume and mice weight were measured as described above. When tumor reached 70–120 mm³ the first treatment of PKHB1-KC (n = 10) or EPI-KC (n = 10) was applied. Killed cells were inoculated subcutaneously in 100 µL of serum-free medium, in the right hind, twice a week for a total of four applications in a 2-week period. Control mice were treated with 100 µL of serum-free medium.

**Long-term antitumor effect evaluation**

Mice in complete remission after prophylactic (n = 9) or therapeutic (n = 9) 4T1-PKHB1-KC application were re-challenged with 5 × 10^5 4T1 viable cells in 100 µL of PBS in the left hind and tumor volume was measured as described above.

**Long-term splenocytes-cytotoxicity**

Mice in complete remission after prophylactic (n = 4) or therapeutic (n = 4) PKHB1-KC application were re-challenged with 5 × 10^5 4T1 viable cells in 100 µL of PBS in the left hind. Three days after tumor inoculation mice were sacrificed, spleens were harvested, filtered through a cell strainer (70 µM) with PBS, and PBMCs were obtained as described above. Splenocytes were recovered and co-cultured with 4T1 cells (previously stained with calcein-AM) at 44:1 ratio (respectively). Finally, calcein positive or negative cells were assessed as described above.

**Statistical Analysis**

Mice were randomly assigned to different groups for all in vivo studies. At least three independent experiments were repeated three independent times. Mann-Whitney tests and two-tailed unpaired Student’s t-tests were performed using GraphPad Prism Software (San Diego CA, USA) and presented as mean values ± SD. The p values were considered significant as follows: p < 0.05.
Results

PKHB1 induces breast cancer cell death

Although the potency of PKHB1 was previously demonstrated in hematopoietic malignancies,9,11 its effectiveness was not yet evaluated for solid tumors in vivo. Thus, after peptide synthesis and characterization (Supplemental material and methods, table sup.1 and figure sup.1), we evaluated here its effect in two types of human breast cancer cell lines, I) MCF-7 (luminal subtype) and II) MDA-MB-231 (triple negative subtype), as well as on the murine 4T1 cell line (mimics triple negative subtype).16 We observed that PKHB1 induces cell death in a concentration-dependent way in MCF-7 (Figure 1(a)), MDA-MB-231 (Figure 1(b)), and 4T1 (Figure 1(c)) cells, as they showed an increase in the percentage of double-positive Annexin V-APC/PI staining (figure sup. 2). We determined that the cytotoxic concentration that induces approximately 50% of cell death (CC50) in MDA-MB-231 and MCF-7 is 200 μM whereas in 4T1 is 300 μM. Next, we evaluated mitochondrial damage and cytosolic Ca2+ augmentation, and observed that PKHB1 (CC50) induced loss of mitochondrial membrane potential (Figure 1(d)), ROS production (Figure 1(e)) and increase of intracellular Ca2+ (Figure 1(f)) in all cell lines. Afterwards, we searched to determine the cell death effectors and evaluated ROS-dependence, also, we used inhibitors of caspases (Z-VAD), necroptosis (Nec-1), and the Ca2+-chelator (BAPTA) and assessed cell death. We used epirubicin (EPI) and H2O2 as controls for inhibition. We found that Z-VAD inhibited the cell death induced by EPI but not PKHB1-induced cell death (Figure 1(g)), while NAC and NEC-1 inhibited the cell death induced by H2O2 but not PKHB1-mediated cell death (Figure 1(h,i)). Finally, we observed that the Ca2+ chelator BAPTA inhibited the Ca2+ augmentation (figure sup. 3) and cell death (Figure 1(j)), induced by PKHB1. This Ca2+ dependence was previously observed for leukemic cells,9,10 suggesting a similar cell death mechanism and common signaling pathways among solid and liquid cancers.

To assess this hypothesis, we used a phospholipase C (PLC) inhibitor (U73122) and ER receptor inhibitors (dantrolene, for ryanodine receptors, and 2-APB, for IP3 receptors). We determined that PKHB1-cell death is significantly inhibited when blocking the ER-Ca2+-channels with U73122, dantrolene and 2-APB (Figure 1(j,k) and Figure 1(l)) in breast cancer cells, confirming a similar cell death pathway induced by PKHB1 in breast cancer cells, as previously found in leukemic cells. PKHB1 has antitumor effects in breast cancer and promotes intratumorally CD8 + T cell infiltration

To evaluate in vivo the potential antitumor effect of PKHB1, 4T1 breast cancer cells were grafted into BALB/c mice. Daily treatments were initiated when tumor volume reached approximately 100 mm3, and 16 days after the cell transplant, tumor volume of the control mice had reached 1500 mm3, requiring the sacrifice of the animals, while the tumor volume of the PKHB1-treated mice reached a maximum volume of 890 mm3 (at day 8) which started to decrease, reaching a volume of 570 mm3 at day 16 (Figure 2(a)) (individual growth curves in figure sup. 4). Also, daily treatment with PKHB1 did not affect mice weight (figure sup. 5). The decrease in tumor volume was correlated with the decrease in tumor weight, going from 1.5 grams in the controls to 0.40 grams in PKHB1-treated mice (Figure 2(b)). The decrease of tumor volume in mice treated with PKHB1, led us to evaluate the involvement of T cells in the observed effect. First, we analyzed histological sections of tumors from control (Figure 2(c)) or PKHB1-treated mice (Figure 2(d)); results revealed that tumors from control mice showed tumor cells (black arrows) with moderate mitotic activity (blue arrows), whereas the PKHB1-treated mice showed sporadic mitotic activity, extensive necrosis with abundant accumulation of cellular debris (green arrows), and abundant inflammatory exudate, composed of polymorphonuclear elements, eosinophils, and lymphoplastacytic cells (red arrows).

Additionally, we evaluated if the cell number and distribution of T lymphocytes in peripheral blood, spleen, lymph nodes, and tumor site, changed after PKHB1 treatment. We observed (Figure 2(e)) that the percentage of CD3+ cells increased in blood, lymph nodes, and tumors of PKHB1-treated mice, while it was maintained in spleen. When we assessed CD4+ cells, the percentage of cells significantly augmented in lymph nodes, whereas it was significantly diminished in the tumor site (Figure 2(f)). Furthermore, CD8 + T cells significantly increased in peripheral blood and specially in tumor site (gating strategy in figure sup. 6), while they significantly diminished in lymph nodes (Figure 2(g)). To extend this analysis, we evaluated the proportion of myeloid-derived suppressor cells (MDSCs) and Tregs in peripheral blood of control and PKHB1-treated mice. We found a significative decrease in the percent of MDSCs (mean from 48% to 23%) and Tregs (mean from 12% to 3%) in mice treated with PKHB1, when compared with control mice (Figure 2(h)). Additionally, we determine a significative increase in the ratio of CD8/Tregs (mean 2% to 14%) in mice treated with PKHB1, when compared with control mice (Figure 2(i)). Finally, we determined whether splenocytes from PKHB1-treated mice could induce an antitumor cell cytotoxicity. For this purpose, we evaluated the calcine negative 4T1 cells after co-culture with splenocytes obtained from control or PKHB1-treated mice. In Figure 2(k), results show that splenocytes from PKHB1-treated mice induced a significant increase in calcine negative 4T1 cells (75%) in comparison with control mice (40%). These results improve the knowledge of the immune system involvement in the antitumor effect mediated by PKHB1 treatment.

PKHB1 induces DAMPs exposure and release in breast cancer cell lines

As we observed that PKHB1 induced cell death in breast cancer cell lines and CD8 + T lymphocyte-recruitment in tumor site and the decrease of immunosuppressive cells, we wondered if cell death induced by PKHB1 was able to induce DAMPs’ exposure/release in breast cancer cells. The first step was to evaluate the exposure of CALR (one of the principal DAMPs related with ICD).17 Our results show that PKHB1-treatment and EPI-treatment were able to induce a significative increase of CALR positive cells in MCF-7 (Figure 3(a)), MDA-MB-231 (Figure 3(b)), and 4T1 (Figure 3(c)) cells. The CALR exposure induced by PKHB1 was confirmed by immunofluorescence microscopy,
where we observed that PKHB1 induced CALR exposure in all the cases Figure 3(d,e) and Figure 3(f). Additionally, PKHB1 treatment induced 24 ± 3, 32 ± 1.3 and 4.23 ± 2-fold of HSP70 exposure (Figure 3(g)), 2.7 ± 0.6, 2 ± 0.6, and 7 ± 1.85-fold of HSP90 exposure (Figure 3(h)) in MCF-7, MDA-MB-231 and 4T1 cells, respectively, when compared with untreated cells.
Finally, we wondered if PKHB1 induced the release of HMGB1 and ATP, two important DAMPs related with ICD. Therefore, the presence of HMGB1 and ATP was assessed in the supernatants of treated and untreated breast cancer cells. In Figure 3, results showed a significant release of HMGB1 (Figure 3(i)) and ATP (Figure 3(j)) in the supernatants of PKHB1-treated cells, when compared with untreated cells.

**PKHB1-KC induce maturation of bone marrow-derived DCs and antitumor T cell responses**

To assess the immunogenicity of the dead cells obtained upon treatment with PKHB1, 4T1 cells were treated with 400 μM of PKHB1, epirubicin (EPI) or freeze and thaw cycles (FT) as positive or negative controls (respectively) of ICD. The PKHB1-killed cells (PKHB1-KC), EPI-KC, or FT-KC were then prepared as described in the methods section, and its ability to induce DCs maturation was evaluated as we show in the schema of Figure 4(a). Thus, bone marrow-derived murine DCs were left untreated (Control) or pulsed for 24 h with the PKHB1-KC, EPI-KC, FT-KC, or LPS (1 μg/mL). After co-culture, DCs pulsed with the different stimulus (killed cells or LPS) maintained the expression of the DCs marker CD11c (Figure 4(b)), while only LPS induced a significative increase of the MHC-II in cell surface (Figure 4(c)). However, the PKHB1-KC, EPI-KC, and LPS induced a significative increase of the co-stimulatory molecule CD86 while no difference was observed in the DCs stimulated with the FT-KC (Figure 4(d)). Additionally, DCs pulsed with PKHB1-KC show a significant increase in CD80 cell surface expression and TNFα release in comparison with unstimulated DCs (figure sup. 7A-C).

Once we determined DCs markers after co-culture with killed cells, we assessed if the DCs pulsed with the different killed cells (PKHB1-KC, EPI-KC or FT-KC) were able to prime T cells. First, primary T lymphocytes (CD3+ cells) were co-
cultured for 96 h with pulsed or unpulsed DCs, and we observed the release of TNFα, IFNγ, and IL-2 in the coculture of CD3+ and DCs-PKHB1-TCL (table sup. 2). Next, primed (co-cultured with pulsed DCs-PKHB1-KC, EPI-KC, or FT-KC) or unprimed (co-cultured with unstimulated DCs) T lymphocytes were collected and co-cultured during 24 h with viable 4T1 cells (previously stained with calcine-AM).

To assess antitumor cell cytotoxicity, we evaluated the increase in calcine negative 4T1 cells after co-culture with primed or unprimed T lymphocytes. Results showed that only T lymphocytes co-cultured with pulsed DCs-PKHB1-KC and DCs-EPI-KC induced a significant increase in calcine negative 4T1 cells (Figure 4(e)), in comparison with the lymphocytes co-cultured with DCs-FT-KC or unpulsed T lymphocytes. Additionally, lymphocytes stimulated with DCs-PKHB1-KC show a significant increase in IFNγ and IL-2 release in the coculture with 4T1 cells in comparison with lymphocytes stimulated with control DCs (figure sup. 7D and E).

**Prophylactic vaccination with PKHB1-KC prevented tumor establishment of 4T1 cells**

Considering that PKHB1 treatment induces tumor decline, infiltration of CD8+ cells into the tumor, DAMPs’ exposure and release, and the antitumor immune response ex vivo, the next step was to carry out the Gold Standard of ICD (prophylactic vaccination)\(^{18,19}\) to confirm whether PKHB1 induced ICD. The vaccine was based in the subcutaneous inoculation of the 4T1-PKHB1-KC, 7 days before the transplantation of viable 4T1 cells, while mice were inoculated with 4T1-EPI-KC used as a positive control, and controls without KC were injected with serum-free medium (Figure 5(a)). Results showed that vaccination with PKHB1-KC prevented tumor establishment in 80% (8/10) of mice compared to 70% (7/10) of mice treated with EPI-KC. No survival (0%) was observed in the Control group inoculated with serum-free medium (Figure 5(b)) (individual growth curves in figure sup. 8 A-C). Additionally, survival rates of mice in each group were consistent with tumor growth, observing, respectively, 80% and 70% of survival in mice vaccinated with PKHB1-KC and EPI-KC by day 60, while control mice perished by day 21 (Figure 5(c)).

**Treatment with PKHB1-KC induces tumor regression**

After ex vivo and in vivo results, we evaluated if the immunogenicity of PKHB1-KC was able to diminish tumor growth and improve overall survival in syngeneic mice bearing 4T1 tumors. First, 4T1 viable cells were inoculated in BALB/c mice. When tumor reached 70–120 mm\(^3\),
a control group was treated with serum-free medium, a second group was treated with PKHB1-KC and the third group was treated with EPI-KC. All mice were treated two times per week for a total of four treatments (Figure 5(d)). Tumor growth measurements show that PKHB1-KC-treated mice had diminished tumor growth after day 10 (7 days after the first treatment), which continued to decrease until no tumor was detected by day 18, in the group of EPI-KC the tumor diminished after day 10 (7 days after the first treatment) which continued to decrease until no tumor was detected by day 16 (Figure 5(e)). Tumor growth diminution was reflected in overall mice survival, as PKHB1-KC-treated mice presented a 78% (7/9) of survival, while the EPI-KC-treated mice presented 67% (6/9) of survival, and all control mice perished by day 23 (Figure 5(f)) (individual growth curves in figure sup. 8D-F).

**PKHB1-KC prophylactic and therapeutic vaccinations induce long-term antitumor effect**

To assess the long-term antitumor response against 4T1 breast cancer cells induced by PKHB1-KC in a prophylactic or therapeutic application, mice in complete remission (tumor free >60 days) were re-challenged with living 4T1 cells. Tumor volume analysis showed that, contrary to naïve mice (Control), which showed a correct 4T1 tumor establishment, mice in remission after PKHB1-KC prophylactic or therapeutic application showed a slight increase in tumor volume at day 3, which immediately disappears by day 6 (Figure 6(a)). These results correlate with mice survival, where we observed that compared to naïve mice, in which a primary 4T1 cells challenge resulted in a 0% (0/9) of survival by day 23, those that were in remission after prophylactic or therapeutic application of PKHB1-KC were completely resistant to a re-challenge with
4T1 cells, resulting in a 100% (9/9) of survival (Figure 6(b)). Furthermore, we determined if splenocytes from re-challenged mice can induce an antitumor cell cytotoxicity. For this purpose, we evaluated the increase in calcein negative 4T1 cells after co-culture with splenocytes obtained from naïve or re-challenged mice. In Figure 6(d), results showed that splenocytes from re-challenged mice induced a significant increase in calcein negative 4T1 cells (60%) in comparison with naïve mice (30%).

**Discussion**

Here, we assessed for the first time the characteristics of the cell death induced by PKHB1 in breast cancer cells, including the triple negative phenotype, which conserves the principal molecular characteristics of cell death (caspase-independent, calcium-dependent, PLC-dependent, and IP3 R and RYR receptor-dependent cell death with the presence of ROS, loss of mitochondrial membrane potential and the intracellular accumulation of Ca2+) reported mainly in
leukemic cells. We recently reported the overexpression of PLCγ1 and its importance in the cell death induced by PKHB1 in CLL cells, but although here we did not assess specially this isoform, it has been demonstrated the overexpression of PLCγ1 in breast cancer patients is correlated with poor clinical outcome, and the overexpression of PLC-β, PLC-ε, and PLC-δ has negative outcomes. Thus, the involvement of PLC in the mechanism of PKHB1-cell death, might have an advantage in the cancer cells that overexpress these proteins.

Our results also revealed the antitumor effect of PKHB1 against 4T1-breast cancer cells in vivo, as PKHB1-treatment diminished tumor volume and weight. Furthermore, we observed the distribution of T cells in PKHB1-treated mice that involves the increase of T cells (in blood, lymph node, and tumor), trafficking of CD4+ cells to lymph nodes, and tumor CD8+ cells infiltration that its associated to an antitumor response. Additionally, PKHB1 induced the decrease of immunosuppressive cells such as MDSCs and Tregs in blood, also, we observed an increase in the
ratio of CD8+/Tregs, which has been related with an enhanced antitumor immune activity.\textsuperscript{23,24} Overall, our results are promising, and indicate that PKHB1 promote a robust antitumor immune response as it has been reported that extensive tumor infiltration by cytotoxic CD8 + T cells, the decrease of MDSCs and Tregs, and the increase in the ratio CD8+/Tregs are strongly associated with patient’s survival and response to therapy, even in different phenotypes of breast cancer.\textsuperscript{23–27} Finally, splenocytes from PKHB1-treated mice were more cytotoxic against breast cancer cells than splenocytes from control mice, probably due to the immunogenicity of the cell death triggered by PKHB1.\textsuperscript{22,28}

The low immunogenicity of tumor cells is a main obstacle of antitumor therapies; therefore, a way to reactivate potent anti-tumor immune responses is through the emission of DAMPs\textsuperscript{39} and dead cells-derived antigens, which can be achieved in the ICD.\textsuperscript{5,50} Here, we demonstrated that PKHB1 is capable of inducing CALR, HSPP70, and HSPP90 exposure, HMGB1 and ATP release, which can promote the uptake of dying cells, and the recruitment, maturation and cross-presentation activity of antigen-presenting cells (APCs).\textsuperscript{28,31} In this sense, we demonstrated that the cell death induced by PKHB1 and epirubicin (our positive control of ICD)\textsuperscript{32–34} are able to promote a mature phenotype of DCs,\textsuperscript{35} both induced a significant increase in the co-stimulatory molecule CD86, such as different ICD inductors.\textsuperscript{36} Additionally, we observed that DCs pulsed with the PKHB1-KC and EPI-KC promote the antitumor specific cytotoxicity of T cells, which confirm the phenotypic and functional maturation of DCs. Additionally, we observed that the freeze and thaw-killed cells do not stimulate the DCs, as other studies have demonstrated that freeze and thaw-killed cells suppress DCs maturation and function.\textsuperscript{37}

However, vaccination assays involving syngeneic models are the gold standard to formally identify ICD inducers, since this demonstrates the tumor rejection capacity of the immunized host.\textsuperscript{29,30} Our results show that the prophylactic application of EPI-KC and PKHB1-KC prevented tumor establishment and increased survival in 70 and 80% (respectively) of the mice, without using adjuvants and with only one vaccination, in comparison with other strategies of prophylactic KC-vaccines.\textsuperscript{39–41} Also, we used Epirubicin, a well-known ICD inductor with major side-effects in human\textsuperscript{32,42} as a positive control, highlighting its ability to prevent the establishment of breast cancer in 70% of the mice. Our results are in line with the protective potential of established ICD inducers including oxaliplatin, doxorubicin, idarubicin, mitoxantrone, and specific forms of radiotherapy (in colon cancer) which presented between 80% and 90% of tumor-free mice\textsuperscript{17,43,44} while the antibody 7A7 (anti-EGFR) induced 50% of survival (lung cancer),\textsuperscript{45} and especially with the fact that an ICD-inductor should display elevated tumor-free survival (>50%).\textsuperscript{19}

The therapeutic application of the dead cells killed by a potential ICD inductor can be used as a confirmatory trial for ICD inductors, to evaluate their ability to mediate therapeutic effects depending on the immune system against established neoplasms.\textsuperscript{38} In this sense, our results show that when we treated tumor bearing mice with only four applications of PKHB1-KC, tumor volume decreased 7 days after the first administration, reaching tumor regression on day 18 in approximately 80% of mice, while in the group of EPI-KC the tumor diminished 7 days after the first treatment which continued to decrease until tumor regression on day 16. Our results highlight the immunogenicity of the PKHB1-induced cell death, because the therapeutic application of the PKHB1-KC induced tumor remission even in the absence of adjuvants. Additionally, we determined the therapeutic potential of the EPI-KC for the first time, as a novel strategy for the application of chemotherapy-ICD inductors. These results differentiate the PKHB1-KC from other therapeutic strategies with tumor lysates against melanoma, prostate and ovarian cancer, which have been poorly evaluated and were mainly used in combination with adjuvants.\textsuperscript{36,47} The success of the therapeutic application of PKHB1-KC in breast cancer was similar to the T-ALL model,\textsuperscript{14} despite their intrinsic molecular differences.\textsuperscript{16,48–50}

The perspectives for cell therapy against cancer are based on the development of T cell responses, resulting in effective rejection of tumors and long-term protection.\textsuperscript{51,52} From this fact, the induction of ICD eventually results in long-lasting protective antitumor immunity.\textsuperscript{53} Our results demonstrate that PKHB1-KC induces long-term antitumor effect since 100% of mice in remission after PKHB1-KC prophylactic or therapeutic application survived at the re-challenge with 4T1 cells. Although immunotherapy with pulsed DCs, primed T lymphocytes or CAR-T cells is the main approach used to stimulate antitumor immune responses, they represent greater technical complexity, higher cost, among other disadvantages regarding the use of crude PKHB1-KC.\textsuperscript{54–56}

Overall, our results demonstrate that PKHB1 is an ICD inductor in breast cancer cells and highlight a new approach for TSP-1 peptides mimic, which could induce ICD as a conserved mechanism of cell death in different types of tumor cells, including solid cancers, additionally, our results provide evidence for a novel strategy in the obtention and application of killed cancer cells against breast cancer.

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Authors’ contributions

KMCR, ACUP and RMR carried out cell death, TMRE, and ROS assessment. KMCR and RMR carried out Ca\textsuperscript{2+} assessment. LGM carried out peptide synthesis. KMCR performed \textit{ex vivo} and \textit{in vivo} experiments. ACMT and PK conceived and supervised the work. KMCR and ACMT prepared the figures and wrote the manuscript. KMCR, LGM, RMR,
ACUP, PK, ACMT, and CRP designed experiments, analyzed and interpreted data, and read and approved the final manuscript.

**Availability of data and material**

The data used to support the findings of this study are available from the corresponding authors upon request.

**Ethics approval**

The Animal Research and Welfare Ethics Committee (CEIBA), of the School of Biological Sciences approved this study; CEIBA-2018-003. All experiments were conducted according to Mexican regulation NOM-062-ZOO-1999.

**Disclosure statement**

The authors declare the following competing financial interest(s): a patent including results from this paper has been filed. The authors declare that no other competing interests exist.

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