2-hexyl-4-pentynoic acid (HPTA), a novel radiosensitizer to breast cancer cells through increasing the instability of DNA repair proteins

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Research

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Abstract

Background

Breast cancer is one of the most common malignant tumors in the world which is the main cause of cancer death for women. Radiotherapy is the main treatment. Although some drugs have been found to enhance the effect of radiotherapy, there are also obvious deficiencies. Therefore, recent applied clinical research has been focusing on locating a suitable radiosensitizer to breast cancer radiotherapy.

Methods

MTT, clonogenic survival assays, comet assays, immunofluorescence and western blot analyses were used to detect the effect of VPA / HPTA on DNA damage induced by radiotherapy for breast cancer through a variety of cell models (MCF7, EUFA423, HCC1937, DMBA-induced rat breast cancer-derived primary culture cell and DMBA-induced transformed human normal breast cell line). At the same time, flow cytometry, immunofluorescence and western blot analyses were used to investigate the effect of VPA / HPTA on DNA damage repair induced by radiation. In vivo experiment, the effect of HPTA as radiosensitizer was investigated by DMBA-induced breast cancer in rats. Finally, the possible mechanism of HPTA acting on target protein was proved by cycloheximide chase experiment.

Results

In this study, a derivative of valproic acid (VPA), 2-hexyl-4-pentynoic acid (HPTA), was demonstrated for the first time that low concentration of HPTA (15 µM) has radiosensitizing properties to breast cancer cells by multiple working models of breast cancer cell lines (in vivo), equivalent to a high concentration of VPA (500 µM). Mechanistic investigations revealed that HPTA induced radiosensitivity through inhibiting the BRCA1-Rad51-mediated homologous recombination pathway. These results were further manifested in breast cancer animal model (in vitro). Most importantly, our study found that HPTA influenced the stability of BRCA1 and Rad51 proteins via shorting their half-life.

Conclusions

Our findings support the proposition HPTA as an alternate, safe and effective radiosensitizer to tumor cells. Targeting BRCA1-Rad51-mediated homologous recombination pathway through HPTA may be a rational strategy to improve the radiotherapeutic efficacy of breast cancer.

Introduction

Breast cancer is one of the most common malignant tumors in the world, with more than one million new cases every year and is the main cause of cancer death for women [1, 2]. Treatment for breast cancer -
surgery, chemotherapy, and radiotherapy - have significantly improved the prognosis of patients and reduced the mortality of breast cancer [3]. Radiotherapy for breast cancer includes adjuvant radiotherapy after breast conserving surgery or after mastectomy, for local recurrence and distant metastasis (e.g. brain and bone). Radiotherapy has been shown to improved survival rate post-mastectomy, reduction in locoregional recurrent rate and ipsilateral breast cancer recurrences [4]. However, radiation to the breast has the potential to cause severe complications due to incidental dosage to heart, lungs and contralateral breast [5], incidence and severity of adverse events is dose and volume dependent [6]. All of these are considered the causes of treatment failure. Thus, acquiring a deeper understanding of the mechanisms related to radiotherapy and identifying a novel and suitable radiosensitizer are crucial events for improving the survival of breast cancer patients.

Valproic acid (VPA) is used in the treatment of epilepsy since the early 1970s [7–10]. In addition to its antiepileptic effect, VPA has been found to induce tumor cell death, inhibit tumor invasion and metastasis, and inhibit angiogenesis in various cancer models. More recent studies have also found VPA to increase the radiosensitivity of solid malignant tumor cells, including esophagus cancer [11], lung cancer [12], prostate cancer [13], and colon cancer [14]. Our previous studies have demonstrated that 500 µM VPA increases the sensitivity of osteosarcoma U2OS cells, breast cancer MCF7 cells and breast cancer tissue-derived primary cells to radiation therapy [15, 16]. The mechanism involves inhibiting BRCA1-Rad51-mediated homologous recombination (HR) and Ku80-mediated non-homologous end joining (NHEJ) repairs, resulting in DSBs accumulation and tumor cell death. A high dose (500 µM) is required for VPA's radiosensitization effects [16], this increases the likelihood of adverse side effects such as hepatotoxicity, nephrotoxicity, and teratogenicity [17–19]. It is therefore imperative to identify alternatives with low toxicity and high potency.

At present, whilst some VPA derivatives have been reported to show similar anti-tumor effects as high-dose VPA, few have radiosensitizing effects. The 2-propyl-pentanoic acid 4-(3H-1,2-dithiole-3-thione-5-yl)-phenyl ester (ACS2) [19], phospho-valproic acid (P-V; MDC-1112) [20], and 2-propyl-pentanoic acid 2-methanesulfonylsulfanyl-ethyl ester (ACS33) [21] have shown some promise, high concentrations of these three derivates were needed to induce tumour cell death [22]. The derivative 2-hexyl-4-pentynoic acid (HPTA) has demonstrated a strong inhibition effect on HDACs activity in HeLa cells and cerebellar granule cells [6, 22]. The IC\textsubscript{50} of its inhibition on HDAC activity is 11–15 µM, lower than VPA (348–448 µM). Whilst HPTA has been reported to be possibly teratogenesis - caused brain malformation in pregnant mouse embryos – a significantly high dose is required (1.25 mmol/kg, equivalent to 227.85 mg/kg [23]; 450 µM). It remains unknown whether a lower dose of HPTA could be as effective as the high dose VPA as a radiosensitizer in breast cancer.

In this study, we discovered for the first time that 15 µM HPTA has a similar radiosensitization effect on breast tumor cells as 500 μM VPA. The mechanism involves disrupting the DNA repair pathway by reducing the half-life of both DNA repair proteins BRCA1 and Rad51. In conclusion, we reveal that HPTA may be an efficacious sensitizer for breast cancer radiotherapy, indicating that BRCA1 and Rad51 may be the promising targets for breast cancer radiotherapy.
Methods And Materials

Cell culture

MCF7 and EUFA423 breast cancer cell lines and U2OS cell lines were purchased from American Type Culture Collection (ATCC) and maintained in DMEM (12100046, Gibco) medium with 10% Fetal Bovine Serum (10270106, Gibco) and 1% Penicillin-Streptomycin (V900929, Sigma). MCF10A cell line was provided by Stem Cell Bank, Chinese Academy of Sciences, cultured with Dulbecco's Modified Eagle's Medium/Nutrient Mixture F-12 Ham (D9785, Sigma) combined with 5% Horse Serum (26050088, Gibco), 100 ng/ml Cholera toxin (C8052, Sigma), 20 ng/ml Epidermal Growth Factor (E5036, Sigma), 0.5µ g/ml Hydrocortisone (614157, Sigma), 10µ g/ml Human Insulin (I9278, Sigma), and 1% Penicillin-Streptomycin (V900929, Sigma). HCC1937 cells were stably transfected with the plasmids of pcDNA3-wild type BRCA1 (wtBRCA1) and vector control (pcDNA3), respectively. Parental HCC1937 cell line was cultured in RPMI 1640 Medium (c11875500bt, Gibco), combined with 10% Fetal Bovine Serum (10270106, Gibco) and 1% Penicillin-Streptomycin (V900929, Sigma). All cells were confirmed to be mycoplasma-free, and maintained at 37°C and 5% CO₂.

Drug treatment and Ionizing Radiation

The cells were treated with VPA (BP452, Sigma) or HPTA (H0964, TCI) and were subjected to IR using X-ray Irradiator (X-RAD225 OptiMAX, Pxi).

MTT assay

Cells were seeded in a 96-well plate at a density of 1 × 10³ cells per well. Following treatments, MTT solution (5 mg/ml, Sigma) was added to the treated cells and incubated for 4 h at 37°C. Then the medium was replaced with dimethyl sulfoxide. The absorbance of the solution was measured using an enzyme immunoassay analyzer at 490 nm.

Clonogenic survival assay

Cells were seeded in P60 dishes followed by the VPA, HPTA and IR treatment and cultured for about two weeks when the visible colonies formed. At the endpoint, cells were fixed with ethanol and stained with 0.1% crystal violet in 20% methanol for 30 min. The detailed method was described in our previous publications [15, 16]. The number of cell colonies was counted and cell survival was presented by the cell survival fraction (SF), with SF= (the number of clones/seeded cells)/plating efficiency (PE). These experiments were performed in triplicate.

Comet assay

The neutral comet assay and alkaline comet assay were performed for the detection of DSBs by using the Comet Assay kit (Trevigen, Gaithersburg, MD) following the manufacturer's instructions and was analyzed using CometScore software (TriTek, Sumerduck, VA).

Immunofluorescence
Cells were seeded in an eight-well chamber at a density of $3 \times 10^4$ cells per well. The methods were described in detail in our previous publications [15, 16]. The primary antibodies used for immunoblotting analysis specific to $\gamma$H$_2$AX (1:500; 05-636, EMD Millipore), 53BP1 (1:1000; NB100-304, Gene Tex), Rad51 (1:100; sc-8349, Santa Cruz), BRCA1 (1:100; sc-6954, Santa Cruz) were used. Secondary antibodies including Alexa Fluor 594 goat anti-mouse IgG(H + L) (1:300; A11032, Molecular probes), Alexa Fluor 488 chicken anti-rabbit IgG(H + L) (1:300; A21441, Molecular probes). Images were captured using a fluorescence microscope (IX71 + DP73, Olympus).

**Immunoblotting**

The protocols were described in our previous publication [24]. The primary antibodies include: $\gamma$H$_2$AX (1:500; 05-636, EMD Millipore), 53BP1 (1:1000; NB100-304, Gene Tex), BRCA1 (1:200; sc-6954, Santa Cruz), BRCA1 (1:250; OP92, Calbiochem), Rad51 (1:200; sc-8349, Santa Cruz), Rad51 (1:2500; PC130, Calbiochem), Ku70 (1:200; sc-5309, Santa Cruz), Ku80 (1:200; sc-5280, Santa Cruz), DNA-PKcs (1:5000; ab32566, abcam), $\beta$-actin (1:2000, TA-09, ZSGB-BIO) and GAPDH (1:2000, TA-08, ZSGB-BIO). The secondary antibodies were goat anti-mouse IgG (H + L) (1:5000, 31430, Thermo Fisher) and goat anti-rabbit IgG (H + L) (1:5000, 31460, Thermo Fisher).

**DSB assay**

For HR assay, the MCF7 cell line expressing a recombination substrate of the pDR–GFP reporter was studied to determine the number of GFP-positive cells in the treated cells for the analysis of the spontaneous HR frequency using flow cytometry as described previously [15, 16]. For NHEJ assay, the U2OS cell line expressing an End Joining reporter (EJ5-GFP) was studied to determine the number of GFP-positive cells in the treated cells for the analysis of the spontaneous NHEJ frequency using flow cytometry as described previously [25].

**Animal keeping and the establishment of breast cancer model**

Female Sprague-Dawley (SD) rats were purchased from Pengyue Laboratory Animal Co. Ltd. Jinan, China. The studies of animal tissue were performed in adherence to national standards on animal care, specifically the Shandong University Human and Animal Ethics Research Committee's requirements (project identification code 81472800, approved on 3 March 2014). All rats were housed in a specific-pathogen-free environment, at a temperature of 23 ± 1 °C. The lights were at a daily rhythm of 12 hrs and the rats were fed fresh food and water ad libitum throughout the experiment. The DMBA (Sigma) was dissolved in purified corn oil and adjusted to the concentration of 20 mg/ml. Intragastric gavage (i.g.) was performed on 50-day-old SD rats at a single dose of 1 ml DMBA-oil solution [26, 27]. At 40–60 days after gavage, primary tumors could be detected through palpation around the breast.

**Tumor observation on rats**

For a period of time after DMBA gavage, rats with appropriate tumor size and location were selected for grouping and then treated. The tumor size of each group was recorded and measured with a vernier.
caliper. Tumor volume was calculated according to the clinical standard formula "Volume V (mm\(^3\)) = \text{Length (L)} \times \text{Width (W)}^2 \times 0.5".

**HE and immunohistochemistry (IHC) staining**

The rats were given chloral hydrate (Sigma) for the anesthesia and breast tissues and cancer tissues were isolated, then fixed overnight in 4% paraformaldehyde solution, embedded in paraffin and serially sectioned 5 µm thick for HE and IHC staining.

The sections for IHC staining were dewaxed with xylene and hydrated with 100–75% ethanol. And then boiled in a 10 mM sodium citrate solution above 92 °C for 20 minutes for antigen retrieval. The sections were cooled to room temperature and then washed with TBS. After washing with TBS, incubate the sections in 3% hydrogen peroxide for 15 minutes to remove endogenous peroxidase and washed again with TBS. Incubate with 10% goat serum for 1 hour to block non-specific binding, then dilute \(\gamma H_2AX\) (1:500,05-636, EMD Millipore) and Rad51 (1:500; PC130, Calbiochem) with solution (TBS + 1% BSA) primary antibody, and the sections were placed at 4 °C overnight. Secondary antibody was diluted by TBST (TBS + 0.25% Triton X-100 + 1% BSA): Biotinylated goat anti-mouse IgG (1:300, BA-9200, Vector), Biotinylated goat anti-rat IgG (1:300, BA-9400, Vector) and incubated sections for 1 hour at room temperature. Wash with TBST and immediately incubated in ABC working solution for 30 minutes, then test with DAB working solution. Subsequently, hematoxylin staining was carried out, and it was dehydrated with ethanol after washing, and then fixed with xylene for 5 minutes, observed with neutral gum.

The IOD density of \(\gamma H_2AX\) and Rad51 staining by IHC staining was further quantified by Image pro plus software (Media Cybernetics).

**Chemical carcinogen-induced rat breast cancer-derived primary cells**

The method was described in our previous publication [28]. After establishment of the breast cancer model, the rats were given chloral hydrate (Sigma) for the anesthesia and breast cancer tissues were isolated in a sterile manner. The breast cancer tissues were further cut into small pieces (2 × 2 mm) and placed on the coated dishes for primary cell culture. The cells from the second and third passage were used for the immunofluorescence analysis of \(\gamma H2AX\) and 53BP1.

**Soft agar colony formation assay**

For the bottom layer, 1.5 ml/well 0.6% low-melting agar (214230, Difco Agar Nobel) was added into a 6-well plate for solidification. The top layer contained cell suspension (1 × 104/well) and 0.3% agar. The cells were cultured for 3–4 weeks until visible colony formed, these were stained with 0.005% crystal violet.

**RI-1 assay**
Cells were seeded onto 60 mm dishes with cell culture media supplemented with 10 µM Rad51 inhibitor 1 (RI-1, S8077, Silleck ) pretreated for 24hrs. Cells were harvested at indicated time points, and lysates were subjected to Western blotting experiments to analyze protein expression at indicated time points. Image J software was used to quantify the relative expression level compared with control treatment.

**Cycloheximide Chase Experiment**

Cells were seeded onto 60 mm dishes with cell culture media supplemented with 40 µg/mL cycloheximide (CHX, HY-12320, MedChemExpress ). Cells were harvested at indicated time points, and lysates were subjected to Western blotting experiments to analyze protein expression at indicated time points. Image J software was used to quantify the relative expression level compared with control treatment.

**Statistical analysis**

All statistical analyses were performed with Student’s t-test on SPSS Statistics for Windows, Version 23.0 (Armonk, NY: IBM Corp; licensed to Shandong University). and represented as mean ± SD. The p values were designated as: *, $P<0.05$; **, $P<0.01$, indicating a statistically significant difference.

**Results**

**HPTA results in the accumulation of more DSBs in response to radiation**

Our previous research showed that 500 µM VPA inhibits the proliferation of multiple breast cancer cells and increases the sensitivity of cells to radiation [16]. To explore whether HPTA has a similar effect on breast cancer cells with VPA, we performed MTT experiments using HPTA in MCF7 cells. Accordingly, we set three concentrations: 15 µM, 30 µM, and 60 µM HPTA against 500 µM VPA (positive control) and subject to 4 Gy of ionizing radiation (IR). We found that cell proliferation in the VPA-/HPTA-treatment groups was inhibited after IR treatment as anticipated ($P<0.01$) (Figure. 1B). There were no statistical differences between the three different concentrations of HPTA as compared to 500 µM VPA, indicating that HPTA at a low concentration (15 µM) possesses similar radiosensitizing effect as 500 µM VPA. Based on the IC$_{50}$ of VPA and HPTA, we opted for 15 µM HPTA for further experiments.

We next performed a clonogenic assay on the MCF7 cells (Figure. 1C). To better compare the effects of HPTA and VPA, we added 15 µM VPA as a negative control. At the same time, to observe cell survival at different IR doses, we also included t 2, 4, and 6 Gy. The results showed that there was no statistical difference between VPA (15 µM or 500 µM) and HPTA treatment (15 µM) in the unirradiated control group (Figure. 1D upper, $P>0.05$). HPTA- treated cells were more sensitive to IR as compared to the untreated control group (Figures. 1C & 1D, $P<0.05$). But VPA was more sensitive only at 500 µM, and there was no difference between the untreated control group and the VPA at a low concentration of 15 µM, which also proved that HPTA was more effective. Similar results were observed in another breast cancer cell line,
EUFA423 (Figure. S1A&1B). These data indicate that the 15 µM HPTA has similar radiosensitizing effects as 500 µM VPA in vitro.

Our previous research [16] has shown that 500 µM VPA could cause significant DSBs following IR treatment, to explore whether the low concentration of HPTA would result in similar DSBs after IR treatment, DSBs were evaluated by two experiments using comet and immunofluorescence assays. To maximize the DSBs in cells, we set the IR dose at 8 Gy. In MCF7 cells, the alkaline comet assay showed no significant change in the VPA- or HPTA- treatment alone compared with the untreated control group pre-IR ($P>0.05$; Figure. 1E). At 0 min post-IR treatment, the relative DSBs in the 500 µM VPA (70.04) and 15 µM HPTA treatment group (69.06) were significantly higher than that of the control group (52.43) ($P<0.01$; Fig. 1E), indicating that VPA or HPTA in combination with IR can further increase intracellular DSBs accumulation. At 30 min post-IR treatment, the comet tail length began to shorten, but in the VPA treatment group (48.52) and the HPTA treatment group (49.70), the comet tail length was still significantly longer than that of the IR treatment alone (36.79) ($P<0.01$; Figure. 1E). At 120 min post-IR treatment, the tail length became shorter and gradually recovered, while the tail length of the VPA treatment group (31.53) and the HPTA treatment group (31.32) was also significantly longer than that of the control alone (20.58) ($P<0.01$). Similar results were obtained from the neutral comet experiments (Figure. 1F). The above observations were further validated in the EUFA423 cells (Figure. S1C&1D).

DNA DSBs biomarkers γH2AX and 53BP1 were employed to determine whether 15 µM HPTA can enhance IR-induced DSBs. Immunofluorescence assay showed that the proportion of MCF7 cells with γH2AX or 53BP1 foci formation in the VPA-/HPTA-treatment alone was not increased compared with the control group ($P>0.05$; Figure. 1G). The proportion of MCF7 cells with γH2AX or 53BP1 foci increased significantly 6 hrs post-IR (81.37% and 87.34% respectively; Figure. 1H). The percentage of γH2AX or 53BP1 foci in the VPA-/HPTA- treatment group was further increased as compared to the control: 98.59%/97.50% (γH2AX) and 98.28%/98.21% (53BP1) ($P<0.01$; Figure. 1H), indicating that 15 µM HPTA can generate a similar amount of DNA DSBs as 500 µM VPA. Next, we observed the clearance of γH2AX and 53BP1 foci. At 24 hr post-IR treatment, the γH2AX and 53BP1 foci disappeared significantly slower in the VPA-/HPTA- treatment group as compared to the control group ($P<0.01$; Figure. 1H), indicating that 15 µM HPTA is equivalent to 500 µM VPA in affecting the ability of cells to repair DNA damage. We analyzed the above data by categorizing the cells containing γH2AX or 53BP1 foci into two groups based on the number of foci in each cell: small foci (n < 20) and big foci (n ≥ 20). The results in Figure. S2A showed that the proportion of cells containing big foci in the VPA-/HPTA- treatment group was significantly higher than that in the control group. We next detected the protein levels of γH2AX and 53BP1 by an immunoblotting assay (Figure. 1I). The levels of γH2AX and 53BP1 were increased by IR treatment, and further increased in the VPA-/HPTA treatment group ($P<0.01$; Figure. 1L). The results from the above experiments indicate that 15 µM HPTA enhances IR-induced DSBs, similar to 500 µM VPA. We further verified the above results in the EUFA423 cells (Figure. S1E-1H).

In summary, in both MCF7 and EUFA423 cells, our data demonstrate that 15 µM HPTA can lead to a similar extent of DSBs as 500 µM VPA in breast cancer cells after IR treatment.
Low dose HPTA dysregulates DNA repair pathway in response to IR-induced DSBs

We next tested the DSB repair process in breast cancer cells to determine the likely mechanism of HPTA's action.

Mammalian cells employ two major DNA repair pathways: homologous recombination (HR) and non-homologous end-joining (NHEJ), to maintain cell survival [29–31]. Error-free HR repair requires a homologous template such as a sister chromatid, whereas NHEJ joins the two ends of a DSB through a process largely independent of homology [32]. Our previous research also showed that 500 µM VPA reduces HR efficiency [16], so we next explore whether 15 µM HPTA has the same effect on the frequency of HR in the above cell models. The MCF7 cells expressing the pDR-GFP recombination reporter for the HR frequency assay were subjected to flow-cytometry after the introduction of I-SceI-induced DSBs (Figure. 2A). The HR frequency decreased in cells treated with 15 µM HPTA by 42.88% as compared to the cells without HPTA treatment (P < 0.01; Figure. 2B), indicating that low concentration of HPTA is sufficient to dysregulate HR repair pathway.

Recombinase Rad51 plays a central role in the HR mechanism and our previous studies have also shown that the effect of VPA on the HR pathway is mediated by Rad51 [16]. Thus, we investigated whether HPTA influences the Rad51-mediated HR pathway. The results from the immunofluorescence assay showed that the percentage of the cells with Rad51 foci formation in the VPA-/HPTA- treatment group was almost the same as the control group pre-IR treatment (P > 0.05; Figure. 2C). At 6 hr post-IR treatment, the percentage of cells with Rad51 foci in the IR control group was 77.91%, and were reduced by 13.31% (P < 0.01) and 14.04% (P < 0.01) in the VPA-/HPTA- treatment groups (Figure. 2C & 2D-upper). At 24 hr post-IR treatment, the Rad51 foci decreased as DNA repairs gradually completed, the VPA-/HPTA- treatment group still showed a significant decrease in Rad51 foci formation, a reduction by 10.39% (P < 0.01) and 9.65% (P < 0.01) respectively as compared with the control group (46.51%) (Figure. 2C & 2D-upper). The results were verified by immunoblotting. At 6 hrs post-IR treatment, Rad51 protein level reduced significantly in both the VPA and HPTA treatment groups as compared to the IR treatment alone (P < 0.01; Figure. 2E & 2F). This indicated that 15 µM HPTA impaired Rad51 activity after IR, and the HPTA inhibited HR pathway is Rad51-dependent.

BRCA1 is another important protein that regulates HR via its interaction with Rad51 [33], so we next determined whether BRCA1 activity was influenced by HPTA. Through immunofluorescence and immunoblotting experiments, we observed similar results as per Rad51(Figure. 2C − 2F). The results were verified in EUFA423 cells (Figure. S3A − 3D).

To next investigate the possibility that HPTA may also influence the NHEJ repair in our cell models, we used U2OS cells expressing the EJ5-GFP reporter to measure the NHEJ frequency [25, 34] using flow-cytometry after the generation of I-SceI-induced DSBs (Figure. S4A). The NHEJ frequency decreased in
cells treated with 15 µM HPTA by 19.48% as compared to the cells without HPTA treatment ($P < 0.05$; Figure. S4B), indicating that HPTA leads to the disruption of the NHEJ pathway.

A number of proteins are involved in the NHEJ repair pathway, such as DNA-PKcs, Ku70 and Ku80 [16]. Since our data demonstrated that HPTA had a suppressive effect on NHEJ, it would be reasonable to detect whether HPTA influences the major NHEJ-associated proteins. We next employed immunoblotting assay to detect DNA-PKcs, Ku70 and Ku80 proteins in our cell models. The immunoblotting assay results in MCF7 cells showed that after 8 Gy IR, no significant changes in the three proteins were observed ($P > 0.05$; Figure.S4C & S4D). Similar results were also noted in the EUFA423 cell line (Figure. S4E). However, the expression of Ku70 and Ku80 proteins was decreased in the VPA treatment group as compared with the IR-control group ($P < 0.05$), but not in the HPTA treatment group (Figure. S4F). These results indicate that HPTA and VPA are different in the role of key proteins in NHEJ pathway. Through flow-cytometry, we found that HPTA could reduce the NHEJ efficiency (19.48%), even if it was not as obvious as the reduction in HR efficiency (42.88%), but the changes of several key proteins in different cells were not consistent, so we next focused on the effect of HPTA on HR.

**HPTA exhibits radiosensitizing properties to IR treatment in DMBA-induced breast cancer in rats in vivo**

To study whether HPTA has a radiosensitization effect *in vivo*, we first investigated the appropriate dose of HPTA on rat breast tumor. The primary model of transformed breast tumor cells in rats was induced by the environmental carcinogen 7,12-dimethylbenz[a]anthracene (DMBA). This was previously described and employed in related studies [28]. The experimental scheme is shown in Fig. 3A. In brief, around 40 days after DMBA gavage to female Sprague Dawley rats, lumps in breast sites could be found. The shape of lumps in the location of mammary glands was irregular (Figure. 3B). By HE staining (Figure. 3C), compared with normal breast tissue, abnormal hyperplasia, fibroadenoma and abnormal proliferation of epithelial cells of the breast were found, indicating that breast cancer in rats was successfully induced.

Reported studies of VPA on glioblastoma utilized intraperitoneal injection of VPA in the range from 150 mg/kg to 600 mg/kg [35]. Therefore, we choose 200 mg/kg as the treatment dose of VPA, which was equivalent to its dose of 500 µM used *in vitro* experiments.

However, the concentration of HPTA in rats has not been reported. In our pilot experiment, four concentrations: 5 mg/kg, 10 mg/kg, 20 mg/kg and 50 mg/kg were used. After intraperitoneal injection, there was no change in the 5 mg/kg dose group, while the 10 mg/kg, 20 mg/kg and 50 mg/kg concentration showed the same performance as that after the 200 mg/kg VPA injection, but only 20 mg/kg dose group was close to 200 mg/kg VPA, and the rest lasted for a shorter or longer time. Therefore, we opted for 20 mg/kg HPTA as the treatment dose for the study, this is equivalent to 15 µM used in the *in vitro* experiments.

To achieve better therapeutic effect, we next experimented with four fractionated doses of 2 Gy IR which more closely mirrored the clinical IR treatment used in breast cancer [36]. VPA or HPTA were administrated before, during, and after IR treatment (Figure. 3D). All the rats were alive over the 32 days
observation period. As shown in Fig. 3E, at the second day post-IR treatment, IR induced a 21% decreased in breast cancer volume, and the addition of VPA or HPTA further reduced the cancer volume by 62% and 54%. During the 32 days of observation, compared with the IR treatment alone, the VPA-/ HPTA- treatment group significantly reduced the tumor volume ($P<0.05$). The tumors in the IR treatment control group had recovered to the volume before IR, while the VPA-/ HPTA- treatment group had recovered to about half of the tumor volume before IR. On the 10th day after treatment, we excised the tumors under general anesthesia (Figure. 3E- right), the tumor size in the VPA-/ HPTA- treatment groups were smaller than that in the IR treatment control group.

The morphological structure of tumors was observed by HE staining (Figure. 3F). VPA or HPTA treatment led to vacuole structures formation in the breast cancer tissue as compared with the untreated group; there were more vacuoles structures and number of necrotic cells after the IR treatment, and large necrotic areas and cells were seen in the tissues in both VPA and HPTA treatment groups. The morphological findings are consistent with the above findings. The results demonstrated that 20 mg/kg HPTA or 200 mg/kg VPA can effectively sensitize breast cancer to IR treatment.

Next, we explored whether DNA damage and repair proteins were influenced by HTPA in vivo using the DSBs marker γH2AX. Whole-cell lysate extracted from the tumor tissue was analyzed for γH2AX using Western blot. The results showed that there were DSBs in both VPA and HPTA treatment groups (Figure. 3G), and the concentration of γH2AX in both groups was higher than that in the untreated control group. The concentration of γH2AX post-IR treatment was significantly increased, significantly higher in VPA treatment group ($P<0.05$) and no difference between the HPTA treatment group and the IR control group. We speculated that the reason why there was no obvious change in the tumor tissue of HPTA combination group might be related to the cell lysate containing the protein components of non-tumor cells (connective tissue cells, such as fibroblast, macrophages, etc.), so we further studied the expression of γH2AX by immunohistochemistry to verify this possibility. We found that compared with other groups, the amount of γH2AX in the tumor areas of the VPA and HPTA treatment groups increased significantly ($P<0.01$; Figure. 3H). The result showed that more DSBs in tumor cells was induced by the combination of HPTA and IR. The level of Rad51 in the VPA-/ HPTA- treatment group was significantly lower than that in IR control group ($P<0.01$; Figure. 3I). We also stained Rad51 with immunohistochemistry, and the results were consistent with those of Western blot (Figure. 4J).

Based on the in vivo and in vitro experiments, we concluded that HPTA, like VPA, can induce DSBs in tumor cells under IR and inhibit Rad51 function in DNA recombination process.

**The role of HPTA on the radiosensitivity of DMBA-induced rat breast cancer-derived primary culture cells**

Based on the model of breast cancer in rats, we designed the experiment of breast cancer-derived primary culture cells according to Fig. 4A. Immunofluorescence assay showed that there were no significant changes in cells with γH2AX or 53BP1 foci formation in the VPA-/HPTA- treatment group compared with the untreated control group (Figure. 4B & 4C). At 6 hr post-IR treatment, the positive rate of cells with
γH2AX or 53BP1 foci formation in the IR control group (90.04%) increased significantly from baseline (Figure. 4C). Importantly, the rate in the VPA-/HPTA- treatment group was further increased to 100% (P < 0.05; Figure. 4C), indicating that 15 µM HPTA induces DSBs in cells similar to 500 µM VPA. Next, we observed the clearance of γH2AX and 53BP1 foci. At 24 hr post-IR treatment, the γH2AX and 53BP1 foci in the IR treatment alone were significantly decreased than in the VPA-/HPTA- treatment group (P < 0.01; Figure. 4C), suggesting that HPTA significantly affect DNA repair activity in cells and delay DNA repair process. Consistent results were obtained from our further analysis (Figure. S2C). Together, the results supported our hypothesis that IR-HPTA combination promotes more DSBs accumulation in primary tumor cells.

**The role of HPTA on the radiosensitivity of DMBA-induced transformed human normal breast cell line**

To triangulate the findings that low concentration of HPTA (15 µM) can promote radiosensitivity, we next test the findings using the DMBA-induced transformed human normal breast cell line, MCF10A. Similar to the rat model mentioned above, the MCF10A cells were treated with 20 µg/ml DMBA for 24 hr then cultured for around 60 days (Figure. 5A-up). The colonies of transformed cells were significantly formed compared to the control group using soft agar-colony formation assay (Figure. 5A-down). The cell survival of the DMBA-induced transformed cells was tested by clonogenic assay (Figure. 5B). The results showed that there was no statistical difference in cell planting efficiency between groups (Figure. 5B). The survival fraction in the VPA-/ HPTA- treatment group was significantly decreased as compared to the IR treatment alone in transformed cells after 4 Gy (P < 0.05) and 6 Gy (P < 0.01) IR treatment (Figure. 5B-right). The survival fraction of untransformed MCF10A human breast cancer cells in the VPA-/HPTA-treatment group did not alter after IR (Figure. S5A & S5B).

Next, we verified HPTA's radiosensitizing effect on DMBA-induced transformed MCF10A cells. DSBs in the tumor cells were observed by immunofluorescence assay using γH2AX and 53BP1 biomarkers. The percentage of cells containing γH2AX or 53BP1 foci significantly increased 6 hr after IR treatment (P < 0.01; Figure. 5C & 5D), the proportion was further increased by VPA-/ HPTA- treatment (P < 0.01). At 24 hr post IR treatment, the γH2AX and 53BP1 foci resolved significantly slower in the VPA-/HPTA- treatment group than in the IR control group (P < 0.01; Figure. 5C & 5D), indicating that 15 µM HPTA affects the ability of transformed cells to repair damaged DNA.

We detected the protein levels of γH2AX and 53BP1 protein by immunoblotting assay. Similarly, the expression of the two proteins in the VPA-/HPTA- treatment group was significantly higher than that of the IR control group (P < 0.05; Figure. 5E & 5F), which was consistent with the immunofluorescence results. In the untransformed MCF10A cells, no statistically significant differences between treatment groups were observed (Figure. S5C - S5F).

To investigate whether HPTA-regulated radiosensitizing effect is associated with DNA repair function in the DMBA-transformed MCF10A cells, the activity of HR associated proteins Rad51 and BRCA1 were
examined. The immunofluorescence assay showed that the number of cells containing Rad51 or BRCA1 foci was significantly decreased in the VPA-/HPTA- treatment group as compared to the IR control group at 6 hr and 24 hr post-IR \((P< 0.01; \text{Figure. 5G & 5H})\). We next detect the protein levels of the Rad51 and BRCA1 in the transformed cells. The protein levels of Rad51 and BRCA1 had decreased in the VPA-/HPTA- treatment group \((P< 0.01; \text{Figure. 5I & 5J})\) as compared to IR control group; though not in untransformed MCF10A cells (Figure. S5G - S5J). These data further indicate that HR function was significantly inhibited in the VPA-/HPTA- treatment group in DMBA-transformed MCF10A cells, consistent with our earlier studies reported above.

**HPTA-mediated radiosensitization to breast cancer cells is dependent on BRCA1 and Rad51 proteins**

We used multiple breast cancer cell lines (MCF7 and EUFA423, primary cultured rat breast cancer cell lines, and DMBA-induced transformed human normal breast cell line) and breast cancer animal model, to investigate whether the radiosensitizability of lower concentration of HPTA (15 µM) is associated with BRCA1-Rad51-mediated HR activity.

The HCC1937 cell line expressing a defective BRCA1 gene was used to establish an isogenic wild type BRCA1 reconstituted cell line (wtBRCA1) (Figure. 6A). The clonogenic assay was employed to analyze the surviving fraction in HCC1937 cells expressing wtBRCA1 in response to IR. The VPA-/HPTA- treatment group did not significantly decrease survival fraction (plating efficiency) in the paired cells as compared with untreated cells pre-IR (Figure. 6B & 6C-up). The survival curve showed that HCC1937 cell expressing defective BRCA1 was more sensitive to IR as compared with BRCA1 cells (Figure. 6B & 6C-down). Furthermore, the survival fraction in VPA-/HPTA-pretreated BRCA1 cells was significantly reduced after 4 Gy and 6 Gy IR, similar to BRCA1 defective cells in response to DNA damage (Figure. 6C-down). The survival fraction in VPA-/HPTA-pretreated BRCA1-deficient cells only moderately decreased compared to wtBRCA1-cells after IR. These results indicated that HPTA treatment led to cell death by inhibiting BRCA1-mediated HR repair pathway in response to IR.

To investigate whether Rad51 is directly involved in the observed HPTA’s radiosensitizing effect, we used RI-1 [37], an inhibitor of Rad51, in the MCF7 cells. After treatment with 10 µM RI-1 for 24 hrs, the expression of Rad51 protein was significantly inhibited (Figure. 6D). The clonogenic assay was employed to analyze the surviving fraction in MCF7 cells in response to IR. We noted that HPTA treatment alone did not decrease survival fraction (plating efficiency) as compared with untreated cells, irrespective of RI-1 treatment pre-IR (Figure. 6E & 6F-up). The survival curve showed that MCF7 untreated with RI-1 was more sensitive to IR treatment as compared with cells treated with RI-1 (Figure. 6E&6F-down). Furthermore, the survival fraction in VPA-/HPTA-pretreated MCF7 cells was significantly reduced after IR, similar to cells treated with RI-1 upon DNA damage, especially under 4 and 6 Gy treatment (Figure. 6F-down). However, in RI-1-treated cells, the survival fraction in VPA-/HPTA-pretreated cells did not decrease compared to MCF7 untreated with RI-1 after IR. The results indicated that HPTA treatment, like and VPA, can lead to cell death via inhibiting Rad51-mediated HR repair pathway in response to IR.
In the established BRCA1 and Rad51 isogenic cell lines, we demonstrated that HPTA- and VPA-induced radiosensitization in breast cancer cells is associated with BRCA1-Rad51 mediated pathway.

**HPTA increases the sensitivity of breast tumor cells to radiotherapy by inhibiting BRCA1 and Rad51 protein synthesis and shortening their half-life**

We next investigate how does HPTA affects sensitize tumor cells to IR. Cycloheximide (CHX) which hinders the translation process by interfering with shifting steps in protein synthesis, is often used to inhibit protein synthesis in eukaryotic cells [38–41]. We designed CHX-chase experiment to measure the half-life of the Rad51 and BRCA1 proteins in MCF7 cells. Immunoblotting was employed to analyze the expression of BRCA1 and Rad51 protein in the treated cells. Prior to IR, the HPTA treatment group and the negative control group showed an inverse association with the duration of CHX treatment and there was no significant difference between them ($P > 0.05$; Figure 7A & 7B). After IR treatment, the protein level of BRCA1 in the control group decreased to 54.45% and 36.61% at 4 and 8 hrs post-CHX treatment respectively, while in the HPTA treatment group was decreased to 55.29% and 31.39% only at 2 and 4 hrs post-CHX treatment respectively (Figure. 7B). The findings suggest that the half-life of BRCA1 protein in the HPTA treatment group was shortened by 2 hrs and that HPTA can inhibit the synthesis of BRCA1 protein. The change of Rad51 protein after CHX treatment was similar to that of BRCA1 (Figure. 7A & 7C). Collectively, HPTA inhibits the synthesis of both BRCA1 and Rad51, shortens their half-life and sensitizes tumor cells to IR.

**Conclusion**

In summary, we discovered for the first time in this study that 15 µM HPTA has a similar radiosensitization effect on breast tumor cells as 500 µM VPA. The mechanism involves disrupting the HR repair pathway by inhibiting both DNA repair proteins BRCA1 and Rad51, as proposed in Fig. 8. Given that HPTA can reduce the half-life of BRCA1 and Rad51 proteins, targeting BRCA1 and Rad51 may be a desirable therapeutic strategy for HPTA to enhance breast cancer radiotherapy.

**Discussion**

The overexpression of HDACs was extensively investigated in many cancers [42, 43]. HDACis inhibit histone acetylation and result in transcriptionally inactive chromatin [26, 44, 45]. Increasing evidence show that many HDACis not only have potential anticancer effects at the cellular and animal level but also are radiosensitizers for some tumors, such as glioblastoma [14] and esophageal squamous cell carcinoma(ESCC) [46]. VPA, a HDACi, increases histone acetylation level, affects chromatin structure and gene regulation by inhibiting HDACs activity [42, 44, 47, 48]. Our previous studies have shown that 500 µM VPA inhibits the proliferation of a variety of breast cancer cells and induce radiosensitivity, suggesting that VPA may be a promising anti-cancer drug and radiosensitizer [15, 16]. However, a large dose (500 µM) of VPA is needed to render such anti-cancer and radiosensitization activities, other studies
reported that high dose VPA can induce adverse events, such as liver, kidney toxicity and teratogenicity [17–19], therefore there is clinical interest to locate a suitable alternative.

Some scholars have studied and compared the effect of VPA and its derivatives on the survival of HepG2 hepatoma cells [49]. It was found that VPA derivatives inhibit the growth of tumor cells at 40 µM. HPTA is a VPA derivative and an effective HDAC inhibitor. Studies have shown that in cerebellar granule cells, HPTA at 5 µM was able to significantly increased acetylated histone H3 (ac-H3) level in a dose-dependent manner [6]. In comparison to VPA which at 100 µM increases the level of ac-H3 by 200%, HPTA at 50–100 µM increases the level of ac-H3 by 600% – 700% [6]. In this study, we demonstrated that HPTA had both anti-cancer and radiosensitizing effects on breast cancer, and that HTPA is more efficacious than VPA since a lower concentration of HPTA is needed to achieve effects as high concentration of VPA. The findings of HPTA's radiosensitizing effect is novel.

A series of our previous studies other group studies have shown that the mechanism of VPA-induced radiosensitization involves the DNA damage repair mechanisms by inducing more DSBs damage to accumulate in the tumor cells, the broken DSBs are unable to be repaired effectively thus lead to the lethality of tumor cells from IR [15, 16]. Our current findings that HPTA acts in a similar mechanism of action to VPA by disrupting BRCA1-Rad51 mediated HR repair pathway (Figure. 8).

For the NHEJ repair, whilst VPA can inhibit Ku70 and Ku80 proteins in EUFA423 cells similar to our previous report [16], HPTA affects the NHEJ independent of the major repair proteins such as Ku70, Ku80 and DNA PKcs in both MCF7 and EUFA423 cells. The data indicated that there is a different mechanism of action between HPTA and VPA in regulating DNA NHEJ repair function. It may be possible that HPTA influences other NHEJ-related proteins and should be subject to further exploration.

We found that HPTA inhibits the synthesis of BRCA1 and Rad51 protein and shortens their half-life indicating that BRCA1 and Rad51 are the key target proteins in mediating HPTA radiosensitization. HPTA can induce the activation of 53BP1, which forms a barrier that inhibits DNA-end resection [50]. Although BRCA1 can promote DNA-end resection by relieving the 53BP1-dependent barrier [51], HPTA inhibits BRCA1 protein synthesis and promotes degradation, which is undoubtedly bad for HR repair. We therefore hypothesize that HPTA may lead to the failure of forming Rad51 nucleoprotein filaments on damaged single-stranded DNA ends in time, since this process requires the presence of a large number of Rad51 proteins [52], and the homology search and strand exchange mediated by Rad51 nucleoprotein filaments are key steps of the homologous recombination process [53]. The inhibition of Rad51 synthesis and the shortening of its half life cycle are fatal to the HR repair of tumor cells, and should be further investigated.

In summary, our work has bridged laboratory research and clinical practice: HPTA is a more efficacious radiosensitizer than VPA, and its clinical application warrants further investigation and promotion.

**Abbreviations**

VPA: valproic acid; HPTA: 2-hexyl-4-pentynoic acid; HR: homologous recombination
Declarations

Availability of data and materials

The analyzed datasets generated during the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate

The studies of animal tissue were performed in accordance with the requirements of the Shandong University Human and Animal Ethics Research Committee (81472800, approved March 2014).

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Authors’ contributions

Zuchao Cai, David Lim and Zhihui Feng designed the study, analyzed the data, and wrote the manuscript. Zuchao Cai and Guochao Liu performed most of the experiments. Wenwen Ding, Zhendong Wang, Zhujun Tian and Junxuan Peng finished the rest part of the experiments in this study, and they analyzed the data and designed the figures. Chao Dong and Fengmei Zhang provided guidance for this work. All authors provided critical feedback on the manuscript. All authors read and approved the final manuscript.
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Figures
**Figure 1**

HPTA results in the accumulation of more DSBs in response to IR. (A) The structural formula of VPA (left) and HPTA (right). (B) The cell growth was tested by the MTT assay in MCF7 cells in VPA and HPTA alone or the combination treatment of VPA and HPTA with 4 Gy of IR. (C) Representative results of clonogenic survival in MCF7 cells treated with VPA and HPTA alone or the combination treatment of VPA and HPTA with different doses of 0, 2, 4 or 6 Gy (upper and down). Cells were treated with VPA or HPTA for 24 h.

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**A**

VPA (2-Heptyl-4-Pentanal) and HPTA (3-Heptyl-4-Pentanal) structural formulas.

**B**

Graph showing relative survival percentages for MCF7 cells treated with VPA or HPTA alone or in combination with 4 Gy of IR. IR-4Gy groups show a decrease in survival compared to NO IR groups.

**C**

Clonogenic survival results for MCF7 cells treated with different doses of IR (0, 2, 4, 6 Gy) in the presence of VPA, HPTA, or their combination.}

**D**

Bar graph showing clonogenic survival as a percentage for different treatment groups.

**E**

Representative images showing cells treated with VPA or HPTA alone or in combination with IR. Cells treated with IR show increased DSBs.

**F**

Similar to **E**, but with an additional time point of 120 min post-IR.

**G**

Representative images showing γH2AX and 53BP1 staining in MCF7 cells treated with VPA, HPTA, or their combination. γH2AX and 53BP1 are markers for DSBs.

**H**

Bar graph showing the percentage of γH2AX- and 53BP1-positive cells in different treatment groups.

**I**

Western blot images showing GAPDH, H2AX, and 53BP1 protein expressions in MCF7 cells treated with VPA, HPTA, or their combination. The blot shows increased expression of H2AX and 53BP1 in IR-treated cells.

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**J**

Bar graph showing the relative density of H2AX and 53BP1 expressions in different treatment groups.
before IR treatment. (D) The planting efficiency (upper) and surviving fraction (down) of the clonogenic survival assay in MCF7 cells. (E) The alkaline comet assay was used for the analysis of DNA DSBs in MCF7 cells (upper) and the relative DSBs were analyzed after normalizing (down). (F) The neutral comet assay was used for the analysis of DNA DSBs in MCF7 cells (upper) and the relative DSBs were analyzed normalizing (down). (G) The dynamic change of IR-induced γH2AX and 53BP1 foci formation in MCF7 cells treated with 500μM VPA and 15μM HPTA was observed after 8Gy treatment. The images represent γH2AX and 53BP1 foci formation in the VPA/HPTA-treated cells at pre-8Gy, 6 h and 24 h post-8Gy, DAPI was used for nuclear staining. (H) The percentage of γH2AX (upper) and 53BP1 (down) foci formation was evaluated. (I) The level of γH2AX and 53BP1 in MCF7 cells was detected by immunoblotting. (J) The density was further analyzed by the Image J in MCF7 cells. Each data point in the graphs was from three independent experiments (mean ± SD). P-values were calculated by Student’s t-test (* P<0.05, ** P<0.01).

Figure 2
Low dose HPTA dysregulates DNA repair pathway in response to IR-induced DSBs (A) The working schematic of the recombination substrate, pDR-GFP, was presented. (B) MCF7 cells with pDR-GFP expression were transfected with plasmid of I-SceI or GFP and then treated with 15μM HPTA for 24 h. GFP was used as a positive control to represent transfection efficiency. The images showed that the results of HR frequency in HPTA treated cells were measured by flow cytometry. (C) The dynamic change of IR-induced Rad51 and BRCA1 foci formation in MCF7 cells treated with 500μM VPA and 15μM HPTA was observed after 8 Gy treatment. The images represent Rad51 and BRCA1 foci formation in the VPA/HPTA-treated cells at pre-8Gy, 6 h and 24 h post-8Gy, DAPI was used for nuclear staining. (D) The percentage of Rad51 (upper) and BRCA1 (down) foci formation was evaluated. (E) The level of Rad51 and BRCA1 in MCF7 cells was detected by immunoblotting. (F) The density was further analyzed by the Image J in MCF7 cells. Each data point in the graph was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).
Figure 3

HPTA exhibits radiosensitizing properties to IR treatment in DMBA-induced breast cancer in rats in vivo
(A) The scheme of experiment in vivo. (B) Normal breast and DMBA-induced breast cancer of rats under gross observation. (C) HE staining for the morphology of normal tissue and DMBA-induced breast cancer. (D) The schedule of VPA or HPTA administration and 2Gy x 4days IR treatment. 200mg/kg VPA or 20mg/kg HPTA injected (i.p.) into rats every 12hrs. VPA or HPTA injection was performed before and after
the daily 2Gy irradiation, totaling 12 times of VPA or HPTA injection. (E) The change of tumor volumes in different groups after treatment, which was normalized by the untreated group. (F) The morphological change of tumor in groups after treatment was outlined in the scheme as was shown by HE staining. (G) The protein level of \( \gamma \)-H2AX was detected by western blot using the whole lysate of treated tumor tissues. Bands were quantified by Image J software. (H) \( \gamma \)-H2AX expression in situ was detected by immunohistochemical staining and the IOD density was analyzed. (I) The protein level of Rad51 was detected by western blot using the whole lysate of treated tumor tissues. Bands were quantified by Image J software. (J) Rad51 expression in situ was detected by immunohistochemical staining and the IOD density was analyzed. Each data point in the graph was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).
Figure 4

The role of HPTA on the radiosensitivity of DMBA-induced rat breast cancer-derived primary cells (A) The culture of DMBA-induced rat breast cancer-derived primary cells. (B) The dynamic change of IR-induced γH2AX and 53BP1 foci formation in primary cells treated with 500μM VPA and 15μM HPTA was observed after 8Gy treatment. The images represent γH2AX and 53BP1 foci formation in the VPA/HPTA-treated cells at pre-8Gy, 6 h and 24 h post-8Gy, DAPI was used for nuclear staining. (C) The percentage of γH2AX
(upper) and 53BP1 (down) foci formation was evaluated. Each data point in the graph was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).

Figure 5

The role of HPTA on the radiosensitivity of DMBA-induced transformed human normal breast cell line (A) Clonogenic survival assay and soft agar assay showed the forming colonies after 4 weeks of culturing to identify cell transforming. (B) Representative results of clonogenic survival in DMBA-induced transformed cell lines.
human normal breast cells treated with VPA and HPTA alone or the combination treatment of VPA and HPTA with different doses of 0, 2, 4, or 6Gy. (C) The dynamic change of IR-induced γH2AX and 53BP1 foci formation in transformed cells treated with 500μM VPA and 15μM HPTA was observed after 8Gy treatment. The images represent γH2AX and 53BP1 foci formation in the VPA/HPTA-treated cells at pre-8Gy, 6 h and 24 h post-8Gy, DAPI was used for nuclear staining. (D) The percentage of γH2AX (upper) and 53BP1 (down) foci formation was evaluated. (E) The level of γH2AX and 53BP1 in transformed cells was detected by immunoblotting. (F) The density was further analyzed by the Image J in transformed cells. (G) The dynamic change of IR-induced Rad51 and BRCA1 foci formation in transformed cells treated with 500μM VPA and 15μM HPTA was observed after 8Gy treatment. The images represent Rad51 and BRCA1 foci formation in the VPA/HPTA-treated cells at pre-8Gy, 6 h and 24 h post-8Gy, DAPI was used for nuclear staining. (H) The percentage of Rad51 (upper) and BRCA1 (down) foci formation was evaluated. (I) The level of Rad51 and BRCA1 in transformed cells was detected by immunoblotting. (J) The density was further analyzed by the Image J in transformed cells. Each data point in the graph was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).
Figure 6

HPTA-mediated radiosensitization of breast cancer cells is dependent on BRCA1 and Rad51 proteins (A) RI-1, an inhibitor of Rad51, was used to down-regulate the expression of Rad51 in MCF7 cells. (B) By colony formation assay, cell survival in the paired cell lines was performed after different treatments. (C) Survival fractions in different groups were showed. (D) The stable expressed cell line of BRCA1-deficient and BRCA1-proficient was successfully established by BRCA1-deficient cell line, HCC1937. (E) By colony
formation assay, cell survival in the paired cell lines was performed after different treatments. (F) Survival fractions in different groups were showed. Each data point in the graphs was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).

**Figure 7.**

(A) After 0, 1, 2, 4, 8 hours treatment with Cycloheximide (CHX), the protein changes of Rad51 and BRCA1 were detected by immunoblotting. (B) Analysis of results in A (after correction). Each data point in the graphs was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).

HPTA increases the sensitivity of breast tumor cells to radiotherapy by inhibiting BRCA1 and Rad51 protein synthesis and shortening their half-life (A) After 0, 1, 2, 4, 8 hours treatment with Cycloheximide (CHX), the protein changes of Rad51 and BRCA1 were detected by immunoblotting. (B) Analysis of results in A (after correction). Each data point in the graphs was from three independent experiments (mean ± SD); P-Values were calculated by t-test (*P<0.05, ** P<0.01).
Figure 8

The molecular model of the combination of HPTA and IR to suppress breast cancer cells through BRCA1-Rad51 mediated HR repair pathway.

Supplementary Files

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- Supplementaryfigures.docx