Research Paper

DNA-PKcs inhibitors sensitize neuroendocrine tumor cells to peptide receptor radionuclide therapy in vitro and in vivo

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Abstract

Background: Peptide receptor radionuclide therapy (PRRT) increases progression-free survival and quality of life of neuroendocrine tumor (NET) patients, however complete cures are rare and dose-limiting toxicity has been reported. PRRT induces DNA damage of which DNA double strand breaks (DSBs) are the most cytotoxic. DNA-dependent protein kinase catalytic subunit (DNA-PKcs) is a key player in DSB repair and its inhibition therefore is a potential way to enhance PRRT efficacy without increasing the dosage.

Methods: We analyzed effects of combining PRRT and DNA-PKcs inhibitor AZD7648 on viability, cell death and clonogenic survival on SSTR2-expressing cell lines BON1-SSTR2, GOT1 and NCI-H69. Therapy-induced DNA damage response was assessed by analyzing DSB foci levels and cell cycle distributions. In vivo efficacy was investigated in BON1-SSTR2 and NCI-H69 xenografted mice and hematologic and renal toxicity were monitored by blood counts, creatinine levels and analyzing renal morphology.

Results: Combining PRRT and AZD7648 significantly decreased viability of BON1-SSTR2, GOT1 and NCI-H69 cells and induced cell death in GOT1 and BON1-SSTR2 cells. A strong effect of AZD7648 on PRRT-induced DSB repair was found. In GOT1 cells, this was accompanied by induction of cell cycle blocks. However, BON1-SSTR2 cells were unable to fully arrest their cell cycle and polyploid cells with high DNA damage levels were detected. In vivo, AZD7648 significantly sensitized BON1-SSTR2 and NCI-H69 xenograft models to PRRT. In addition, combination therapy did not induce significant changes in body weight, blood composition, plasma creatinine levels and renal morphology, indicating the absence of severe acute hematologic and renal toxicity.

Conclusion: These results highlight that the potentiation of the therapeutic effect of PRRT by DNA-PKcs inhibition is a highly effective and well-tolerated therapeutic strategy. Based on our findings, we recommend initiation of phase I/II studies in patients to find a safe and effective combination regimen.

Keywords: Neuroendocrine tumors, peptide receptor radionuclide therapy, DDR inhibitors, DNA-PKcs, ionizing radiation

Introduction

NETs are a relatively rare form of cancer originating from the diffuse neuroendocrine system [1]. PRRT using [177Lu]Lu-[DOTA-Tyr³]octreotate ([177Lu-DOTA-TATE], was approved as a treatment for somatostatin receptor subtype 2 (SSTR2)-positive advanced gastroenteropancreatic NETs (GEP-NETs).
Unfortunately, complete cure with the current regimen of PRRT is rare and dose-limiting kidney and bone marrow toxicity has been reported [2, 3]. These challenges have sparked research interests into PRRT radiobiology to better understand cellular effects and eventually improve therapy [4]. After systemic administration, \(^{177}\)Lu-DOTA-TATE binds to SSTR2 on NET cells, after which radioactive decay of lutetium-177 induces DNA damage leading to cancer cell death [5, 6]. However, cells can repair this damage by employing a tightly orchestrated network of proteins called the DNA damage response (DDR), which counteracts the anti-cancer effects of PRRT and consequently can lead to cancer cell survival [7]. An attractive avenue of improvement is therefore to combine PRRT with targeted inhibitors of DDR proteins, as has been demonstrated using inhibitors of poly (ADP-ribose) polymerase 1/2 (PARP1/2), a protein central in the repair of DNA single-strand breaks [8, 9]. The strategy of combining radiation with other DDR inhibitors has been used extensively for external beam radiation therapy (EBRT), but due to the inherent differences between the physical aspects of EBRT and PRRT (e.g. timing, dose rate, radiation source), these methods cannot readily be transferred to PRRT [10].

One of the most lethal DNA lesions induced by radiation therapies are double-strand breaks (DSBs). Therefore, inhibition of the DSB repair machinery has been evaluated in a vast panel of cancer types as radiosensitization strategy. The major DSB repair pathway is non-homologous end-joining (NHEJ) and a central protein in this pathway is DNA-PKcs [11]. Cells derived from DNA-PKcs-deficient mice display severe radiosensitivity, highlighting the importance of this protein in genomic stability and DNA repair [12-14]. Moreover, upregulation of DNA-PKcs has been shown to correlate with radioresistance in cancer cells [15-17]. Interestingly, functions other than in DSB repair have been found for DNA-PKcs, including transcription regulation, telomere maintenance and cell cycle progression, and those may play additional roles in the response to radiation therapies [18]. For these reasons, different small molecule inhibitors of DNA-PKcs have been developed the past years, with varying selectivity and potency [19]. These have been shown to significantly sensitize different tumor types to EBRT [20-22]. However, data on the effect of these inhibitors combined with PRRT is still lacking.

In this paper we investigated PRRT radiosensitization of NET in \textit{in vitro} and \textit{in vivo} models by DNA-PKcs inhibition and showed that selective DNA-PKcs inhibition is a feasible strategy to potentiate the cancer-killing effect of PRRT.

### Materials & Methods

#### Cell culture and reagents

BON1-SSTR2 cells were generated by transfecting wildtype BON1 cells with the pcDNA3.1-SSTR2 vector as described previously [23], using Lipofectamine Transfection Reagent (Thermo-Fisher Scientific). Selection was done for 1 week using 2 mg/mL G418 (Invivogen), after which cells were cultured in presence of 500 μg/mL G418 every other passage. BON1-SSTR2 cells were cultured in DMEM/F12 medium (Gibco), supplemented with 10% fetal calf serum (FCS; Biowest) and 1% penicillin-streptomycin (PS; Sigma-Aldrich). GOT1 cells were maintained in RPMI 1640 (Gibco), supplemented with 10% FCS, 1% PS, 5 mM L-glutamine (Stemcell Technologies), 5 μg/mL insulin (Sigma) and 5 μg/mL human holo-transferrin (Sigma). NCI-H69 cells (ATCC) were cultured in RPMI 1640, supplemented with 10% FCS and 1% PS. All cells were cultured at 37 °C and 5% CO₂.

#### Radiolabeling

Lu-Mark was purchased from IDB Holland and \(^{177}\)Lu-DOTA-TATE for \textit{in vitro} studies was synthesized in-house according to a standard labeling procedure as used for patient treatment (molar activity 53 MBq/nmol, radiochemical yield > 98% and radiochemical purity > 95%). For \textit{in vivo} studies, \(^{177}\)Lu-DOTA-TATE was synthesized in-house at a molar activity of 86 MBq/nmol, radiochemical yield of > 95% and radiochemical purity of > 90%.

#### Antibodies

For immunofluorescent (IF) and immunohistochemical (IHC) stainings, primary antibodies were used for SSTR2 (Abcam, ab134152, 1:250) and 53BP1 (Novus Biologicals, NB100-904, 1:1000). The used secondary antibodies were goat-anti-rabbit IgG Alexa Fluor 488 (Thermo Fisher, A-11066, 1:1000) for IF and peroxidase-conjugated AffiniPure donkey-anti-rabbit IgG (Jackson ImmunoResearch, 705-035-147, 1:100) for IHC.

For western blotting, primary antibodies were used for DNA-PKcs (homemade [24], 1:1000), and phospho-DNA-PKcs (Ser2056) (CST, E9J4G, 1:1000). The used secondary antibody was peroxidase-conjugated AffiniPure donkey-anti-rabbit IgG (Jackson ImmunoResearch, 705-035-147, 1:1000).

#### Viability assay

Cells were treated with \(^{177}\)Lu-DOTA-TATE or vehicle in suspension (1 MBq/mL (1.9x10⁻⁸ M) in culture medium, 2x10⁵ cells/mL) and plated in triplicate in white-walled, round, flat bottom poly-
styrene 96-well plates (Corning) in the presence of a concentration range of AZD7648 in 200 μL culture medium per well (BON1-SSTR2: 500 cells/well; GOT1: 3x10^4 cells/well; NCI-H69: 1.5x10^4 cells/well). Samples were incubated for 7 days before readout. At the time of readout, 100 μL of CellTiterGlo® 2.0 assay reagent (Promega) was added to each well, after which plates were incubated for 10 min at RT. Luminescence was recorded using a Spectramax iD3x microplate reader (Molecular Devices) without wavelength selection. All raw luminescence values were normalized to the values of cells untreated with AZD7648 to calculate cell viability relative to vehicle- or PRRT-treated controls. Experiments were performed as 3 independent replicates.

**Colony survival assay**

Following ^177^Lu-DOTA-TATE (1 MBq/mL (1.9×10^-8 M) in culture medium, 6x10^5 cells in 60mm-dish) or vehicle treatment for 2 h, BON1-SSTR2 cells were seeded in 6-well plates at a density of 300 cells/well in triplicate in the continuous presence of an AZD7648 concentration range. Plates were incubated for 14 days to allow colony formation. After incubation, the medium was removed, wells were washed in phosphate-buffered saline (PBS; Lonza), and colonies were fixed and stained in 50% (v/v) methanol (Sigma), 7% (v/v) acetic acid (Sigma) and 0.1% (m/v) Brilliant Blue R (Sigma) in dH₂O. Colonies were quantified manually. Experiments were performed as 3 independent replicates.

**Cell death detection**

GOT1 and BON1-SSTR2 cells were treated with ^177^Lu-DOTA-TATE or vehicle in suspension (1 MBq/mL (1.9×10^-8 M) in culture medium, 2x10^5 cells/mL) and seeded with and without 1 μM AZD7648 in 6-well plates. After 5 days, cell death detection was done using the CellEvent™ Caspase-3/7 Green Flow Cytometry Assay Kit (Invitrogen), following the manufacturer’s instructions. In short, cells were incubated with CellEvent Caspase3/7 Green Detection Reagent (500 nM) in culture medium for 55 min at RT in the dark, after which SYTOX™ AADvanced™ Dead Cell Stain (1 μM) was added for an additional 5 min. Fluorescent signal of both dyes was detected by flow cytometry on a LSRFortessa flow cytometer (BD Biosciences). Gating and data analysis were performed using FlowJo™ software (BD Biosciences). Experiments were performed as 3 independent replicates.

**Cell cycle distributions**

GOT1 and BON1-SSTR2 cells were treated with ^177^Lu-DOTA-TATE or vehicle in suspension (1 MBq/mL (1.9×10^-8 M) in culture medium, 2x10^5 cells/mL) and incubated with and without 1 μM AZD7648 for 3 days, after which 5-ethynyl-2'-deoxyuridine (EdU; ThermoFisher Scientific) was added for 3 h at 30 μM until fixation in ice-cold ethanol. Cells were permeabilized in PBS + 0.1% Triton X-100 (Merck) on ice. For EdU detection, samples were incubated with a reaction mix consisting of 40 mM Tris buffer, 4 mM CuSO₄ (Sigma), 30 μM Atto488 azide (ATTO-TEC) and 4 mM ascorbic acid (Sigma) for 30 min at RT in the dark. Samples were washed in PBS + 1% bovine serum albumin (BSA; Sigma) and afterwards resuspended in 1 μg/mL DAPI (Thermo Fisher Scientific) in PBS until analysis. EdU- and DAPI-intensities were recorded on a LSRFortessa flow cytometer. Data analysis was done using FlowJo™ software. Experiments were performed as 3 independent replicates.

**DNA damage analysis and nuclear area quantification**

After ^177^Lu-DOTA-TATE or vehicle treatment in suspension (1 MBq/mL (1.9×10^-8 M) in culture medium, 2x10^5 cells/mL), GOT1 and BON1-SSTR2 cells were incubated with and without 1 μM AZD7648 for 24, 72 and 120 h. For GOT1, cells were plated in 24-well plates, trypsinized and cytospun on glass coverslips using a Rotofix 32A centrifuge (Hettich). For BON1-SSTR2, after PRRT cells were plated on glass coverslips. Cells were fixed in 2% paraformaldehyde (PFA) for 15 min at RT at the indicated timepoints.

To stain cells for 53BP1, samples were permeabilized in PBS + 0.1% Triton X-100. Cells were washed in blocking buffer (PBS + 0.5% BSA) and incubated with primary antibody (diluted in blocking buffer) for 1 h at RT in the dark. Subsequently, samples were incubated with secondary antibody (diluted in blocking buffer) for 1 h at RT in the dark. Coverslips were mounted on microscope slides using Vectashield containing DAPI (Vector Laboratories).

Imaging of 53BP1 foci was done using 405 nm (DAPI) and 488 nm (53BP1) lasers on a TCS SP5 confocal microscope (Leica) with an oil immersion 40x objective. At least four fields of view were captured and 50 cells were analyzed per condition. Images were acquired as Z-stacks. Creation of maximum Z-projections of each image, nuclei segmentation and analysis of the number of 53BP1 foci per nucleus was done in FIJI using a homemade macro. In short, nuclei were segmented in the maximum Z-projections based on thresholding the DAPI channel with a defined minimum and maximum nucleus size. Clusters of cells that could not be segmented correctly were removed from the analysis. Additionally, nuclear size was determined based on this segmentation method.
53BP1 foci were segmented per nucleus by thresholding the Alexa Fluor 488-channel, again with a defined minimum and maximum focus size. Experiments were performed as 3 independent replicates.

**In vivo experiments**

Animal experiments were approved by the Animal Welfare Committee of the Erasmus Medical Center Rotterdam and were conducted in compliance with European guidelines. 6-week-old Rj:NMRI-Foxn1nu/nu mice (Janvier Labs) were subcutaneously injected in the right flank with 5x10⁶ BON1-SSTR2 cells in Matrigel (33% v/v; Corning) in Hank’s Balanced Salt Solution (HBSS; Gibco) or 5x10⁶ NCI-H69 cells in HBSS. Tumor size was determined by caliper measurements. When tumors reached an average size of 200-400 mm³, mice were randomly distributed in treatment groups and intravenously injected in the tail vein with 30 MBq (0.35 nmol) ¹⁷⁷Lu-DOTA-TATE or with vehicle (PBS + 0.1% BSA) (N=9 per group). Starting 2 h before ¹⁷⁷Lu-DOTA-TATE treatment, mice received AZD7648 (50 mg/kg or 100 mg/kg) or vehicle (0.5% hydroxypropyl methylcellulose (HPMC; Colorcon)/ 0.1% Tween-80 (Sigma-Aldrich) by oral gavage daily for 7 consecutive days. Tumor volume and body weight were determined twice per week. When the tumor volume reached 2000 mm³, blood was collected by orbital puncture and mice were euthanized by cervical dislocation.

At the timepoint of euthanization, whole blood samples were collected in EDTA tubes, after which white and red blood cell and platelet counts were performed using an abc Plus counter (scil Vet). Blood plasma was isolated in lithium-heparin tubes and plasma creatinine levels were determined using a Cobas 8000 analyzer (Roche). Tumor and kidneys were collected immediately after euthanization and processed as described below.

**Tumor and kidney SSTR2- and histology assessment**

Tumor and kidneys were collected and fixed in 10% formalin (J.T.Baker) for 24 h at RT and embedded in paraffin. Subsequently, 4 μm sections were generated. For the tumors, hematoxylin & eosin (H&E) staining was performed using standard procedures [6]. For IHC analysis of SSTR2, sections were deparaffinized and rehydrated and antigen retrieval was performed in Tris-EDTA buffer (pH 9.0). Slides were incubated with 3% H₂O₂ in methanol and subsequently blocked in 5% protifar (Nutricia) in washing buffer (PBS + 0.1% Triton X-100). Sections were incubated with primary antibody in blocking buffer at 4 °C O/N, washed and subsequently incubated with secondary antibody in blocking buffer for 30 min. Antibodies were detected using the DAB+ Substrate Chromogen Kit (DAKO). Samples were mounted using Pertex Mounting Medium.

For the kidneys, periodic acid-schiff (PAS) staining was carried out on a Ventana BenchMark Special Stains machine (Roche) as previously described [25]. Assessment of kidney damage was performed in a blinded manner by scoring acute tubular necrosis (ATN) on 10x magnification. Every section was scored for tubular dilatation, cast deposition, brush border loss and necrosis. Scoring was done by a 5-point scale with a score of 0 reflecting virtually no damage, and 5 reflecting severe damage.

Visualization of all stainings described in this section (H&E, IHC, PAS) was done by a NanoZoomer slice scanner (Hamamatsu Photonics).

**Statistical analyses**

Statistical analyses for all experiments were performed in Graphpad Prism (version 8). Used statistical tests include one-way ANOVA with Tukey’s test for multiple comparisons. All statistical analyses yielding a p-value < 0.05 were considered significant.

**Results**

**SSTR2-overexpression in BON1 cells increases ¹⁷⁷Lu-DOTA-TATE binding**

As availability of human cell lines of neuroendocrine origin with sufficiently high SSTR2-expression is limited [26], we generated a SSTR2-overexpressing clone of the neuroendocrine cell line BON1. Compared to the relatively low SSTR2-expression of wildtype BON1 cells, a sharp increase in SSTR2-levels was confirmed by microscopy and flow cytometry in BON1-SSTR2 cells (Figure S1A-B). Functional analysis was done by uptake studies, which confirmed an approximately 100-fold increase in binding of ¹⁷⁷Lu-DOTA-TATE to BON1-SSTR2 cells compared to wildtype (Figure S1C). The majority of the radioluclide, 84%, was internalized after a 2 h-incubation period, as expected for an agonist.

**Inhibition of DNA-PKcs by AZD7648 sensitizes SSTR2-expressing NET in vitro models to PRRT**

For our studies on DNA-PKcs inhibition, we used AZD7648, a recently developed potent and selective inhibitor of DNA-PKcs catalytic activity [22]. Inhibition of DNA-PKcs autophosphorylation at Ser2056 (p-DNA-PKcs), which is induced by ionizing radiation and required for DSB repair [27], was confirmed after AZD7648 treatment in BON1-SSTR2...
cells (Figure S2A). Compared to PRRT monotherapy, AZD7648 treatment did not decrease p-DNA-PKcs levels after combination therapy. An explanation for this is that a high amount of DNA-PKcs is activated upon increased levels of DNA damage during combination therapy, and that this activation is only partially inhibited at 1 μM AZD7648. To further confirm the effects of AZD7648 on DNA-PKcs activity, we measured the balance between classical non-homologous end joining (C-NHEJ) and microhomology-mediated end-joining (MMEJ) in BON1-SSTR2 cells using an end-joining assay (Figure S2B). Inhibition of C-NHEJ is expected to lead to an increase of MMEJ as one of the compensatory repair mechanisms and indeed, we found that 1 μM of AZD7648 led to a severe shift to MMEJ (91%) in BON1-SSTR2, compared to 24% of relative MMEJ activity in untreated cells, confirming the inhibitory effects of AZD7648 on C-NHEJ repair.

To determine the effect of combining PRRT and AZD7648 on cell viability, we treated SSTR2-positive cell lines BON1-SSTR2, GOT1 and NCI-H69 with PRRT or vehicle and a concentration range of AZD7648. For all three cell lines, AZD7648 induced a notable concentration-dependent decrease in cell viability in PRRT-treated cells compared to vehicle-treated cells, indicating that DNA-PKcs inhibition significantly sensitizes these cells to PRRT (Figure 1A). To validate if the enhanced effect of combination therapy on cell viability was due to cell death, apoptosis and necrosis induction was measured at 5 days post-PRRT in GOT1 and BON1-SSTR2 cells, two cell lines from our panel that are derived from neuroendocrine tumors (Figure 1B; Figure S3). PRRT + AZD7648 significantly increased cell death levels compared to control in GOT1 cells (1.6-fold) compared to PRRT alone (1.2-fold). In BON1-SSTR2 cells, a massive induction of cell death was detected after combination therapy (15-fold increase compared to control) as compared to PRRT monotherapy (2.2-fold). No induction of cell death was seen in the AZD7648 monotherapy-treated cells. Finally, to further confirm the observed therapy effects on cell viability, we assessed clonogenic survival after

![Figure 1](https://www.thno.org)

**Figure 1.** In vitro effects of PRRT + AZD7648 on SSTR2-positive cell lines. A, Effect of AZD7648 on cell viability of BON1-SSTR2, GOT1 and NCI-H69 cells administered as monotherapy (blue line) or combined with 1 MBq/mL PRRT for 4 h (red line). Both curves are normalized to their respective viability without AZD7648. Relative viability (mean ± SD) after PRRT monotherapy, as compared to vehicle, is indicated below the corresponding graph for each cell line. B, Cell death (apoptosis/necrosis) induction at 5 days post-PRRT for BON1-SSTR2 and GOT1 cells, expressed as fold change from untreated control. Cells were treated with 1 μM AZD7648. Ns = not significant; *p > 0.05; ****p < 0.0001. C, Effect of AZD7648 on clonogenic survival of BON1-SSTR2 cells administered as monotherapy (blue line) or combined with 1 MBq/mL PRRT (red line). Both curves are normalized to their respective survival without AZD7648. Relative clonogenic survival (mean ± SD) after PRRT monotherapy, as compared to vehicle, is indicated below the corresponding graph. All data points represent the mean of 3 independent replicates. All error bars represent SD.
therapy in BON1-SSTR2 cells (Figure 1C). Due to its low proliferation rate, this assay was not feasible for GOT1 cells. Again, AZD7648 dose-dependently decreased the number of BON1-SSTR2 colonies after PRRT versus vehicle-treated cells.

Combination of PRRT and AZD7648 results in high levels of DNA damage and induces cell type-dependent effects on cell cycle

As DNA-PKcs plays a central role in DSB repair, we measured the number of 53BP1 foci (a marker for DSBs [28]) per nucleus in GOT1 and BON1-SSTR2 cells after treatment. Quantification of the number of foci in GOT1 showed a small but significant increase for PRRT alone, as compared to control (Figure 2A-B), while only at 72 h post-PRRT a significant increase in number of foci could be detected in BON1-SSTR2 (Figure 2C-D). However, when PRRT was combined with AZD7648, a significant increase in foci number was seen for both cell lines at 24, 72 and 120 h after PRRT. For BON1-SSTR2, this was accompanied by the presence of large nuclei containing a massive number of foci (Figure 2C-D). No increase in number of 53BP1 foci was seen for AZD7648 monotherapy.

To investigate the observed effects on the nucleus in more detail, the nuclear area was quantified for both cell lines. For GOT1, no trend in differences in nuclear size could be detected after both monotherapies or combination therapy over time post-PRRT (Figure 3A), while for BON1-SSTR2, it was confirmed that PRRT + AZD7648 induced a sharp and significant increase in nuclear size (Figure 3B), which further increased over time. Subsequently, we assessed therapy-induced cell cycle distribution by DNA content and EdU-incorporation analysis in both cell lines (Figure 3C; Figure S4). In GOT1 cells, PRRT alone slightly decreased the percentage of cells with EdU-incorporation, indicating a decrease of S-phase cells and induction of cell cycle blocks. PRRT + AZD7648 further abolished proliferating cells in S-phase compared to PRRT alone. In contrast, neither AZD7648 nor PRRT monotherapy significantly changed the cell cycle distribution of BON1-SSTR2 cells. However, PRRT + AZD7648 induced cells with large DNA content, while a relatively high number of S-phase cells could still be detected. This indicated that BON1-SSTR2 cells could only partially induce cell cycle blocks and that polyploidy was induced after combination therapy, confirming a difference in cellular response compared to GOT1.

AZD7648 potentiates PRRT anti-tumor effects in SSTR2-expressing xenograft models

To assess the efficacy and safety of combining PRRT and DNA-PKcs inhibition in vivo, BON1-SSTR2-xenografted mice were injected with a subtherapeutic dose of 177Lu-DOTA-TATE (30 MBq/mouse) and subjected to two different doses of AZD7648 (50 or 100 mg/kg/day for 7 consecutive days) (Figure 4A). Administered as monotherapy, both doses of AZD7648 did not have an effect on tumor growth, while PRRT monotherapy had a minor inhibitory effect (Figure 4B; Figure 55A). However, when PRRT was co-administered with AZD7648, a notable dose-dependent delay in tumor growth could be observed. Consequently, median survival in combination therapy groups increased to 41 days (PRRT + 50 mg/kg AZD7648) and 62 days (PRRT + 100 mg/kg AZD7648) after PRRT administration, compared to 34 days for PRRT monotherapy (Figure 4C). Importantly, none of the treatments had a significant effect on body weight, except for an individual mouse with severe weight loss in the 100 mg/kg AZD7648 monotherapy group, of which the cause is unknown (Figure S6).

PRRT combined with 100 mg/kg AZD7648 showed the highest efficacy for the BON1-SSTR2 tumors with no signs of acute toxicity, and we therefore tested this regimen in a second xenograft model with NCI-H69 tumors. No effect on tumor growth was seen for either AZD7648 or PRRT monotherapy, but the combination treatment resulted in strong tumor growth inhibition (Figure 4D, Figure S5B). Here, median survival after combination therapy increased to 32 days post-therapy initiation, compared to 18 days after either vehicle treatment, AZD7648 or PRRT monotherapy (Figure 4E).

After euthanization, BON1-SSTR2 xenografts were collected and analyzed for tissue morphology and SSTR2 expression. No morphological difference was observed between therapy groups and all analyzed tumors stained homogeneously positive for SSTR2 (Figure S7). This indicated that tumor recurrence after the initial regression upon administering PRRT + AZD7648 was not initiated from SSTR2-negative cells present in the tumor, but rather due to a different mechanism of therapy resistance.

Combination treatment of PRRT and AZD7648 does not induce acute hematologic and renal toxicity

As the bone marrow and kidneys are the main organs at risk during PRRT, we investigated whether co-treatment with AZD7648 would increase toxicity at these sites. For the mice bearing BON1-SSTR2 xenografts, hematologic toxicity was assessed at the timepoint of animal sacrifice by measuring white blood cell, red blood cell and platelet counts in whole blood samples (Figure 5A).
**Figure 2. Effect of PRRT + AZD7648 on DSB repair foci.** Cells were treated with 1 MBq/mL PRRT or vehicle and subsequently with 1 μM AZD7648 or vehicle. **A+C,** Representative images of IF staining for 53BP1 (green) as a marker for DSBs in GOT1 cells (A) and BON1-SSTR2 cells (C) after vehicle treatment, AZD7648 or PRRT monotherapy and PRRT + AZD7648. DAPI was used as a nuclear counterstain. Images are shown for 120 h post-PRRT.

**B+D,** Quantification of the number of 53BP1 foci per nucleus for GOT1 (B) and BON1-SSTR2 (D) cells at 24, 72 and 120 h post-PRRT treatment. At least 50 nuclei in 3 independent experiments were analyzed per condition. Graphs show all data points from these 3 experiments merged. Grey horizontal bars represent the mean. Ns = not significant; *p > 0.05; ****p > 0.0001.
Both PRRT and AZD7648 monotherapy and PRRT + AZD7648 did not induce a significant difference in the level of any of these blood components compared to vehicle-treated mice, indicating absence of severe hematologic toxicity at the point of euthanization. For white blood cells, increased intra-group variability was seen after PRRT treatment (monotherapy and combination), but the cause of this is unknown. Subsequently, nephrotoxicity was assessed by scoring kidney histology for acute tubular necrosis (ATN) (Figure 5B-C). ATN scores were low in general (max. 1 on a 5-point scale) and similar between treatment groups. In addition, creatinine levels in blood plasma were determined (Figure 5D). No significant difference was observed for both PRRT monotherapy and combination therapy, compared to vehicle treatment. These data combined indicate absence of severe nephrotoxicity after any of the applied treatments at the timepoint of euthanization.
Figure 4. In vivo efficacy of PRRT + AZD7648 in xenografted mice. A, Experimental setup for in vivo therapy study. After inoculation with BON1-SSTR2 or NCI-H69 cells and tumor growth, mice received a daily dose of AZD7648. At 2 h after the first AZD7648 administration, mice received a single dose of 177Lu-DOTA-TATE. B, Average absolute tumor volume (in mm^3) for each treatment group of BON1-SSTR2 xenografted mice. Error bars represent SD. C, Kaplan-Meier curve visualizing survival of different treatment groups of BON1-SSTR2 xenografted mice. D, Average absolute tumor volume (in mm^3) for each treatment group of NCI-H69 xenografted mice. AZD7648 was only administered at 100 mg/kg as this was the optimal dose in the BON1-SSTR2 xenografts. Error bars represent SD. E, Kaplan-Meier curve visualizing survival of different treatment groups of NCI-H69 xenografted mice.

Discussion

Interfering with the DDR machinery to sensitize cancer cells to ionizing radiation is a well-known strategy for EBRT, but evidence on combination regimens with PRRT is still lagging behind. Due to its central function in DSB repair, DNA-PKcs has been deemed a promising DDR target for this purpose. Importantly, in the past years various selective DNA-PKcs inhibitors have been developed, which have been tested extensively preclinically as monotherapy and in combination with ionizing radiation [19]. Overall, targeting DNA-PKcs has been shown to significantly sensitize a plethora of cancer types in vitro and in vivo to different radiation types, such as photon and proton beams [17, 20-22, 29, 30].
Here, we showed strong and well-tolerated radiosensitization of PRRT in SSTR2-expressing preclinical models by the potent DNA-PKcs inhibitor AZD7648. This inhibitor targets DNA-PKcs kinase activity and exhibits high selectivity versus structurally related kinases such as ATM, ATR and PI3K isoforms, rendering it a suitable compound for potential clinical implementation [22]. Our in vitro treatments in a panel of 3 SSTR2-expressing cell lines showed that AZD7648 dose-dependently and...
significantly reduced cell viability of PRRT-treated cells, as opposed to a relatively small effect of AZD7648 and PRRT monotherapy. This indicates a strong synergistic effect of PRRT and AZD7648, which was evident at submicromolar doses. In GOT1 and BON1-SSTR2 cell lines, two neuroendocrine tumor models with different genetic backgrounds, this was accompanied by a sharp increase in cell death compared to both monotherapies, indicating that at least part of the reduction in viability can be attributed to cancer-eradicating cellular fates such as apoptosis. The strong potentiation of PRRT by AZD7648 was confirmed in our in vivo xenograft experiments, as the selected doses of both AZD7648 and PRRT monotherapy had a minimal effect on tumor growth and survival, while combination therapy yielded significant tumor inhibition and corresponding increase in survival. However, especially in NCI-H69 xenografts, it was surprising that after PRRT monotherapy no difference in tumor control was detected compared to vehicle-treated mice, as previous in vivo experiments employing this model showed significant tumor-inhibitory activity of PRRT [6]. When comparing SSTR2-expression of our NCI-H69 line with a different clonal lineage of the same cell type, our cell line showed, on average, a reduced SSTR2-expression by flow cytometric assessment (data not shown), most likely leading to lower PRRT uptake. Nevertheless, the magnitude of the combination therapy effect as compared to monotherapies indicates that a strong anti-tumor effect can be reached by combining subtherapeutic doses of PRRT and AZD7648. This means that patients with relative low level SSTR2-positive tumors that are not responding to PRRT could potentially benefit from this combination therapy regimen.

In GOT1 and BON1-SSTR2 cells, a significant increase in the number of DSBs was detected after combination therapy compared to PRRT monotherapy. This is in line with the central function of DNA-PKcs in the NHEJ pathway. However, both cell lines responded differently to the combination therapy. PRRT inhibited cell proliferation and induced a cell cycle block in GOT1 cells and this effect was increased upon combination therapy. In contrast, while PRRT had no detectable effect on BON1-SSTR2 nuclear size or cell cycle, these cells were unable to fully arrest cell cycle progression upon combination therapy, accompanied by polyploidy. It is likely that this is caused by therapy-induced endoreplication. It has been reported that p53 status is an important determinant for cellular effects after IR and DNA-PKcs inhibition, where p53 mutant cell lines were unable to induce cell cycle blocks and entered mitosis with high levels of DNA damage, leading to the formation of polyploid cells [31]. Indeed, the wildtype BON1 model has been reported to harbor a mutation in the p53 gene [32]. Consistently, for GOT1, a p53 wildtype model [32], no effect on nuclear size was detected. In addition, it has been shown that p53 status influences cellular fate after ionizing radiation and DNA-PKcs inhibitor treatment, with the induction of premature senescence in p53 wildtype cells and apoptosis in p53 mutant cells [31]. However, in our experiments we detected significant cell death induction both in GOT1 and BON1-SSTR2 after PRRT + AZD7648, meaning that, although different cellular response mechanisms are involved, eventually cell death is induced in both p53 wildtype and mutant cells. The latter might be highly dependent on specific, yet unidentified cellular characteristics.

Importantly, in the case of p53 mutant cells, induction of polyploidy by endoreplication has been debated to be a potential resistance mechanism for tumors after therapy [33]. In addition, it has been shown that polyploid cells rewire their DDR to cope with cellular stresses during endoreplication [34]. In this study, under AZD7648 treatment we detected high levels of unresolved DSBs in polyploid nuclei, indicating that DNA-PKcs inhibition might exceed the threshold of cellular stress in these cells and consequently induces cell death. For example, it is likely that these treated cells are subjected to high levels of replication stress after endoreplication. When analyzing tumor morphology after PRRT + AZD7648 in our in vivo studies, the presence of large polyploid nuclei was not detected. This leaves the question if the observed endoreplication in vitro also occurs in vivo and if yes, these cells could not be detected because they had already been killed after therapy. Nevertheless, BON1 has been shown to harbor multiple other genomic lesions in genes associated with cell cycle progression, such as CDKN2A and CHEK2 [32], which also might play important roles in the formation of polyploid cells after PRRT + AZD7648. In addition, it must be noted that mutations in the p53 gene are rare in NETs [35], which is a limitation of using BON1-SSTR2 for preclinical studies for this specific tumor type. Thus, exact genomic determinants of therapy response and (long-term) consequences for tumor control remain unclear and require additional research.

In addition to the function of DNA-PKcs in DSB repair, other cellular roles of DNA-PKcs might play a role in the described radiosensitizing effects. For example, it has been shown that DNA-PKcs is involved in mitotic progression, which might add to the observed phenotype after PRRT + AZD7648 in BON1-SSTR2 cells [36]. However, another possibility is that these effects are the result of an impaired DNA
damage response upon AZD648 treatment and elevated levels of DNA damage per se and not of DNA repair-independent functions of DNA-PKcs, which warrants further investigation.

One of the major concerns when combining ionizing radiation and DDR inhibitors is overlapping toxicity profiles [37]. Specifically for DNA-PKcs inhibitors, systemic administration might lead to the impairment of radiation-induced DSB repair in healthy tissues throughout the body. In our xenograft studies, we assessed hematologic and renal toxicity as these are the main dose-limiting organs for PRRT and DNA-PKcs inhibition could potentially increase PRRT-induced damage here [2, 3]. These experiments indicated that combination of PRRT and AZD7648 induced no severe acute toxicity at these sites. Previous research on combining radiation with AZD7648 showed increased normal tissue toxicity after EBRT and AZD7648 combination therapy in vivo, along with significant body weight loss [38]. Not only are the non-target organs at risk different between EBRT and PRRT, but also a different toxicity profile can be expected based on differences in radiobiological characteristics. Compared to EBRT, a higher tumor selectivity of PRRT might render a more favorable toxicity profile to combine with systemic DNA-PKcs inhibition. Moreover, the total administered radiation dose between the aforementioned study assessing AZD7648 with EBRT and our study are not matched. Nevertheless, a drawback of our approach might be that we only assessed acute toxicity at a single timepoint upon euthanization. In addition, the use of the immunodeficient mouse strain Rj:NuF1-Foxn1nu limits conclusions that can be drawn on therapy-induced lymphotoxicity. However, the fact that we did not detect any signs of acute hematologic and renal toxicity, along with no significant effects on body weight along the course of the experiment, strongly supports the hypothesis that combining PRRT and AZD7648 in this regimen is well-tolerated in mice. In the future, long-term studies are needed to investigate late toxicity effects, most importantly on kidneys and bone marrow, as well as the effect of different combination therapy scheduling on efficacy and toxicity. For example, the effect of starting DNA-PKcs inhibitor treatment after PRRT administration, when the radionuclide has been cleared from the dose-limiting organs but is still present in the tumor, might be considered.

Previous work by us and others has evaluated other combinatorial strategies of DDR inhibitors and PRRT, such as poly (ADP-ribose) polymerase (PARP)-inhibition [9, 39, 40]. Here, inhibitors of PARP1/2 were used to sensitize several SSTR2-expressing cell lines to PRRT. Inhibition of PARP1/2 attenuates its role in single strand break repair and these breaks can be converted to DSBs during DNA replication [41]. In this way, PARP1/2 inhibition will preferentially target replicating cells, while DNA-PKcs inhibition targets NHEJ as the major DSB repair pathway after IR, which is active throughout the cell cycle. Compared to PARP1/2 inhibition, DNA-PKcs inhibition thus might show better efficacy in slow-growing tumors, which is often the case for NETs, especially those of lower grades.

In conclusion, we have shown strong and tolerable potentiation of PRRT by DNA-PKcs inhibitor AZD7648 in preclinical neuroendocrine tumor in vitro and in vivo models. Further research on dosing and scheduling of PRRT co-administrated with new generation DNA-PKcs inhibitors such as AZD7648, along with additional toxicity studies, should pave the way to clinical trials in humans, as this might be very different from combination therapies with EBRT. Currently, DNA-PKcs inhibitors are evaluated as monotherapy in advanced clinical trials, showing good tolerability but limited efficacy [19]. In addition, a few clinical trials are underway evaluating the effects of DNA-PKcs inhibition and EBRT, and results on efficacy and safety in humans are expected soon (e.g. NCT05116254, NCT04555577, NCT04068194). The development of next generation DNA-PKcs inhibitors could potentially further improve therapy outcomes. This might include inhibitors with a different mechanism of action, such as blocking specific interaction sites of DNA-PKcs and downstream proteins, enabling targeting of specific DNA-PKcs functions important in tumor progression after ionizing radiation. Furthermore, combination of DNA-PKcs inhibitors with α-emitters might yield significantly enhanced responses, considering the increased probability of DSB induction compared to β-emitters such as lutetium-177 [42]. These developments, together with an improved understanding of PRRT radiobiology, hold the promise to stimulate implementation of PRRT-based combination regimens in the clinic and thus further improve responses to radionuclide therapies in neuroendocrine tumors and other cancer types.

**Abbreviations**

ATN: acute tubular necrosis; C-NHEJ: classical non-homologous end joining; DDR: DNA damage response; DNA-PKcs: DNA-dependent protein kinase catalytic subunit; DSB: double-strand break; EBRT: external beam radiotherapy; EdU: 5-ethynyl-2'-deoxyuridine; H&E: hematoxylin & eosin; HBSS: Hank’s Balanced Salt Solution; IF: immunofluorescence; IHC: immunohistochemistry; MMEJ: microhomology-mediated end joining; NET: neuroendocrine
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Supplementary methods and figures.
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Competing Interests
The authors have declared that no competing interest exists.

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