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*J Clin Invest.* 2018;128(5):2116-2126. [https://doi.org/10.1172/JCI97992](https://doi.org/10.1172/JCI97992).

**BACKGROUND.** Poly(ADP-ribose) polymerase (PARP) inhibitors are effective in a broad population of patients with ovarian cancer; however, resistance caused by low enzyme expression of the drug target PARP-1 remains to be clinically evaluated in this context. We hypothesize that PARP-1 expression is variable in ovarian cancer and can be quantified in primary and metastatic disease using a novel PET imaging agent.

**METHODS.** We used a translational approach to describe the significance of PET imaging of PARP-1 in ovarian cancer. First, we produced *PARP1*-KO ovarian cancer cell lines using CRISPR/Cas9 gene editing to test the loss of PARP-1 as a resistance mechanism to all clinically used PARP inhibitors. Next, we performed preclinical microPET imaging studies using ovarian cancer patient–derived xenografts in mouse models. Finally, in a phase I PET imaging clinical trial we explored PET imaging as a regional marker of PARP-1 expression in primary and metastatic disease through correlative tissue histology.

**RESULTS.** We found that deletion of *PARP1* causes resistance to all PARP inhibitors in vitro, and microPET imaging provides proof of concept as an approach to quantify PARP-1 in vivo. Clinically, we observed a spectrum of standard uptake values […]

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RESULTS. We found that deletion of PARP1 causes resistance to all PARP inhibitors in vitro, and microPET imaging provides proof of concept as an approach to quantify PARP-1 in vivo. Clinically, we observed a spectrum of standard uptake values (SUVs) ranging from 2–12 for PARP-1 in tumors. In addition, we found a positive correlation between PET SUVs and fluorescent immunohistochemistry for PARP-1 (r² = 0.60).

CONCLUSION. This work confirms the translational potential of a PARP-1 PET imaging agent and supports future clinical trials to test PARP-1 expression as a method to stratify patients for PARP inhibitor therapy.

TRIAL REGISTRATION. Clinicaltrials.gov NCT02637934.

FUNDING. Research reported in this publication was supported by the Department of Defense OC160269, a Basser Center team science grant, NIH National Cancer Institute R01CA174904, a Department of Energy training grant DE-SC0012476, Abramson Cancer Center Radiation Oncology pilot grants, the Marsha Rivkin Foundation, Kaleidoscope of Hope Foundation, and Paul Calabresi K12 Career Development Award SK12CA076931.

Introduction

Epithelial ovarian cancer (EOC) is the fifth deadliest malignancy in women, with more than 70% of patients presenting with advanced disease (1). Until the recent emergence of targeted therapeutics, treatment has relied heavily on cytoreductive surgery and platinum/taxane-based chemotherapy (2). Poly(ADP-ribose) polymerase inhibitors (PARPis) have emerged as promising drug candidates for the treatment of EOC. In current clinical practice, patients are commonly selected for PARPi therapy by testing positive for genetic mutations in genes encoding DNA repair proteins that result in homologous recombination deficiency (HRD), most notably mutations in breast and ovarian cancer susceptibility genes 1 and 2 (BRCA1 and BRCA2) (3, 4). However, only approximately 50% of patients with HRD respond to PARPi therapy (3). Moreover, patients without known HRD have also shown a clinical benefit from PARPis, as seen in recent trials assessing niraparib, olaparib, or rucaparib, as maintenance therapy in platinum-sensitive recurrent ovarian cancer (5–8). Given that not all patients will respond to PARPi therapy, improved clinical tools for predicting which patients will respond are urgently needed.

Numerous clinical trials have led to FDA approval of 3 PARPis since 2014 and there is continued development of 2 additional drugs within this class (9–13). Despite growth in the
enhanced sensitivity to a DNA alkylating agent in WT cells, suggesting that loss of catalytic function could not be the only reason for PARPi efficacy (17–19). Next, Murai et al. demonstrated that all clinically used PARPis have differential abilities to trap PARP-1 on DNA, but equally lack cytotoxicity in PARP1-KO −/− cells (22, 23). While some PARPis are potent PARP-1 trappers, all PARPis target the PARP-1 enzyme, making it a potential prerequisite biomarker for efficacy (22–24). Furthermore, PARP-1 has been development and application of PARPis, the primary drug target poly(ADP-ribose) polymerase 1 (PARP-1) has never been evaluated in vivo, even though loss of expression in vitro is a well-characterized resistance mechanism (3, 14–19). It was first hypothesized that PARPis work primarily through a synthetic lethality pathway where loss of BRCA1 or BRCA2 combined with chemical inhibition of PARP-1 results in cell death (20, 21). However, it was later shown that deletion of PARP1 did not result in enhanced sensitivity to a DNA alkylating agent in WT cells, suggesting that loss of catalytic function could not be the only reason for PARPi efficacy (17–19). Next, Murai et al. demonstrated that all clinically used PARPis have differential abilities to trap PARP-1 on DNA, but equally lack cytotoxicity in PARP1-KO −/− cells (22, 23). While some PARPis are potent PARP-1 trappers, all PARPis target the PARP-1 enzyme, making it a potential prerequisite biomarker for efficacy (22–24). Furthermore, PARP-1 has been
Effect that stems from their primary pharmacological mechanism of action (16, 35). As such, [18F]FTT and [125I]KX1 quantify PARP-1 expression and have the ability to measure drug-target engagement of clinical PARPi by competing with one another for the NAD+ binding pocket on the catalytic subdomain of PARP-1. Contrary to current methodologies that measure the biochemical product of PARP-1, poly(ADP-ribose) (PAR), this is a direct measurement of drug-target engagement.

In this work, we validate the preclinical rationale for measuring PARP-1 expression as a predictive biomarker of response to PARPi and report the first clinical trial studying PARP-1 expression with [18F]FTT PET in EOC.

**Results**

CRISPR/Cas9 deletion of PARP1 in ovarian cancer cells. Using CRISPR/Cas9 gene editing, we mediated the deletion of PARP1 in 2 ovarian cancer cell lines, one with a BRCA1 mutation and another with BRCA1 promoter methylation (UWB1.289 and OVCAR8). OVCAR8 cells have been previously shown to have reduced BRCA-1 expression attributed to promoter methylation and are sensitive to DNA-damaging agents (36–38). Strikingly, the genetic deletion of PARP1 in EOC cells with a BRCA1 mutation or BRCA1 promoter methylation did not result in synthetic lethality, in that the cells were viable and grew in culture (21, 39). Identifying the mechanism of viability was beyond the scope of this work, but is being pursued. Using this system for PARP1 deletion, we achieved more than 90% reduction of PARP-1 expression through a receptor-ligand effect that stems from their primary pharmacological mechanism of action (16, 35). As such, [18F]FTT and [125I]KX1 quantify PARP-1 expression and have the ability to measure drug-target engagement of clinical PARPi by competing with one another for the NAD+ binding pocket on the catalytic subdomain of PARP-1. Contrary to current methodologies that measure the biochemical product of PARP-1, poly(ADP-ribose) (PAR), this is a direct measurement of drug-target engagement.

In this work, we validate the preclinical rationale for measuring PARP-1 expression as a predictive biomarker of response to PARPi and report the first clinical trial studying PARP-1 expression with [18F]FTT PET in EOC.
damage and sensitivity to PARPi in vitro. First, we examined isogenic UWB1.289 cells by IF after treatment with olaparib for 24 hours. We found no change in DNA damage measured by γH2AX foci formation following olaparib treatment in both BRCA1-restored and PARP1-KO cells compared with DMSO controls. In contrast, γH2AX levels increased 2.6 times in olaparib-treated UWB1.289 cells compared with DMSO controls (Figure 1D). Similarly, the same effect was observed in OVCAR8 cells (Figure 1D). Next, the cytotoxicity evaluated for clinically used PARPi showed that the loss of PARP1 resulted in equal or greater resistance compared with restoring BRCA1 in UWB1.289 cells (Figure 1E, Supplemental Figure 2D, and Supplemental Table 3, A and B). Strikingly, a greater resistance in PARP1-KO cells compared with BRCA1-restored cells from parent controls was observed for the potent PARP trappers, niraparib and talazoparib. The loss of PARP1 resulted in greater than a 1,000-fold decrease in sensitivity to talazoparib, whereas restoring BRCA1 had a lesser effect in UWB1.289 cells. OVCAR8 cells also became resistant to PARPi after loss of PARP1. No differences in sensitivity to cisplatin were observed in cell lines after loss of PARP1, indicating there was no change in overall sensitivity to DNA-damaging drugs. These data provide direct support for PARP-1 expression as a requirement for PARPi sensitivity in vitro.

Preclinical imaging of PARP-1 expression. The aforementioned findings demonstrate the need for dynamic, noninvasive, and quantitative monitoring of PARP-1 expression in patients selected for PARPi therapy. To this end, proof-of-concept studies were conducted in preclinical mouse patient-derived xenograft (PDX) models to confirm the in vitro radioligand binding studies and demonstrate the specificity of [18F]FTT for in vivo quantitative imaging of PARP1. Significant differences were observed in radiotracer uptake in tumors before and after treatment with...
sue specimens were collected for in vitro analysis. Of the 10 patients included in the study, 8 underwent PET/CT imaging (Figure 3). One of the 8 patients had a negative [18F]FTT scan and was excluded from the [18F]FTT PET versus PARP-1 IF correlation.

PARP-1 immunohistochemistry and autoradiography on clinical specimens. Radioligand binding with [125I]KX1 was found to colocalize with PARP-1 colormetric IHC (c-IHC) (Figure 4A) on adjacent tissue sections, supporting the specificity of the iodinated radiotracer for PARP-1. In addition, a spectrum of PARP-1 expression was observed by fluorescent IHC (f-IHC) and autoradiography (Figure 4B and Supplemental Figure 4), providing direct evidence for the wide range of PARP-1 expression in ovarian cancer. Interestingly, PARP-1 colocalized with both tumor cells and lymphocytes in the tumor microenvironment (Supplemental Table 4 and Supplemental Figure 5). PARP-1 expression was also positive in 2 lymph node samples, with residual disease on pathology, and costained with γH2AX. Finally, positive staining for PARP-1 and p53 co-occurred in 77% (10/13) of tissue samples, whereas positive staining for PARP-1 and γH2AX had 100% (13/13) co-occurrence (Supplemental Table 4 and Supplemental Figure 5).

Clinical observations with [18F]FTT PET imaging. PET imaging demonstrated [18F]FTT localized to areas of known EOC based on clinical [18F]FDG PET/CT imaging with low bladder uptake, allowing for clear visualization of lesions within the pelvic region (Figure 5). Despite hepatobiliary excretion, [18F]FTT uptake was discernible in omental disease in the abdomen (Figure 6). Differences in radiotracer uptake were seen between [18F]FTT and [18F]FDG PET, which is commonly used for evaluating tumor response and recurrence in EOC as well as identifying extrapelvic metastases and diagnosis. Such differences suggest these radiotracers provide different and complementary information, concordant with their distinct molecular targets. For example, patient 2, who had completed 4 cycles of carboplatin and paclitaxel 2 weeks prior to imaging, showed multiple omental lesions by [18F]FTT PET imaging with maximum SUVs ranging from 5–8 that were low on [18F]FDG PET (Figure 6A, Figure 6B, and Supplemental Table 5). In follow-up, patient 2 showed disease progression within 4 months of therapy and was platinum resistant. In contrast, patient 11 had a BRCA1 mutation and showed low uptake on both [18F]FTT and [18F]FDG PET after 4 cycles of therapy (carboplatin and paclitaxel) and was platinum sensitive (Figure 6, C and D). These results suggest that [18F]FTT PET provides unique molecular information as a biomarker of viable tumor tissue and could be used in the future to guide clinical management in addition to current biomarkers.

olaparib. By administering olaparib (50 mg/kg) we observed a reduction in [18F]FTT tumor-to-muscle ratios, a semiquantitative measure of tracer binding in vivo (Figure 2A and Supplemental Figure 3A). In addition, tumor tissue analyzed from untreated and olaparib-treated mice by ex vivo autoradiography was used to confirm microPET findings (Figure 2B and Supplemental Figure 3B). Results between microPET imaging and ex vivo autoradiography were similar. Tumor-to-muscle ratios from microPET imaging before and after olaparib treatment were 4.2 ± 0.32 vs. 2.5 ± 0.11 (paired t test, P < 0.0025, n = 4) (Figure 2C). In agreement with microPET results, ex vivo autoradiographs showed a significant difference between untreated and olaparib-treated groups (5.14 ± 0.13 vs. 2.41 ± 0.18, unpaired t test, P < 0.0001, n = 2, 10 sections/tumor) (Figure 2D). Autoradiographs also showed significant differences in tumor and muscle of untreated and olaparib-treated groups (unpaired t test, P < 0.0001) (Supplemental Figure 3C).

Clinical trial enrollment. In this trial, 20 patients were enrolled and 10 patients who underwent surgical debulking or biopsy were included in the study. From the 10 patients, 13 tissue samples represent 275 μM. (A) HE, PARP-1, and [125I]KX1 autoradiograph on adjacent tissue sections showed colocalization between [125I]KX1 and PARP-1 c-IHC. (B) In vitro autoradiography showed a difference in PARP-1 expression that was also confirmed by PARP-1 f-IHC. Max intensity by autoradiograph was 0.28 vs. 0.20 μCi/mg and fluorescent intensity of PARP-1 from whole-section f-IHC was 9.6 vs. 6.7 RFU.
of metabolic activity. Intertumor heterogeneity of [$^{18}$F]FTT was observed in patients (Supplemental Figure 6), further emphasizing a beneficial role of PET imaging compared with standard biopsy approaches for quantifying PARP-1 in local and disseminated disease.

Clinical [$^{18}$F]FTT PET imaging positively correlates with PARP-1 expression in patient tumors. A spectrum of [$^{18}$F]FTT uptake was seen in tumors, indicative of varying levels of PARP-1 expression (Supplemental Table 5). Maximum SUVs of tumors ranged from as low as 2 (background) to as high as 12 and a similar range of PARP-1 expression was also observed by immunofluorescence and autoradiography (Figure 7, A and B, and Supplemental Table 5). Since [$^{18}$F]FTT uptake is a marker of PARP-1 expression, this could translate to some tumors expressing up to 6 times as much PARP-1 as other tumors, with consequently higher drug-target densities for PARPi (Figure 7A). Further characterizing [$^{18}$F]FTT as a measurement of PARP-1 expression, we found positive correlations between PARP-1 f-IHC and [$^{18}$F]FTT PET imaging using linear regression analysis and calculating Pearson coefficients, in addition to PARP-1 f-IHC and [$^{125}$I]KX1 autoradiography ($r^2 = 0.60, 0.79, r = 0.77, 0.89$) (Figure 7B and Supplemental Table 5). As expected, there was no correlation observed among PARP-1 f-IHC, [$^{18}$F]FTT, or [$^{125}$I]KX1 autoradiography with [$^{18}$F]FDG, supporting distinct molecular imaging targets for [$^{18}$F]FTT and [$^{18}$F]FDG.

Discussion
This translational work further supports PARP-1 as the primary target for PARPi and describes the development of [$^{18}$F]FTT as a noninvasive measure of regional tumor PARP-1 expression in patients with ovarian cancer. In vitro radioligand binding of [$^{125}$I]KX1, the radio-iodinated analog of [$^{18}$F]FTT, was previously reported (16) as a method to assay PARP-1 expression and was further confirmed in the present work. The genetic deletion of PARP1 was explored as a mechanism of PARPi resistance in a BRCA1 mutant and a BRCA1 methylated ovarian cancer cell line, and our findings support the notion that PARP-1 expression is necessary for PARPi sensitivity in vitro. Preclinical imaging of PARP-1 in mice with [$^{18}$F]FTT demonstrated the specificity of the radiotracer for its molecular target. Lastly, [$^{18}$F]FTT was studied in patients with EOC involved in a phase 1 trial. [$^{18}$F]FTT demonstrated a wide dynamic range of uptake correlating with in vitro measures of PARP-1 expression and the absence of correlation with [$^{18}$F]FDG. In addition, we confirmed the feasibility of [$^{18}$F]FTT as a practical radiotracer to image lesions in disseminated omental disease despite bowel uptake. Given our small sample size, it is too early to remark on imaging nodal disease involvement with [$^{18}$F]FTT although a recent preclinical study suggests uptake in malignant nodes is greater than normal lymph nodes in a diffuse large B cell lymphoma model (41). It is difficult to extrapolate between diseases, but we did observe a single case that showed nodal uptake on [$^{18}$F]FDG but not [$^{18}$F]FTT and was negative on pathology, which is consistent with inflammation. Together, these data are important first steps in establishing [$^{18}$F]FTT as a novel noninvasive biomarker of PARP-1 expression and lay the foundation for future testing of [$^{18}$F]FTT PET in parallel with PARPi clinical trials.

We explored the loss of drug-target expression as a mechanism of PARPi resistance by studying the effects of the genetic deletion of PARP1 in BRCA1 mutant and BRCA1 methylated cell
lines. While loss of PARP1 in BRCA1 mutant or BRCA1 methylated cell lines has the potential to be synthetically lethal, we found that stable deletion of PARP1 by CRISPR/Cas9 resulted in polyclonal populations of cells in which more than 90% of clones lacked PARP-1 expression and grew in culture. It has been recently shown that OVCAR8 cells are capable of forming RAD51 foci, indicating functional homologous recombination, which could explain why loss of PARP1 was not synthetically lethal in this cell line (42). The observed viability with UWB1.289 cells is most likely due to the biological redundancy between PARP-1 and PARP-2 or PARP-3. Future studies are ongoing to assess PARP1/PARP2-KO and PARP1/PARP3-KO models to test this hypothesis. In addition, we intend to evaluate loss of PARP1 in isogenic BRCA2 mutants with and without reversion mutations, such as the PE01/PE04 cell lines, to test whether similar results are found. We also found that lack of PARP-1 expression prevented PARPi from inducing DNA damage, as measured by γH2AX. These results are consistent with previous reports (17–19, 22, 23). Loss of PARP1 most likely prevents the DNA damage caused by PARPi-induced PARP-1 trapping. This mechanistic understanding is highly relevant to PARPi therapy and our findings are consistent with what has been previously reported, showing PARP-1 as the primary target for PARPi-induced DNA damage (3, 14, 15, 17–19, 23).

In further support, we also observed a greater reduction in PARPi sensitivity with loss of PARP-1 expression compared with BRCA1 restoration in the UWB1.289 isogenic cell lines when treated with the potent PARP trappers, niraparib and talazoparib. This observation may have translational implications, since olaparib, rucaparib, and niraparib were recently approved by the FDA as maintenance therapy in platinum-sensitive patients, regard-

Figure 6. Patients 2 and 11 underwent [18F]FTT PET/CT imaging within 2 weeks of completing 4 cycles of carboplatin and paclitaxel. Omental metastases in patient 2 showed higher uptake of [18F]FTT than [18F]FDG with maximum SUVs of (A) 7.8 vs. 3.4 and (B) 5.1 vs. 2.0. Patient 2 was platinum resistant and relapsed within 4 months of therapy. Patient 11 showed low uptake on both [18F]FTT and [18F]FDG with maximum SUVs of (C) 2.4 vs. 3.7 and (D) 2.3 vs. 2.9. Patient 11 received 2 additional cycles of chemotherapy and was platinum sensitive. Yellow arrows indicate sites of disease.
PARP-1 expression may also play a role in predicting inherited or incurred DNA damage burden to identify DNA repair deficiencies or even treatment response as a functional biomarker of HRD (44, 45). We are currently entering several clinical trials in both breast and ovarian cancer at the University of Pennsylvania to directly evaluate [18F]FTT as a predictive and pharmacodynamic biomarker of PARPi therapy and other DNA-damaging chemotherapy. These studies should provide the impetus for multicenter clinical trials to fully evaluate the importance of PARP-1 expression as a biomarker for cancer therapy and the technological advantages of [18F]FTT for measuring intratumoral and intertumoral PARP-1 expression versus conventional IHC approaches.

**Methods**

**Study design.** The objective of this study was to highlight the significance of PARP-1 expression for PARP inhibitor therapy using preclinical models of ovarian cancer and perform the first PET imaging study of PARP-1 in ovarian cancer using [18F]FTT with histological correlates to confirm radiotracer uptake as a biomarker of PARP-1 expression. This goal was addressed by using CRISPR/Cas9 to delete PARP1 in models of ovarian cancer, which were then used to test PARPi efficacy in vitro. In addition, we used patient-derived orthotopic xenograft models of ovarian cancer to show proof of concept for [18F]FTT microPET imaging of PARP-1 and PARPi drug-target engagement in
In vitro. [18F]FTT PET imaging of PARP-1 was performed in patients with ovarian cancer, and tumor specimens were retrieved at surgical debulking or biopsy for in vitro correlative analysis of PARP-1 using [125I]KX1 autoradiography and PARP-1 f-IHC.

All in vitro experiments were carried out using a standard cell culture technique and repeated 3 independent times. The deletion of PARP1 with CRISPR/Cas9 was performed in 2 ovarian cancer cell lines. In addition, 3 unique guide RNAs were used to mediate the deletion of PARP1. Polyclonal populations of PARP1-KO cell lines were used to test PARPi efficacy in vitro. A sample size of 4 was selected for preclinical imaging studies to prevent bias greater than 10% for interpreting and analyzing microPET image results. No data from preclinical models was excluded from analyses.

For the clinical portion of this study, 20 patients with EOC were enrolled, 10 of whom had tissue available for in vitro analysis. Eight of those 10 underwent [18F]FTT PET imaging prior to subsequent standard clinical management (Figure 3). One of the 8 patients who underwent [18F]FTT had a negative PET image and was excluded from correlational analysis. A total of 13 tissue specimens was available from the 10 patients and these specimens were used for in vitro assays including PARP-1 f-IHC and [125I]KX1 autoradiography (Supplemental Table S5). All tissue specimens from patients who underwent [18F]FTT PET imaging were included in correlations among [18F]FTT PET, [18F]FDG, [125I]KX1 autoradiography, and PARP-1 f-IHC. All tissue specimens collected from consenting patients were included in correlations between [125I]KX1 and PARP-1 f-IHC. Patients with and without BRCA1 mutations were enrolled in this study (Supplemental Table S6). All experimental procedures for preclinical and clinical work are described in detail in the supplemental information.

CRISPR/Cas9 deletion of PARP1 in ovarian cancer cells. OVCAR8 cells were a gift from David M. Livingston (Dana Farber Cancer Institute, Boston, Massachusetts, USA). All other cell lines were available through ATCC or the Basser Center for BRCA (University of Pennsylvania, Philadelphia, Pennsylvania, USA) (Supplemental Table 1). PARP1 was deleted in UWB1.289 and OVCAR8 EOC cell lines that have reduced BRCA1 expression through a deletional mutation (UWB1.289) or promoter methylation (OVCAR8). These mutations are characterized elsewhere in the literature (46, 47). Using 3 unique guide RNAs (Supplemental Material), PARP1 deletion was mediated by Cas9 in both cell lines, effectively producing 3 unique polyclonal PARP1-KO cell lines for each. Loss of PARP1 expression was confirmed by IF cell microscopy and Western blot in each polyclonal population. Polyclonal populations were used in all experiments.

[125I]KX1 radioligand binding measures differences in PARP-1 expression. Using multiple ovarian cancer cell lines, we performed in vitro pharmacological PARP-1 saturation radioligand binding studies to quantify PARP-1 expression. [125I]KX1 was synthesized and in vitro radioligand binding studies were performed as previously described (16). Briefly, increasing concentrations of [125I]KX1 from 0.01-10 nM were added to a 96-well plate in quadruplicate for each respective cell line and experiments were completed in triplicate.

Loss of PARP1 promotes PARPi resistance in ovarian cancer cell lines. Immunofluorescence was used to determine DNA damage induced by olaparib or DMSO control in ovarian cancer cell lines. Using 5 clinically used PARPis, we performed in vitro cell viability assays to determine changes in drug sensitivity based on loss of PARP-1 expression. The experimental procedures were adopted from previous work (16). Cisplatin was also examined as a control to evaluate changes in sensitivity of DNA-damaging agents that occur after loss of PARP1. Cell viability was assessed using CellTiterGlo (Promega). Experiments were completed in duplicate 3 independent times.

Preclinical microPET imaging of PARP-1. Through serial [18F]FTT PET imaging studies, we evaluated PARP-1 drug-target engagement using the FDA-approved PARPi olaparib. PET imaging of PARP-1 drug-target engagement is a noninvasive methodology to directly assess PARP inhibition. Using 2 ovarian cancer patient-derived xenograft models in 15- to 18-week-old female NOD SCID mice provided from the Simpkins laboratory at the University of Pennsylvania (48), including a BRCA1 mutant and WT, we performed microPET imaging on a Phillips Medical Systems scanner (n = 4 mice). Mice were imaged on day 1 and then olaparib was orally administered daily by gavage at a dose of 50 mg/kg. Mice were then reimaged on day 4, 1 hour after the fourth dose of olaparib. A total of 4 doses of olaparib was administered between imaging studies to achieve steady state. Ex vivo autoradiography studies were performed on mice from PDX model WO-12-2 with and without olaparib treatment (n = 6). The differences in tumor-to-muscle ratios were measured and were representative of the percentage of PARP1 drug-target engagement.

PARP-1 immunohistochemistry and autoradiography on clinical specimens. Colorimetric immunohistochemistry was performed at the University of Pennsylvania Pathology Core for H&E and biomarkers, including PARP-1, H2AX, p53, LCA, and AE1/3. These biomarkers were chosen due to their relevance to EOC. AE1/3 is a keratin marker and was used to identify tumor cells by c-IHC. In order to determine where in the tumor microenvironment PARP-1 was expressed, we stained with LCA, which is a pan lymphocyte marker. In addition, we analyzed common biomarkers present in high-grade serous EOC, including p53 and H2AX. Lastly, PARP-1 was evaluated by f-IHC and [125I]KX1 autoradiography on adjacent sections to determine whether the 2 methods colocalize.

Clinical [18F]FTT PET imaging. The clinical trial included 2 imaging cohorts: a biodistribution cohort and a dynamic cohort. The patients included in this trial with pathologic correlates were all enrolled in the dynamic cohort. As such, only that protocol is reported in this section. Dynamic scanning was performed for approximately 60 minutes from the time of injection with a field-of-view over the abdomen/pelvis or to include a known site of tumor involvement. Static images from the skull base to mid-thigh were performed starting approximately 90 and 180 minutes after [18F]FTT injection; these static scans were optional at the discretion of the investigator or study physician. Static images obtained at approximately 90 minutes were used in the analysis. [18F]FTT PET/CT imaging was performed on a Phillips Ingenuity PET/CT scanner (Phillips Medical Systems). Images were reconstructed using standard reconstruction techniques and interpreted by trained radiology readers. Lesions were first located on a contemporaneous clinical [18F]FDG PET/CT and then located on the [18F]FTT PET/CT. The maximum SUV was recorded for each scan.

Statistics. Data presented are mean ± SEM unless otherwise noted. All graphs were produced and statistically analyzed using Prism version 6.0, GraphPad. The maximum number of PARP binding sites (B max) was calculated by an exponential nonlinear curve-fit 1-site binding hyperbola function. All statistical tests comparing the mean of 2 groups were 2-sided. The comparisons of the means from more than 2 groups were performed using 1-way ordinary ANOVA.
Dose-response curves were fitted using a nonlinear sigmoidal dose-response curve, and effective calculations for 50% reduction in cell viability were calculated. Image analysis was performed using PMOD software, and tumor-to-muscle ratios were calculated. Preclinical microPET imaging studies comparing tumor uptake of radiotracer were analyzed using a paired, 2-tailed t test. Autoradiography studies were analyzed using an unpaired 2-tailed t test. We chose an α of 0.05 (95% confidence interval) and data were deemed significant if \( P < 0.05 \). For correlations, the Pearson correlation coefficient was calculated and the data were considered statistically significant if \( P < 0.05 \). To determine the ability of \([^{18}F]FTT\) PET maximum SUV to predict PARP-1 expression measured by f-IHC, we used a linear regression and results were considered statistically significant if different from zero.

Study approval. All animal studies were performed under protocols approved by the University of Pennsylvania Institutional Animal Care and Use Committee, Philadelphia, Pennsylvania, USA. Studies were conducted under guidelines for animal welfare provided by the NIH. This was an open-label, phase I trial of \([^{18}F]FTT\) PET conducted under an approved IRB at the University of Pennsylvania, Philadelphia, PA, USA, and registered with the FDA as an expanded investigational new drug (clinical trial identifier: NCT02637934). All subjects consented to PET imaging and providing clinical tissue specimens for in vitro analysis.

Author contributions

MM designed experiments, performed experiments, collected data, analyzed data, and composed the manuscript. AP read PET scans, determined SUV values in lesions for PET tracers, analyzed data, and composed the manuscript. LS interpreted tissue histology and evaluated biomarker positivity. ES wrote the clinical protocol. KX synthesized precursors for radiotracers used. CJI analyzed preclinical PET imaging data. CH performed in vivo preclinical PET imaging studies. HK developed and produced radiotracer to patients and interpreted PET images. RAG radiotracer to patients and interpreted PET images. DAP administered the radiotracer to patients and interpreted PET images. RHM designed radiotracers and experiments and composed the manuscript. LLL designed experiments, analyzed data, recruited patients, led the clinical trial, and composed the manuscript.

Acknowledgments

The authors would like to acknowledge the University of Pennsylvania PET center and cyclotron facility for clinical support and manufacturing of radiopharmaceuticals used in this study. The authors would like to specifically thank Mark A. Sellmyer and Elizabeth S. McDonald (Perelman School of Medicine, University of Pennsylvania) for their input regarding this work.

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