Role of PET imaging for biochemical recurrence following primary treatment for prostate cancer

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Abstract: Prostate cancer is one of the most common cancers in men worldwide, and primary prostate cancer is typically treated with surgery, radiation, androgen deprivation, or a combination of these therapeutic modalities. Despite technical advances, approximately 30% of men will experience biochemical recurrent within 10 years of definitive treatment. Upon detection of a rise in serum prostate specific antigen (PSA), there is great need to accurately stage these patients to help guide further therapy. As a result, there are considerable efforts underway to establish the role of positron emission tomography (PET) in the diagnostic algorithm of biochemically recurrent prostate cancer. This manuscript provides an overview of PET tracers used for the detection and localization of prostate cancer in the setting of biochemical recurrence with a focus on PET tracers that are currently being used in clinical practice in the United States.

Keywords: Biochemical recurrence; positron emission tomography (PET)/computed tomography (CT); PET/magnetic resonance imaging (MRI); prostate cancer

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Introduction

Prostate cancer is one of the most common cancers in men worldwide (1). Radical prostatectomy can cure appropriately selected patients with localized disease, as can radiation therapy. Imaging for staging at the time of diagnosis varies slightly depending on the patient’s overall risk of extraprostatic disease, but patients classified as intermediate risk or higher will undergo skeletal scintigraphy and either an abdominopelvic CT or pelvic MRI for comprehensive staging per the National Comprehensive Center Network (NCCN) clinical practice guidelines (2). Locally advanced prostate cancer is defined by the NCCN as T3b-T4 tumors, tumors with primary Gleason pattern of 5, or tumors with greater than four cores of Gleason grade group 4 or 5 (2). Patients are deemed metastatic at initial staging if any distant or nodal metastasis is detected on conventional anatomic imaging (2).

Following definitive treatment, patients will undergo serial serum PSA sampling to evaluate for response to treatment and to detect recurrence. Approximately 20–40% of patients undergoing radical prostatectomy and 30–50% of patients undergoing radiation therapy will experience biochemical recurrence within 10 years (3-5). The American Urological Association defines biochemical recurrence in patients who have undergone prostatectomy as an initial PSA value of at least 0.2 ng/mL followed by a subsequent confirmatory PSA value of 0.2 ng/mL or more (6). In patients who have undergone definitive primary radiation therapy, biochemical recurrence is defined as a rise of 2 ng/mL or more above the nadir PSA level (7). Common locations of local recurrence are within the prostate gland, prostatectomy surgical bed, regional nodal metastases, and osseous metastases. Computed tomography (CT) and magnetic resonance imaging (MRI) are non-invasive modalities that have been utilized for
Methods

A search was conducted through PubMed to assess the existing literature regarding the use of PET imaging in the setting of biochemically recurrent prostate cancer. Terms searched included “prostate cancer”, “PET prostate cancer”, and “PET biochemical recurrence”. These searches were conducted on 12/4/2017. Articles were reviewed and included in the paper if they were scientific articles regarding PET imaging of biochemically recurrent prostate cancer, provided needed statistics on prostate cancer recurrence rates and/or mortality, or provided a useful summary or meta-analysis of the prostate cancer PET literature. Articles were excluded if they did not discuss PET imaging of biochemically recurrent prostate cancer or focused on non-PET imaging modalities of prostate cancer.

Basics of prostate cancer PET imaging and available tracers

PET is currently performed in combination with CT or MR for attenuation correction and anatomic localization of PET findings. The vast majority of clinical PET scanners currently are PET/CT systems, although PET/MRI is in clinical use with an increasing installation base. In PET imaging for prostate cancer, a positron-emitting radiopharmaceutical is administered intravenously, and images are acquired through the detection of coincident gamma rays of 511 keV that result from the annihilation of a positron with an electron in a tissue (11). PET images can provide quantitative localization of radioactivity throughout the body at a single or multiple time points after tracer injection.

The timing of imaging acquisition after PET tracer administration varies depending on the pharmacokinetics of the tracer. In some cases, image acquisition within 5 minutes after PET tracer administration is desirable to minimize the concentration of radioactivity in the urinary bladder and ureters. PET provides higher spatial and temporal resolution than conventional single photon computed tomography (SPECT). However, one of the limitations of PET is resolution, with decreased sensitivity of the detection and characterization of PET tracer uptake in lesions less than 8 mm. Additionally, PET tracers are expensive compared to CT and MRI contrast agents, and reimbursement for PET studies can be challenging even when the PET tracers are approved by the Food and Drug Administration (FDA).

A wide range of radionuclides have been used to label PET tracers for prostate cancer imaging. The radionuclides that have been used most commonly for PET tracers
PET, positron emission tomography.

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Table 1 Radionuclides commonly used for PET imaging of prostate cancer

| Radionuclide               | Half-life (min) | Production method |
|----------------------------|-----------------|-------------------|
| Carbon-11[^{11}C]         | 20              | Cyclotron         |
| Gallium-68[^{68}Ga]       | 68              | Generator         |
| Fluorine-18[^{18}F]       | 110             | Cyclotron         |

Table 2 Selected PET radiotracers used for prostate cancer imaging

| PET radiotracers           | Mechanism of action                  |
|----------------------------|--------------------------------------|
| [^{18}F]FDG                | Glucose metabolism                   |
| Na[^{18}F]F               | Bone chemisorption                   |
| [^{18}F]choline[^{11}C]choline | Cell membrane metabolism             |
| [^{18}F]DCFPyl,[^{68}Ga]PSMA-11 | PSMA binding                        |
| [^{11}C]acetate           | Fatty acid metabolism                |
| [^{18}F]fluciclovine      | Amino acid transport                 |
| [^{68} Ga]DOTA-bombesin   | Androgen receptor binding            |
| [^{18}F]FDHT              |                                       |

[^{18}F]FDG is the most commonly used PET agent for clinical oncologic imaging but has limited clinical utility for the staging of prostate cancer at the time of diagnosis and in the setting of initial biochemical recurrence. In particular, there is an inverse relationship between levels of [^{18}F]FDG uptake and tumoral differentiation with well-differentiated tumors demonstrating low-level [^{18}F]FDG uptake and poorly-differentiated tumors demonstrating increasing [^{18}F]FDG uptake (20,21). Another significant limitation of [^{18}F]FDG-PET is the urinary excretion of the radiotracer, which can limit evaluation of the pelvic lymph nodes and prostate surgical bed (22). [^{18}F]FDG may have a limited role in staging and monitoring response to castrate-resistant metastatic prostate cancer or biopsy-proven poorly differentiated recurrent prostate cancer (23).

[^{18}F]FDG and [^{18}F]Sodium fluoride

[^{18}F] FDG is the most commonly used PET agent for clinical oncologic imaging but has limited clinical utility for the staging of prostate cancer at the time of diagnosis and in the setting of initial biochemical recurrence. However, more recent studies demonstrate increased sensitivity of PSMA ligands and [^{18}F] fluciclovine when compared to choline ligands (17,18). There is little current head-to-head comparison between PSMA ligands and [^{18}F] fluciclovine in the literature, but an early case series suggests possible superiority of [^{68} Ga]PSMA-11 over [^{18}F] fluciclovine for lesion detection (19).

PET tracers that have been developed for the evaluation of prostate cancer recurrence include 2-deoxy-2-[^{18}F] fluoro-D-glucose (FDG), [^{11}C] acetate, [^{18}F] DCFPyl and [^{68} Ga] PSMA-11 (also known as HBED-CC), [^{68} Ga] DOTA-bombesin, [^{18}F] fluoro-5a-dihydrotestosterone (FDHT), [^{11}C] choline, and [^{18}F] fluciclovine (Table 2). Currently in the United States, [^{18}F] FDG, sodium [^{18}F] fluoride, [^{18}F] fluciclovine, and [^{11}C] choline are FDA-approved for use in the setting of biochemical recurrence, although [^{18}F] FDG does not have an established role at the time of initial biochemical recurrence. The use of sodium [^{18}F] fluoride, [^{18}F] fluciclovine, and [^{11}C] choline in prostate cancer is included in the National Comprehensive Cancer Network Guidelines version 1.2018 (2).
evaluation of response to treatment. When compared to $^{99m}$Tc-labelled diphosphonates, sodium $[^{18}F]$fluoride offers shorter scanning times, higher spatial resolution, and better image quality (24). A study has reported a sensitivity and specificity of 100% for sodium $[^{18}F]$fluoride-PET/CT compared to 70% sensitivity and 57% specificity for planar bone scintigraphy (25), although it is not clear that $[^{18}F]$fluoride-PET/CT reaches this high degree of specificity in clinical practice. Sodium $[^{18}F]$fluoride-PET/CT also allows for earlier detection of osseous metastases compared to conventional bone scintigraphy and detection of occult osseous metastases when compared to conventional imaging (26,27). Additionally, sodium $[^{18}F]$fluoride-PET/CT positivity tends to associate with increasing PSA levels in men who have undergone prostatectomy and may occur at lower levels of PSA than expected (27). Although the NCCN guidelines state that sodium $[^{18}F]$fluoride can be used when planar bone scans are negative but there is high suspicion for osseous metastases or when there is disease progression, there is concern about its use in the treatment response setting. This concern is largely due to established guidelines being based on planar bone scintigraphy. The impact of sodium $[^{18}F]$fluoride-PET/CT has been demonstrated through the initial results of the National Oncologic PET Registry, where 40% of treatment plans were revised following sodium $[^{18}F]$fluoride-PET/CT (28). Despite this, there are limitations of $[^{18}F]$sodium fluoride-PET/CT, including cost, availability of PET/CT scanners, and lack of soft tissue evaluation which can be performed with other PET tracers.

$[^{11}C]$Choline

Choline is a substrate for the synthesis of phosphatidylcholine, which is the major phospholipid in the cell membrane (29). Choline is incorporated into tumor cells after transport into prostate cancer cells and phosphorylation by choline kinase, which is upregulated in prostate cancer (30). $[^{11}C]$Choline has been approved by the FDA for use in the detection and localization of suspected biochemically recurrent prostate cancer. $[^{11}C]$Choline offers an advantage over $^{18}F$-labeled choline derivatives in that the metabolite of $[^{11}C]$choline demonstrates less urinary excretion than fluorinated choline compounds, which is advantageous in the evaluation of the prostatectomy bed and pelvic lymph nodes (Figure 2). However, a comparative analysis between $[^{11}C]$choline

Figure 1 Sodium $[^{18}F]$fluoride-PET/CT images from a patient with prostate cancer status post external beam radiation therapy, androgen-deprivation therapy, and brachytherapy presenting with elevated PSA and biochemical recurrence. The maximum intensity projection image (MIP, panel A) demonstrates numerous foci of increased tracer activity in several ribs and the bony pelvis. The fused PET/CT (B) and CT only (C) images demonstrate a right sacral metastasis (arrow) with a sclerotic correlate.
and $[^{18}F]$choline demonstrated comparable results in the detection of disease (31). Fanti et al. performed a meta-analysis and critical review of literature consisting of 12 studies including 1,270 patients, specifically looking at the ability of $[^{11}C]$choline detect prostate cancer in the setting of biochemical recurrence after definitive treatment, with a derived pooled sensitivity and specificity of 89% (32). This derived pool data is similar to other published meta-analyses of $[^{11}C]$choline-PET/CT (33-35). The performance of $[^{11}C]$choline-PET/CT is better for larger lymph nodes, but studies have shown decreased sensitivity in lymph nodes measuring less than 7 mm (36). A meta-analysis of $[^{11}C]$choline-PET/CT demonstrated a pooled rate of detection of 36% for nodal disease and 25% for osseous metastases (32). The same meta-analysis demonstrated a pooled detection rate for locally recurrent disease of 27%, with a pooled sensitivity of 61% and a pooled specificity of 97% (32). This result is consistent with comparative studies that have shown that multiparametric MRI with endorectal coil is superior to $[^{11}C]$choline for detection of local recurrence, while $[^{11}C]$choline-PET/CT was shown to be superior to MRI for pelvic lymph node metastases and equal with respect to bone metastases (37). The NCCN guidelines 1.2018 state that $[^{11}C]$choline-PET/CT or PET/MRI can be considered for recurrence or disease progression after definitive therapy or for disease progression during systemic therapy (2).

$[^{18}F]$Fluciclovine

Amino acid transport is another important target for the imaging of prostate cancer due to upregulation of...
transmembrane amino acid transport (38) (Figures 3,4).

The most work with radiolabeled amino acids for prostate cancer imaging has been with $^{18}$Ffluciclovine, a non-natural alicyclic amino acid. $^{18}$Ffluciclovine targets the LAT1 and ACST2 transmembrane transporters, both of which are overexpressed in prostate cancer cells (39,40). $^{18}$Ffluciclovine was approved by the FDA in 2016 for use in biochemically recurrent prostate cancer. An advantage of $^{18}$Ffluciclovine is relatively little urinary excretion of radiotracer, which leads to improved evaluation of potential recurrent disease in the pelvis (41,42). However, a limitation of $^{18}$Ffluciclovine is that it is unable to reliably differentiate malignant intraprostatic lesions from nodules due to benign prostatic hypertrophy, although new research suggests that delayed PET of the pelvis on PET/MRI shows promise for the differentiation between malignant and benign tissue (43,44). Additionally, the transport of $^{18}$Ffluciclovine is bidirectional which can lead to washout from prostate cancer at later time points after injection (45).

$^{18}$Ffluciclovine has been shown to have an overall detection rate of approximately 68% in the setting of biochemical recurrence, including in patients with low PSA levels (detection rate 41% for PSA <0.79 ng/mL) (46). When compared to conventional imaging (CT and bone scan), $^{18}$Ffluciclovine has demonstrated the ability to detect subcentimeter lymph node metastases and osseous metastases that were not detected on conventional imaging (47). Schuster et al. reported specificities of 40% and 97% for prostate bed and extraprostatic lesions, respectively (48). Odewole et al. demonstrated similar specificities of 56% and 100% for prostate bed and extraprostatic lesions, respectively (49). A meta-analysis evaluated $^{18}$Ffluciclovine PET, including 6 studies and 251 patients with biochemical recurrence (50). The pooled sensitivity and specificity on a per-patient analysis was 87% and 66%, respectively (50). One study examining the effect of $^{18}$Ffluciclovine-PET/CT on consideration of salvage radiation therapy in the setting of biochemical recurrence demonstrated a change in management in 41% of patients (51).

A study comparing the diagnostic performance of $^{18}$Ffluciclovine-PET/CT to $^{111}$In capromab pendetide-SPECT/CT demonstrated superior performance for $^{18}$F fluciclovine-PET/CT in the detection of both intraprostatic and extraprostatic disease (48). Additionally, a study comparing the diagnostic performance of $^{18}$Ffluciclovine-PET/CT to $^{11}$Ccholine-PET/CT in biochemical recurrence following definitive therapy revealed superior performance of $^{18}$Ffluciclovine-PET/CT (17). In this study, $^{18}$Ffluciclovine demonstrated a sensitivity of 37% compared to 32% for $^{11}$Ccholine and a specificity of 67%.
compared to 40% for $^{11}$Ccholine. A small study in 10 patients with recurrent prostate cancer comparing $^{68}$Ga PSMA-11 to $^{18}$Ffluciclovine suggested that $^{68}$GaPSMA-11 provides higher sensitivity for prostate cancer detection (19). However, larger studies are needed to determine the diagnostic accuracy of PSMA-PET versus $^{18}$Ffluciclovine for initial staging and the detection of recurrent prostate cancer. The NCCN guidelines 1.2018 for $^{18}$Ffluciclovine are very similar to those for $^{11}$Ccholine and state that $^{18}$Ffluciclovine-PET/CT or PET/MRI can be considered for recurrence or disease progression after definitive therapy or for disease progression during systemic therapy (2).

$^{11}$CAcetate

Acetate is a naturally occurring substance that can enter the fatty acid metabolic pathway, which is overexpressed in prostate cancer cells (52) (Figure 5). Currently, $^{11}$Cacetate-PET/CT is used at fewer sites than choline radiotracers (53). $^{11}$Cacetate-PET/CT has demonstrated good performance in the evaluation of recurrence in the prostatectomy bed, with 15 of the 18 patients with biopsy-proven recurrent disease demonstrating positivity on $^{11}$Cacetate-PET/CT and none of the patients with negative biopsies demonstrating $^{11}$Cacetate-PET/CT positivity (54). The same study also demonstrated comparable results between $^{11}$Ccholine-PET/CT and $^{11}$Cacetate-PET/CT (54). At this point in time, no conclusions have been reached about the superiority of $^{11}$Cacetate-PET/CT versus $^{11}$C choline-PET/CT with additional studies demonstrating similar results between the two radiotracers (55,56). A similar limitation between the two radiotracers is that the sensitivity for detection of disease is correlated with PSA levels, with decreased performance for PSA levels of <2 ng/mL (53,57,58).

Prostate specific membrane antigen (PSMA) ligands

PSMA is a transmembrane protein expressed by the prostate and overexpressed in prostate cancer (59) (Figure 6). PSMA has long been a target for imaging patients with metastatic prostate cancer and was originally the target of $^{111}$Inindium capromab pendetide (tradename Prostascint), a radiolabeled monoclonal antibody targeting the intracellular portion of the transmembrane PSMA protein. Although this was an improvement at the time over existing imaging techniques, the sensitivity and specificity of the examination were
Figure 5  
$[^{11}C]$acetate-PET/CT in a patient with elevated PSA of 0.38 ng/mL status post prostatectomy. CT images (A) and fused PET/CT images (B) demonstrate focal soft tissue with increased tracer activity in the prostatectomy bed (arrow). Whole body MIP image (C) demonstrates no distant metastases. Patient was subsequently referred to radiation oncology for salvage pelvic radiotherapy.

Figure 6 $[^{68}Ga]$PSMA-11-PET/CT in a patient with prostate cancer initially treated with radiation therapy in 2013 with subsequent biochemical recurrence treated with salvage pelvic radiation therapy. Patient PSA continued to be elevated to 7.7 ng/mL. CT image (A), fused $[^{68}Ga]$PSMA-11-PET/CT (B), $[^{68}Ga]$PSMA-11-PET (C), and whole body MIP (D) images demonstrate increased tracer activity in several enlarged left internal mammary lymph nodes (arrow). (Images courtesy of Tom Hope, MD, University of California San Francisco).
significantly limited due to the targeting of the intracellular portion of the PSMA protein which permitted radiotracer binding only in the setting of cellular apoptosis or necrosis (36,59-62). An additional practical limitation was that the kinetics of \[^{111}\text{In}\]jindium capromab pendetide mandated imaging at 5–7 days following radiotracer injection.

New small molecular imaging PSMA ligands have been developed such as \[^{18}\text{F}\]DCFBC, \[^{18}\text{F}\]DCFPyl, and \[^{68}\text{Ga}\]PSMA-11, which bind irreversibly to the extracellular component of PSMA and have been shown to improve detection of metastatic prostate cancer (63,64). The most commonly used PSMA ligands in Europe is \[^{68}\text{Ga}\]PSMA-11. It is important to note that less than 10% of prostate cancers have no uptake on PSMA PET (65). In addition, the agent is rapidly cleared from non-target tissue. In the setting of biochemical recurrence, \[^{68}\text{Ga}\]PSMA-11-PET/CT demonstrated higher tumor-to-background and higher detections rates than \[^{18}\text{F}\]choline-PET/CT, particularly at lower PSA levels (66). A large retrospective study demonstrated that \[^{68}\text{Ga}\]PSMA-11-PET/CT demonstrated uptake in 82% of patients with evidence of biochemical recurrence (67). Among the lesions that underwent surgical intervention or biopsy, 30 false-negative results were observed in a total of 4 patients with the rest of the lesion (n=416) classified as either true-positive or true-negative (67). Of note, this study also demonstrated a 50% detection rate in patients with serum PSA of less than 0.5 ng/mL, a significant improvement compared to the existing literature for choline and acetate and a finding that has been replicated in additional studies (67,68). A recent meta-analysis was performed by Perera et al., in which 16 articles including 1,309 patients were evaluated (69). When evaluating on a per-patient basis, the summary sensitivity and specificity were identical at 86% (69). When analyzed on a per-lesion basis, summary sensitivity was 80% and specificity was 97% (69).

A current limitation of widespread use of \[^{68}\text{Ga}\]PSMA-11-PET/CT in the United States is that it is not currently approved by the FDA. Additionally, production of \[^{68}\text{Ga}\]PSMA ligands currently requires an onsite generator for most sites as the radius of distribution of \[^{68}\text{Ga}\]labeled compounds is limited by the relatively short half-life. However, research is currently ongoing regarding the development of PSMA analogues labeled with \[^{18}\text{F}\]fluorine, which has potential to significantly reduce the cost and increase the availability of PSMA ligand through batch production and remote distribution (70). A potential future in targeted molecular imaging of metastatic prostate carcinoma is the use of PSMA for targeted therapy with alpha or beta-emitters (including \[^{90}\text{Y}\] and \[^{177}\text{Lu}\]), currently being utilized in clinical trials for patients with biochemical recurrence, but not currently approved for use in the United States (71).

Emerging and experimental PET radiotracers for prostate cancer

Gastrin-releasing peptide receptor (GRPR) is a current investigational target for prostate cancer imaging (Figure 7). GRPR is overexpressed in prostate cancer cells, but demonstrates lower levels of expression in benign prostate tissue (72). Increases in GRPR expression have been shown to be present in 63–100% of intraprostatic prostate cancers and 50–85% of nodal and osseous metastases (73). To target and image GRPR overexpression, several peptide-based bombesin and gastrin-related peptide (GRP) analogs have been developed and labeled with a number of radioisotopes (74-76). A recent study examined the use of \[^{68}\text{Ga}\]RM2-PET/CT (GRPR antagonist) in patients with known biochemical recurrence and negative or equivocal \[^{18}\text{F}\]fluoroethylcholine-PET/CT and demonstrated that \[^{68}\text{Ga}\]RM2-PET/CT was helpful in localizing the recurrence in a majority of the cases (77). An additional study examining the use of a \[^{68}\text{Ga}\]labeled bombesin analog for PET/CT in patients with both initial diagnosis of prostate cancer and biochemical recurrence demonstrated a sensitivity of 88%, specificity of 81%, and accuracy of 83% for primary prostate cancer and 70% sensitivity for detection of metastatic lymph nodes (78). The same bombesin analog also correctly identified two of three cases of local recurrence both in the prostatectomy surgical bed and regional lymph nodes when compared to \[^{13}\text{C}\]acetate-PET/CT (78). Similar to PSMA ligands, bombesin agents are being examined for their potentialtheranostic capabilities in delivering targeted radiation via alpha-emitters or beta-emitters (79).

Therapies targeting androgen receptor signalling are a cornerstone of the treatment algorithm of prostate carcinoma, both for treatment-naïve patients and patients diagnosed with biochemical recurrence. Although not currently utilized in the clinical setting, PET imaging targeting the androgen receptor would therefore make sense in the setting of prostate cancer. The majority of preclinical and initial in-human work for PET imaging of androgen receptors has been with 16β-\[^{18}\text{F}\]DFHT (Figure 8). An initial study examining the \[^{18}\text{F}\]FDHT in patients...
Figure 7 Whole body MIP (A), MR (B), $^{68}$GaRM2-PET (C), and fused $^{68}$GaRM2-PET/MRI images (D) in a 73-year-old patient with biochemical recurrence status post radiation therapy (PSA = 1.32 ng/mL) demonstrate activity in the right seminal vesicle, suspicious for local recurrence (arrow). (Images courtesy of Andrei Iagaru, MD, Stanford University).

Figure 8 $^{18}$FDFHT-PET images of a patient with biochemically recurrent metastatic prostate cancer (PSA = 62 ng/mL) before (A) and after (B) initiation of flutamide demonstrate osseous and lymph node metastases (arrows) and subsequent treatment response. (Images courtesy of Farrokh Dehdashti, MD, Washington University in St. Louis).
with progressive metastatic prostate cancer demonstrated uptake is 46/59 lesions compared with $^{18}$FFDG uptake in 57/59 (80). An additional pilot study demonstrated $^{18}$FDHT PET positivity in 63% of patients with advanced prostate cancer and demonstrated an additional 17 unsuspected lesions in a total of 10 patients (81). Interestingly, when the patients underwent a repeat $^{18}$FDHT-PET/CT one day following the initiation of flutamide, there was an immediate decrease in the radiotracer activity of greater than 50% that suggests $^{18}$FDHT-PET may be useful in tracking response to androgen deprivation therapy (81). This finding was later replicated in a phase 1/2 study evaluating the use of $^{18}$FDHT-PET to assess androgen blockade in patients with castration-resistant prostate cancer (82). The optimal use of androgen receptor PET imaging in prostate cancer remains to be established, but the evidence is promising that certain niche applications of $^{18}$FDHT-PET may emerge in the future.

PET/CT vs. PET/MRI for prostate cancer

PET/MRI scanners are being implemented throughout the world for routine clinical applications with increasing frequency. These scanners are capable of acquiring PET and MRI data simultaneously with the potential for more accurate image co-registration. In the setting of biochemical recurrence following both prostatectomy or definitive radiation therapy, MRI of the pelvis remains a cornerstone of evaluation for potential sites of local recurrence owing to its superior soft tissue contrast. Given the development of multiple PET radiotracers that show excellent sensitivity and specificity for recurrent prostate cancer [e.g., $^{18}$F] fluciclovine, $^{68}$GaPSMA-11], it is logical that PET/MRI may become the optimal imaging modality for patients with biochemically recurrent prostate cancer. Several studies have shown a high detection rate of PET/MRI for pelvic recurrence in the setting of biochemical recurrence (83,84). As PET/MRI scanners become increasingly common, further research is needed to demonstrate the added value of PET/MRI both in the pretreatment and posttreatment settings.

Conclusions

Molecular imaging of prostate cancer continues to evolve as new radiotracers are studied and put into clinical practice. The use of both PET/CT and PET/MRI are an important diagnostic consideration for patients with biochemically recurrent prostate cancer, particularly if locoregional therapy is being considered. As newer PET radiotracers become approved, it will be important for larger prospective head-to-head studies to be performed if an optimal molecular imaging algorithm is to be developed. At this point in time, it appears that both $^{18}$Ffluciclovine and PSMA tracers are superior to the older PET tracers, but little existing data exists directly comparing the two. Given the existing research in theranostics with both PSMA and GRPR tracers, these imaging agents may be the optimal agent if consideration is being given to targeted molecular therapy with alpha-emitters or beta-emitters.

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Footnote

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