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Molecular Targeted Positron Emission Tomography Imaging and Radionuclide Therapy of Pancreatic Ductal Adenocarcinoma

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Simple Summary: Pancreatic ductal adenocarcinoma (PDAC) has a dismal prognosis, mainly due to difficulty in early detection of the disease by current imaging modalities. In this review, we discuss the more specific diagnostic imaging modality that evaluates the presence of specific tumour tracers via positron emission tomography. In addition, we review the available therapeutic applications of these tumour-specific tracers.

Abstract: Pancreatic ductal adenocarcinoma (PDAC) has an inauspicious prognosis, mainly due to difficulty in early detection of the disease by the current imaging modalities. The upcoming development of tumour-specific tracers provides an alternative solution for more accurate diagnostic imaging techniques for staging and therapy response monitoring. The future goal to strive for, in a patient with PDAC, should definitely be first to receive a diagnostic dose of an antibody labelled with a radionuclide and to subsequently receive a therapeutic dose of the same labelled antibody with curative intent. In the first part of this paper, we summarise the available evidence on tumour-targeted diagnostic tracers for molecular positron emission tomography (PET) imaging that have been tested in humans, together with their clinical indications. Tracers such as radiolabelled prostate-specific membrane antigen (PSMA)—in particular, ¹⁸F-labelled PSMA—already validated and successfully implemented in clinical practice for prostate cancer, also seem promising for PDAC. In the second part, we discuss the theranostic applications of these tumour-specific tracers. Although targeted radionuclide therapy is still in its infancy, lessons can already be learned from early publications focusing on dose fractioning and adding a radiosensitiser, such as genecitabine.

Keywords: pancreatic ductal adenocarcinoma; positron emission tomography; radionuclide; tumour tracer

1. Introduction

Pancreatic ductal adenocarcinoma (PDAC) is the most frequent type of all pancreatic cancers and has an inauspicious prognosis, with a five-year survival rate of less than 5% [1,2]. This extremely low survival rate is mainly due to difficulty in early detection of the disease by the current imaging modalities. Staging and, hence, rational use of treatment are highly dependent on information yielded from conventional imaging modalities (i.e., computed tomography (CT), magnetic resonance imaging (MRI), or endoscopic ultrasound



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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). (EUS) [3]. However, almost 50% of surgeries are performed without patient benefit (i.e., due to benign diagnoses, undetected metastases, or rapid recurrence <6 months), indicating that these imaging modalities are lacking diagnostic precision and therapy response evaluation accuracy.

During surgery for PDAC, 10% of the patients already present with metastases at laparoscopy, and approximately half of the patients undergoing a resection will have microscopically positive resection margins (R1), of whom 25% will develop disease recurrence within six months after surgery. Furthermore, the imaging in patients with borderline resectable or locally advanced PDAC who started chemotherapy is unreliable due to the difficulty in distinguishing between fibrosis and stroma in PDAC [4]. Additionally, ¹⁸F-fluorodeoxyglucose (¹⁸F-FDG) PET-CT, the most commonly used tracer in oncology, has a variable and debatable role in the routine pancreatic work up, mainly due to the large number of false positive findings by also identifying pancreatitis, potentially resulting in futile resections of the pancreas. ¹⁸F-FDG PET-CT is therefore only reserved on indication for the individual patient [4].

The upcoming development of tumour-specific tracers provides an alternative solution for more accurate diagnostic techniques, staging, and therapy response monitoring. Targeted radionuclides such as radiolabelled peptides, which bind to the receptors overexpressed by cancer cells, and radiolabelled antibodies to tumour-specific antigens can provide a more specific diagnosis [5–7]. Additionally, this development offers new possibilities to maximally capitalise on the theranostic applicability, i.e., the possibility to use the tracer both for imaging purposes as well as a targeting binder for radionuclide therapy.

In the first part of this review, we summarise the available evidence on tumourtargeted imaging tracers for molecular PET-CT imaging that have been tested on humans, together with their clinical indications, and in the second part, we discuss the theranostic applications of these tumour-specific tracers.

For this narrative review, our search strategy for both the diagnostic and therapeutic sections consisted of a general search of diagnostic and therapeutic tracers in pancreatic cancer, followed by a search of specific tracers and, finally, reviewing the papers for leads to other—not yet included—tracers.

2. Part 1: Tumour-Targeted Tracers for the Detection of Pancreatic Cancer

Early detection is important for the treatment of PDAC. It is believed there are two main precursors for PDAC, namely pancreatic intraepithelial neoplasia (PanIN) and intraductal papillary mucinous neoplasm (IPMN).

Three grades can be distinguished in PanIN. PanIN-1 and PanIN-2 are commonly found in patients over the age of 40 or in chronic pancreatitis. PanIN-3 is more exclusively found in the pancreas with PDAC and is the stage prior to invasion [8–10].

IPMNs develop in the cells lining the pancreatic ducts and contribute to mucin production, cystic dilatation of the pancreatic ducts, and intraductal papillary growth. IPMNs are at risk of developing into malignancy 20% of the time over a period of 10 years [8,11,12].

PanINs are challenging to diagnose, as all types are under the resolution of conventional imaging, due to their limited size. EUS may help in detecting identifiable parenchymal changes, such as acinar cell loss, the proliferation of small ductular structures, and fibrosis. This combination of changes, labelled as lobulocentric atrophy (LCA), however, are not specific to PanIN [8,13].

In diagnosing PDAC, an important challenge is to distinguish PDAC from pancreatitis, as both entities have abundant stroma. Additionally, neoadjuvant treatment such as FOLFIRINOX makes it difficult to discriminate between viable tumour and chemoradiation-induced tumour necrosis and fibrosis [14–16].

Tumour-targeted molecular imaging could provide essential knowledge in these situations by adding metabolic molecular imaging information to the anatomical changes.

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2.1. ¹⁸F-Fluorodeoxyglucose—¹⁸F-FDG

PET imaging for the diagnosis of PDAC uses ¹⁸F-FDG, a radiolabelled glucose [17]. ¹⁸F-FDG PET imaging relies on the property that a normal pancreas tissue has low glucose usage compared to PDAC. In PDAC, a KRAS mutation induces an overexpression of hexokinase-2 and the glucose cell membrane transporter, GLUT-1 [18]. ¹⁸F-FDG is accumulated by PDAC, where it is phosphorylated and consequently goes into metabolic arrest [17,19] (Figure 1 and Table 1).



Figure 1. Available tracers and their properties. **1.** ¹⁸F-FDG tumour cells with the property of using glucose; GLUT-1 and hexokinase are upregulated in tumours. ¹⁸F-labelled FDG accumulation. **2.** ¹⁸F-FLT: cell proliferation in tumour cells; thymidine kinase is upregulated. ¹⁸F-labelled (FLT) \rightarrow FLTTP accumulation. **3.** $\alpha\nu\beta6$ overexpressed in tumour cells; targeted by labelled peptides: ¹⁸F-FP-R01-MG-F2 or ⁶⁸Ga-NODAGA-R01-MG. **4.** PSMA expressed in tumour cells; targeted by inhibitor molecules labelled with ¹⁸F or ⁶⁸Ga. **5.** Expression of FAP (fibroblast activation protein) by CAF (cancer-associated fibroblasts) targeted by FAPI (FAP inhibitor) labelled with ⁶⁸Ga or ¹⁸F. Abbreviations: $\alpha\nu\beta6$ = integrin $\alpha\nu\beta6$; DNA = deoxyribonucleic acid; FAP= fibroblast activation protein; ¹⁸F = ¹⁸Fluorodeoxyglucose-labelled; ¹⁸F-FDG = ¹⁸F-FDG -6-P = ¹⁸F-FDG -6-P hosphate; ¹⁸F-FLT = ¹⁸F-fluorothymidine; ¹⁸F-FLTDP = ¹⁸F-FLT diphosphate; ¹⁸F-FLTMP = ¹⁸F-FLT monophosphate; ¹⁸F-FLTTP = ¹⁸F-FLT triphosphate; ¹⁸F-FP-R01-MG-F2 = ¹⁸F-labelled rintegrin tracer; ¹⁸F-PSMA = ¹⁸F-labelled PSMA; ⁶⁸Ga = ⁶⁸Gallium-labelled; ⁶⁸Ga-FAPI = ⁶⁸Ga-labelled fibroblast activation protein inhibitor; ⁶⁸Ga-NODAGA-R01-MG = ⁶⁸Ga-labelled integrin tracer; ⁶⁸Ga-NODAGA-R01-MG = ⁶⁸Ga-labelled integrin tracer; ⁶⁸Ga-PSMA = ⁶⁸Ga-labelled PSMA; Glucose-6-P = Glucose-6-Phosphate; TTP = thymidine triphosphate.

The average sensitivity and specificity for detecting PDAC by ¹⁸F-FDG is reported to be superior to CT, with a sensitivity of 94% and specificity of 90% for ¹⁸F-FDG compared to 82% and 75%, respectively, for CT [14,18,20].

A major limitation of PET imaging with ¹⁸F-FDG is that glucose uptake can also be seen in inflammation, leading to similar appearances of pancreatitis and PDAC [14,21]. However, when the diagnosis of PDAC is correct, the degree of ¹⁸F-FDG uptake can predict tumour aggressiveness and survival [22,23].

Tracer	Properties	Localisation	Main Advantage	Main Disadvantage
¹⁸ F-FDG	Marker of glucose consumption	Intracellular	High-glucose-use of malignant cells	High-glucose-using cells in inflammation
¹⁸ F-FLT	Marker of cell proliferation	Intracellular	Cell proliferation in malignancies	
⁶⁸ Ga-FAPI; ⁶⁸ Ga-DOTA-FAPI-04	Expression of FAP by CAF targeted by FAPI labelled with ⁶⁸ Ga	Cell membrane of cancer-associated fibroblast	After multiple time points, PDAC and pancreatitis show a trend for differential uptake kinetics	Can be false positive in pancreatitis
¹⁸ F-FP-R01-MG-F2; ⁶⁸ Ga-NODAGA-R01-MG; ⁶⁸ Ga-Trivehexin	Labelled peptides targeting $\alpha v \beta 6$ overexpressed in tumour cells	Cell membrane	Distinguishment between PDAC and pancreatitis. Additional uptake in lymph node metastases	
Radiolabelled PSMA (i.e., ¹⁸ F-PSMA; ⁶⁸ GA-PSMA)	Inhibitor molecules labelled with ¹⁸ F or ⁶⁸ Ga targeting PSMA expressed in tumour cells	Cell membrane	Very high diagnostic accuracy between PDAC and pancreatitis	

Table 1. Available tracers and their properties for the diagnosis of the primary disease.

Abbreviations: $\alpha \nu \beta 6$ = integrin $\alpha \nu \beta 6$; CAF = cancer associated fibroblast; FAP = fibroblast activation protein; FAPI = FAP inhibitor; ¹⁸F = ¹⁸Fluorodeoxyglucose-labelled; ¹⁸F-FDG = ¹⁸F-fluorodeoxyglucose; ¹⁸F-FLT = ¹⁸F-fluorothymidine; ¹⁸F-FP-R01-MG-F2 = ¹⁸F-labelled integrin tracer; ¹⁸F-PSMA = ¹⁸F-labelled PSMA; ⁶⁸Ga = ⁶⁸Gallium-labelled; ⁶⁸Ga-DOTA-FAPI-04 = ⁶⁸Ga-labelled (macrocyclic chelator) 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid-FAPI-04; ⁶⁸Ga-FAPI = ⁶⁸Ga-labelled fibroblast activation protein inhibitor; ⁶⁸Ga-NODAGA-R01-MG = ⁶⁸Ga-labelled integrin tracer; ⁶⁸Ga-Trivehexin = ⁶⁸Ga-labelled Trivehexin; ⁶⁸Ga-PSMA = ⁶⁸Ga-labelled PSMA; PSMA = prostate-specific membrane antigen.

In clinical practice, differentiation between pancreatitis and PDAC is possible by performing a dual-phase PET scan. This method consists of performing a PET scan at two different time intervals after the injection of the tracer. Pancreatic masses on PET images in pancreatitis have lower standardised uptake values (SUV), which further decrease in the delayed phase. However, there can be overlap in SUV values between inflammation and PDAC. Furthermore, dual-phase ¹⁸F-FDG PET imaging is very time-consuming and, therefore, not often feasible in daily practice [17,24].

The specificity of PET imaging for the diagnosis of PDAC could be improved by using more disease-specific imaging agents compared to ¹⁸F-FDG. Several other radiotracers have been used for the evaluation of PDAC [17,25,26]. These include radiotracers such as ¹⁸F-Fluorothymidine (¹⁸F-FLT), ⁶⁸Gallium-labelled fibroblast activation protein inhibitor (⁶⁸Ga-FAPI), ⁶⁸Ga-labelled 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid-FAPI-04 (⁶⁸Ga-DOTA-FAPI-04), ¹⁸Fluorodeoxyglucose-labelled PSMA (¹⁸F-PSMA), ⁶⁸Ga-labelled PSMA (⁶⁸Ga-PSMA), and Integrin $\alpha\nu\beta6$ tracers. These tracers are amply discussed in the following sections.

2.2. ¹⁸F-Fluorothymidine—¹⁸F-FLT

¹⁸F-fluorothymidine is a marker of cell proliferation due to tracer accumulation in proliferating cells. Thymidine kinase activity is upregulated during proliferation, subsequently phosphorylating ¹⁸F-FLT, which gets trapped intracellularly (Figure 1 and Table 1).

¹⁸F-FLT PET imaging has shown a good correlation with histological Ki-67 expression, a marker of cell proliferation [17,27]. Furthermore, ¹⁸F-FLT PET imaging is potentially superior to ¹⁸F-FDG PET, as ¹⁸F-FLT uptake is not affected by inflammation or hypergly-caemia [17].

2.3. Fibroblast Activation Protein Inhibitor-FAPI

In PDAC, more than 90% of the tumour volume consists of cancer-associated fibroblasts (CAF). CAFs are associated with the promotion of tumour growth, tissue invasion, metastasis developing, and therapy resistance [28–31]. CAFs express Fibroblast Activation Proteins (FAP) on the cell surface, a type II membrane-bound glycoprotein [32,33]. FAP can be detected by performing a PET-CT with ⁶⁸Ga-labelled FAP inhibitors (Figure 1 and Table 1).

Röhrich et al. showed in a small study including 19 PDAC patients (seven primary and 12 progressive/recurrent) that ⁶⁸Ga-FAPI PET-CT led to restaging in half of the patients with PDAC and, also, in most patients with recurrent disease compared to standard of care imaging. Differentiation from pancreatitis was challenging but significantly improved with imaging at multiple time points after the injection of ⁶⁸Ga-FAPI [28].

Chen et al. compared the use of ⁶⁸Ga-DOTA-FAPI-04 to ¹⁸F-FDG PET for the diagnosis of the primary disease and metastatic lesions for various types of cancer. Four patients with pancreatic cancer were included. In one patient, pancreatic cancer was not visualised due to uptake throughout the pancreas caused by tumour-induced pancreatitis [34]. Identical findings of uptake of ⁶⁸Ga-DOTA-FAPI-04 in patients with Ig-G4-related disease have been reported by others [34–38].

The study of Chen et al. did show a significantly lower uptake of ⁶⁸Ga-DOTA-FAPI-04 than ¹⁸F-FDG, thus facilitating an improved detection of possible liver metastases [34].

2.4. Integrin αvβ6

Integrins are proteins that facilitate the adhesion of cells to the extracellular matrix (ECM) of polypeptides. Integrins play a crucial role in the signalling pathway for the regulation of cell differentiation, migration, proliferation, and apoptosis [39,40]. In many cancers, the expression of specific integrins can become dysregulated, such as $\alpha\nu\beta3$ and $\alpha\nu\beta6$. The overexpression of $\alpha\nu\beta3$ results in overpromotion of the angiogenesis pathway [39,41].

Integrin $\alpha\nu\beta6$ promotes PDAC by modulating the proliferation, survival, migration, and invasion of both the cancer cells and its microenvironment [42]. Studies have shown higher expressions of $\alpha\nu\beta6$ in PDAC compared to other type of cancers [39,43,44]; additionally, the differentiation of PDAC from pancreatitis was possible [45]. Tumour-positive lymph nodes also showed elevated levels of $\alpha\nu\beta6$ [45].

As $\alpha\nu\beta6$ seems to be an important integrin for the detection of PDAC and distinguished from pancreatitis, numerous research groups have been developing PET tracers [46–48].

Kimura et al. used ¹⁸F-FPR01-MG-F2 to target $\alpha\nu\beta6$ (Figure 1 and Table 1). The study group demonstrated that the targeting peptide was able to penetrate the pancreatic tumour rapidly and also showed an improved uptake compared to ¹⁸F-FDG, reflecting the difference in the peptide's target, namely glucose metabolism versus expression of the ECM protein. In addition, it was observed that uptake only occurred in the viable part of the tumour compared to the parts with significant necrosis [39].

A recent small study from Quigley et al. showed the first promising results for ⁶⁸Galabelled trimerized $\alpha v\beta 6$ -integrin-selective nonapeptide (⁶⁸Ga-Trivehexin)-enabled PET-CT imaging. One patient, out of a total of four, was included with PDAC, showing a high tracer uptake in the pancreatic tumour, including multiple liver metastases [49].

2.5. Prostate-Specific Membrane Antigen Targeted PET-CT Imaging

Currently, different types of radiolabelled PSMA tracers exist (i.e., ¹⁸F-PSMA and ⁶⁸Ga-PSMA) with different biodistributions, as largely described in the literature [50].

Prostate-specific membrane antigen (PSMA) is a type II transmembrane glycoprotein highly expressed on the surfaces of prostate cancer cells. The expression of PSMA in the tumour-associated (neo)vasculature of prostate cancer, breast cancer and primary gliomas has been reported and has also been proven to be high in PDAC [51–53].

The immunohistochemical experiments from our group showed high expression of PSMA in four out of five patients with PDAC de novo, as well as in 32 out of 33 PDAC patients after neoadjuvant treatment (mean tumour H-score of 99 (maximum 300)). These experiments also showed no expression in adjacent normal and pancreatitis tissue (H-score 0), thus yielding a high tumour contrast with the background and improved tumour detection [54].

Radiolabelled PSMA-targeted PET-CT has proven highly successful for the primary staging and restaging of prostate cancer patients and is currently being implemented worldwide [55,56]. PSMA expression can be imaged by labelling small inhibitor molecules with PET radionuclides, i.e., ¹⁸F or ⁶⁸Ga [57] (Figure 1 and Table 1).

2.6. ¹⁸F-labelled Prostate-Specific Membrane Antigen—¹⁸F–PSMA

¹⁸F-radiolabelled PSMA PET has been recently technically validated and successfully implemented in clinical practice for prostate cancer [54,55,58].

One of the more commonly used variants, ¹⁸F-DCFPyL, a second-generation ¹⁸F-fluorinated PSMA ligand, has advantages over ⁶⁸Ga-labelled PSMA tracers. It provides a higher spatial resolution, along with a longer half-life, which may result in more accurate staging due to the detection of small local tumour deposits [59].

2.7. ⁶⁸Ga-labelled Prostate-specific Membrane Antigen—⁶⁸Ga-PSMA

Krishnaraju et al. showed improved diagnostic accuracy with ⁶⁸Ga-PSMA compared to ¹⁸F-FDG in a study among 40 patients with pancreatic lesions—positive predictive value 90.5% versus 65.4% for ⁶⁸Ga-PSMA compared to ¹⁸F-FDG; accuracy 92.5% versus 72.5%, respectively [57].

3. Part 2: Targeted Radionuclide Therapy of Pancreatic Ductal Adenocarcinoma

Locally advanced and metastatic pancreatic cancer have poor prognoses. The current standard of care treatments such as gemcitabine or FOLFIRINOX provide minimal survival benefits. Targeted radionuclide therapy may provide improved survival in addition to less systemic toxicity than seen with the current chemotherapy [60].

In patients with cancer, the ultimate goal is, first, to identify the receptor expression (by using a diagnostic scan with a diagnostic tracer) and, then, in the case of an adequate expression of the receptor, to use the tracer, radiolabelled with alpha or beta-particles, for therapeutic purposes, with curative or sometimes palliative intent [61].

In addition to therapy with radiolabelled antibodies, there is a possibility to add gemcitabine, serving as a radiosensitiser, which is generally well-tolerated in combination with external radiotherapy [62–66].

In this second part, we will discuss the theranostic applications of tumour-specific tracers. To our knowledge, there have only been two different types of antibodies that have been reported on for humans as a targeted radionuclide therapy of PDAC: ¹³¹I-KAb201 antibody and ⁹⁰Y-clivayuzumab tetraxetan antibody (⁹⁰Y-labelled hPAM4). Furthermore, there is only one single study registered at the popular databases that is currently recruiting patients and focusing on the theranostic pair of ⁶⁸Ga-DOTA-5G /¹⁷⁷Lu-DOTA-ABM-5G [67].

3.1. ¹³¹I-labelled KAb201 Antibody—¹³¹I-KAb201

Carcinoembryonic antigen (CEA) is expressed in most patients with pancreatic cancer and therefore serves as an interesting target for antibodies. One potential antibody is KAb201, an anti-CEA antibody labelled with ¹³¹Iodine [68].

Sultana et al. performed a randomised phase I/II trial assessing the safety and efficacy of ¹³¹I-KAb201 in patients with inoperable PDAC. Patients were randomised to receive ¹³¹I-KAb201 via either the intra-arterial (gastroduodenal artery) or intravenous (standard intravenous line) delivery route. The hypothesis of including an intra-arterial delivery route is expected to have a higher concentration of the radiolabelled drug at the target site and, thus, increased effectiveness with reduced toxicity [68].

In total, nineteen patients were randomised (nine in the intravenous arm, 10 in the intra-arterial arm), of whom one patient was excluded from the intra-arterial arm, as there was no uptake on the pretherapy scan. The overall response rate was 6% (one out of 18 patients). Dose-limiting toxicity was only reached in the intra-arterial route (at 50 mCi).

Both anti-chimeric antibodies (HACA) and anti-sheep antibodies (HASA) were developed for the entire study population, thus limiting the possibility of repeat dosing, as this could lead to either hypersensitivity reactions or to complexing with circulating antibodies, creating a challenge to maintain effective therapeutic levels [68,69].

The median overall survival was 5.2 months (95% CI = 3.3–9.0 months), with no significant difference between either delivery arm (log-rank test p = 0.79) [70]. Survival and efficacy data were comparable with a single agent therapy of gemcitabine [71–73].

Future improvements can be found in the ability to predict the occurrence and type (I or II) of antibody response, thus aiding the possibility of repeat dosing. Additionally, humanisation of the antibody may reduce the immunogenicity [68].

3.2. ⁹⁰Y-labelled Clivazutumab Tetraxetan Antibody—⁹⁰Y-hPAM4

Preclinical studies in nude mice have shown that ⁹⁰Y-labelled PAM4 decelerates tumour growth [70,74]. PAM4 is a monoclonal antibody that binds to a mucin produced primarily in PDAC [75–78].

Gulec et al. performed a phase I single-dose escalation trial among 21 patients with PDAC (four stage III—locally advanced; 17 stage IV—metastatic) with the primary aim of determining the dose-limiting toxicity and the maximum tolerated dose. Patients first received ¹¹¹In-hPAM4 for diagnostic imaging and, finally, ⁹⁰Y-hPAM4 for therapy [79].

The drug-related toxicities among the study group were grade 3/4 neutropenia and thrombocytopenia, which both increased with the ⁹⁰Y dose. Fourteen patients progressed rapidly; however, seven patients remained progression-free for 2–6 months, with three patients showing a partial response with tumour shrinkage. The combination with gemcitabine showed further improvements [79].

Fractionated dosing of ⁹⁰Y-hPAM4 (for an increased total radiation dose) in combination with gencitabine acting as a radiosensitiser (for an increased potency of the radiation) could be a promising treatment regimen [38,79].

Ocean et al. showed in a phase 1 trial among 38 untreated patients with pancreatic cancer (five stage III—locally advanced, 33 stage IV—metastatic) that the fractionated dosing of ⁹⁰Y-hPAM4 in combination with gemcitabine in repeated cycles (the number of cycles variated among the patients) allowed for double the radioimmunotherapy dose [80].

The drug-related toxicities among the study group were grade 3/4 thrombocytopenia and neutropenia in 28 patients. Sixteen patients showed stabilisation and six patients a partial response. The median overall survival was 7.7 months for all patients, with improved survival up to 11.8 months with repeated cycles [80].

An important terminated and unpublished study is the PANCRIT-1 trial. This was an international, multi-centre, double-blinded, randomised phase III trial of ⁹⁰Y-labelled hPAM4 in combination with gemcitabine versus a placebo in combination with gemcitabine in patients with metastatic PDAC who had progressed despite receiving at least two prior therapies for metastatic disease. After the enrolment of 334 patients, an interim analysis on overall survival was performed, showing that the treatment arm did not demonstrate a sufficient improvement in the overall survival [81]. One major flaw of the study seemed to be the lack of a pre-treatment evaluation of the receptor expression, thus not applying the theranostic concept.

3.3. ⁶⁸Ga-DOTA-5G /¹⁷⁷Lu-DOTA-ABM-5G Theranostic Pair

(⁶⁸Gallium-labelled—DOTA—G/¹⁷⁷labelled Lutetium—DOTA–ABM-5G; DOTA = 1,4, 7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid).

The only study listed in a clinical study database (ClinicalTrials.gov) is from Sutcliffe et al. from the University of California. This is a phase I study evaluating the safety and efficacy of the theranostic pair of ⁶⁸Ga-DOTA-5G /¹⁷⁷Lu-DOTA-ABM-5G in patients with locally advanced or metastatic PDAC. Patients will first receive a diagnostic ⁶⁸Ga-DOTA-5G PET scan; subsequently, only the patients that show any uptake will receive ¹⁷⁷Lu-DOTA-ABM-5G as a therapy.

The primary objective is to identify the dose limiting toxicity and the recommend phase 2 dose. The objective is to enrol 30 participants, with an expected completion in 2023 [67].

In summary, the results from the study with ¹³¹I-KAb201 demonstrated the importance of further investigation into the type of antibody response and the ability to predict this adverse event for the possibility of repeat dosing. Future research will learn if humanisation of the antibody is able to reduce immunogenicity. Additionally, the route of delivery (intraarterial versus intravenous) did not show any difference in survival benefit or reduction in toxicity. Studies using ⁹⁰Y-labelled hPAM4 proved that dose fractioning could be successful in increasing the total radiation dose without increasing the adverse events. In addition, a combination with gemcitabine acting as a radiosensitiser can increase the potency of the radiation. Future research, including randomised controlled trials, will need to confirm these results.

We are looking forward to the first results from the ⁶⁸Ga-DOTA-5G/¹⁷⁷Lu-DOTA-ABM-5G theranostic pair, as an evaluation of the quantifiable antibody localisation at the site of disease before administering a therapeutic dose seems to be the best tailor-made medicine.

4. Summary and Challenges for the Future

Impressive efforts have been made in improving tumour-specific tracers for the detection of PDAC. In patients with PDAC, the ultimate goal is to firstly identify the radiolabelled peptide expression and, then, in the case of an adequate expression of the peptide, to use the tracer, radiolabelled with alpha or beta-particles, for therapeutic purposes, with curative or sometimes palliative intent.

18F-FDG is a well-known radiotracer that already is being used in PDAC. An important limitation, however, is that increased glucose metabolism is not specific for malignant processes only but can also be found in inflammatory and infectious disease sites. The specificity of PET imaging for the diagnosis of PDAC could be improved by a more diseasespecific imaging agent.

In general, there are currently three types of tracers: tracers that accumulate in the proliferating cell, such as ¹⁸F-FLT, and the second type of tracers target highly expressed integrin receptors or PSMA on the surfaces of cells in PDAC. In this group, ¹⁸F-PSMA is a promising tracer (Figure 2) that has already been validated and successfully implemented in clinical practice for prostate cancer. The last type of tracer targets fibroblast activation protein (FAP inhibitor), which is expressed by CAF (cancer-associated fibroblast).

Targeted radionuclide therapy is still in its infancy. The effectiveness of targeted radionuclide therapy has been limited by poor delivery to tumours. There have only been two different types of antibodies that have been reported on in humans as targeted radionuclide therapy of pancreatic ductal adenocarcinoma: ¹³¹I-KAb201 antibody and ⁹⁰Y-clivayuzumab tetraxetan antibody (⁹⁰Y-labelled hPAM4). Besides selecting the correct radionuclide antibody, the important contributing factors for successful therapy are dose fractioning and the addition of a radiosensitiser, such as gemcitabine.

These novel diagnostic and therapeutic approaches, in populations often characterised by poor outcomes and a decreased quality of life, have the potential to add a new chapter to patient's lives.



Figure 2. CT versus MRI versus ¹⁸F-FDG versus ¹⁸F-PSMA PET and the corresponding pathology and HE and immunohistochemistry tests in a patient with pancreatic adenocarcinoma. (**A**) CT image with an arrow pointing towards a pancreatic lesion. (**B**) MRI image with an arrow pointing towards a pancreatic lesion. (**C**) ¹⁸F-FDG PET image with an arrow pointing towards a pancreatic lesion. (**D**) ¹⁸F-PSMA PET image with an arrow pointing towards a pancreatic lesion. Note the more specific PSMA uptake compared to the ¹⁸F-FDG PET scan. (**E**) Haematoxylin and eosin staining (HE) image. Adenocarcinoma is not visible on this staining. (**F**) Immunohistochemistry staining of PSMA. The arrow points towards stained PSMA. CT, computed tomography; MRI, magnetic resonance imaging; PET, positron emitting tomography; HE, haematoxylin–eosin; IHC, immunohistochemistry. (The "HE" image has previously been published by our study group [54] and is licensed under a Creative Commons Attribution 4.0 International—http://creativecommons.org/licenses/by/4.0, accessed on 1 June 2021).

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Abbreviations

ανβ6	integrin αvβ6
CT	computed tomography
CAF	cancer-associated fibroblast
CEA	carcinoembryonic antigen

DOTA	1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid
DNA	Deoxyribonucleic acid
ECM	extracellular matrix
EUS	endoscopic ultrasound
¹⁸ F	¹⁸ Fluorodeoxyglucose-labelled
FAP	fibroblast activation protein
FAPI	fibroblast activation protein inhibitor
¹⁸ F-DCFPyL	¹⁸ Fluorodeoxyglucose-labelled PSMA ligand
¹⁸ F-FDG	¹⁸ Fluorodeoxyglucose-labelled fluorodeoxyglucose
¹⁸ F-FDG-6-P	¹⁸ Fluorodeoxyglucose-labelled FDG -6-phosphate
¹⁸ F-FLT	¹⁸ Fluorodeoxyglucose-labelled fluorothymidine
¹⁸ F-FLTDP	¹⁸ Fluorodeoxyglucose-labelled FLT diphosphate
¹⁸ F-FLTMP	¹⁸ Fluorodeoxyglucose-labelled FLT monophosphate
¹⁸ F-FLTTP	¹⁸ Fluorodeoxyglucose-labelled FLT triphosphate
¹⁸ F-FP-R01-MG-F2	¹⁸ Fluorodeoxyglucose-labelled integrin tracer
¹⁸ F-PSMA	¹⁸ Fluorodeoxyglucose-labelled PSMA
⁶⁸ Ga	⁶⁸ Gallium-labelled
⁶⁸ Ga-FAPI	⁶⁸ Gallium-labelled fibroblast activation protein inhibitor
⁶⁸ Ga-DOTA-5G	⁶⁸ Gallium-labelled—DOTA—5G
68C DOTA FADI 04	⁶⁸ Ga-labelled 1,4,7,10-tetraazacyclododecane-
Ga-DOTA-FAPI-04	1,4,7,10-tetraacetic acid-FAPI-04
⁶⁸ Ga-NODAGA-R01-MG	⁶⁸ Ga-labelled integrin tracer
⁶⁸ Ga-PSMA	⁶⁸ Ga-labelled PSMA
⁶⁸ Ga-Trivehexin	⁶⁸ Gallium-labelled Trivehexin
Glucose-6-P	Glucose-6-Phosphate
GLUT-1	glucose transporter type 1
HACA	anti-chimeric antibodies
HASA	anti-sheep antibodies
HE	haematoxylin eosin
IHC	immunohistochemistry
¹³¹ I-KAb201	¹³¹ Iodine-labelled KAb201 antibody
IPMN	intraductal papillary mucinous neoplasm
KRAS	Kirsten rat sarcoma virus
LCA	lobulocentric atrophy
¹⁷⁷ Lu-DOTA-ABM-5G	¹⁷⁷ labelled Lutetium—DOTA-ABM-5G
MRI	magnetic resonance imaging
SUV	standardised uptake values
PanIN	pancreatic intraepithelial neoplasia
PDAC	pancreatic ductal adenocarcinoma
PET	positron emission tomography
PSMA	prostate-specific membrane antigen
TDP	thymidine diphosphate
TMP	thymidine monophosphate
TTP	thymidine triphosphate
⁹⁰ Y-hPAM4	⁹⁰ Yttrium-labelled clivazutumab tetraxetan antibody

References

- 1. Vincent, A.; Herman, J.; Schulick, R.; Hruban, R.H.; Goggins, M. Pancreatic cancer. Lancet 2011, 378, 607–620. [CrossRef]
- Latenstein, A.E.; Van Der Geest, L.G.; Bonsing, B.A.; Koerkamp, B.G.; Mohammad, N.H.; De Hingh, I.H.; De Meijer, V.E.; Molenaar, I.Q.; Van Santvoort, H.C.; Van Tienhoven, G.; et al. Nationwide trends in incidence, treatment and survival of pancreatic ductal adenocarcinoma. *Eur. J. Cancer* 2020, 125, 83–93. [CrossRef] [PubMed]
- Gerritsen, A.; Dutch Pancreatic Cancer Group; Molenaar, I.Q.; Bollen, T.L.; Nio, C.Y.; Dijkgraaf, M.G.; Van Santvoort, H.C.; Offerhaus, G.J.; Brosens, L.A.; Biermann, K.; et al. Preoperative Characteristics of Patients with Presumed Pancreatic Cancer but Ultimately Benign Disease: A Multicenter Series of 344 Pancreatoduodenectomies. *Ann. Surg. Oncol.* 2014, *21*, 3999–4006. [CrossRef]
- Van Veldhuisen, E.; van den Oord, C.; Brada, L.J.; Walma, M.S.; Vogel, J.A.; Wilmink, J.W.; Del Chiaro, M.; Van Lienden, K.P.; Meijerink, M.R.; Van Tienhoven, G.; et al. Locally advanced pancreatic cancer: Work-up, staging, and local intervention strategies. *Cancers* 2019, 11, 976. [CrossRef] [PubMed]

- Orlando, L.A.; Kulasingam, S.L.; Matchar, D. Meta-analysis: The detection of pancreatic malignancy with positron emission tomography. *Aliment. Pharmacol. Ther.* 2004, 20, 1063–1070. [CrossRef]
- Montemagno, C.; Cassim, S.; De Leiris, N.; Durivault, J.; Faraggi, M.; Pagès, G. Pancreatic Ductal Adenocarcinoma: The Dawn of the Era of Nuclear Medicine? *Int. J. Mol. Sci.* 2021, 22, 6413. [CrossRef] [PubMed]
- Ren, H.; Zhang, H.; Wang, X.; Liu, J.; Yuan, Z.; Hao, J. Prostate-specific membrane antigen as a marker of pancreatic cancer cells. *Med. Oncol.* 2014, *31*, 1–6. [CrossRef]
- Petrone, M.C.; Arcidiacono, P.G. New strategies for the early detection of pancreatic cancer. *Expert Rev. Gastroenterol. Hepatol.* 2016, 10, 157–159. [CrossRef] [PubMed]
- Canto, M.I.; Goggins, M.; Hruban, R.H.; Petersen, G.M.; Giardiello, F.M.; Yeo, C.; Fishman, E.K.; Brune, K.; Axilbund, J.; Griffin, C.; et al. Screening for Early Pancreatic Neoplasia in High-Risk Individuals: A Prospective Controlled Study. *Clin. Gastroenterol. Hepatol.* 2006, 4, 766–781. [CrossRef]
- 10. Maire, F.; Couvelard, A.; Palazzo, L.; Aubert, A.; Vullierme, M.-P.; Rebours, V.; Hammel, P.; Sauvanet, A.; Levy, P.; Ruszniewski, P. Author Index for American Pancreatic Association Abstracts: Pancreas. *Pancreas* **2013**, *42*, 1392–1394. [CrossRef]
- 11. Del Chiaro, M.; Verbeke, C.; Salvia, R.; Klöppel, G.; Werner, J.; McKay, C.; Friess, H.; Manfredi, R.; Van Cutsem, E.; Löhr, M.; et al. European experts consensus statement on cystic tumours of the pancreas. *Dig. Liver Dis.* **2013**, *45*, 703–711. [CrossRef]
- Tanaka, M.; Fernández-del Castillo, C.; Adsay, V.; Chari, S.; Falconi, M.; Jang, J.-Y.; Kimura, W.; Levy, P.; Pitman, M.B.; Schmidt, C.M.; et al. International consensus guidelines 2012 for the management of IPMN and MCN of the pancreas. *Pancreatology* 2012, 12, 183–197. [CrossRef]
- Del Chiaro, M.; Segersvärd, R.; Löhr, M.; Verbeke, C. Early detection and prevention of pancreatic cancer: Is it really possible today? World J. Gastroenterol. 2014, 20, 12118–12131. [CrossRef] [PubMed]
- 14. Tummers, W.S.; Willmann, J.K.; Bonsing, B.A.; Vahrmeijer, A.L.; Gambhir, S.S.; Swijnenburg, R.-J. Advances in Diagnostic and Intraoperative Molecular Imaging of Pancreatic Cancer. *Pancreas* **2018**, *47*, 675–689. [CrossRef]
- Conroy, T.; Desseigne, F.; Ychou, M.; Bouché, O.; Guimbaud, R.; Bécouarn, Y.; Adenis, A.; Raoul, J.-L.; Gourgou-Bourgade, S.; De La Fouchardière, C.; et al. FOLFIRINOX versus Gemcitabine for Metastatic Pancreatic Cancer. *N. Engl. J. Med.* 2011, 364, 1817–1825. [CrossRef]
- 16. Ferrone, C.R.; Marchegiani, G.; Hong, T.S.; Ryan, D.P.; Deshpande, V.; McDonnell, E.I.; Sabbatino, F.; Santos, D.D.; Allen, J.N.; Blaszkowsky, L.S.; et al. Radiological and Surgical Implications of Neoadjuvant Treatment With FOLFIRINOX for Locally Advanced and Borderline Resectable Pancreatic Cancer. *Ann. Surg.* **2015**, *261*, 12–17. [CrossRef]
- 17. Panda, A.; Garg, I.; Johnson, G.B.; Truty, M.J.; Halfdanarson, T.R.; Goenka, A.H. Molecular radionuclide imaging of pancreatic neoplasms. *Lancet Gastroenterol. Hepatol.* 2019, 4, 559–570. [CrossRef]
- Ghaneh, P.; Hanson, R.; Titman, A.; Lancaster, G.; Plumpton, C.; Lloyd-Williams, H.; Yeo, S.T.; Edwards, R.T.; Johnson, C.; Abu Hilal, M.; et al. PET-PANC: Multicentre prospective diagnostic accuracy and health economic analysis study of the impact of combined modality 18fluorine-2-fluoro-2-deoxy-d-glucose positron emission tomography with computed tomography scanning in the diagnosis and management of pancreatic cancer. *Health Technol. Assess.* 2018, 22, 1–114. [CrossRef] [PubMed]
- Samuel, N.; Hudson, T.J. The molecular and cellular heterogeneity of pancreatic ductal adenocarcinoma. *Nat. Rev. Gastroenterol. Hepatol.* 2011, 9, 77–87. [CrossRef]
- Gambhir, S.S.; Czernin, J.; Schwimmer, J.; Silverman, D.H.; Coleman, R.E.; Phelps, M.E. A tabulated summary of the FDG PET literature. J. Nucl. Med. 2001, 42, 1S–935. [PubMed]
- 21. Yokoyama, Y.; Nagino, M.; Hiromatsu, T.; Yuasa, N.; Oda, K.; Arai, T.; Nishio, H.; Ebata, T.; Nimura, Y. Intense PET Signal in the Degenerative Necrosis Superimposed on Chronic Pancreatitis. *Pancreas* **2005**, *31*, 192–194. [CrossRef]
- Epelbaum, R.; Frenkel, A.; Haddad, R.; Sikorski, N.; Strauss, L.G.; Israel, O.; Dimitrakopoulou-Strauss, A. Tumor Aggressiveness and Patient Outcome in Cancer of the Pancreas Assessed by Dynamic 18F-FDG PET/CT. J. Nucl. Med. 2013, 54, 12–18. [CrossRef] [PubMed]
- Lee, J.W.; Kang, C.M.; Choi, H.J.; Lee, W.J.; Song, S.Y.; Lee, J.-H.; Lee, J.D. Prognostic Value of Metabolic Tumor Volume and Total Lesion Glycolysis on Preoperative 18F-FDG PET/CT in Patients with Pancreatic Cancer. J. Nucl. Med. 2014, 55, 898–904. [CrossRef] [PubMed]
- Kato, K.; Nihashi, T.; Ikeda, M.; Abe, S.; Iwano, S.; Itoh, S.; Shimamoto, K.; Naganawa, S. Limited Efficacy of 18F-FDG PET/CT for Differentiation Between Metastasis-Free Pancreatic Cancer and Mass-Forming Pancreatitis. *Clin. Nucl. Med.* 2013, 38, 417–421. [CrossRef] [PubMed]
- Sahani, D.V.; Bonaffini, P.A.; Catalano, O.A.; Guimaraes, A.R.; Blake, M.A. State-of-the-Art PET/CT of the Pancreas: Current Role and Emerging Indications. *Radiographics* 2012, 32, 1133–1158. [CrossRef]
- Chang, J.; Schomer, D.; Dragovich, T. Anatomical, Physiological, and Molecular Imaging for Pancreatic Cancer: Current Clinical Use and Future Implications. *BioMed Res. Int.* 2015, 2015, 1–10. [CrossRef] [PubMed]
- 27. Chalkidou, A.; Landau, D.; Odell, E.; Cornelius, V.; O'Doherty, M.; Marsden, P. Correlation between Ki-67 immunohistochemistry and 18F-Fluorothymidine uptake in patients with cancer: A systematic review and meta-analysis. *Eur. J. Cancer* **2012**, *48*, 3499–3513. [CrossRef]
- Röhrich, M.; Naumann, P.; Giesel, F.L.; Choyke, P.L.; Staudinger, F.; Wefers, A.; Liew, D.P.; Kratochwil, C.; Rathke, H.; Liermann, J.; et al. Impact of 68Ga-FAPI PET/CT Imaging on the Therapeutic Management of Primary and Recurrent Pancreatic Ductal Adenocarcinomas. J. Nucl. Med. 2021, 62, 779–786. [CrossRef]

- 29. Von Ahrens, D.; Bhagat, T.D.; Nagrath, D.; Maitra, A.; Verma, A. The role of stromal cancer-associated fibroblasts in pancreatic cancer. *J. Hematol. Oncol.* 2017, 10, 1–8. [CrossRef]
- 30. Sun, Q.; Zhang, B.; Hu, Q.; Qin, Y.; Xu, W.; Liu, W.; Yu, X.; Xu, J. The impact of cancer-associated fibroblasts on major hallmarks of pancreatic cancer. *Theranostics* 2018, *8*, 5072–5087. [CrossRef]
- 31. Kalluri, R. The biology and function of fibroblasts in cancer. Nat. Rev. Cancer 2016, 16, 582–598. [CrossRef] [PubMed]
- 32. Sahai, E.; Astsaturov, I.; Cukierman, E.; DeNardo, D.G.; Egeblad, M.; Evans, R.M.; Fearon, D.; Greten, F.R.; Hingorani, S.R.; Hunter, T.; et al. A framework for advancing our understanding of cancer-associated fibroblasts. *Nat. Rev. Cancer* 2020, *20*, 174–186. [CrossRef] [PubMed]
- McCarthy, J.B.; El-Ashry, D.; Turley, E.A. Hyaluronan, Cancer-Associated Fibroblasts and the Tumor Microenvironment in Malignant Progression. *Front. Cell Dev. Biol.* 2018, 6, 48. [CrossRef]
- 34. Chen, H.; Pang, Y.; Wu, J.; Zhao, L.; Hao, B.; Wu, J.; Wei, J.; Wu, S.; Zhao, L.; Luo, Z.; et al. Comparison of [68Ga] Ga-DOTA-FAPI-04 and [18F] FDG PET/CT for the diagnosis of primary and metastatic lesions in patients with various types of cancer. *Eur. J. Nucl. Med. Mol. Imaging* 2020, 47, 1820–1832. [CrossRef] [PubMed]
- 35. Luo, Y.; Pan, Q.; Zhang, W.; Li, F. Intense FAPI Uptake in Inflammation May Mask the Tumor Activity of Pancreatic Cancer in 68Ga-FAPI PET/CT. *Clin. Nucl. Med.* **2020**, *45*, 310–311. [CrossRef]
- 36. Luo, Y.; Pan, Q.; Zhang, W. IgG4-related disease revealed by 68Ga-FAPI and 18F-FDG PET/CT. *Eur. J. Nucl. Med. Mol. Imaging* 2019, 46, 2625–2626. [CrossRef]
- 37. Pan, Q.; Luo, Y.; Zhang, W. Recurrent Immunoglobulin G4–Related Disease Shown on 18F-FDG and 68Ga-FAPI PET/CT. *Clin. Nucl. Med.* **2020**, *45*, 312–313. [CrossRef]
- Ocean, A.J.; Guarino, M.J.; Pennington, K.L.; O'Neil, B.H.; Rocha Lima, C.S.; Bekaii-Saab, T.S.; Gulec, S.A.; Gold, D.V.; We-gener, W.A.; Goldenberg, D.M. Therapeutic effects of fractionated radioimmu-notherapy (RAIT) with clivatuzumab tetraxetan combined with low-dose gemcitabine (Gem) in advanced pancreatic cancer (APC). J. Clin. Oncol. 2010, 28, 7S. [CrossRef]
- Kimura, R.H.; Wang, L.; Shen, B.; Huo, L.; Tummers, W.; Filipp, F.V.; Guo, H.H.; Haywood, T.; Abou-Elkacem, L.; Baratto, L.; et al. Evaluation of integrin αvβ6 cystine knot PET tracers to detect cancer and idiopathic pulmonary fibrosis. *Nat. Commun.* 2019, 10, 1–18. [CrossRef]
- 40. Hynes, R.O. Integrins: Bidirectional, Allosteric Signaling Machines. Cell 2002, 110, 673–687. [CrossRef]
- 41. Desgrosellier, J.S.; Cheresh, D.A. Integrins in cancer: Biological implications and therapeutic opportunities. *Nat. Rev. Cancer* 2010, 10, 9–22. [CrossRef] [PubMed]
- Reader, C.S.; Vallath, S.; Steele, C.W.; Haider, S.; Brentnall, A.; Desai, A.; Moore, K.M.; Jamieson, N.; Chang, D.; Bailey, P.; et al. The integrin αvβ6 drives pancreatic cancer through diverse mechanisms and represents an effective target for therapy. *J. Pathol.* 2019, 249, 332–342. [CrossRef] [PubMed]
- 43. Bandyopadhyay, A. Defining the Role of Integrin αvβ6 in Cancer. Curr. Drug Targets 2009, 10, 645–652. [CrossRef] [PubMed]
- Sipos, B.; Hahn, D.; Carceller, A.; Piulats, J.; Hedderich, J.; Kalthoff, H.; Goodman, S.L.; Kosmahl, M.; Klöppel, G. Immunohistochemical screening for beta6-integrin subunit expression in adenocarcinomas using a novel monoclonal antibody reveals strong up-regulation in pancreatic ductal adenocarcinomas in vivo and in vitro. *Histopathology* 2004, 45, 226–236. [CrossRef]
- Vachiranubhap, B.; Kim, Y.H.; Balci, N.C.; Semelka, R.C. Magnetic Resonance Imaging of Adenocarcinoma of the Pancreas. *Top. Magn. Reson. Imaging* 2009, 20, 3–9. [CrossRef] [PubMed]
- 46. Hausner, S.H.; DiCara, D.; Marik, J.; Marshall, J.F.; Sutcliffe, J.L. Use of a Peptide Derived from Foot-and-Mouth Disease Virus for the Noninvasive Imaging of Human Cancer: Generation and Evaluation of 4-[18F] Fluorobenzoyl A20FMDV2 forIn vivoImaging of Integrin αvβ6Expression with Positron Emission Tomography. *Cancer Res.* 2007, 67, 7833–7840. [CrossRef] [PubMed]
- Kimura, R.H.; Teed, R.; Hackel, B.J.; Pysz, M.A.; Chuang, C.Z.; Sathirachinda, A.; Willmann, J.; Gambhir, S.S. Pharmacokinetically Stabilized Cystine Knot Peptides That Bind Alpha-v-Beta-6 Integrin with Single-Digit Nanomolar Affinities for Detection of Pancreatic Cancer. *Clin. Cancer Res.* 2012, *18*, 839–849. [CrossRef]
- Altmann, A.; Sauter, M.; Roesch, S.; Mier, W.; Warta, R.; Debus, J.; Dyckhoff, G.; Herold-Mende, C.; Haberkorn, U. Identification of a Novel ITGαvβ6-Binding Peptide Using Protein Separation and Phage Display. *Clin. Cancer Res.* 2017, 23, 4170–4180. [CrossRef]
- Quigley, N.G.; Steiger, K.; Hoberück, S.; Czech, N.; Zierke, M.A.; Kossatz, S.; Pretze, M.; Richter, F.; Weichert, W.; Pox, C.; et al. PET/CT imaging of head-and-neck and pancreatic cancer in humans by targeting the "Cancer Integrin" αvβ6 with Ga-68-Trivehexin. *Eur. J. Nucl. Med. Mol. Imaging* 2021, 1–12. [CrossRef] [PubMed]
- Ceci, F.; Oprea-Lager, D.E.; Emmett, L.; Adam, J.A.; Bomanji, J.; Czernin, J.; Eiber, M.; Haberkorn, U.; Hofman, M.S.; Hope, T.A.; et al. E-PSMA: The EANM standardized reporting guidelines v1.0 for PSMA-PET. *Eur. J. Nucl. Med. Mol. Imaging* 2021, 48, 1626–1638. [CrossRef]
- Stock, K.; Steinestel, K.; Wiesch, R.; Mikesch, J.-H.; Hansmeier, A.; Trautmann, M.; Beller, N.; Rehkämper, J.; Wardelmann, E.; Heitkötter, B.; et al. Neovascular Prostate-Specific Membrane Antigen Expression Is Associated with Improved Overall Survival under Palliative Chemotherapy in Patients with Pancreatic Ductal Adenocarcinoma. *BioMed Res. Int.* 2017, 2017, 1–8. [CrossRef]
- Chang, S.S.; Reuter, V.E.; Heston, W.D.; Bander, N.H.; Grauer, L.S.; Gaudin, P.B. Five different anti-prostate-specific membrane antigen (PSMA) antibodies confirm PSMA expression in tumor-associated neovasculature. *Cancer Res.* 1999, *59*, 3192–3198. [PubMed]
- 53. Sahbai, S.; Rieping, P.; Pfannenberg, C.; la Fougère, C.; Reimold, M. Pancreatic Ductal Adenocarcinoma with High Radiotracer Uptake in 68Ga–Prostate-Specific Membrane Antigen PET/CT. *Clin. Nucl. Med.* **2017**, *42*, 717–718. [CrossRef] [PubMed]

- 54. Vuijk, F.A.; De Muynck, L.D.A.N.; Franken, L.C.; Busch, O.R.; Wilmink, J.W.; Besselink, M.G.; Bonsing, B.A.; Bhairosingh, S.S.; Kuppen, P.J.K.; Mieog, J.S.D.; et al. Molecular targets for diagnostic and intraoperative imaging of pancreatic ductal adenocarcinoma after neoadjuvant FOLFIRINOX treatment. *Sci. Rep.* **2020**, *10*, 1–9. [CrossRef]
- Mottet, N.; van den Bergh, R.C.; Briers, E.; van den Broeck, T.; Cumberbatch, M.G.; De Santis, M.; Fanti, S.; Fossati, N.; Gandaglia, G.; Gillessen, S.; et al. EAU-EANM-ESTRO-ESUR-SIOG Guidelines on Prostate Cancer—2020 Update. Part 1: Screening, Diagnosis, and Local Treatment with Curative Intent. *Eur. Urol.* 2010, *79*, 243–262. [CrossRef]
- Jansen, B.H.; Cysouw, M.C.; Vis, A.N.; Van Moorselaar, R.J.; Voortman, J.; Bodar, Y.J.; Schober, P.R.; Hendrikse, N.H.; Hoekstra, O.S.; Boellaard, R.; et al. Repeatability of Quantitative 18F-DCFPyL PET/CT Measurements in Metastatic Prostate Cancer. J. Nucl. Med. 2020, 61, 1320–1325. [CrossRef]
- 57. Krishnaraju, V.S.; Kumar, R.; Mittal, B.R.; Sharma, V.; Singh, H.; Nada, R.; Bal, A.; Rohilla, M.; Singh, H.; Rana, S.S. Differentiating benign and malignant pancreatic masses: Ga-68 PSMA PET/CT as a new diagnostic avenue. *Eur. Radiol.* 2021, 31, 2199–2208. [CrossRef] [PubMed]
- Jansen, B.H.; Yaqub, M.; Voortman, J.; Cysouw, M.C.; Windhorst, A.D.; Schuit, R.C.; Kramer, G.M.; Eertwegh, A.J.V.D.; Schwarte, L.A.; Hendrikse, H.N.; et al. Simplified Methods for Quantification of 18F-DCFPyL Uptake in Patients with Prostate Cancer. J. Nucl. Med. 2019, 60, 1730–1735. [CrossRef] [PubMed]
- Rowe, S.P.; Gorin, M.A.; Allaf, M.E.; Pienta, K.J.; Tran, P.T.; Pomper, M.G.; Ross, A.E.; Cho, S.Y. PET imaging of prostate-specific membrane antigen in prostate cancer: Current state of the art and future challenges. *Prostate Cancer Prostatic Dis.* 2016, 19, 223–230. [CrossRef]
- 60. Aigner, K.R.; Gailhofer, S.; Kopp, S. Regional versus systemic chemotherapy for advanced pancreatic cancer: A randomized study. *Hepatogastroenterology* **1998**, *45*, 1125–1129. [PubMed]
- 61. Shah, M.; Da Silva, R.; Gravekamp, C.; Libutti, S.K.; Abraham, T.; Dadachova, E. Targeted radionuclide therapies for pancreatic cancer. *Cancer Gene Ther.* 2015, 22, 375–379. [CrossRef]
- 62. Maitra, A.; Hruban, R.H. Pancreatic Cancer. Annu. Rev. Pathol. Mech. Dis. 2008, 3, 157–188. [CrossRef] [PubMed]
- 63. Moore, M.J.; Goldstein, D.; Hamm, J.; Figer, A.; Hecht, J.R.; Gallinger, S.; Au, H.J.; Murawa, P.; Walde, D.; Wolff, R.A.; et al. Erlotinib Plus Gemcitabine Compared With Gemcitabine Alone in Patients With Advanced Pancreatic Cancer: A Phase III Trial of the National Cancer Institute of Canada Clinical Trials Group. *J. Clin. Oncol.* **2007**, *25*, 1960–1966. [CrossRef]
- 64. Kulke, M.H.; Blaszkowsky, L.S.; Ryan, D.P.; Clark, J.W.; Meyerhardt, J.A.; Zhu, A.X.; Enzinger, P.C.; Kwak, E.L.; Muzikansky, A.; Lawrence, C.; et al. Capecitabine Plus Erlotinib in Gemcitabine-Refractory Advanced Pancreatic Cancer. *J. Clin. Oncol.* 2007, 25, 4787–4792. [CrossRef] [PubMed]
- Morgan, M.A.; Parsels, L.A.; Maybaum, J.; Lawrence, T.S. Improving Gemcitabine-Mediated Radiosensitization Using Molecularly Targeted Therapy: A Review. *Clin. Cancer Res.* 2008, 14, 6744–6750. [CrossRef]
- Pauwels, B.; Korst, A.E.; Lardon, F.; Vermorken, J.B. Combined Modality Therapy of Gemcitabine and Radiation. Oncology 2005, 10, 34–51. [CrossRef]
- 67. Available online: https://clinicaltrials.gov/ct2/show/NCT04665947 (accessed on 1 June 2021).
- Sultana, A.; Shore, S.; Raraty, M.G.; Vinjamuri, S.; Evans, J.E.; Smith, C.T.; Lane, S.; Chauhan, S.; Bosonnet, L.; Garvey, C.; et al. Randomised Phase I/II trial assessing the safety and efficacy of radiolabelled anti-carcinoembryonic antigen I131KAb201 antibodies given intra-arterially or intravenously in patients with unresectable pancreatic adenocarcinoma. *BMC Cancer* 2009, *9*, 66. [CrossRef]
- 69. Adams, G.P.; Weiner, L.M. Monoclonal antibody therapy of cancer. Nat. Biotechnol. 2005, 23, 1147–1157. [CrossRef] [PubMed]
- 70. Cardillo, T.M.; Ying, Z.; Gold, D.V. Therapeutic advantage of (90)yttrium-versus (131)iodine-labeled PAM4 anti-body in experimental pancreatic cancer. *Clin. Cancer Res.* **2001**, *7*, 3186.
- Berlin, J.D.; Catalano, P.; Thomas, J.P.; Kugler, J.W.; Haller, D.G.; Benson, A.B. Phase III Study of Gemcitabine in Combination With Fluorouracil Versus Gemcitabine Alone in Patients With Advanced Pancreatic Carcinoma: Eastern Cooperative Oncology Group Trial E. J. Clin. Oncol. 2002, 20, 3270–3275. [CrossRef]
- 72. Colucci, G.; Giuliani, F.; Gebbia, V.; Biglietto, M.; Rabitti, P.; Uomo, G.; Cigolari, S.; Testa, A.; Maiello, E.; Lopez, M. Gem-citabine alone or with cisplatin for the treatment of patients with locally advanced and/or metastatic pancre-atic carcinoma: A prospective, randomized phase III study of the Gruppo Oncologia dell'Italia Meridionale. *Cancer* **2002**, *94*, 902. [CrossRef] [PubMed]
- 73. Burris, H.A., III; Moore, M.J.; Andersen, J.; Green, M.R.; Rothenberg, M.L.; Modiano, M.R.; Cripps, M.C.; Portenoy, R.K.; Storniolo, A.M.; Tarassoff, P.; et al. Improvements in survival and clinical benefit with gemcitabine as first-line therapy for patients with advanced pancreas cancer: A randomized trial. *J. Clin. Oncol.* **1997**, *15*, 2403–2413. [CrossRef] [PubMed]
- 74. Gold, D.V.; Cardillo, T.; Vardi, Y.; Blumenthal, R. Radioimmunotherapy of experimental pancreatic cancer with 131I-labeled monoclonal antibody PAM. *Int. J. Cancer* **1997**, *71*, 660. [CrossRef] [PubMed]
- 75. Gold, D.V.; Karanjawala, Z.; Modrak, D.E.; Goldenberg, D.M.; Hruban, R.H. PAM4-Reactive MUC1 Is a Biomarker for Early Pancreatic Adenocarcinoma. *Clin. Cancer Res.* 2007, *13*, 7380–7387. [CrossRef]
- 76. Gold, D.V.; Lew, K.; Maliniak, R.; Hernandez, M.; Cardillo, T. Characterization of monoclonal antibody PAM4 reactive with a pancreatic cancer mucin. *Int. J. Cancer* **1994**, *57*, 204–210. [CrossRef] [PubMed]
- 77. Gold, D.V.; Modrak, D.E.; Ying, Z.; Cardillo, T.M.; Sharkey, R.M.; Goldenberg, D.M. New MUC1 Serum Immunoassay Differentiates Pancreatic Cancer From Pancreatitis. *J. Clin. Oncol.* **2006**, *24*, 252–258. [CrossRef] [PubMed]

- 78. Gold, D.V.; Goggins, M.; Modrak, D.E.; Newsome, G.; Liu, M.; Shi, C.; Hruban, R.H.; Goldenberg, D.M. Detection of Early-Stage Pancreatic Adenocarcinoma. *Cancer Epidemiol. Biomark. Prev.* **2010**, *19*, 2786–2794. [CrossRef]
- 79. Gulec, S.A.; Cohen, S.J.; Pennington, K.L.; Zuckier, L.S.; Hauke, R.J.; Horne, H.; Wegener, W.A.; Teoh, N.; Gold, D.V.; Sharkey, R.M.; et al. Treatment of Advanced Pancreatic Carcinoma with 90Y-Clivatuzumab Tetraxetan: A Phase I Single-Dose Escalation Trial. *Clin. Cancer Res.* 2011, *17*, 4091–4100. [CrossRef]
- 80. Ocean, A.J.; Pennington, K.L.; Guarino, M.J.; Sheikh, A.; Bekaii-Saab, T.; Serafini, A.N.; Lee, D.; Sung, M.W.; Gulec, S.A.; Goldsmith, S.J.; et al. Fractionated radioimmunotherapy with90Y-clivatuzumab tetraxetan and low-dose gemcitabine is active in advanced pancreatic cancer: A Phase 1 Trial. *Cancer* 2012, *118*, 5497–5506. [CrossRef]
- 81. Available online: https://clinicaltrials.gov/ct2/show/NCT01956812 (accessed on 1 June 2021).