Review

Radiolabeled Bombesin Analogs

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Simple Summary: Recent medical advancements have strived for a personalized medicine approach to patients, aimed at optimizing therapy outcomes with minimum toxicity. In this respect, nuclear medicine methodologies have been playing increasingly important roles. For example, the overexpression of peptide receptors, such as the gastrin-releasing peptide receptor (GRPR), on tumor cells as opposed to their lack of expression in healthy surrounding tissues can be elegantly exploited with the aid of “smart” peptide carriers, such as the analogs of the amphibian 14-peptide bombesin (BBN). These molecules can bring clinically attractive radionuclides to malignant lesions in prostate, breast, and other human cancers, sparing healthy tissues. Depending upon the radionuclide in question, diagnostic imaging with single-photon emission computed tomography (SPECT) or positron emission tomography (PET) has been pursued, identifying patients who are eligible for peptide radionuclide receptor therapy (PRRT) in an integrated “theranostic” approach. In the present review, we (i) discuss the major steps taken in the development of anti-GRPR theranostic radioligands, with a focus on those selected for clinical testing; (ii) comment on the present status in this field of research; and (iii) reflect on the current limitations as well as on new opportunities for their broader and more successful clinical applications.

Abstract: The gastrin-releasing peptide receptor (GRPR) is expressed in high numbers in a variety of human tumors, including the frequently occurring prostate and breast cancers, and therefore provides the rationale for directing diagnostic or therapeutic radionuclides on cancer lesions after administration of anti-GRPR peptide analogs. This concept has been initially explored with analogs of the frog 14-peptide bombesin, suitably modified at the N-terminus with a number of radiometal chelates. Radiotracers that were selected for clinical testing revealed inherent problems associated with these GRPR agonists, related to low metabolic stability, unfavorable abdominal accumulation, and adverse effects. A shift toward GRPR antagonists soon followed, with safer analogs becoming available, whereby, metabolic stability and background clearance issues were gradually improved. Clinical testing of three main major antagonist types led to promising outcomes, but at the same time brought to light several limitations of this concept, partly related to the variation of GRPR expression levels across cancer types, stages, previous treatments, and other factors. Currently, these parameters are being rigorously addressed by cell biologists, chemists, nuclear medicine physicians, and other discipline practitioners in a common effort to make available more effective and safe state-of-the-art molecular tools to combat GRPR-positive tumors. In the present review, we present the background, current status, and future perspectives of this endeavor.

Keywords: gastrin-releasing peptide receptor; radiolabeled bombesin; cancer theranostics
1. Introduction

The latest trends in medicine involve a personalized approach to patients. Such a patient-tailored approach can be well served by nuclear medicine techniques in oncological patients, following the so-called “theranostic” concept. This concept can be well exemplified in the case of bombesin and its analogs. Bombesin (BBN) is a 14-peptide (Figure 1) first isolated from the skin of the European fire-bellied toad Bombina bombina and exerting a series of actions on mammalian tissues through binding to distinct G-protein coupled receptors (GPCRs), known as bombesin receptors (BB-Rs) [1,2]. Interestingly, a number of BBN-like peptides have been identified in humans: the 27-mer gastrin-releasing peptide (GRP) and two other C-terminal decapeptide versions, namely neuromedin B (NMB) and neuromedin C (NMC, GRP(18–7)), all with a high sequence homology (Figure 1). The bombesin receptor family comprises three receptor subtypes in mammals, which are pharmacologically distinguished by their distinct affinities to native peptides. BB1-R, also known as NMBR, shows high affinity for NMB; BB2-R, also known as GRPR, strongly binds GRP and NMC; BB3-R is an orphan receptor with no native ligand identified thus far; BBN shows high binding affinities for both NMBR and GRPR, but not for BB3-R.

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Figure 1. Amino acid sequence of frog BBN and the respective GRP, NMC, and NMB peptides, native in humans; conserved residues are indicated by red color.

The GRPR, in particular, has attracted much attention in oncology and in nuclear medicine by virtue of its high-density expression in major human cancers, such as prostate [3–5], breast [6–9], and small-cell lung cancer [10], as well as in gastrinoma, gastrointestinal stromal tumors [11,12], and other cancer types [13]. This finding in combination with the lack of physiological GRPR expression in healthy tissues (with the exception of the GRPR-rich pancreas and gastrointestinal tract [14]) has provided the rationale for using properly designed BBN-like peptide carriers of radionuclides in malignant lesions for cancer theranostics [2,15,16]. BBN and/or GRP, but mostly their C-terminal fragments that still retain affinity for the GRPR, have been employed as motifs for anti-GRPR radioligand development. For such a purpose, suitable radiometal-binding chelators are covalently attached in their N-terminus, either directly or via a linker, and “theranostic radiopeptide pairs” may become available. Accordingly, the peptide conjugate is used for labeling with diagnostic gamma emitting radiometals (e.g., Tc-99m and In-111) for single-photon emission computed tomography (SPECT), or a positron emitter (e.g., Ga-68 and Cu-64) for positron emitter tomography (PET) imaging. The imaging will elucidate disease spread and monitor disease progress as well as post-therapy responses. Moreover, the imaging will identify patients eligible for peptide receptor radionuclide therapy (PRRT), using the same peptide conjugate labeled, instead, with a particle emitter, either a beta (e.g., Lu-177 and Y-90) or an alpha (e.g., Ac-225 and Bi-213) emitter. The concept of combining diagnostic and therapeutic tools in the same vector has already been successfully applied in nuclear medicine in the case of somatostatin and represents a valid example of personalized and precision medicine [17,18].

In the present review, we briefly discuss the development of GRPR radioligands, from its early days to its present status, with an emphasis on analogs involved in translational studies in breast and prostate cancer. Keeping a strong focus on theranostics of GRPR-positive cancer, we limit this review to peptide analogs labeled with diagnostic
or therapeutic radiometals, excluding the F-18 analogs suitable for PET imaging only. Likewise, delivery platforms loaded with BBN-like domains, such as nanoparticles, are not included in this review. The shift from GRPR agonists to antagonists is more thoroughly discussed, in view of the plethora of preclinical and clinical studies on this topic. After briefly outlining the issue of metabolic stability, the new trends in GRPR-targeted cancer theranostics are described, revealing the challenges and new opportunities in the field. Of particular interest is the final part of this review, in which we present the views of a clinician on the limitations, strengths, and future opportunities in prostate cancer theranostics using GRPR-targeted radiolabeled bombesin analogs.

2. First BBN-like Radioligands in Nuclear Oncology: GRPR Agonists

Most efforts involving BBN-like radioligands have been initially directed to the diagnosis and treatment of prostate cancer, a major cause of death worldwide [19]. While GRPR is highly expressed in prostatic intraepithelial neoplasia, as well as primary and invasive prostatic carcinoma, it is practically absent in normal prostate tissue and in benign prostate hyperplasia [3,4]. Several authors have correlated the expression of the receptor with Gleason score, i.e., high GRPR expression at low Gleason scores and low expression at high Gleason scores [5,20–22]. Accordingly, the application of anti-GRPR radioligands may represent a valid complementary approach to prostate-specific membrane antigen (PSMA) targeting for diagnosis and treatment of prostate cancer [23]. As mentioned above, the first introduced BBN-like radioligands were based on native motifs, and hence possessed agonistic properties at the GRPR. This implies that they internalized into target cells after receptor binding. This feature was previously considered to be an important prerequisite for high and prolonged tumor uptake [15,18]. During this initial period, several SPECT and PET GRPR-directed agents were designed for tumor diagnosis with a strong focus on the choice of linker and chelator to optimize pharmacokinetics, and a few of them were tested in patients (Table 1). It should be noted that information on peptide doses used in these studies, along with dosimetry data (where available) are not included or discussed herein, and the reader is advised to seek more detailed information in the references provided. The same holds true for Table 2 (vide infra).

Table 1. Bombesin-based agonists tested in the clinic. BBN(7–14), H-Gln-Trp-Ala-Val-Gly-His-Leu-Met-NH₂.

| BBN Analog | Radioconjugate Sequence | Clinical Indication | Ref. |
|------------|-------------------------|---------------------|-----|
| SPECT Tracers |
| [⁹⁹mTc]Tc-RP527 | [⁹⁹mTc]Tc-N₂S-Gly-5aVa⁴-BBN(7–14) | Metastasized PC⁵ or BC⁶ | [24] |
| [⁹⁹mTc]Tc-DB4 | [⁹⁹mTc]Tc-N₄-[Pro¹, Nle¹⁴]BBN | PC | [25] |
| PET Tracers |
| [⁶⁸Ga]Ga-AMBA | [⁶⁸Ga]Ga-DOTA-Gly-4-aminobenzyl-BBN(7–14) | Different malignancies | [26] |
| PRRT Agents |
| [¹⁷⁷Lu]Lu-AMBA | [¹⁷⁷Lu]Lu-DOTA-Gly-4-aminobenzyl-BBN(7–14) | Metastatic castration-resistant PC | [27] |

⁴5aVa, 5-aminovaleric acid; ⁵PC, prostate cancer; ⁶BC, breast cancer.
Table 2. GRPR-antagonist-based peptide radioligands tested in the clinic. BBN(7–14), H-Gln-Trp-Ala-Val-Gly-His-Leu-Met-NH₂.

| Radioligand | Radioconjugate Sequence | Clinical Indication | Ref. |
|-------------|-------------------------|---------------------|------|
| **SPECT Tracers** | | | |
| [99mTc]Tc-DB15 | [99mTc]Tc-N₃S-AMA¹-DGA³-[DPhe⁶,Sar¹⁹,Leu₁⁵,NH₂]BBN(6–13) | Advanced BC | [28] |
| [⁶⁸Ga]Ga-SB3 | [⁶⁸Ga]Ga-DOTA-AMA¹-DGA³-[DPhe⁶,Leu₁⁵,NH₂]BBN(6–13) | Disseminated PC | [21,29] |
| [⁶⁸Ga]Ga-RM2 | [⁶⁸Ga]Ga-DOTA-Pip⁶-[DPhe⁶,Sta₁⁵,Leu₁⁵,NH₂]BBN(6–14) | PC⁺, BC | [30,31] |
| [⁶⁸Ga]Ga-RM26 | [⁶⁸Ga]Ga-DOTA-AMA¹-DGA³-[DPhe⁶,Sta₁⁵,Leu₁⁵,NH₂]BBN(6–14) | PC⁺, BC | [32,33] |
| [⁶⁸Ga]Ga-NeoBOMB1 | [⁶⁸Ga]Ga-DOTA-AMA¹-DGA³-[DPhe⁶,Gln₂⁰,His₂⁵,NH₂]GRP(20–25)⁶ | Different malignancies | [34,35] |
| [⁶⁹Cu]Cu-CB-TE2A | [⁶⁹Cu]Cu-CB-TE2A-Peg⁴-[DPhe⁶,Sta₁⁵,Leu₁⁵,NH₂]BBN(6–14)⁶ | Newly diagnosed PC | [36] |
| **PET Tracers** | | | |
| [¹⁷⁷Lu]Lu-RM2 | [¹⁷⁷Lu]Lu-DOTA-Pip⁶-[DPhe⁶,Sta₁⁵,Leu₁⁵,NH₂]BBN(6–14) | Castration-resistant PC | [37] |
| [¹⁷⁷Lu]Lu-NeoBOMB1 | [¹⁷⁷Lu]Lu-DOTA-AMA¹-DGA³-[DPhe⁶,Gln₂⁰,His₂⁵,NH₂]GRP(20–25)⁶ | Advanced solid tumors | (NCT03872778) |

¹ AMA, p-aminomethylane; ²DGA, diglycolic acid; ³Pip, 4-amino-1-carboxymethyl-piperidine; ⁴R.-CH(CH₂)₂CH(CHOH)₂; ⁵PC, prostate cancer; ⁶CB-TE2A, 4,11-bis(carboxymethyl)-1,4,8,11-tetraazabicyclo(6.6.2)hexadecane; ⁷ BC, breast cancer.

One of the first SPECT tracers developed for such purposes was [⁹⁹ᵐTc]Tc-RP527 (RP527, N₃S-Gly-5-Ava-BBN(7–14)), whereby, the N₃S donor atom set, attached at the N-terminal via a Gly-5-aminovaleric acid linker, was used for binding Tc-99m under formation of a stable, neutral [⁹⁹ᵐTc]Tc⁵⁺O(N₃S) complex in square pyramidal configuration [38]. The tracer was assessed in a small cohort of prostate and breast cancer patients and successfully visualized four out of six breast cancers and one out of four androgen-resistant bone-metastasized prostate carcinomas [24]. Hence, the feasibility of GRPR-targeted imaging of human tumors was established for the first time, despite the shortcomings of the tracer with regards to pharmacokinetics.

Several more GRPR-specific tracers have been developed for SPECT imaging, but very few have been evaluated in the clinic. Suboptimal tumor uptake, in part as a result of poor metabolic stability, and excessive hepatobiliary excretion, due to high lipophilicity, have been the most frequent problems of these compounds [15]. Excretion of [⁹⁹ᵐTc]Tc⁶⁺ complexes rapidly through the kidneys into urine to minimize radioactivity levels in the abdomen could be promoted by using hydrophilic metal chelates. For such purposes, open chain tetraamines were soon coupled at the N-terminus via a Gly-5-aminovaleric acid linker, was used for binding Tc-99m under formation of a stable, neutral [⁹⁹ᵐTc]Tc⁶⁺O(N₃S) complex in square pyramidal configuration [38]. The tracer was assessed in a small cohort of prostate and breast cancer patients and successfully visualized four out of six breast cancers and one out of four androgen-resistant bone-metastasized prostate carcinomas [24]. Hence, the feasibility of GRPR-targeted imaging of human tumors was established for the first time, despite the shortcomings of the tracer with regards to pharmacokinetics.

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the α-therapy as compared with the β−-therapy and, among the two [213Bi]Bi-radioligands, a better safety profile for [213Bi]Bi-DOTA-PESIN than for [213Bi]Bi-AMBA.

Nevertheless, AMBA continues to be the bombesin analog most extensively tested in registered clinical trials, either radiolabeled with Ga-68 as a PET tracer, or radiolabeled with Lu-177 as radiotherapeutic treatment. Interestingly, in a small cohort of patients with different cancer types, [68Ga]Ga-AMBA was reported to be tolerated with minor adverse effects (abdominal discomfort and tachycardia), revealing several pathological lesions, especially in breast and prostate cancer patients, and displaying significant uptake mainly in the pancreas and intestinal tract as well as in the esophago-gastric junction [26]. However, a phase I escalation study conducted in patients with metastatic castration-resistant prostate cancer was discontinued due to the severe adverse effects induced by GRPR activation after injection of therapeutic doses of [177Lu]Lu-AMBA [27].

In conclusion, the application of radiolabeled BBN-like agonists in human, especially during PRRT requiring higher peptide doses administered to patients, has been linked to biosafety hurdles. Although receptor activation following agonist binding triggers internalization of the receptor-(radio)peptide complex into target cells, at the same time severe side effects can be released, predominantly in the gastrointestinal system [14,44,45]. Furthermore, bombesin and its analogs have been reported for their mitogenic actions [46,47]. Consequently, another paradigm was urgently needed for theranostic management of GRPR-positive cancers with this issue being promptly addressed by the advent of radiolabeled GRPR antagonists (vide infra).

3. From Agonists to Antagonists in GRPR-Targeting Radioligand Development

Several GRPR antagonists have been developed in previous years in order to better understand the pharmacology and roles of bombesin and its receptors in various physiological and pathophysiological conditions. The growth stimulatory effects of bombesin agonists in several human cancers have triggered systematic efforts to develop GRPR antagonists as antitumoral drugs, capable of binding to the receptor with a high affinity without it being activated [48–50]. As summarized in Figure 2, antagonists are not expected to elicit any adverse effects after injection in humans, possessing better biosafety [20,51].

![Figure 2](image-url)

**Figure 2.** After binding, agonists activate the GRPR on the target-cell membrane, whereas antagonists do not. Antagonists compete with agonists for GRPR binding, and thereby (partially) ban the agonist-induced activation of the GRPR.

Extensive structure-activity relationship studies have resulted in different classes of GRPR-antagonists, generated by modifying the peptide backbone of native BBN or GRP motifs, and especially the C-terminal Leu-Met-NH$_2$ dipeptide [1]. The most prominent modifications comprised truncation of the last C-terminal Met and alkyl-amidation or esterification of the exposed carboxy group of the penultimate residue, reduction of peptide bonds, substitution of key amino acids by their D-counterparts or other residues, all leading to a plethora of potent GRPR antagonists, displaying antiproliferative activity in GRPR-expressing cells as well as in GRPR-expressing tumors in mice [52,53]. This
rich library of well-characterized compounds has provided a multitude of motifs for the
development of radiolabeled GRPR antagonists (GRPR radioantagonists) for application in
nuclear medicine. Again, in addition to selecting the most promising peptide motif, other
modifications become crucial, such as the choice of the appropriate chelator and radiometal
for the intended application and the use or not of a suitable linker, for making available
new radioligands with high potential for clinical translation [51]. Representative families of
GRPR antagonist-based radioligands, classified according to the peptide motif with clinical
interest are discussed below, with relevant structures shown in Figure 3. A list of GRPR
antagonist-based radioligands tested in patients is included in Table 2.

Figure 3. Major BBN- and GRP-based motifs used in the GRPR antagonist design.

3.1. Radioligands Based on the GRPR Antagonist [DPhe6,Leu13-NHEt]BBN(6–13) Motif

Despite the increasing use of PET tracers in clinical practice, medical diagnostic imaging
techniques using [99mTc]Tc-based radiopharmaceuticals still account for approximately
80% of all nuclear medicine procedures. This radionuclide is widely available in phar-
aceutical grade in many hospitals via commercial [98Mo/99mTc]Tc-generators, in low-cost
and high specific activity and has ideal physical properties with a half-life of 6 h, an
optimal energy of 140 keV for imaging with currently used instrumentation combined
with low radiation exposure of patients and personnel [39]. Therefore, it is not surpris-
ing that the first attempts were devoted to obtaining GRPR antagonists, radiolabeled
with Tc-99m and suitable for SPECT imaging. The potent GRPR antagonist [DPhe6,Leu13,
NHEt]BBN(6–13), obtained by ethylamidation of C-terminal Leu13 after removal of the
last amino acid ([des-Met14]-class of GRPR antagonists) [54] was conjugated to an acyclic
tetraamine chelator via the p-aminomethylamine-diglycolic acid (AMA-DGA) linker. The
forming radioligand [99mTc]Tc-Demobesin 1 ([99mTc]Tc-DB1) was the first radiolabeled
GRPR antagonist tested in PC-3 tumor-bearing mice showing excellent and specific recep-
tor targeting properties, combining high accumulation in the experimental tumor with a
fast background clearance via the kidneys into urine. Rapid washout was observed even
from GRPR-rich organs, such as the pancreas, resulting in improved tumor-to-background
ratios over time, a feature characteristic for most GRPR radioantagonists thereafter [55]. In
a following side-by-side comparison of [99mTc]Tc-DB1 with the agonist-based [99mTc]Tc-
DB4, [99mTc]Tc-DB1, despite the lack of internalization, achieved higher tumor uptake and
faster washout from GRPR-positive tissues [56]. It should be noted that efforts to prolong
tumor retention involved the introduction of different linkers and amino acid replacements,
eventually revealing the Sar11-analog [99mTc]Tc-DB15 as a promising candidate for clinical
translation [28,57]. The first results in patients with advanced breast cancer revealed an
excellent pharmacokinetic profile of [99mTc]Tc-DB15, whereby, the tracer visualized several
bone and soft tissue metastases on SPECT/CT, while being well tolerated [28].
With the aim to obtain a theranostic tool labeled with either diagnostic or therapeutic radiometals, the above GRPR antagonist was coupled to the universal chelator DOTA, affording SB3 (SB3, DOTA-AMA-DGA-[DPhe\textsuperscript{6},Leu\textsuperscript{13}-NHEt]BBN(6–13), Table 2). During preclinical testing, in tumor-bearing mice, \[^{67}\text{Ga}]\text{Ga-SB3}, serving as a surrogate of \[^{68}\text{Ga}]\text{Ga-SB3}, compared well with \[^{99m}\text{Tc}]\text{Tc-DB1}, with regards to tumor uptake and retention and background clearance. \[^{68}\text{Ga}]\text{Ga-SB3} was selected for clinical testing in a small cohort of patients with disseminated prostate and breast cancers, whereby it was able to visualize lesions in about 50% of the patients on PET/CT [29]. A much higher detection accuracy was subsequently documented in therapy-naïve primary prostate cancer patients. Furthermore, an excellent correlation could be established between imaging findings and GRPR expression levels by histopathology of excised lesions [21]. Unfortunately, both the SPECT \[^{111}\text{In}]\text{In-SB3} counterpart and the therapeutic \[^{177}\text{Lu}]\text{Lu-SB3} version were shown to be rapidly catabolized in vivo by neutral endopeptidase. This handicap, compromising their clinical translation perspectives, could be, however, overcome by means of specific inhibitors (vide infra) [58]. Indeed, \[^{111}\text{In}]\text{In-SB3} administered in combination with such an inhibitor in mice, successfully visualized PC-3 tumors using SPECT/MRI, revealing its potential for preoperative imaging of prostate cancer [59].

3.2. Radioligands Based on the GRPR Antagonist \([\text{DPhe}^6,\text{Sta}^{13},\text{Leu}^{14}\text{NH}_2]\text{BBN(6–14)}\) Motif

Replacement of the C-terminal Leu\textsuperscript{13}-Met\textsuperscript{14}-NH\textsubscript{2} by the dipeptide Sta\textsuperscript{13}-Leu\textsuperscript{14}-NH\textsubscript{2} (Sta or statine, [3S, 4S]-4-amino-3-hydroxy-6-methylheptanoic acid) in the truncated BBN(7–14) motif, led to the potent “statine-based” GRPR antagonist JMV594 [60]. A variety of chelators and linkers have been introduced at its N-terminal to modulate the receptor binding affinities, metabolic stability, hydrophilicity, and hence the endopharmacokinetics of resulting GRPR radioantagonists. Initially, \[^{111}\text{In}]\text{In-RM1} (RM1, DOTA-Gly-4-aminobenzoyl-JMV594) was developed with DOTA coupled to JMV594 via the same linker used in \[^{111}\text{In}]\text{In-AMBA} to allow for a side-by-side comparison [61]. The antagonist showed higher accumulation in PC-3 xenografts and faster washout from the GRPR-positive organs in mice as compared with the agonist, confirming the superior performance of the GRPR antagonist. Intensive structure-activity studies have led to the development of a considerable number of radiolabeled JMV594 derivatives.

A positive charge at the N-terminus, introduced either by a positively charged linker or by the radiometal chelate per se, turned out to favor GRPR affinity as well as in vivo pharmacokinetics of resulting radioligands [62–65]. Thus, by using a positively charged 4-amino-1-carboxymethyl-piperidine (Pip) linker, instead, RM2 (DOTA-Pip-[DPhe\textsuperscript{6},\text{Sta}\textsuperscript{13},\text{Leu}\textsuperscript{14}, \text{NH}_2]BBN(6–14), Table 2) was suitable for labeling with Ga-68. \[^{68}\text{Ga}]\text{Ga-RM2} has been widely studied in different prostate cancer cell lines and tumor models and has shown improved binding affinity to the GRPR and metabolic stability [63]. \[^{68}\text{Ga}]\text{Ga-RM2} was the most clinically tested PET tracer based on this class of GRPR antagonists, studied in a small cohort of healthy volunteers [66], breast [30], and prostate cancer patients [31]. In these two latter studies, \[^{68}\text{Ga}]\text{Ga-RM2} was well tolerated by all patients and showed high sensitivity, specificity, and accuracy for the detection of primary cancer. Its feasibility for detecting most metastatic lesions highlights that it has high potential as a diagnostic PET tracer for GRPR-positive tumors. In another approach, NOTA (1,4,7-triazacyclononane-1,4,7-triacetic acid) was attached at the N-terminus of JMV594 either directly or via a PEGx linker, allowing labeling with Ga-68 and the formation of the PET tracer \[^{68}\text{Ga}]\text{Ga-RM26} (RM26, NOTA-PEG\textsubscript{x}-[DPhe\textsuperscript{6},\text{Sta}\textsuperscript{13},\text{Leu}\textsuperscript{14},\text{NH}_2]BBN(6–14), Table 2) [67]. The tracer successfully (85\%) visualized pathological lesions in breast cancer patients, showing significantly more intensive uptake in estrogen receptor (ER+) patients. Furthermore, an excellent correlation could be established between lesion uptake and GRPR expression status [32]. In patients with primary prostate cancer, the tracer detected prostate confined pathological lesions with an excellent correlation with GRPR expression status in the excised specimens. The tracer was also able to visualize several bone and lymph node metastases in patients with recurring disease [33].
Recently, the therapeutic counterpart $^{[177]}$Lu-Lu-RM2 has been studied in patients with metastatic castration-resistant prostate cancer, showing promising features for clinical use. Thus, absorbed doses in tumor lesions were found to be therapeutically relevant, whereas rapid clearance from normal GRPR-rich organs was confirmed, such as the pancreas, considered to be the dose-limiting organ due to its high radioligand uptake [37]. In addition, positively charged radiometal chelates, formed after stable binding of specific radiometals of clinical interest to chelators other than DOTA, were also considered [62,68]. Thus, in an interesting study, the GRPR antagonist AR (H-PEG$_4$-JM594) carrying four different radiometal chelates on its N-terminus ($[^{111}$In-DOTA, $[^{99m}$Tc]Tc-N$_4$, $[^{68}$Ga]Ga-NODAGA, and $[^{64}$Cu]Cu-CB-TE2A), confirmed the superiority of positively charged conjugates ($[^{99m}$Tc]Tc-N$_4$-AR and $[^{64}$Cu]Cu-CB-TE2A-AR) in terms of GRPR affinity and tumor uptake and retention. Notably, $[^{68}$Ga]Cu-CB-TE2A-AR (CB-TE2A, 4,11-bis(carboxymethyl)-1,4,8,11-tetraazabicyclo(6.6.2)hexadecane, Table 2) was finally chosen for further clinical evaluation also owing to the attractive nuclear properties of Cu-64. Indeed, the half-life of 12.7 h and its decay scheme render this theranostic radiometal an attractive alternative to the usually applied PET radionuclides F-18 and Ga-68 [69]. In a first proof-of-concept study in a small number of prostate cancer patients $[^{64}$Cu]Cu-CB-TE2A-AR showed long tumor retention and fast clearance from other organs [36].

3.3. Radioligands Based on the GRPR Antagonist [Ac-His$_{20}$,His$_{25}$-NHR]GRP(20–25) Motif

Alkylamide derivatives of the human [(N-acetyl)His$_{20}$,His$_{25}$-NHR,des-(Leu$_{26}$, Met$_{27}$)NH$_2$]GRP(20–27) motif (especially when NHR = NHCH(CH$_2$CH(CH$_3$)$_2$)$_2$) represent a successful class of very potent GRPR antagonists, characterized by high receptor affinity and in vivo enzymatic stability. Accordingly, the [DPhe$_{19}$,Gln$_{20}$,His$_{25}$-NHCH(CH$_2$CH(CH$_3$)$_2$)$_2$]GRP(20–25) sequence was modified by DOTA-AMA-DGA at the N-terminus affording NeoBOMB1. NeoBOMB1, labeled with Ga-68 (for PET), In-111 (for SPECT), or Lu-177 (for PRRT) showed promising theranostic profiles during preclinical evaluation, characterized by high receptor affinity and cell-uptake efficiency in vitro together with high metabolic stability and high and prolonged tumor uptake in GRPR-expressing tumors in vivo [34,70,71]. First clinical data with $[^{68}$Ga]Ga-NeoBOMB1 in a group of prostate cancer patients highlighted its ability to visualize primary tumors as well as liver metastases and bone lesions (example in Figure 4). These promising outcomes justifi- ed the assessment of the theranostic pair $[^{68}$Ga]Ga/$[^{177}$Lu]Lu-NebOMB1 in an ongoing multicenter clinical trial (NCT03872778) in patients with advanced solid tumors known to overexpress GRPR [35].

![Figure 4](image_url) Patient with PC, post radical prostatovesiculectomy with pelvic lymphadenectomy, IMRT, and androgen deprivation therapy (PSA on date of scan, 21.77 ng/mL). $[^{68}$Ga]Ga-NeoBOMB1 PET/CT: (A) PET MIP; (B) serial PET transverse; (C) corresponding CT transverse; (D) fusion PET/CT images. Multiple mediastinal, abdominal, paraoesophageal and pelvic lymph node metastases (arrows and crossbars). This research was originally published in JNM 2017 [34].
4. Metabolic Stability Issues

A major handicap in the development of peptide-based radioligands for cancer theranostics is the notorious susceptibility of peptides to proteolytic degradation [72–75]. Omnipresent peptidases, i.e., enzymes breaking down peptide bonds, can drastically reduce the number of intact radiopeptide molecules eventually reaching the tumor-situated target. As a result, tumor uptake drops and, consequently, diagnostic accuracy and therapeutic efficacy are compromised [76]. It should be noted that amongst the >550 proteases comprising the human protease degradome [77], the circulating radiopeptide will encounter quite a few to which it may potentially be a substrate on its quick trip to the target. Another point to keep in mind is that most peptide radioligands carry the radiometal chelate at the N-terminus (either directly or via a spacer), and therefore are resistant to the proteolytic action of ubiquitous N-aminopeptidases.

The performance of BBN-like radioligands is likewise affected by rapidly degrading peptidases in the blood stream (Figure 5a). Previous studies have revealed neprilysin (NEP) as the major player in the degradation of radiolabeled BBN-like analogs [76,78,79]. NEP is an ectoenzyme found in high local concentrations anchored on the membrane of epithelial cells of the vasculature and major organs of the body, such as the kidneys, the lungs, and the intestinal track. Typically, NEP hydrolyzes peptide bonds at the amino side of hydrophobic residues and shows a wide substrate repertoire, including BBN-like peptides [80,81]. Four cleavage sites have been, thus far, identified in the BBN(6/7–14) motif most often present in anti-GRPR radioligands, namely the His12-Leu13, Ala9-Val10, Trp8-Ala9, and Gln17-Trp8 bonds [58,79].

![Figure 5. (a) The radiolabeled peptide conjugate (based on a GRPR agonist or antagonist) after entering the blood (top) is attacked by NEP (indicated by the two descending green arrows) and is degraded. As a result, only a few of the initial molecules arrive intact at the tumor cells and interact with the GRPR (orange receptors) located at the cell membrane (bottom); (b) following co-administration of a NEP inhibitor, the degrading action of NEP is suspended, and thus the number of intact molecules arriving at the tumor and binding to GRPR sites is higher, leading to higher tumor uptake.](image-url)
In yet another promising approach, NEP inhibitor administration was proposed to ban the action of the major degrading protease (Figure 5b). The resulting in situ stabilization of circulating BBN-like radioligands maximized their delivery to tumor-associated GRPR sites, thereby improving tumor uptake. This strategy turned out to be particularly effective in enhancing the GRPR-targeting capabilities of biodegradable analogs without interfering with other important biological properties, such as cell uptake or pharmacokinetics, but unsurprisingly, was less evident in metabolically more robust analogs [58,76,82,88–92]. Although GRPR radioantagonists turned out to be typically more stable than their agonist-based counterparts, notable tumor uptake improvements could still be observed in most cases, providing new exciting prospects for better diagnosis and therapy [59,93]. In most early preclinical studies testing this approach, phosphoramidon (PA) was used as an NEP inhibitor [76,94,95]. Translation of this methodology in the clinic needs to overcome a series of safety and regulatory hurdles and may be facilitated by the availability of registered and orally administered NEP inhibitors used as anti-diarrhea (Hidrasec™, in vivo releasing the potent and selective NEP-inhibitor thiorphan) or as anti-hypertensive drugs (Entresto®, containing the prodrug sacubitril (AHU377), in vivo releasing the active substance sacubitrilat (LBQ657), a potent NEP inhibitor) [96–101].

5. Newer Trends in the Application of Anti-GRPR Theranostic Radiopeptides

In view of the above, it is fair to conclude that molecular imaging and PRRT of human GRPR-positive tumors has been gaining momentum in radiopharmacy and nuclear medicine research with a few tracers currently being evaluated by multicenter clinical trials for entering the clinical arena. Recent breakthroughs include, on the one hand, the advent of GRPR radioantagonists competently addressing biosafety concerns and, on the other hand, the application of clinically established protease inhibitors or the “smart” design of protease-resistant analogs to improve metabolic stability. Nevertheless, a few critical questions still need to be tackled before drastically improving the diagnostic accuracy and, most importantly, the therapeutic efficacy of this approach. Hence, it is essential to acquire a better understanding of the pathophysiology of GRPR-positive cancers and, especially, the dependence of GRPR expression density on disease type, stage, coexpression of other biomolecular targets, individual patient biochemical background, and previous or even concomitant therapies.

For example, an alternative biomolecular target in prostate cancer is PSMA, prompting the development of several radiolabeled PSMA inhibitors dynamically entering the field of nuclear medicine as novel diagnostic and therapeutic tools [102]. Heterogeneity of GRPR and/or PSMA expression in primary and metastatic cancer lesions through the various stages of the disease has been documented [9,22,51,103,104]. This finding has triggered the development of heterodimeric radioligands, which are able to target both the GRPR and PSMA in prostate cancer lesions, thereby, increasing diagnostic sensitivity and therapeutic efficacy [84,105–107]. Likewise, in another approach, the GRPR and integrin αVβ3 associated with tumor angiogenesis have been concomitantly targeted by a dual PET tracer [68Ga]Ga-BBN-RGD enhancing diagnostic accuracy in breast and prostate cancer patients [108,109].

How the above strategies applied in PET imaging can be adopted in PRRT remains to be explored, given that the issue of inadvertent accumulation of radioactivity in excretory organs or organs with high physiological GRPR expression levels (e.g., the pancreas) have to be competently addressed first. This issue becomes even more relevant in the case of PRRT with alpha emitters [43]. Higher tumor-to-background ratios have been achieved, in preclinical studies, with the application of higher peptide doses, an approach potentially applied in humans only for GRPR antagonists owing to biosafety concerns [70]. Interestingly, an extra benefit of administering higher peptide amounts was recently shown to favor penetration of radioactivity into the whole tumor mass in mice, thereby, improving therapeutic efficacy [110]. In another interesting approach depicted in Figure 6, a pretargeting strategy has been proposed to circumvent pancreatic uptake. Pretargeting strategies have been
applied with success to lower tumor-to-background radioactivity ratios of long-circulating high molecular weight molecules, for example, antibodies [111,112]. Preliminary studies in mice with derivatives of NeoBOMB1, carrying a clickable residue, have been applied in combination with the radiolabeled complementary small molecule [111In]In-Tz, and thus far, have failed to improve tumor-to-pancreas ratios, most likely due to undesired folding of the targeting vector in the binding pocket of the GRPR [113]. Therefore, targeting vectors with longer linkers are currently being developed and preclinically tested, aiming to prevent pancreatic radiotoxicity, and thus enabling higher radioactivity doses to the tumor.

Figure 6. The pretargeting concept for safe and effective GRPR-mediated radionuclide therapy: (a) The targeting vector adapted with a click moiety is injected first. It binds to the GRPR on the cells of the pancreas and on GRPR-expressing tumor cells. When the targeting vector is (mostly) cleared from the pancreas but still present on the tumor cells, a radiolabeled small molecule containing a complementary click element is injected. This radiolabeled small molecule couples the targeting vector still present on the tumor cells, hereby, specifically delivering radioactivity to these cells; (b) graph depicting uptake of the targeting vector and radioactivity/radiation dose to the pancreas and tumor using the pretargeting strategy.

Several other efforts have been directed to enhance the therapeutic efficacy of PRRT in GRPR-positive tumors via adjuvant therapeutic schemes. Thus, the efficacy of monotherapy with [177Lu]Lu-RM26 alone in mice models has been reported to further improve when combined with the mTOR inhibitor rapamycin [114]. Interestingly, mTOR inhibitors were reported to enhance the expression levels of other GPCRs in experimental tumors, revealing the need for systematic studies in this promising area of research [115]. The combination of PRRT and hormone therapy in breast and prostate cancer may provide another promising route toward higher therapeutic responses [116]. Immunotherapy may also enhance therapeutic efficacy of PRRT, as recently shown for the [177Lu]Lu-RM26 and trastuzumab combination in prostate cancer mice models [110]. In a recent innovative approach, prolonged tumor retention could be achieved in mice models when using GRPR peptide radioligands, either agonists or antagonists, modified with cysteine cathepsin inhibitors, via an endolysosomal trapping mechanism [117].

Last but not least, one should report recent breakthroughs in the production of clinically appealing therapeutic radionuclides, including the Tb and Sc therapeutic isotopes and alpha emitters (Ac-225 and Bi-213), dynamically entering radiopharmaceutical development, and expected to soon upgrade the arsenal of anti-GRPR therapeutics with new powerful molecular tools [118–120].

6. The Clinician’s View: GRPR-Targeted Theranostics with a Focus on Prostate Cancer

In the last decade, new developments in imaging of prostate cancer have had a major impact on the daily practice of prostate cancer specialists. To put things in perspective, it is important to realize that guidelines are still heavily based on results of two large
prostate cancer screening trials (ERSPC and PLCO), starting in the late 1980s with only PSA, rectal examination, low-resolution ultrasound, and six random biopsies as screening tools [121]. Two big game changers can be depicted that have significantly reformed the management of prostate cancer, i.e., MRI and (PSMA-) PET (or PET/CT), improving the average prognosis [122,123]. However, PSMA is not as specific as we wish and several other malignant, benign, and non-clinically relevant conditions might be mistaken for a prostate cancer lesion and can lead to false conclusions [124,125]. Furthermore, limitations of PSMA are also apparent with respect to the relative low sensitivity for early stage, low-grade tumors, or early recurrent disease, thereby impairing decision making in patients with oligo-metastatic disease, whereby the exact number and localization of metastatic lesions is of great importance [126]. Accordingly, PET imaging of other molecular targets, such as GRPR, may well contribute to this clinical need of refining decision making [104].

GRPR-targeted imaging may constitute a relevant addition for those patients with PSMA-negative tumors, estimated to constitute 74% of patients, or those with low-grade tumors that do not show on MRI nor PSMA PET scans [127]. Indeed, retrospective studies have revealed that PSMA expression was inversely correlated with GRPR, underscoring the potential value of their combined use [22]. Recent phase I studies have confirmed GRPR PET imaging to detect (primary) prostate tumors with high sensitivity and specificity comparable to those observed for PSMA [59]. Further reports have shown that PSMA and GRPR PET/CT may have added value in evaluating biochemical recurrence of prostate cancer [103,104]. In the direct comparison of [68Ga]Ga-RM2 and [68Ga]Ga-PSMA-11 on PET/CT in seven patients with biochemically recurring prostate cancer, shown in Figure 7, the heterogeneity of GRPR and PSMA expression was evident, highlighting the benefit of using both targets in a complementary fashion to increase diagnostic accuracy [103]. Promising new developments are directed towards bispecific radioligands of PSMA and GRPR to reach maximal detection rates and to capture all prostate cancer lesions [84,105–107].

![Figure 7. Maximum intensity projection [68Ga]Ga-RM2 and [68Ga]Ga-PSMA-11 images of the 7 enrolled prostate cancer patients. This research was originally published in JNM 2016 [103].](image)

Importantly, GRPR and PSMA expression are both affected by standard of care treatments applied in prostate cancer, including targeting the androgen receptor pathway and taxane-based chemotherapy [29]. Although reported data are contradictory on the effect size, this clearly constitutes a serious issue for clinicians. As a consequence of the multitude of options for treatment of disseminated prostate cancer [128], late-stage patients have highly variable treatment histories that jeopardize good clinical evaluation of the most optimal timing of GRPR and PSMA imaging (and therapy) in these patients.

In addition to their use in diagnostics, PSMA and GRPR hold promise for the application of radionuclide therapy in the management of prostate cancer. Clinical application
of this theranostic approach has just started, and reports are still scarce [129]. Although several spectacular cases have been presented with seemingly complete remissions after treatment with PSMA or GRPR, as well as promising data achieved in several retrospective studies, robust scientific proof of efficacy and long-term safety in randomized clinical trials are yet to be completed [37,130,131]. A meta-analysis of 12 studies in late-stage/end-stage patients failing on standard of care treatments, revealed transient effectiveness of PSMA radiotherapy, although overall survival benefit was not significantly improved as compared with any other third-line treatment [132–134]. From these studies, short-term side effects appear to be minimal for PSMA, but administration of these treatments will not be approved in an early metastasized stage until long term safety is scientifically proven. Likewise, GRPR expression is also not unique to the prostate nor to prostate cancer; the high absorbed pancreatic dose and diffuse low uptake throughout the gastrointestinal tract constitute a serious concern for therapy [21,135]. With new developments in alpha radionuclide therapy, toxicity in surrounding tissues will become even more critical. Despite major consequences of PSMA radiotherapy with loss of saliva, taste, speech, and dry eyes, \[^{225}\text{Ac}\]Ac-PSMA therapy is already being evaluated in the clinic with promising biochemical responses and low treatment-related toxicity [136]. Finally, tandem therapy with both \[^{177}\text{Lu}\]Lu- and \[^{225}\text{Ac}\]Ac-PSMA have been applied that suggest improving PSMA-radiotherapy response while minimizing toxicity of the salivary glands [133]. Alternatively, combination strategies of radiotherapy with both PSMA and GRPR are being considered as a solution to mitigate toxicity while optimizing antitumor efficacy [137].

The fast introduction of PSMA radiotracers for imaging and therapy has radically shifted the clinical perspectives of prostate cancer patients. But clearly, clinical evaluation has just started, and well-designed prospective randomized trials are needed to learn how to use these new applications in daily practice: (1) how to interpret PSMA PET/CT (e.g., the clinical relevance of lesions, how to deal with PSMA-negative tumors); (2) how to optimally position PSMA radiotherapy in the sequence of standard of care therapies; and most importantly, (3) how to alleviate radiotoxicity without losing therapeutic efficacy. GRPR-targeted radioligands may help to solve, at least in part, these clinical issues by complementing the PSMA PET/CT and radioligand therapy and improving the prostate cancer detection rate as well as therapeutic efficacy.

7. Conclusions

In view of the above, it becomes evident that research in the field of radiolabeled bombesin analogs for application in the theranostic management of GRPR-positive human tumors has been thriving with major breakthroughs already achieved. Firstly, the paradigm shift from agonist- to antagonist-based radioligands has competently addressed the issue of biosafety, a very crucial point during PRRT with higher peptide amounts administered to patients. Secondly, the challenge of rapid degradation of BBN-like tracers entering the circulation has been “elegantly” met with the application of NEP inhibitors. The latter may be directly applied for in situ stabilization of biodegradable radioligands, but at the same time have been instrumental in a smarter design of metabolically robust analogs. Extended and systematic structure-activity relationship studies have pinpointed candidates eligible for clinical translation in breast, prostate, and other GRPR-positive cancer patients, applying SPECT/CT, or PET/CT imaging and PRRT.

Such pioneer clinical studies have been eye-openers with regards to the strengths and limitations of GRPR-positive cancer theranostics, revealing new challenges to be met. Our incomplete understanding of cancer pathology needs to be improved in order to provide solid information on critical issues, such as GRPR expression status in different types and stages of cancer, as well as following previous therapies in cases of advanced disease. This critical knowledge will certainly provide new opportunities for the successful application of multitargeted theranostic agents and, most importantly, of adjuvant therapies. Accordingly, combining PRRT with the aid of anti-GRPR radioligands with other therapies, such as hormone therapy, immunotherapy, and other therapeutic schemes, should be adopted to
individual patients, aiming at maximum efficacy and low toxicity. The seeds for these innovative developments have already been planted and will most certainly lead to new exciting breakthroughs in the near future.

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