Theranostics in nuclear medicine practice

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Abstract: The importance of personalized medicine has been growing, mainly due to a more urgent need to avoid unnecessary and expensive treatments. In nuclear medicine, the theranostic approach is an established tool for specific molecular targeting, both for diagnostics and therapy. The visualization of potential targets can help predict if a patient will benefit from a particular treatment. Thanks to the quick development of radiopharmaceuticals and diagnostic techniques, the use of theranostic agents has been continually increasing. In this article, important milestones of nuclear therapies and diagnostics in the context of theranostics are highlighted. It begins with a well-known radioiodine therapy in patients with thyroid cancer and then progresses through various approaches for the treatment of advanced cancer with targeted therapies. The aim of this review was to provide a summary of background knowledge and current applications, and to identify the advantages of targeted therapies and imaging in nuclear medicine practices.

Keywords: theranostics, nuclear medicine, personalized medicine, PET/CT, therapy, diagnostics

Introduction: goals of theranostics in nuclear medicine
Challenges in modern oncology include the fact that patients are getting older and are typically unfit for conventional chemotherapy regimens because of comorbidities or poor performance status.¹ Furthermore, the occurrence of side effects may aggravate treatment compliance in both young and elderly patients.² To manage these problems, it is important to improve patient selection, reduce side effects, and enhance therapeutic efficacy. Taking these factors into consideration, the combination of targeted cancer imaging and therapy is a considerable achievement for personalized medicine.

The theranostic approach in nuclear medicine couples diagnostic imaging and therapy using the same molecule or at least very similar molecules (Figure 1), which are either radionuclide labeled differently or given in different dosages. For example, iodine-131 and lutetium-177 are gamma and beta emitters; thus, these agents can be used for both imaging and therapy. Furthermore, different isotopes of the same element, for example, iodine-123 (gamma emitter) and iodine-131 (gamma and beta emitters), can also be used for theranostic purposes.³⁴ Newer examples are yttrium-86/yttrium-90 or terbium isotopes (Tb): ¹⁵²Tb (beta plus emitter), ¹⁵⁵⁵Tb (gamma emitter), ¹⁴⁹Tb (alpha emitter), and ¹⁶⁴Tb (beta minus particle).⁵⁶

The detection of potential targets can help predict whether a patient will benefit from a particular treatment. Theranostics can be useful for estimating the potential response and eventual toxicity. During the treatment, theranostics can be applied in monitoring the therapy course. However, one cause of concern is the safety of high cumulative doses of radioactive agents after multiple repeated cycles. For instance, reports concerning irreversible high-grade toxicity followed soon after the first treatments with radioiodine (iodine-131).⁷ However, there have been remarkable advances in nuclear
The first theranostic radiopharmaceutical in nuclear medicine history was radioiodine, which was used for therapy and imaging in thyroid diseases. Since then, the use of theranostic agents has been consistently increasing. Nuclear targeted therapies play an essential role, especially in patients with advanced neuroendocrine tumors, such as gastroentero-pancreatic (GEP) tumors, bronchopulmonary neuroendocrine tumors, pheochromocytoma, and neuroblastoma. Furthermore, there are positive results with radioligand therapies in metastatic PC and metastatic melanoma.

The aim of this review was to discuss the most important milestones of nuclear theranostics in current practice (Table 1), and to provide a summarized background and overview of current applications and advantages of targeted therapies and imaging.

**Radioiodine therapy: “the gold standard” in thyroid diseases**

Iodine (stable isotope: iodine-127) is taken up by the thyroid gland for the production of thyroid hormones, namely, thyroxine (T4) and triiodothyronine (T3). Thyroid hormones are vital for the embryonic and neonatal development of the brain, normal growth, and metabolic balance. In 1936, Dr Saul Hertz, director of the Thyroid Clinic in Massachusetts (1931–1943), developed the idea of administering radioactive iodine in patients with thyroid diseases. Iodine and external beam radiation were well-known tools in thyroid disease therapy, but the combination of these was a considerably innovative approach. This idea followed a few years of preclinical studies in collaboration with the Massachusetts Institute of Technology (MIT), where the first cyclotron for medical use had been built. The MIT Cyclotron produced 90% iodine-130 (I-130, half-life: 12 hours) and 10% iodine-131 (I-131, half-life: 8 days). Subsequently, on March 31, 1941, Dr Hertz treated the first patient with radioiodine (I-131).

The first radioiodine therapy (RAI) with I-131 in patients with thyroid cancer (TC) was undertaken by Seidlin et al in 1946. This group studied the use of RAI in patients with metastatic thyroid carcinomas.

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the TC and the thyroid remnant from the inside and, at the same time, targeted lesions can be visualized using a gamma camera or SPECT.9,32,33 Another radioisotope is $^{131}$I, which is a pure gamma emitter and is used for pre- and post-therapeutic diagnostics. The advantages of imaging with $^{131}$I include a higher quality of whole-body scans, which improves the sensitivity in detecting thyroid remnants.3,4

Figure 2A shows the initial $^{131}$I-planar images of a 74-year-old female with metastatic TC (lung, bone, and intracranial soft-tissue metastases). The level of tumor marker thyroglobulin (Tg) before RAI was 572 ng/mL. After two administrations of RAI (cumulative activity: 14.3 GBq), the patient was in complete remission with a Tg level of 0.2 ng/mL (Figure 2B).

**Table 1** Overview of theranostic agents

<table>
<thead>
<tr>
<th>Theranostic molecule</th>
<th>Iodine</th>
<th>mIBG</th>
<th>SSA</th>
<th>PSMA-ligands</th>
<th>Benzamide/arylcarboxamide</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target</td>
<td>Thyroid cancer cells</td>
<td>Neurosecretory granules</td>
<td>SSRT, especially the subtype SSTR2</td>
<td>PSMA</td>
<td>Melanin</td>
</tr>
<tr>
<td>Planar imaging/ SPECT or PET</td>
<td>$^{131}$I and $^{123}$I</td>
<td>$[^{123}]$I-miBG, $[^{123}]$I-miBG</td>
<td>$[^{123}]$I-miBG</td>
<td>$[^{123}]$I-MIP-1072</td>
<td>$[^{123}]$I-BA52</td>
</tr>
<tr>
<td>Therapeutic agent</td>
<td>$^{123}$I</td>
<td>$[^{123}]$I-miBG</td>
<td>$[^{123}]$I-miBG</td>
<td>$[^{123}]$I-PSMA-11</td>
<td>$[^{123}]$I-ICF15002</td>
</tr>
<tr>
<td>Indication</td>
<td>Thyroid cancer</td>
<td>Neuroblastomas, pheochromocytomas, paragangliomas, medullary thyroid carcinomas, and other NEN</td>
<td>SSRT, especially the subtype SSTR2</td>
<td>PSMA</td>
<td>Melanin</td>
</tr>
</tbody>
</table>

**Abbreviations:** mIBG, metaiodobenzylguanidine; SSA, somatostatin analogs; SSRT, somatostatin receptors; NEN, neuroendocrine neoplasia; GEP, gastrointestinal pancreatic system; SPECT, single photon emission computed tomography; PET, positron emission tomography.

Diagnostics and therapy with metaiodobenzylguanidine

Metaiodobenzylguanidine (mIBG), or iobenguane, is a molecule similar to noradrenaline and enters neuroendocrine cells from the sympathetic nervous system, either by endocytosis or passive diffusion before being stored in neurosecretory granules.39 Among the used radiolabeled molecules, $[^{123}]$I-miBG has a lower gamma energy than $[^{123}]$I-miBG (159 keV vs 360 keV), which makes it more suitable for planar imaging/SPECT. Furthermore, the pure gamma emitter $[^{123}]$I-miBG consists of a shorter half-time of 13 hours compared to 8 days for the combined beta and gamma emitter $[^{123}]$I-miBG, leading to a smaller radiation burden. Thus, higher quantities of $[^{123}]$I-miBG can be injected.40 Both $[^{113}]$I-miBG and $[^{123}]$I-miBG are used in mIBG scintigraphy for the detection of neuroendocrine tumors, such as neuroblastomas, pheochromocytomas, paragangliomas, medullary thyroid carcinomas, and other neuroendocrine neoplasias.41 In patients with inoperable or advanced tumors with distant metastases, mIBG imaging plays an essential role in response assessment after therapy and in the evaluation of potential $[^{131}]$I-miBG therapy.32,43

In patients with neuroblastoma and pheochromocytoma, $[^{123}]$I-miBG has high sensitivity (97% and 94%) and specificity (up to 96% and 92%), respectively.44,46 If available, $[^{124}]$I-miBG-PET can be equally used for the planning of mIBG targeted therapy.47,48

Targeted therapy with $[^{113}]$I-miBG presents an encouraging efficacy with tolerable toxicity in relapsed or refractory neuroblastomas with response rates between 20% and 40% being used alone or in combination with high-dose
chemotherapy followed by autologous stem cell transplantation.\textsuperscript{23–25,49} Recently, the NB2004 Trial for Risk Adapted Treatment of Children with Neuroblastoma closed, and further analysis of the usage of mIBG therapy as a first-line therapy is outstanding.\textsuperscript{23,25}

**Palliation of bone metastases**

Radiolabeled phosphonates have a high bone affinity and can be used for imaging and palliation of painful bone metastases. Depending on the degree of osseous metabolism, the tracer accumulates via adhesion to bones and, preferably, to osteoblastic bone metastases.\textsuperscript{8}

Therapy planning requires a bone scintigraphy with technetium-99m-hydroxyethylidene diphosphonate (HEDP) to estimate the metabolism and the extent of the metastases involvement. Bisphosphonate HEDP can be labeled for therapy either with rhenium-186 (beta-emitter, half-life: 89 hours, 1.1 MeV maximal energy, maximal range: 4.6 mm) or rhenium-188 (beta-emitter [to 85%, 2.1 MeV] and gamma-emitter [to 15%, 155 keV], half-life: 16.8 hours, maximal range in soft tissue: 10 mm).\textsuperscript{50}

Both agents induce pain relief in ≥90% of patients.\textsuperscript{51} However, rhenium-188 is of particular interest because it can be cost-effectively produced using a tungsten-188/rhenium-188 generator. The gamma component of the agent allows post-therapeutic imaging of good quality.\textsuperscript{5} Furthermore, patients with prostatic cancer have shown, after two [\textsuperscript{188}Re]Re-HEDP injections, a significantly higher response rate (92%), a prostate-specific antigen (PSA) decline >50% (38%), and a longer PFS (7 months) compared to patients who have had a single injection.\textsuperscript{52,55} Biersack et al showed an improved OS of 15.7 months in patients after repeated injections (≥3 injections) compared to patients with a single therapy.\textsuperscript{54} Besides transient bone marrow toxicity (thrombocytopenia grade III), the reported toxicity is low to moderate.\textsuperscript{55}

New promising radiopharmaceuticals for bone palliation therapy include radiolabeled complexes of zoledronic acid. Zoledronic acid belongs to a new, most potent generation of bisphosphonates with cyclic side chains. The bone affinity of zoledronic acid labeled with scandium-46 or lutetium-177 has shown excellent absorption (98% for [\textsuperscript{177}Lu]Lu-zoledronate and 82% for [\textsuperscript{89}Sc]Sc-zoledronate), which is much higher than of bisphosphonates labeled with samarium-153 (maximum: 67%).\textsuperscript{55}

**Radiolabeled somatostatin analogs**

Neuroendocrine neoplasia (NEN) of the GEP system originates most frequently from the pancreas, jejunum, ileum, cecum, rectum, appendix, and colon. The common characteristic of all GEP-NEN is the compound features of endocrine and nerve cells.\textsuperscript{56–58} Well-differentiated NEN overexpresses somatostatin receptors (SSTRs), especially the SSTR-2 subtype. Thus, SSTRs are theranostic targets in NEN that have been known for almost three decades and have become well established.\textsuperscript{59–61}

SSTR imaging is necessary for staging, therapy planning, and follow-up. In the PET diagnostics, there are three routine somatostatin analog (SSA) tracers labelled with gallium-68: DOTATATE, DOTA-TOC, and DOTA-NOC. All three tracers bind specifically with SSTRs.\textsuperscript{62,63} Gallium-68-labeled SSTRs have high sensitivity (82%–97%) and specificity (80%–92%) in the detection of small primary tumors or metastases of GEP-NEN.\textsuperscript{64–66} Alternatively, a conventional whole-body scintigraphy (planat plus SPECT) or SPECT alone can be performed using indium-111-labeled SSTR, which is available at many institutions. Even if the \textsuperscript{111}In imaging does not have as good a quality as PET, this method has been shown to be superior to chromogranin A measurements alone for the management of GEP-NET patients and is better than computed tomography or ultrasound.\textsuperscript{67,68}

Peptide receptor radionuclide therapy (PRRT) is a systemic therapy in patients with advanced metastatic NEN. PRRT requires a good tumor uptake in the SSTR imaging. For therapeutic purposes, the peptides DOTA-TATE and DOTA-TOC can be labeled with either \textsuperscript{90}Y or \textsuperscript{177}Lu. Because of the renal excretion and the proximal tubular reabsorption of these tracers, the kidneys are one of the primary limiting organs for this therapy.\textsuperscript{69} Due to the smaller range (2 mm) and lower energy, \textsuperscript{177}Lu is less nephrotoxic than \textsuperscript{90}Y (range maximum: 11 mm, pure beta emitter, higher energy). \textsuperscript{177}Lu is also less hematotoxic.\textsuperscript{70} This allows for the performance of several PRRT cycles without any relevant toxicity. Furthermore, the overall survival is not significantly different between the therapies. For that reason, [\textsuperscript{177}Lu]Lu-DOTA-TATE and DOTA-TOC are, in many centers, the preferred agents for NEN therapy.\textsuperscript{10,11,71,72}

PRRT has been increasingly gaining attention. The first randomized controlled phase III study, NETTER-1 (started 2012), compared the standard therapy of Sandostatin® LAR (somatostatin analog) with PRRT in patients with midgut NEN. The study showed a significant clinical benefit from the therapy, achieving a prolonged progression-free survival (median not reached, approximately 40 months, p<0.001), an overall response of 18%, and a presumably longer overall survival (median not reached, p=0.004).\textsuperscript{11}

Figure 3 represents the image of a male patient with a neuroendocrine tumor (unknown origin), with a recurrence of
the disease nearly 2 years after the baseline PRRT. After a salvage therapy of PRRT, the patient had a partial response.

Radiolabeled PSMA-ligands

PC is the most common cancer in men in Western countries. Hormone- and chemotherapy-refractory patients with metastatic PC have a poor prognosis. The main cause of death in these patients is progression to the androgen-independent stage.

PC cells overexpress prostate-specific membrane antigen (PSMA) on the cell surface. There are several available radiopharmaceuticals that target PSMA including [54Ga]Ga-PSMA-HBED-CC (also known as [54Ga]Ga-PSMA-11 [PET]), a monoclonal antibody (mAb) [177Lu]Lu[58Y]Y-J591 (therapy), [123I]-MIP-1072 (planar/SPECT), [131I]-MIP-1095 (therapy), and the theranostic agents PSMA-I&T and DFKZ-PSMA-617 (PSMA-617), which are labeled with 54Ga for PET or with 177Lu for therapy.

The specificity of the two commercially available PET-tracers, [54Ga]Ga-PSMA-617 and [54Ga]Ga-PSMA-11, is similar – 99% and 100%, respectively. However, due to slower kinetics, PSMA-617 was suggested for labeling with the long-half-life Lutetium-177 for therapy and, therefore, there is sparse systematic evaluation of [54Ga]Ga-PSMA-617 for diagnostics. The high kidney uptake of PSMA-11 in the kidneys makes it unsuitable for therapy and does not fit the definition “theranostics.”

Still, it is at issue if molecules with affinity to identical target structures might already present a “theranostic surrogate” even if more than the isotope differs between the diagnostics and the therapeutics.

The latest studies have shown that treatment with [177Lu] Lu-PSMA-617 is effective and well tolerated. In fact, nearly 70% of patients have benefited from this therapy. Dosimetry studies have shown that the most critical organs are the kidneys, with a maximum kidney radiation dose of 0.88 Gy/GBq for [177Lu]Lu-PSMA-617 and 0.93 Gy/GBq for [177Lu]Lu-PSMA-I&T. Okamoto et al stated that a cumulative activity of 40 GBq [177Lu]Lu-PSMA-I&T could be safely applied in patients. Furthermore, [177Lu]Lu-PSMA-617 has been shown to produce no relevant increase in renal toxicity in the salvage setting or in patients with kidney radiation doses >19 Gy.

Figure 4A shows the pre-therapeutic [54Ga]Ga-PSMA-11 images of a patient with multiple lymph node, peritoneal, and bone metastases (arrows), and a history of chemotherapy (first and second line), enzalutamide, and abiraterone. After three cycles of [177Lu]Lu-PSMA-617, the follow-up images show a very good response with a substantial PSA decline (Figure 4B).

Melanin targeting in patients with metastatic melanoma

A promising approach in patients with metastatic melanoma is the specific targeting of melanin. The newly developed theranostic agents include [123I]-[131I]-BA52 and [18F]-[131I]-ICF15002, which may play a considerable role in the future.

Abnormal melanin targeting in patients with metastatic melanoma

Figure 4B shows the [18F]-[131I]-ICF15002 images of a patient with a known primary skin melanoma of the cheek. After 3 cycles of [177Lu]Lu-PSMA-617, the patient had a partial response with a reduction in the size of the primary tumor (unknown origin) 23 months after the baseline PRRT (three cycles, cumulative: 40 GBq). The patient had a recurrence of the disease with multiple metastases in the bone, lung, liver, and lymph nodes (marked with arrows). The latest studies have shown that treatment with [177Lu] Lu-PSMA-617 is effective and well tolerated. In fact, nearly 70% of patients have benefited from this therapy.
ICF15002 is an arylcarboxamide derivative and, as a small molecule, can passively enter the cell and bind to melanin. The PET tracer is radiolabeled with $^{18}$F and $^{[131]}$I-ICF15002 is designed for the therapy. However, both tracers are still in the preclinical phase of their studies. One major problem may be the absorbed dose in melanin-rich tissues, such as skin, dark eyes, and the brain. For instance, in a murine model, there was a 30% decrease of the retinal thickness after two cycles of $^{[131]}$I-ICF15002.

Conclusion

In nuclear medicine, theranostics combine diagnostic imaging and therapy with the same, but differently radiolabeled, molecule, or the same agent in different dosages. The visualization of potential targets can help predict if a patient would benefit from a particular treatment. In properly preselected patients, targeted nuclear therapies have proven to be effective with a favorable safety profile. To conclude, the combination of targeted cancer imaging and therapy is a considerably contribution to personalized medicine and may play an increasingly important role in the future.

Disclosure

The authors report no conflicts of interest in this work.

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