With the recent approval of 177Lu-DOTATATE for use in gastroenteropancreatic neuroendocrine tumors, access to peptide receptor radionuclide therapy is increasing. Representatives from the North American Neuroendocrine Tumor Society and the Society of Nuclear Medicine and Molecular Imaging collaborated to develop a practical consensus guideline for the administration of 177Lu-DOTATATE. In this paper, we discuss patient screening, maintenance somatostatin analog therapy requirements, treatment location and room preparation, drug administration, and patient release as well as strategies for radiation safety, toxicity monitoring, management of potential complications, and follow-up. Controversies regarding the role of radiation dosimetry are discussed as well. This document is designed to provide practical guidance on how to safely treat patients with this therapy.

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Preamble
The present guidelines/standards were developed collaboratively by THE North American Neuroendocrine Tumor Society (NANETS) and the Society of Nuclear Medicine and Molecular Imaging (SNMMI). NANETS is a multidisciplinary professional society of neuroendocrine specialists in North America that was founded in 2005. NANETS’ mission is to improve neuroendocrine tumor disease management through increased research and educational opportunities. NANETS is committed to a multidisciplinary approach and consists of doctors and scientists involved in different specialties of neuroendocrine tumors (NETs). SNMMI is an international scientific and professional organization founded in 1954 to promote the science, technology and practical application of nuclear medicine. In addition to publishing journals, newsletters, and books, the SNMMI also sponsors international meetings and workshops designed to increase the competencies of nuclear medicine practitioners and to promote new advances in the science of nuclear medicine. This collaboration between NANETS and SNMMI aims to define new procedure standards for peptide receptor radionuclide therapy (PRRT) to improve the quality of service to patients. Existing practice guidelines will be reviewed for revision or renewal as appropriate, on their fifth anniversary or sooner, if indicated. Each practice standard has undergone a thorough consensus process in which it has been subjected to extensive review. Much of the content within the guidelines are based on working group experience. These procedure standards are intended to assist practitioners in providing appropriate nuclear medicine care for patients. They are not inflexible rules or requirements of practice and are not intended, nor should they be used, to establish a legal standard of care. For these reasons and those set forth below, NANETS and SNMMI caution against the use of these guidelines in litigation in which the clinical decisions of a practitioner are called into question. The ultimate judgment regarding the propriety of any specific procedure or course of action must be made by medical professionals taking into account the unique circumstances of each case. Thus, there is no implication that an approach differing from the guidelines, standing alone, is below the standard of care. To the contrary, a conscientious practitioner may responsibly adopt a course of action different from that set forth in the guidelines when, in the reasonable judgment of the practitioner, such course of action is indicated by the condition of the patient, limitations of available resources, or advances in knowledge or technology subsequent to publication of the guidelines. The variety and complexity of human conditions make it impossible to always reach the most appropriate diagnosis or to predict with certainty a particular response to treatment. Therefore, it should be recognized that adherence to these guidelines will not ensure an accurate diagnosis or a successful outcome. All that should be expected is that the practitioner will follow a reasonable course of action based on current knowledge, available resources, and the needs of the patient to deliver effective and safe medical care. The sole purpose of these procedure standards is to assist practitioners in achieving this objective.

Background
NETS are a heterogeneous group of malignancies that frequently overexpress somatostatin receptors (SSTRs) (1). NETs can be imaged using somatostatin analogs (SSAs) labeled with \(^{68}\text{Ga}\) (\(^{68}\text{Ga-DOTATATE}\) and \(^{68}\text{Ga-DOTATOC}\) (2)). \(\beta\)-emitting radionuclides such as \(^{177}\text{Lu}\) (\(^{177}\text{Lu}\)) can be used for PRRT (3). The NETTER-1 trial demonstrated prolonged progression-free survival in mid-gut NET patients treated with four cycles of \(^{177}\text{Lu-DOTATATE}\), which subsequently led to the approval of this therapy (4).

Treatment Overview
\(^{177}\text{Lu-DOTATATE}\) is administered at an activity of 7.4 GBq (200 mCi) every 8±1 wk for four cycles. Combined with prophylactic amino acid infusions and antiemetics, each treatment visit can last approximately 5 –8 hours (Fig. 1). Before starting PRRT treatments, each treatment site must ensure that \(^{177}\text{Lu}\) is included in their institutional radioactive materials license. A detailed review of procedures surrounding \(^{177}\text{Lu-DOTATATE}\) therapy is provided below.

Patient Screening
Patients should be evaluated by a multidisciplinary NET team, including a cancer specialist with expertise in the medical management of NETs as well as a nuclear medicine physician or appropriate authorized user to decide on the appropriateness and timing of PRRT in individual patients. Potential candidates should undergo a SSTR PET scan or a SSTR scintigraphy (\(^{111}\text{In-pentetreotide}\)) to demonstrate adequate SSTR expression (2). Traditionally, SSTR expression on \(^{111}\text{In-pentetreotide}\) greater than background hepatic uptake has been considered an eligibility requirement for PRRT (5). Necessary levels of SSTR expression on \(^{68}\text{Ga}\)-based SSTR PET have not been clearly defined, but lesion uptake should exceed background hepatic uptake. Laboratory values should be checked shortly before the treatment is ordered (typically 2 wk prior to each cycle). These should include blood urea nitrogen, creatinine, albumin,
alkaline phosphatase, aspartate aminotransferase, alanine aminotransferase, total bilirubin, white blood cell with differential, hemoglobin, and platelet counts. The threshold values provided in Table 1 should be taken as general eligibility guidelines for therapy.

Somatostatin Analog Therapy
SSA therapy is used frequently to treat NET patients, and SSAs are typically administered in depot formulations every 4 wk. It is recommended that PRRT treatments be scheduled at least 4 wk after the last long-acting SSA therapy to prevent interference with SSTR binding. For symptomatic patients, short-acting SSAs being used as a bridge should be stopped at least 24 h before treatment. Subsequent SSA doses can be administered as soon as several hours after the completion of the radiopharmaceutical therapy. During and after completion of PRRT, it is generally agreed that syndromic patients should remain on SSA therapy. It is unclear whether patients with non-functional tumors should remain on SSA treatment regardless of whether or not they had progressed on SSA therapy before initiation of PRRT. In the NETTER-1 study, all patients remained on long-acting release octreotide despite prior progression on this drug, and the $^{177}$Lu-DOTATATE package insert suggests that patients should remain on SSAs for up to 18 mo after treatment (6). However, there are no clear data to support or refute this recommendation.

Table 1

<table>
<thead>
<tr>
<th>Laboratory</th>
<th>Acceptable value before first treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin (HGB)</td>
<td>&gt;8 g/dL</td>
</tr>
<tr>
<td>White blood cell count (WBC)</td>
<td>&gt;2K/mm³</td>
</tr>
<tr>
<td>Platelet count (PLT)</td>
<td>&gt;70K/mm³</td>
</tr>
<tr>
<td>Estimated glomerular filtration rate (eGFR)</td>
<td>&gt;50 mL/min</td>
</tr>
<tr>
<td>Total bilirubin</td>
<td>≤3 x ULN</td>
</tr>
<tr>
<td>Serum albumin</td>
<td>&gt;3.0 g/dL</td>
</tr>
</tbody>
</table>

ULN = upper limits of normal.

during administration of radioactivity. If a combination approach is taken (i.e., the patient is transported between departments for specific components of the procedure), extra caution regarding radiation protection is required for patient travel. When treated as an outpatient, a patient should be forewarned of the uncommon possibility of an overnight hospital stay should a complication such as a neuroendocrine hormonal crisis or severe emesis occur (6).

Room Preparation
Because body fluids (primarily urine) are radioactive after $^{177}$Lu-DOTATATE administration, room preparation is essential to reduce potential contamination. For example, patient stretchers, chairs, floors, and lower walls can be covered with a prophylactic protective covering (Fig. 2). Furthermore, whereas a treatment suite with an attached toilet is ideal, having a dedicated toilet nearby is acceptable as the patient will need to void frequently on the completion of the $^{177}$Lu-DOTATATE infusion and may need assistance. Local rules related to radioactive waste materials should be followed under the guidance of a local radiation safety officer.

Patient Preparation
It may be helpful to have patients change into hospital scrubs or gowns on arrival in order to avoid potential contamination of personal belongings. If stress urinary incontinence is a concern, disposable undergarments are also recommended. For some patients, a Foley catheter with acrylic shielding of the Foley bag may be necessary. However, routine bladder catheterization is not recommended. A peripheral vein in the antecubital fossa is the preferred location for venous access, and intravenous lines in both arms are preferred, one to administer the radioactivity and one to administer the amino acid solution. If 2 lines
are not possible, the amino acids and radiotherapy may be infused through the same line. Central lines may be used for the administration of the premedications and the amino acid solution in the case of patients with difficult peripheral venous access; however, a peripheral line is the recommended route for the administration of the $^{177}$Lu-DOTATATE infusion, as the use of a central line has not been studied. Intermittent assessment of ongoing vascular access is necessary throughout the procedure due to the high osmolarity of the amino acid solutions, and local reactions at the infusion site may occur. Additionally, patency of the peripheral intravenous site should be monitored continuously throughout the infusion of the $^{177}$Lu-DOTATATE.

### Amino Acid Solutions

Administration of an amino acid solution with the appropriate lysine and arginine concentration (Table 2) before, during, and after the $^{177}$Lu-DOTATATE infusion is required to decrease reabsorption of $^{177}$Lu-DOTATATE via the proximal renal tubules and thereby decrease the radiation dose to the kidneys (7). Several high concentration commercial amino acid solutions are currently available with the correct concentration of arginine and lysine (Table 3); however, they also include additional amino acids, which raise the osmolality of the solution and are associated with significant nausea and vomiting during infusion. The target infusion rate of the commercial amino acid solutions should reach 320 mL/h. $^{177}$Lu-DOTATATE should generally not be administered until this rate is reached or until one eighth of the total volume of amino acid solution has been infused. The amino acid solution should infuse concurrently with the $^{177}$Lu-DOTATATE, and continue at 320 mL/h or greater until the total volume has been administered. Commencing at a low rate of 100 mL/h for the high concentration amino acid solution and increasing slowly (e.g., by 20–50 mL/h every 15–20 min) has been somewhat successful at reducing side effects relating to nausea or vomiting.

An alternative compounded amino acid formulation consisting solely of 25 g of lysine and 25 g of arginine diluted in 1L of normal saline for injection should be considered (8). The compounded two amino acid solution is substantially less emetogenic and can generally be infused over a shorter period of time. The use of arginine-lysine formulations may be preferred due to improved tolerability; however, due to licensing requirements and compounding regulations, compounded arginine-lysine formulations may not be available at many institutions. When using the compounded arginine-lysine solution, it should be infused at a rate of 250 mL/h for 4 h, commencing 30 min before the treatment with the radiopharmaceutical.

### Antiemetic Medications

Nausea and vomiting are common during the administration of commercial amino acid solutions when infusion rates are above 250 mL/h. Therefore, it is recommended to use an intravenous premedication regime consisting of a 5-HT3 antagonist (e.g., granisetron, ondansetron, or palonosetron), an NK1 receptor antagonist (e.g., fosaprepitant), and an H2 receptor antagonist (e.g., famotidine). When increasing the amino acid rate to the target of 320 mL/h, additional doses of the 5-HT3 antagonist may be required with the addition of a D2 receptor antagonist (e.g., prochlorperazine). Benzodiazepines may also be required for anticipatory nausea and vomiting (9). Steroids, such as dexamethasone, can also be administered after infusion of $^{177}$Lu-DOTATATE.

Additionally, cooling and pressure aids may also be beneficial to help with the possible side effects of the amino acid solution infusion. Patient education is highly important at the beginning of the procedure to ensure the patient understands the importance of where and how to contain emesis under these circumstances.

Patients receiving compounded arginine/lysine generally require only a 5-HT3 antagonist as prophylactic medication.

### Radiopharmaceutical Administration

Treatment centers can choose among different infusion methods for $^{177}$Lu-DOTATATE administration, including the gravity method, the saline infusion method, the pump method with a vial, and the pump method with a
syringe (Fig. 3) (6). Supplemental Table 1 (supplemental materials are available at http://jnm.snmmjournals.org) provides step-by-step instructions for each method (10). The saline infusion method may result in leakage from the vial, and care should be taken to prevent contamination. The pump method with a syringe can result in extra radiation exposure to the technologists when drawing the activity from the vial and higher residual values. The pump method with the vial requires the availability of an automated infusion pump. Regardless of the method used, one should use appropriate radiation shielding and aseptic technique when preparing and administering the radiopeptide solution, including wearing appropriate personal protective equipment, using tongs when handling the vial to minimize radiation exposure, confirming the amount of the radioactivity of the radiopeptide vial (and syringe if applicable) with an appropriate dose calibrator dose administration, and inspecting the product visually for particulate matter and discoloration under a shielded screen (the vial should not be used if particulates or discoloration are present). The therapy dosage should be administered over 30 min and should not be administered as a bolus.

### Dosage Modifications for Adverse Reactions

In patients who have baseline renal, liver, or bone marrow dysfunction, or in those who develop toxicity while on treatment, modification of the administered $^{177}$Lu-DOTATATE activity can be considered (6). In patients with preexisting toxicity, decreased administered activity or longer intervals between administrations can be considered. Bone marrow toxicity is the most common adverse event to develop during treatment. In patients experiencing myelosuppression greater than grade 1, one can delay treatment, allowing bone marrow function to recover, administer a lower administered activity (i.e., 3.7 GBq [100 mCi]) during the next treatment, or permanently stop therapy. Oftentimes thrombocytopenia will resolve with a delay. Renal and liver toxicity rarely occur during treatment, but if patients develop toxicity that is attributable to $^{177}$Lu-DOTATATE (e.g., elevated bilirubin or reduction in kidney function), therapy should be withheld until toxicity resolves. Because of issues related to differential reimbursement based on the administered activity, it may be difficult to administer at half the normal activity. Therefore, it may only be feasible to prolong the delay between treatments.

### Patient Monitoring and Potential Reactions

#### Hormonal Crisis (Carcinoid Crisis)

Neuroendocrine hormonal crises due to excessive release of hormones or bioactive substances develop in 1% of patients and typically occur during treatment or within 2 d after the initial treatment (6,11). Typical clinical manifestations include cutaneous flushing, diarrhea, bronchospasm, and hypertension. Hormonal crises can be treated with intravenous high-dose SSAs, intravenous fluids, corticosteroids, and correcting of electrolyte disturbances in patients with diarrhea or vomiting. Pretreating patients at high risk for crisis has been suggested, although this is not done at the majority of centers (12).

#### Infiltration of $^{177}$Lu-DOTATATE

Prevention of infiltration is critical and includes testing the intravenous line patency before administration of the radiopharmaceutical, direct observation of the site during administration, and use of protective tongs to minimize radiation exposure. If infiltration occurs, the site should be allowed to drain freely for 24 h and then covered with a sterile dressing. If swelling persists or increases, the site should be immobilized, elevated, and wrapped in a cold pack. The patient should be observed for signs of compartment syndrome.

### TABLE 2

**Content Requirements for the Amino Acid Solution**

<table>
<thead>
<tr>
<th>Item</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lysine HCl content</td>
<td>Between 18 and 24 g</td>
</tr>
<tr>
<td>Arginine HCl content</td>
<td>Between 18 and 24 g</td>
</tr>
<tr>
<td>Volume</td>
<td>1.5 to 2.2 L for commercial</td>
</tr>
<tr>
<td></td>
<td>(1.0 L for compounded)</td>
</tr>
<tr>
<td>Osmolarity</td>
<td>&lt;1,050 mOsmol</td>
</tr>
</tbody>
</table>

### TABLE 3

**Currently Available Commercial Amino Acid Solutions**

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Amino acid concentration</th>
<th>Dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aminosyn II 10%</td>
<td>21 g of lysine, 20.4 g of arginine in 2 L</td>
<td>Dilute to approximately 2 L</td>
</tr>
<tr>
<td>Aminosyn II 15%</td>
<td>23.6 g of lysine, 22.9 g of arginine in 1.5 L</td>
<td>Dilute to approximately 2.2 L</td>
</tr>
<tr>
<td>Clinisol 15%</td>
<td>18 g of lysine, 18 g of arginine in 1.6 L</td>
<td>Dilute to approximately 2.1 L</td>
</tr>
<tr>
<td>Plenamine 15%</td>
<td>18.8 g of lysine, 23.5 g of arginine in 1.6 L</td>
<td>Dilute to approximately 2.1 L</td>
</tr>
<tr>
<td>Trophamine 10%</td>
<td>18 g of lysine, 26 g of arginine in 2.2 L</td>
<td>Consult*</td>
</tr>
<tr>
<td>Premosol 10%</td>
<td>16.4 g of lysine, 24 g of arginine in 2 L</td>
<td>Consult*</td>
</tr>
</tbody>
</table>

*Consult with treating physician as lysine–arginine concentration may be outside specifications, which may increase adverse events.
swelling or pain develop. If infiltration occurs, clearance of the radiotracer from the site can be facilitated with warm packs, compression, and elevation (13). Infiltration must be reported to radiation safety for monitoring and calculation of skin dose.

Radiation Safety

Contamination

With a half-life of 6.6 d, 177Lu raises the possibility of prolonged contamination. Blood and urine are the main sources of contamination during and after radionuclide administration. Because 177Lu-DOTATATE is primarily excreted in urine, with a cumulative excretion of 44% within 5 h, 58% within 24 h and 65% within 48 h AFTER administration, the main focus for individual patients for the first 3 d after therapy is on preventing urinary contamination (6). Of note, when produced by neutron activation, there is a long-lived 177mLu contaminant, which has a half-life of 160 d and needs to be surveyed for before disposal of radioactive waste from the treatment center. This issue is of greater importance in locations that treat patients in an inpatient setting. Emesis can contain small amounts of radioactive material and should be treated as contaminated and disposed of appropriately.

Preparation for Inpatient Therapy

For patients who require an overnight stay, typically due to medical complications, the recommendations largely follow those already in place for 131I inpatients, noting that the external dose rate from 177Lu is significantly lower than 131I given the lower energy and abundance of its γ emissions. Nursing personnel will need to be instructed in pertinent radiation safety precautions (i.e., potential for contamination related primarily to the patient’s blood and urine), but also be advised that nausea and vomiting related to the administration of the amino acid solution may occur. Universal precautions (e.g., gloves, gowns, shoe covers) should be used to avoid contact with patient bodily fluids. If blood or urine specimens are needed for laboratory testing, nursing staff should be advised to collect the smallest amount necessary for testing. Nursing staff should be provided with radiation monitoring devices (passive dosimeter, direct-reading dosimeter). Ideally, radiation safety staff should help prepare the patient’s bed, floor, and bathroom to minimize potential radioactive contamination from patient bodily fluids. Institutional radiation safety guidelines should be developed for general nursing care of 177Lu-DOTATATE inpatients including approaches to medical emergencies or patient deaths before a 177Lu-DOTATATE therapy program is established.

Release Criteria

To ensure that radiation dose to members of the public remains less than 5 mSv (500 mrem), the patient should be provided with instructions (Table 4) at discharge. Each center should determine its own recommendations based each patient’s specific circumstances and local regulations, but we have provided a basic guidance in Table 4 for minimizing exposure to others and potential urine contamination (14). The average exposure at one m immediately after treatment is 1.8 ± 0.5 mrem/h, and at time of discharge is 0.9 ± 0.4 mrem/h (combined institutional experiences from more than 100 therapies). This exposure is less than that from 131I therapy and below release criteria from published Nuclear Regulatory Commission guidelines (15). The time periods for following various instructions will vary, but 3 d should be sufficient for resumption of most public activities, given the physical half-life of 6.7 d, the mean effective blood elimination half-life of 3.5 ± 1.4 h and the mean terminal blood half-life of 71 ± 28 h (6). Extra precautions should be taken to minimize exposure of young children and pregnant women. The mathematics of release criteria for any nuclear medicine therapy patients have been addressed comprehensively.
The RADAR website has an online tool that allows for calculation of cumulative doses to family members or members of the public from exposure to patients treated with $^{177}$Lu-DOTATATE (16).

**Pregnancy**
Radiopharmaceutical therapy is almost universally contraindicated during pregnancy. Therapy must be delayed until childbirth or termination of pregnancy. Breast feeding should be stopped for treatment and not be restarted until 2.5 mo after the final therapy. Future children may be breastfed. Contraception should be used for 6 mo after completion of the final treatment.

**Issues with Cremation and Patient Death**
It is important to notify the local radiation safety officer of a death involving a radioactive patient. Deceased patients should be appropriately labeled and the death certificate should note that the patient is radioactive. If possible the radiation safety officer should appropriately train the medical examiners and the mortuary personnel, as well as perform radiation surveys. Please refer to the National Council on Radiation Protection and Measurements (NCRP) Report No. 161 (NCRP 2008) for additional guidance on the management of radiation accident victims, regarding guidelines for the medical examiner and mortuary personnel (17).

**Dosimetry and Post-Treatment Imaging**
Although dosimetry was not a part of the Phase III NETTER-1 trial, there may be a role for patient specific dosimetry when considering cumulative renal and bone marrow dose. There is a large variability in tumor and organ uptake of radiolabeled SSAs across patients, which suggests that tailored dosimetry may be useful for $^{177}$Lu-DOTATATE therapy (18). For example, 1 study showed a biologically effective dose to the kidneys ranging from 9 to more than 40 Gy (19). Please see Supplemental Table 2 for standardized dose estimates for the reference adult for $^{177}$Lu-DOTATATE (20). Although $^{68}$Ga-DOTATATE may be used for diagnostic evaluation of tracer uptake, it cannot be used for dosimetry planning due to its short (68 min) half-life. However, $^{177}$Lu gamma-rays are suitable for gamma camera imaging (Supplemental Table 3), and imaging for dosimetry may be performed after the initial therapy cycle (Fig. 4).

Further details about potential imaging protocols for dosimetry can be found in the supplemental materials.

**Toxicity**
Rates of toxicity vary between patients, and heavily pretreated patients have higher rates of PRRT-associated toxicity. In particular, liver toxicity has been reported in patients who have had extensive prior liver-directed therapy or who are treated with $^{90}$Y-based PRRT (21). The following sections focus on bone marrow and kidney toxicity in more detail.

**Bone Marrow Toxicity**
Grade 3 and 4 thrombocytopenia and neutropenia occur in ≤5% of patients and resolve within 8 wk (4,22). Grade 3 and 4 lymphopenia is more common, but is rarely of clinical significance since opportunistic infections are not observed in association with $^{177}$Lu-DOTATATE. The most significant long-term hematological risk is myelodysplastic syndrome or acute leukemia, which occurs in roughly 2%
- 3% of patients at a median of 2 y after therapy. Although little can be done during treatment to minimize marrow toxicity, there are possible risk factors for the development of toxicity such as prior chemotherapy (6,23,24).

**Renal Toxicity**

Due to the high exposure to the kidneys from the renal excretion of the radiotracer, renal toxicity is possible. With the introduction of concurrent amino acids for renal protection and the use of $^{177}$Lu-labeled compounds in place of $^{90}$Y-labeled compounds, the rates of renal toxicity are low, with long-term grade 3-4 renal toxicity less than 2% (4,23). Variations in individual kidney sensitivity to radiation may explain the variability seen between renal dose and the subsequent development of renal toxicity. For example, there is evidence that patients with long-standing diabetes or hypertension may be at higher risk for renal dysfunction after $^{177}$Lu-DOTATATE treatment (19). Nevertheless, severe renal toxicity is rare (<5%) when using the current administration guidelines (4,19).

**Follow-Up**

Monitoring of patients after PRRT treatment is an essential part of the treatment plan. The recommended monitoring should include clinical evaluation to assess symptoms and detection of possible treatment sequelae, laboratory, and imaging tests (Table 5).

**Clinical Evaluation**

Clinical evaluation by the treating or primary team should ideally be conducted at 1 mo, 3 mo, 6 mo, and 12 mo after PRRT. If there are no laboratory abnormalities or clinical symptoms concerning for posttreatment sequelae, patients can resume clinical follow-up per the primary team. National Comprehensive Cancer Network (NCCN) guidelines recommend follow up intervals of 3-12 mo based on clinical presentation (25). Clinical symptoms and presentations that could reflect possible progression, increased symptoms from carcinoid syndrome, or posttreatment sequelae warrant closer monitoring.

**Laboratory Tests and Markers**

Blood tests including complete blood count with differential, aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase, total bilirubin, albumin, and serum creatinine/GFR should be monitored at roughly 1 mo, 3 mo, 6 mo, and 12 mo after treatment. If all blood tests are within normal limits, a complete blood count and serum creatinine should be monitored at least yearly or sooner if clinically indicated or per primary team. If there are any abnormalities in blood tests, more frequent monitoring is recommended.

Both secondary myelodysplastic syndrome and acute leukemia are known toxicities of PRRT. Increased monitoring is recommended for those patients with persistent cytopenias with blood tests until recovery. In addition, hematology consult should be considered for patients with persistent cytopenias.

Patients with mild to moderate renal impairment after PRRT treatment warrant closer monitoring with serum creatinine measurements. In addition, nephrology consult should be considered for patients with persistent or worsening renal impairment.

There are no formal recommendations for following tumor markers after PRRT. NCCN NET guidelines recommend following markers if clinically applicable every 3-12 mo (25).
Diagnostic Imaging Evaluation

In most cases, diagnostic imaging should be done at 1 – 3 mo, 6 mo, and 12 mo after the completion of all treatment cycles. Thereafter, patients should undergo diagnostic imaging based on treatment response. It is important to remain consistent with imaging modalities. NCCN guidelines recommend following patients with contrast-enhanced abdominal and pelvic CT or MRI with contrast along with chest CT with or without contrast (if clinically indicated) every 3-12 mo (25). MRI of the liver with gadoxetate disodium (Eovist) should be considered for those patients with liver dominant disease (26,27). Increased monitoring is recommended for clinical presentation concerning for possible progressive disease, clinical worsening, or possible posttreatment sequel. If there is evidence of progression or equivocal findings on CT or MRI, SSTR-based imaging with 68Ga-DOTATATE PET should be considered and is preferred over SSTR scintigraphy (111In-pentetreotide) (2).

It should be noted that the first imaging study performed after the completion of PRRT can be complicated by pseudoprogression (28). In nearly 10% of patients with stable disease, metastatic disease can increase in volume transiently, presumably due to edema from radiation to the tumor. Diagnostic imaging during the course of PRRT can be considered in patients with relatively aggressive tumors, or patients with clinical evidence of progression.

Conclusion

177Lu-DOTATATE is an effective treatment for patients with NETs. The above guidelines provide information on how to safely administer this novel treatment to patients.
REFERENCES