ACR-ACNM-ARS-SNMMI PRACTICE PARAMETER FOR LUTETIUM-177 (Lu-177) DOTATATE THERAPY

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PREAMBLE

This document is an educational tool designed to assist practitioners in providing appropriate radiologic care for patients. Practice Parameters and Technical Standards are not inflexible rules or requirements of practice and are not intended, nor should they be used, to establish a legal standard of care¹. For these reasons and those set forth below, the American College of Radiology and our collaborating medical specialty societies caution against the use of these documents 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 the practitioner considering all the circumstances presented. Thus, an approach that differs from the guidance in this document, standing alone, does not necessarily imply that the approach was below the standard of care. To the contrary, a conscientious practitioner may responsibly adopt a course of action different from that set forth in this document when, in the reasonable judgment of the practitioner, such course of action is indicated by variables such as the condition of the patient, limitations of available resources, or advances in knowledge or technology after publication of this document. However, a practitioner who employs an approach substantially different from the guidance in this document may consider documenting in the patient record information sufficient to explain the approach taken.

The practice of medicine involves the science, and the art of dealing with the prevention, diagnosis, alleviation, and treatment of disease. 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 the guidance in this document will not assure 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 purpose of this document is to assist practitioners in achieving this objective.

I. INTRODUCTION

¹ Iowa Medical Society and Iowa Society of Anesthesiologists v. Iowa Board of Nursing, 831 N.W.2d 826 (Iowa 2013) Iowa Supreme Court refuses to find that the "ACR Technical Standard for Management of the Use of Radiation in Fluoroscopic Procedures (Revised 2008)" sets a national standard for who may perform fluoroscopic procedures in light of the standard's stated purpose that ACR standards are educational tools and not intended to establish a legal standard of care. See also, <u>Stanley v. McCarver</u>, 63 P.3d 1076 (Ariz. App. 2003) where in a concurring opinion the Court stated that "published standards or guidelines of specialty medical organizations are useful in determining the duty owed or the standard of care applicable in a given situation" even though ACR standards themselves do not establish the standard of care.

This practice parameter was developed collaboratively by the American College of Radiology (ACR), the American College of Nuclear Medicine (ACNM), the American Society for Radiation Oncology (ASTRO), and the Society of Nuclear Medicine and Molecular Imaging (SNMMI).

This practice parameter is intended to guide appropriately trained and licensed physicians performing therapy with lutetium-177 (Lu-177) DOTATATE. Such therapy requires close cooperation and communication between the physicians who are responsible for the clinical management of the patient, those who administer radiopharmaceutical therapy and those who manage the attendant side effects. Adherence to this parameter should help to maximize the efficacious use of these procedures, maintain safe conditions, and ensure compliance with applicable regulations.

Application of this parameter should be in accordance with the <u>ACR–AAPM–SPR Technical Standard for</u> <u>Therapeutic Procedures Using Radiopharmaceuticals [1]</u>, in so far as that standard relates to the handling of radiopharmaceuticals, radiation safety, and radiation protection of patients, personnel, and the public. There must also be compliance with applicable laws and regulations.

The goal of therapy with Lu-177 DOTATATE is to slow disease progression, to palliate symptoms, or even to extend life, while minimizing untoward side effects and complications.

Neuroendocrine tumors (NETs) are relatively rare and typically slow-growing neoplasms that originate in neuroendocrine tissue distributed throughout the body. They secrete bioactive amines and hormones, giving rise to variable clinical presentations [2]. Surgical resection of the tumor is the preferred initial therapy; however, because of the indolent course and nonspecific presentation of the disease, many patients are diagnosed with locally advanced or metastatic disease, making curative resection difficult or impossible. Alternative conventional treatments include use of nonradioactive somatostatin analogues that take advantage of the overexpression of somatostatin receptors (SSRs) by these NETs. Use of other agents, including cytotoxic chemotherapy, can be limited because of the unwanted side effects and minimal effectiveness in certain grades of tumor. Despite use of these currently available conventional treatments, many patients continue to progress with life-altering signs and symptoms, such as unrelenting diarrhea, flushing, or right-sided heart disease [3,4].

Lu-177 DOTATATE is an effective therapy for patients with inoperable, locally advanced, or metastatic NETs that progress on conventional treatments [3-6]. Improvement in disease control rates, progression-free survival, overall survival, and quality of life have advanced this radiopharmaceutical agent to a place of primary consideration in advanced disease management [7]. Lu-177 DOTATATE specifically binds to the SSRs that are overexpressed on the cell surfaces of most NETs, with highest affinity for subtype 2. The complex formed is chemically stable and is internalized into the cell resulting in a favorable position to irradiate the nucleus to induce DNA damage–related inhibition of growth and death [8]. This treatment process is called peptide receptor radionuclide therapy (PRRT). Beta (B-) emission from Lu-177 has a maximum energy (Q-value) of 0.5 MeV, range in soft tissues of 2 mm, and half-life of 6.7 days. Lu-177 also emits low-energy gamma rays at 208 keV (11% abundance) and 113 keV (6.4% abundance) that can be used for gamma camera imaging and dosimetry if desired [9]. Although PRRT with Lu-177 DOTATATE has been proven to be effective in NET, there are adverse side effects and safety issues that must be understood and taken into consideration by the treating physicians so that appropriate plan and required interventions can be instituted [5].

Side effects associated with PRRT with Lu-177 DOTATATE can be categorized as acute, subacute, or delayed [5]. It is highly advisable that a multidisciplinary team coordinate the care of a patient being considered for treatment with Lu-177 DOTATATE [6].

<u>General</u>: Abdominal pain, nausea, and vomiting can occur typically within 24 hours of treatment. In addition, patients can also experience fatigue, diarrhea, alopecia and cough [7]. In most cases, these symptoms are self-limiting and rarely require more than supportive therapy.

<u>Nephrotoxicity:</u> Lu-177 DOTATATE is excreted by the kidneys through glomerular filtration and is reabsorbed by the proximal tubules where radiation damage can occur [9]. Reduction of proximal tubular reabsorption has been effectively achieved with use of other ligands that can competitively bind to the receptors in the proximal tubular

cells without affecting the SSR targets of Lu-177 DOTATATE in the circulation [10]. The most efficacious solution to date to reduce renal uptake of somatostatin analogues consists of a combination of basic amino acids lysine (25 g) and arginine (25 g) [9,11]. Renal toxicity is generally mild and well-tolerated with amino acid co-infusions. However, grade 1 nephrotoxicity in 20% and grade 2 nephrotoxicity in 4% of patients has been reported [5,7]. Higher-grade toxicities are rare (0% to 0.4%) [4,5]. Many studies have shown improvement of renal function over time, but long-term renal impairment remains a clinical concern, with some studies reporting an annual decrease in creatinine clearance of 3.4% to 3.8% [12,13]. Details on administration are provided in the "Specific Procedures" section of this document (IV.B).

<u>Hematologic:</u> The bone marrow is a rapidly dividing organ and is thus radiosensitive. Mild subacute myelosuppression can be seen in the first days to weeks after treatment and typically reverses within weeks after cessation of treatment [5]. The most frequently seen effects include anemia, thrombocytopenia, and leukopenia. Grade 3 and 4 bone marrow toxicity are seen less frequently and are generally reversible without intervention within 2 to 3 months but may take up to 12 months [4,14,15]. Bone metastases can increase the likelihood of myelotoxicity [15,16]. Rarely, 1% to 2% of patients can develop leukemias and myelodysplastic syndrome (MDS), which can lead to a fatal outcome in patients heavily pretreated with myelosuppressive therapies prior to receiving Lu-177 DOTATATE [4,5,13,14].

<u>Hepatic:</u> Liver dysfunction has been noted with increase in bilirubin and transaminases. A few patients have developed grade 3 toxicity that progressed to liver failure and death within one year after PRRT [<u>5</u>].

<u>Hormonal Crisis</u>: This is a rare complication that presents as flushing and significant diarrhea and, less frequently, heart failure, emesis, and bronchoconstriction. It typically occurs within 48 hours of infusion, with greater likelihood in patients with large tumor burden [<u>17,18</u>]. This is a serious adverse side effect requiring prompt inhospital care for continuous somatostatin analogue infusion and supportive care.

<u>Risk of Infertility</u>: The recommended cumulative activity of 800 mCi (29.6 GBq) Lu-177 DOTATATE results in radiation absorbed dose to the testis and ovaries within the range where temporary or permanent infertility may ensue, such as seen following pelvic external-beam radiotherapy [4,6,7,19].

Facilities and their responsible staff should consult with their radiation safety officer (RSO) to ensure that there are policies and procedures specific to Lu-177 DOTATATE that address 1) required instrumentation, calibration, and calibration frequency and 2) ordering and receiving, recordkeeping, safe use, and waste disposal in compliance with the applicable laws and regulations as described in <u>ACR–AAPM Radiation Safety Officer</u> <u>Resources [19]</u>.

II. INDICATIONS

Lu-177 DOTATATE is indicated for the treatment of SSR-bearing gastroenteropancreatic NETs (GEP-NETs), including foregut, midgut, and hindgut NETs in adults [6].

III. QUALIFICATIONS AND RESPONSIBILITIES OF PERSONNEL

The qualifications and responsibilities of physicians and other personnel performing these therapeutic procedures should be in accordance with the <u>ACR–AAPM–SPR Technical Standard for Therapeutic Procedures Using</u> <u>Radiopharmaceuticals</u> and/or the <u>ACR–ASTRO Practice Parameter for Radiation Oncology</u> [1,20].

In addition, training and experience must be in compliance with the applicable laws and regulations.

IV. SPECIFICATIONS OF THE EXAMINATION AND TREATMENT

The written or electronic request for a Lu-177 DOTATATE procedure should provide sufficient information to demonstrate the medical necessity of the examination and allow for its proper performance. Documentation that satisfies medical necessity includes 1) signs and symptoms and/or 2) relevant history (including known diagnoses). Additional information regarding the specific reason for the procedure or diagnosis would be helpful and may at times be needed to allow for the proper performance of the procedure.

The request for the procedure must be originated by a physician or other appropriately licensed health care provider. The accompanying clinical information should be provided by a physician or other appropriately licensed health care provider familiar with the patient's clinical problem or question and consistent with the state scope of practice requirements. (ACR Resolution 35, adopted in 2006 - revised in 2016, Resolution 12-b)

A. Clinical Evaluation:

The clinical evaluation should be in concordance with the <u>ACR–ASTRO Practice Parameter for Radiation</u> <u>Oncology</u> and the <u>ACR–ASTRO Practice Parameter for Communication: Radiation Oncology</u> [20,21]. The treating physician's initial evaluation of the patient must include review of the patient's history, physical examination, pertinent diagnostic studies, laboratory reports, and complete history of all available records of previous pertinent therapies, including, but not limited to, myelosuppressive systemic therapy and/or radiotherapy.

- <u>Verification of Pathology and Indication for Therapy</u>: A pathology report confirming diagnosis of GEP-NET should be reviewed and included in the patient's record. Efficacy of Lu-177 DOTATATE is well documented, particularly in well-differentiated NET often with Ki-67 index of <20% [22]. Because Lu-177 DOTATATE localizes to NET expressing SSR, it is of paramount importance to confirm that the NET being treated expresses the required SSR through positive indium-111 octreotide scan or gallium-68 DOTATATE PET/CT (see <u>ACR-ACNM-SNMMI-SPR Practice Parameter for the Performance of Neuroendocrine Tumor Scintigraphy</u> and <u>ACR Practice Parameter for the Performance of Gallium-68 DOTATATE PET/CT for Neuroendocrine Tumors</u>) [23,24].
- 2. <u>Discontinuation of Somatostatin Analogue Therapy with Baseline Laboratory Evaluation</u>: If the patient is being treated with long-acting somatostatin analogue, this should be stopped for 4 weeks prior to Lu-177 DOTATATE infusion. Short-acting analogues can be stopped 24 hours prior to PRRT. In anticipation of possible side effects, each patient should have a complete blood count with differentials and metabolic panel including renal and hepatic function tests. Such monitoring should be performed before each infusion and as needed for hematologic monitoring in between treatments. Dose reduction based upon laboratories is discussed in Section IV.B.2. Although institution and patient-specific considerations take precedence, a creatinine clearance >50 mL/min and grade 1 to 2 or less hepatic enzyme elevation or myelosuppression is sufficient to allow therapy. Women of childbearing age should undergo pregnancy testing [6].
- 3. <u>Special Populations:</u> Lu-177 DOTATATE has not been tested in lactating patients, and these patients should be advised to stop breastfeeding while receiving treatment and for 2.5 months after the last treatment fraction, as effects on infants have not been determined. For patients of reproductive potential, discussion should be carried out to use effective contraceptive measures during and after PRRT. For female patients, because of the possibility of fetal harm, effective contraception should be continued for 7 months following the last treatment fraction of PRRT. For male patients with female partners, contraception should be continued until 4 months following the last treatment fraction [6]. Sexual activity should be avoided following therapy for 7 days. The radiopharmaceutical has not been tested in pediatric (<18 years old) and pregnant patients. Caution should be exercised in these patient populations, with extensive discussion regarding risk of radiation.
- 4. <u>Quality Management</u>: In order to use radiopharmaceuticals as unsealed sources for therapy, including Lu-177 DOTATATE, a "quality management" program must be in place as required by applicable state and federal regulations. (An Agreement State is any state with which the Nuclear Regulatory Commission (NRC) or the US Atomic Energy Commission has entered into an effective agreement under subsection 274.b of the Atomic Energy Act of 1954 as amended, 73 Stat, 689). Key elements of such a program include written directives, duplicative procedures for identifying patients, careful record keeping to ensure prescribed administered activity, minimization of the possibility of infiltration for radiopharmaceuticals that are administered intravenously (IV), procedures for minimizing radiation exposure or radiopharmaceutical contamination of personnel, family members of patients, and the public (eg, alerts regarding possible current or future pregnancy), procedures for containment of radioactivity; and an audit mechanism to ensure

compliance with the program.

- 5. <u>Informed Consent:</u> Informed consent must be obtained and documented. See the <u>ACR Practice</u> <u>Parameter on Informed Consent – Radiation Oncology [25]</u>.
- 6. <u>Treatment</u>: The procedure and follow-up should be performed according to an established system of procedural steps unique for Lu-177 DOTATATE [26].
- 7. <u>Radiation Precautions:</u> Radiation precautions and patient release criteria may be regulated federally by the NRC in many states or by the state (with regulations that are closely patterned on the federal regulations and may be more restrictive). The radiation safety officer, medical physicist, or health physicist for the local facility can provide information on the applicable regulations. Details on the federal regulations can be obtained at the NRC website (<u>nrc.gov</u>).

Under the guidelines of federal code 10 of the Code of Federal Regulations (CFR) 35.75 [27] and key sections of NUREG-1556 [28], a patient may be released to the public if the total effective dose equivalent to any other individual (including any caregiver or family member) who is exposed to the patient is not likely to exceed 5 mSv (0.5 rem). If the total effective dose equivalent is likely to exceed 1 mSv (0.1 rem) to any individual, instructions (including written instructions) must be provided to the patient on actions to limit radiation exposure to others by utilizing the "as low as reasonably achievable" (ALARA) principle. Some states may have specific rules and regulations regarding release of patients with significant residual activity.

The dose limits specified by the National Council on Radiation Protection and Measurements (NCRP) differ somewhat from the NRC regulations [29]. Because the fetus and children are more sensitive to radiation injury than adults, the NCRP specifies that children and pregnant women, whether or not they are members of the patient's household, should be limited to 1 mSv (0.1 rem). Any individual who has no familial connection to the patient and whose presence offers no emotional benefit should also be limited to 1 mSv, which is also the NRC dose limit to a member of the public.

Many radiation meters measure exposure rates in milliroentgens per hour (mR/h). For purposes of radiation protection and for low linear energy transfer (LET) radiation (including beta particles and most x-rays and gamma rays), the authors of this document accept the approximation that 1 mR, 0.01 mSv, and 1 mrem are equivalent. Thus, an exposure rate of 7 mR/h at 1 m is an adequate approximation to the dose rate, 0.07 mSv/h (7.0 mrem/h) at 1 m.

Specific Considerations During Lu-177 DOTATATE Therapy and Patient Release:

According to radiation exposure calculations based on whole-body clearance data, patients may need to be kept in radiation isolation for 4 to 5 hours following the administration of the typical dose of 200 mCi (7.4 GBq) Lu-177 DOTATATE [30]. Postinfusion survey by physics or other radiation safety is performed to determine an acceptable maximum exposure rate that conforms to the 10 CFR 35.75 requirement of <5 mSv exposure anticipated to other individuals. An established protocol for documenting this survey result should be used and available. Until the patient has been released, the patient must be kept in an area with suitable radiation shielding to protect others from unnecessary exposure. An administration of 200 mCi Lu-177 DOTATE typically results in exposure levels on the order of 2 mR/h at 1 m immediately after administration, declining to 1 mR/h after 24 hours, allowing outpatient treatment in most cases with appropriate training, protocols, infrastructure, and patient counseling. The procedures and practical example guidance for instruction of patients upon discharge have been reviewed in published literature [31]. For further information, see Appendix A.

Modeling per the NUREG-1556 assumption of 0.25 occupancy factor estimates 1.8 mSv exposure dose to other individuals, thus requiring written instructions be given to the patient on ALARA principles. During therapy, involvement of trained radiation safety personnel qualified in safe management of unsealed sources, waste, accidental contamination, and counseling of patients is

important. The patient and, as relevant, caregivers should be compliant with all radiation safety precautions and instructions. Education should occur before treatment, preferably at the time of consultation so that the patient and caregivers can plan ahead. Inability to comply with the precautions may require an admission or other special accommodations to account for the realities of patient life at home, as determined by the authorized user. The specific instructions and considerations for admission or other special accommodations will vary from institution to institution, but key features are summarized below.

Urinary Contamination:

Specific concern is paid to disposal of urine as the most common potential source of contamination. During therapy, a dedicated toilet is preferred, and although lead shielding is not needed because of the short range of beta emission, disposable lining of the floors and toilet/sink surfaces is recommended to contain radioactive urine or other contamination [31]. Urinary incontinence, if present, would require catheterization prior to administration and for at least 2 days thereafter to minimize radiation contamination. Other simple measures used to minimize urinary contamination upon discharge include:

Use of private room with its own bathroom Washing of hands for 20 seconds after each use of the restroom Instructing the patient to urinate while seated Flushing 2 to 3 times with the toilet lid closed Rinsing of sinks and showers after use Cleanup of urinary spills with damp toilet paper that can be flushed down the toilet (to minimize accumulation of waste product trash requiring long-term storage).

Other Potential Sources of Contamination:

Peritoneal and hemodialysis are not contraindications for treatment, but they may impact the administered activity of Lu-177 given the prolonged residence time within the patient and complicate handling of hemodialysis machines because of the likelihood of retained radioactivity after use, thus requiring logistics planning with dialysis facilities and the patient. Another infrequent but special consideration for Lu-177 DOTATE therapy given its target population is in patients with indwelling drains, such as biliary drains, which require confirmation of ability of caregivers to safely manage disposal of waste with the same precautions applied to urine. When possible, these sources of waste should be flushed down the toilet similar to urine, with use of disposable gloves by the caregiver when handling and cleaning drain equipment and collection bags.

Release to Health Care Facility/Admission to Hospital Considerations:

If confinement in a health care facility is needed, it is not usually necessary to store body effluents, such as urine, stool, or vomitus. In general, for patients who have been released to the public, precautions for the patient should be according to ALARA principles and universal precautions. A discussion should be had in such cases with a facility or hospital's radiation safety department and/or involved parties (clinical leadership) to determine any additional precautions that will be taken for care workers. Furthermore, should a patient receiving Lu-177 DOTATATE require admission to a hospital or transfer to an emergency department, it is highly recommended that the administering team contact the receiving personnel for a "signout."

Although not explicitly required, examples of "extra" precautions include the following. For effluent disposal where acceptable under state or federal regulations, the toilet can be flushed two or three times after each use to ensure sufficient dilution of radioactivity. Food trays, linens, and all other contaminated products may be stored in the patient's room until monitored and cleared by radiation safety staff. The patient must stay in the room except in a medical or nonmedical (eg, fire) emergency, and access by personnel and visitors can be limited. All trash and residual nondisposable items can be monitored after the patient's release and stored until radiation levels reach the statutory level defined for safe disposal or reuse. In some jurisdictions, items in decay storage must

reside there for 10 half-lives (67 days for Lu-177) or until radiation levels are indistinguishable from background. Once all known contamination is removed from the room, the room must be surveyed to verify that the radiation levels and removable contamination are sufficiently low to permit its general use. The room may not be used until this survey is performed and safe level documented. Individual institution's radiation safety procedures may vary somewhat.

If the admitting physician is different from the physician who administers the radiopharmaceutical, there must be a mechanism to prevent premature discharge or release of the patient from confinement.

Waste Disposal:

As above, trash and nondisposable items contaminated by patient fluids must be stored and monitored until their radiation levels reach safe disposal limits, which may vary between institutions and jurisdictions, with one prominent guideline being 10 half-lives (67 days for Lu-177).

Distance of Caregivers and Considerations for Travel:

There is no specific regulation on required distance of caregivers following discharge. However, to meet guidance from NUREG-1556's use of a 0.25 occupancy factor for estimating exposure of public allowing safe discharge of patients after administration, it is assumed that exposed persons will maintain a distance of 1 m (3 ft) for at least 3 days and not sleep in the same bed as the patient for 7 days. There is a further assumption of following ALARA principles to minimize exposure to potential contamination, such as may occur during use of the same toilet facilities.

Prolonged use of personal or public transportation (bus, train, etc) in the company of others for more than one hour is discouraged for the first 3 days following therapy. Although Title 10 of the CFR, part 35.75, does not expressly prohibit release of a radioactive patient to a location other than a private residence such as a hotel, the NRC strongly discourages this practice because it can result in radiation exposure to members of the public for which the licensee may not be able to assess full compliance with the code.

Nonetheless, when travel is unavoidable in the first 3 days after therapy, the patient should be instructed to discuss the matter with treating personnel.

Furthermore, although patients are recommended to travel immediately home, it is acknowledged that some patients may need to reside in a hotel or other public facility. Again, precautions to maximize distance from other members of the public are recommended (>1 m at a minimum) in the 3 days after Lu-177 DOTATATE administration.

B. Treatment Procedures for Infusion of Lu-177 DOTATATE

1. Preparation:

Before ordering Lu-177 DOTATATE solution for PRRT, confirm that treatment with Octreotide analogues has been discontinued for at least 4 weeks for long-acting preparation and for 24 hours for short-acting preparation before scheduled therapy.

Lu-177 DOTATATE is a radiopharmaceutical that requires effective radiation shielding before handling. The vial containing the radiopharmaceutical is delivered in a lead- or Plexiglas-shielded container. It is highly advised that the personnel assigned to prepare or infuse the radiopharmaceutical wear double gloves.

Before the actual administration of Lu-177 DOTATATE, patients should be started on a renoprotective amino acid infusion and may be premedicated with antiemetics according to institutional/physician preference. Depending on institutional preferences and resources, coordination should be made between all involved staff, including the referring physician and the physician administering the radiopharmaceutical to ensure that the steps and processes involved with PRRT are carried out. Two separate IV access sites are preferred: one for the amino acid infusion

and one for Lu-177 DOTATATE infusion.

2. Dosage:

The recommended dosage is 200 mCi (7.4 GBq) Lu-177 DOTATATE, administered every 8 weeks for a total of 4 doses as tolerated. Dosage can be halved, according to the US Food and Drug Administration (FDA)–approved clinical notes, in special clinical situations, such as hematological toxicity [<u>32</u>].

Prophylaxis: Amino Acid Solution and Antiemetics:

The Lu-177 DOTATATE solution needs to be administered with concomitant amino acid infusion to reduce radiation absorbed dose and toxicity to the kidneys. Amino acid infusion should be initiated 30 minutes prior to infusion of Lu-177 DOTATATE and continued for at least 4 to 5 hours after completion of PRRT. There are different amino acid formulations available. The extemporaneously compounded formulation contains only 25 g lysine HCl and 25 g arginine HCl with 1 L of appropriate sterile solvent (eg, water for injection). This formulation has lower osmolality and less patient emetic effects. The commercially available amino acid solutions have a lysine content between 18 and 24 g and arginine content between 18 and 25 g in a volume of 1.5 - 2.2 L of solution having <1,050 mOsmol/L. Aminosyn II 10% used in clinical trials in the United States contained additional essential and nonessential amino acids as well as electrolytes resulting in osmolality of 1,040 mOsmol/L. This preparation was associated with a high incidence of nausea and vomiting. Choice of amino acid formulations depends on institutional resources.

Due to nausea with or without vomiting observed in some patients receiving amino acid infusion, it is advised that use of prophylactic antiemetic medications be considered, as used in each institution with any chemotherapy, 30 minutes prior to commencing amino acid solution administration. Other adjunct treatment for persistent vomiting is reasonable depending on physicians' experiences.

3. Infusion Methods:

It is highly preferred that the IV access for administration of Lu-177 DOTATATE solution be separate from IV access for amino acid infusion. Separate access allows removal of the radiopharmaceutical access materials from the patient after PRRT, ensuring no radioactive medical line leaves the confines of the administering facility. Prior to infusion, measure the source activity to confirm prescribed activity. In some centers, a double lumen peripherally inserted central catheter (PICC) line can be used for infusion to avoid delivery failures.

Lu-177 DOTATATE is delivered in a vial under positive pressure. It can be administered via gravity method, infusion pump method or via automated syringe pump injector, as detailed with illustrative figure at the available link: <u>http://jnm.snmjournals.org/content/60/7/937/F3.expansion.html</u> [26]. Each institution can choose the best technique of radiopharmaceutical administration.

Gravity Method:

- Insert a 2.5-cm-long, 20-gauge needle (short needle) into the Lu-177 DOTATATE vial, ensuring that the beveled tip inside the vial does not touch the solution at any time during the infusion. The hub of the short needle is fastened to the IV tubing of a previously prepared 500-mL sterile 0.9% sodium chloride solution. Keep the IV tubing clamp closed until the entire setup has been completed and is ready for infusion.
- Insert a second needle that is 9 cm long, 18 gauge (long needle) into the Lu-177 DOTATATE vial, ensuring that the beveled tip of this long needle touches and is secured to the bottom of the vial during the entire infusion. Fasten a connecting tube prefilled with sterile 0.9% sodium chloride to the hub of the long needle, ensuring that there are no air bubbles inside the plastic tubing. Check the designated IV access for Lu-177 DOTATATE to ensure patency; once confirmed, fasten the male Lauer lock of the connecting tube to the IV access, keeping clamp closed.
- Do not remove the needles to reposition once the seal is punctured, as this may make the seal

ineffective and prevent dose delivery by this method.

- Open the clamp in the connecting tube from the vial to the patient, and then open the clamp of the tubing from the bag of normal saline solution. Regulate the flow of the sodium chloride solution via the short needle into the Lu-177 DOTATATE vial at a rate of 50 mL/h to 100 mL/h for 5 to 10 minutes and then 200 mL/h to 300 mL/h for an additional 25 to 30 minutes. During infusion, ensure that the level of solution in the Lu-177 DOTATATE vial remains constant and that the vial does not fill up completely. Total duration of infusion is about 30 to 40 minutes.
- Do not administer Lu-177 DOTATATE as an IV bolus.
- Clamp the saline line once the level of radioactivity is stable for at least five minutes.
- Clamp the connecting line from the vial and disconnect from the long needle, taking care that no fluid spills out. Open the connecting tube again and flush with 25 mL of 0.9% sterile sodium chloride to wash off any radiopharmaceutical remaining within the tubing into the patient.
- Remove the IV access used. Measure the remaining activity in the setup, including the vial, and subtract from the measured preinfusion activity to obtain the net activity administered.

Infusion Pump Method:

For the infusion pump method, the short and long needles are also used. The tubing that connects to the long needle should be primed with normal saline solution before attachment to an infusion pump. The other end of this tubing is attached to the IV access of the patient. A 3-way stopcock is connected to the hub of the short needle before it is inserted into the vial with a filter attached to the vent tip. Again, the tip of the short needle should stay above the fluid level, whereas the tip of the long needle is at the bottom of the vial. The positive pressure within the Lu-177 DOTATATE vial drives fluid into the patient and is controlled by the infusion pump, which is usually programmed to deliver 0.8 to 0.9 mL/min for total infusion time of 25 to 30 minutes. Remove the IV access used. Measure the remaining activity in the setup, including the vial, and subtract from the measured preinfusion activity to obtain the net activity administered.

Automated Syringe Pump Injector Method:

Another method involves drawing the Lu-177 DOTATATE solution from inside the vial into a sterile lead-shielded syringe that is then mounted on an automated syringe pump injector to administer the Lu-177 DOTATATE. This method exposes the individual drawing the solution to radiation risk. A connecting tube prefilled with sterile 0.9% sodium chloride solution is used to connect the syringe containing the radiopharmaceutical to the IV access of the patient. Before starting the infusion, confirm patency of patient's IV access. The pump is programmed to deliver the contents of the syringe over 30 minutes, eg, 30 mL at 60 mL/h . Once infusion is completed, the connecting tube can be flushed with 25 mL of 0.9% sterile sodium chloride to wash off any radiopharmaceutical remaining within the tubing into the patient. Attention is required to safely handle the setup to avoid spillage as well as minimize radiation exposure by using tongs. Remove IV access, use and measure remaining radioactivity in the setup and vial, and subtract it from preinfusion activity to determine net activity administered.

C. Post-therapy Management

All personnel involved with Lu-177 DOTATATE therapy should perform a survey of their hands and clothing for any contamination, and appropriate measures should be performed if such contamination is discovered. The room used for infusion should be surveyed for contamination before releasing the room to another patient. All medical wastes associated with the PRRT should be stored as required by radiation safety procedures, making sure that they are separated from other wastes associated with short-acting radiopharmaceuticals.

Care of the patient after Lu-177 DOTATATE therapy follows established institutional protocol for care of patient after radionuclide therapy with special consideration to ALARA principles. Therapy with octreotide LAR or lanreotide is usually given 4 to 24 hours after Lu-177 DOTATATE at the discretion of the attending oncology physician and stopped 4 weeks prior to subsequent PRRT. Short-acting octreotide maybe given for symptomatic management during PRRT cycles and withheld 24 hours prior to next dose of Lu-177

DOTATATE after determination by treating team of physicians.

If desired, posttherapy 3-D imaging may be obtained for the purposes of dosimetry. Personalized dosimetry may be used to assess and estimate potential risk to organs for the individual patient, as data collection for correlative studies seeking to establish maximum organ dose thresholds or lesion treatment efficacy thresholds, or for dose reporting in case of future radiation treatments [26].

V. DOCUMENTATION

Reporting should be in accordance with the <u>ACR–ASTRO Practice Parameter for Communication: Radiation</u> <u>Oncology</u> [21].

A summary of the patient's history, pathologic findings, imaging results and laboratory findings should be included in the report to document the indication and tolerability for treatment with Lu-177 DOTATATE. The report should include the radiopharmaceutical used, the administered activity, site and route of administration, safety precautions for other staff involved in the patient's care, and any associated incident encountered during therapy. If dosimetry is performed, salient organ absorbed dose values, both in directly calculated dose and in equivalent dose (EQD2), should be reported, and, if available, a dose map in DICOM format with the associated CT. On subsequent PRRT, interval history should include a summary of prior Lu-177 DOTATATE treatments, interval imaging to assess treatment efficacy, and pertinent laboratory findings to determine and confirm appropriateness and safety of additional therapy [<u>26</u>].

VI. STATEMENT OF THERAPEUTIC USE OF UNSEALED RADIOPHARMACEUTICAL SOURCES

On the basis of their education, training pathway(s), initial board certification(s), and maintenance of certification(s), NRC or Agreement State Authorized User (AU) status, and clinical work experience, diagnostic radiologists (DRs), nuclear radiologists (NRs), nuclear medicine physicians (NMs), and radiation oncologists (ROs) may have the qualifications to supervise and perform therapy with Lu-177 DOTATATE. Although it is recognized that individual physician variations and state and federal regulatory requirements may, of necessity, dictate site-specific practice patterns, these physicians may best participate in the practice according to their special interests and qualifications. In most clinical settings, one of the following common practice paradigms generally applies:

- Physicians who are NRC and/or Agreement State recognized, board-eligible or board-certified in DR, NR, NM, or RO but do not hold AU status. These physicians may participate in the practice of PRRT under the supervision of an AU. Although they may not issue written directives for Lu-177 DOTATATE, they may administer such a dosage as designated and supervised by an AU.
- Physicians who are NRC and/or Agreement State—recognized and board certified in DR, NR, NM, or RO and hold AU status based on that certification and site-specific credentialing: These physicians may administer Lu-177 DOTATATE therapy under their own AU qualifications and licensure.
- Physicians who are NRC and/or Agreement State—recognized and board certified in DR, NR, NM, or RO and hold the appropriate AU statuses and site-specific credentialing. These physicians may practice parenteral Lu-177 DOTATATE therapy as permitted by their own specific training leading to such AU statuses.

VII. EQUIPMENT SPECIFICATIONS

Equipment performance monitoring should be in accordance with the <u>ACR–AAPM Technical Standard for Nuclear</u> <u>Medical Physics Performance Monitoring of Gamma Cameras [33]</u>.

VIII. RADIATION SAFETY

Radiologists, medical physicists, non-physician radiology providers, radiologic technologists, and all supervising physicians have a responsibility for safety in the workplace by keeping radiation exposure to staff, and to society as a whole, "as low as reasonably achievable" (ALARA) and to assure that radiation doses to individual patients are appropriate, taking into account the possible risk from radiation exposure and the diagnostic image quality necessary to achieve the clinical objective. All personnel who work with ionizing radiation must understand the key principles of occupational and public radiation protection (justification, optimization of protection, application of dose constraints and limits) and the principles of proper management of radiation

dose to patients (justification, optimization including the use of dose reference levels). <u>https://www-pub.iaea.org/MTCD/Publications/PDF/PUB1775_web.pdf</u>

Facilities and their responsible staff should consult with the radiation safety officer to ensure that there are policies and procedures for the safe handling and administration of radiopharmaceuticals in accordance with ALARA principles. These policies and procedures must comply with all applicable radiation safety regulations and conditions of licensure imposed by the Nuclear Regulatory Commission (NRC) and by applicable state, local, or other relevant regulatory agencies and accrediting bodies, as appropriate. Quantities of radiopharmaceuticals should be tailored to the individual patient by prescription or protocol, using body habitus or other customized method when such guidance is available.

Nationally developed guidelines, such as the <u>ACR's Appropriateness Criteria</u>[®], should be used to help choose the most appropriate imaging procedures to prevent unnecessary radiation exposure.

Additional information regarding patient radiation safety in imaging is available from the following websites – Image Gently[®] for children (<u>www.imagegently.org</u>) and Image Wisely[®] for adults (<u>www.imagewisely.org</u>). These advocacy and awareness campaigns provide free educational materials for all stakeholders involved in imaging (patients, technologists, referring providers, medical physicists, and radiologists).

Radiation exposures or other dose indices should be periodically measured by a Qualified Medical Physicist in accordance with the applicable ACR Technical Standards. Monitoring or regular review of dose indices from patient imaging should be performed by comparing the facility's dose information with national benchmarks, such as the ACR Dose Index Registry and relevant publications relying on its data, applicable ACR Practice Parameters, NCRP Report No. 172, Reference Levels and Achievable Doses in Medical and Dental Imaging: Recommendations for the United States or the Conference of Radiation Control Program Director's National Evaluation of X-ray Trends; 2006, 2009, amended 2013, revised 2023 (Res. 2d).

IX. QUALITY CONTROL AND IMPROVEMENT, SAFETY, INFECTION CONTROL, AND PATIENT EDUCATION

Policies and procedures related to quality control and improvement, patient education, infection control, and safety should be developed and implemented in accordance with the ACR Policy on Quality Control and Improvement, Safety, Infection Control, and Patient Education appearing under the heading *ACR Position Statement on Quality Control and Improvement, Safety, Infection Control and Patient Education* on the ACR website (https://www.acr.org/Advocacy-and-Economics/ACR-Position-Statements/Quality-Control-and-Improvement).

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Writing Committee – members represent their societies in the initial and final revision of this practice parameter

ACR	ACNM
Neil B. Desai, MD, Co-Chair	Lorraine E. DeBlanche, MD, FACNM
Charito Love, MD, Co-Chair	Tony Francis, MD

Tony Abraham, DO, MPA	Bital Savir-Baruch, MD	
Erin C. Grady, MD	Ila Sethi, MD	
Jeffrey S. Kempf, MD, FACR		
Rathan M. Subramaniam, MD, PhD, MPH		
Ying Xiao, PhD		
<u>ASTRO</u>	<u>SNMMI</u>	
Thomas Boike, MD	Lisa Bodei, MD	
Michael M. Dominello, DO	David L. Bushnell, MD	
Robert F. Hobbs, PhD	Thomas Hope, MD	
William Rule, MD	Daniel A. Pryma, MD	
Committee on Practice Parameters and Technical Standards – Nuclear Medicine and Molecular Imaging		
(ACR Committee responsible for sponsoring the draft through the process)		
Kevin P. Banks, MD, Co-Chair	Andrew Kaiser, MD	
Richard K. J. Brown, MD, FACR, Co-Chair	Jeffrey S. Kempf, MD, FACR	
Munir V. Ghesani, MD, FACR, Co-Chair Vice Chair	Jennifer J. Kwak, MD	
Rathan M. Subramaniam, MD, PhD, MPH, Co-Chair Vice Chair	Justin G. Peacock, MD	
Esma A. Akin, MD, FACR	Syam P. Reddy, MD	
Alexandru C. Bageac, MD, MBA	Eric M. Rohren, MD, PhD	

Committee on Practice Parameters and Technical Standards – Nuclear Medicine and Molecular Imaging

(ACR Committee responsible for sponsoring the draft through the process)

Twyla B. Bartel, DO, MBA	Levi Sokol, MD	
Elizabeth H. Dibble, MD	Andrew T. Trout, MD	
K. Elizabeth Hawk, MD, MS, PhD	Stephanie P. Yen, MD	
Eric Hu, MD		
Committee on Practice Parameters – Radiation Oncology		
(ACR Committee responsible for sponsoring the draft through the process)		
Naomi R. Schechter, MD, Chair	Join Y. Luh, MD	
Nathan H. J. Bittner, MD	Matthew Poggi, MD	
Samuel T. Chao, MD	Helen A. Shih, MD	
Neil B. Desai, MD	Paul E. Wallner, DO, FACR	
Beth A Erickson-Wittmann, MD	Kristina L. Woodhouse, MD	
Matthew Harkenrider, MD	Ying Xiao, PhD	
Mark Hurwitz, MD	Sue S. Yom, MD, PhD	
Don C. Yoo, MD, FACR, Chair, Commission Nuclear Medicine and Nuclear Medicine		
William Small, Jr., MD, FACR, Chair, Commission on Radiation Oncology		
Jacqueline Anne Bello, MD, FACR, Chair, Commission on Quality and Safety		
Mary S. Newell, MD, FACR, Chair, Committee on Practice Parameters and Technical Standards		

Comments Reconciliation Committee

K. Elizabeth Hawk, MD, MS, PhD, Chair	Erin C. Grady, MD
Ralph P. Lieto, MS, FACR, Co-Chair	Robert F. Hobbs, PhD
Tony Abraham, DO, MPA	Thomas Hope, MD
Kevin P. Banks, MD	Jeffrey S. Kempf, MD, FACR
Jacqueline Anne Bello, MD, FACR	Hyun Kim, MD
Pradeep Bhambhvani, MD	Amy L. Kotsenas, MD, FACR
Lisa Bodei, MD	Paul A. Larson, MD, FACR
Thomas Boike, MD	Charito Love, MD
David L. Bushnell, MD	Mary S. Newell, MD, FACR
Richard K.J. Brown, MD, FACR	Zoubir Ouhib, MS, FACR
Lorraine E. De Blanche, MD, FACNM	Daniel A. Pryma, MD
Neil B. Desai, MD	William Rule, MD
Michael M. Dominello, DO	Naomi R. Schechter, MD
Richard Duszak Jr., MD, FACR	lla Sethi, MD
Samuel A. Einstein, PhD	William Small, Jr., MD, FACR
Saeed Elojeimy, MD, PhD	Rathan M. Subramaniam, MD, PhD, MPH
Tony Francis, MD	Ying Xiao, PhD
Munir V. Ghesani, MD, FACR	Don C. Yoo, MD, FACR

Comments Reconciliation Committee

Michael Goris, MD

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*Practice parameters and technical standards are published annually with an effective date of October 1 in the year in which amended, revised, or approved by the ACR Council. For practice parameters and technical standards published before 1999, the effective date was January 1 following the year in which the practice parameter or technical standard was amended, revised, or approved by the ACR Council.

2020 (Resolution 17)

Amended 2023 (Resolution 2c, 2d)

Appendix A

Post Treatment Instructions to Patient Following Lu-177 DOTATATE Therapy

Name of Patient: ______ Medical Record Number: _____

Last name, First name

Date of Treatment: ______ Isotope: Lu-177 Activity: _____

_____, please show this form to every physician, healthcare worker or emergency Before this date: personnel that provide you care.

Special Precautions

- 1. Maintain a distance of at least 3 feet (1 meter) from others for 3 days since radiation exposure decreases with distance, the further away you are from others, the less radiation they get.
- 2. Minimize visits by family or friends for 3 days. If you have visitors, try to stay at least 3 feet away.
- 3. Minimize close contact with others at work for 3 days.
- 4. For woman of childbearing age (<55 years old), pregnancy must be excluded before initiating the treatment. Both men and women of child-bearing potential must refrain from procreation by using effective contraceptive methods during the treatment and for 6 months after.
- 5. For women who are breastfeeding, discontinue breast feeding for this child.
- 6. Sleep alone for at least 7 nights. Sleeping together with another adult exposes them to the radiation coming from you. Sexual activity is not advised for 7 days after LUTATHERA administration.
- 7. No children should sleep with you for 7 days. No pregnant person should sleep with you for 15 days. Children and fetuses are much more sensitive to radiation than adults, and special care is needed. Limit close contact to brief periods for 7 days.
- 8. Take particular care when urinating for 3 days. Toilets must be used in a seated position, even for men. Use private bathroom or flush 2 to 3 times and clean any spills with disposable gloves and damp cloth after each use. Wash hands thoroughly. The radiation leaves your body mainly from your urine.
- 9. Wash dishware and utensils and bathroom accessories separately for 3 days.
- 10. Do not travel by public transportation (bus, train, plane) for more than 1 hour for 3 days. If you are planning to travel while radiation safety precautions are in effect, please inform Nuclear Medicine or Radiation Facility personnel at (area code) ______.

**For flights, travelers may call TSA Cares toll free at 1-855-787-2227 prior to traveling with questions about screening policies, procedures, and what to expect at the security checkpoint. For more information, visit https://www.tsa.gov/travel/special-procedures.

- 1. No prolonged car trip (more than 1 hour) with others for 3 days.
- 2. Drink plenty of fluid (4-8 glasses) per day for 3 days.

If you are admitted to the ER or hospitalized while radiation safety precautions are in effect, inform the hospital staff to notify the above contact person immediately. During off-hours, contact Nuclear Medicine or Radiation Facility via the operator at _

Instructions for Radioactive Trash Generated by patient

Please be aware that the following items that may be contaminated with urine and blood cannot be disposed into regular trash:

- 1. Pads, tampons
- 2. Toilet papers, tissue
- 3. Towels, linens, sheets
- 4. Any other items that are contaminated with urine, blood, and wound or drainage secretions for the first 3 days post treatment

Towels, linens and sheets can be washed separately and reused.

Toilet paper and tissue need to be flushed down the toilet.

Any other contaminated items that cannot be washed or flushed down the toilet needs to be kept for at least 70 days or bring it to the Nuclear Medicine or Radiation Oncology Facility to be stored.

Rbaisede2022th@aboveipnetautions and instructions and have spoken with the Nuclear Medicine or Radiation Facility personnel and agree to comply.

Patient (print name):		
Signature:	Date/time:	
Witness (print name)		
Signature:	Date/time:	
Authorized User or Supervised Designee (print name)		
Signature:	Date/time:	