Reimbursement Approaches for Radiopharmaceutical Dosimetry: Current Status and Future Opportunities

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Interest in performing dosimetry for clinical radiopharmaceutical therapy procedures has grown in recent years. Several approved therapies include dosimetry in the Food and Drug Administration–approved label instructions, and other therapies are best used under a patient-tailored paradigm. This paper, which is a product of the Society of Nuclear Medicine and Molecular Imaging Dosimetry Task Force, presents motivations and general workflows for radiopharmaceutical therapy dosimetry, as well as existing strategies for obtaining reimbursement for clinical activities related to dosimetry. Several specific patient examples are provided, including suggested codes for reimbursement. In addition to current reimbursement approaches, key dosimetry services that are not supported under the current coding structure are presented and suggested as areas of focus in the coming years.

Key Words: RPT; dosimetry; SPECT/CT; PET/CT

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Radiopharmaceutical therapy (RPT) is a rapidly growing oncologic intervention whereby electron- or α-emitting radionuclides, formulated for accumulation within or near cancer cells, are administered by intravenous, intraarterial, or interstitial injection. The mechanism for accumulation within or near cancer cells can be physical in nature, such as 90Y-labeled microspheres that become trapped in the arterioles of hypervascular lesions, or biochemical in nature, such as the binding of a radiolabeled peptide or antibody to a biologic receptor.

In most cases, the mass of radiolabeled compound administered for therapy is below any threshold for pharmacologic effects, and it is primarily the energy imparted into tissue by radioactive decay that effects a therapeutic response. The interactions between radiation and human biology—including biologic effects—have been extensively investigated over the last approximately 125 y (1,2). The primary endpoints of radiation therapy are so-called deterministic effects in target and nontarget tissue. (Stochastic effects, such as secondary hematologic malignancies, have also been shown to result from radiation exposure and chemotherapy. Current models suggest that these effects are not associated with a dose threshold, but rather the effect risk is thought to increase with increasing cumulative treatment. Rather than individualized dosimetry for toxicity avoidance, stochastic effects are better informed by population-level dosimetry data for risk modeling.) Examples of deterministic effects include radiation-induced nephropathy (kidney damage) and radiation-induced tumor shrinkage. Deterministic effects, which are the product of cell killing, are associated with a dose threshold, below which no effect is observed. Beyond the radiation dose threshold, the severity or magnitude of a deterministic effect is expected to increase with increasing dose. These dose-dependent effects for various biologic endpoints, tissue types, radiation types, and dose rates have been described in literature.

Radiation dose from RPT is therefore a measure that is expected to correlate with tumor control probability and normal-tissue complication probability. Indeed, there is a growing body of evidence showing that dose–response relationships are observed in RPT (3–10). Although not covered in this paper, the current state of knowledge regarding normal-tissue toxicity relationships and dose–response relationships within the context of RPT is thoroughly described in 2 other papers within this dosimetry supplement. Dose to tumors and normal tissues can vary widely among patients for a given administered activity level due to differences in tissue mass, pharmacokinetics, tissue geometries, and tumor phenotype (11–13). It is therefore critical to monitor patient-specific radiation-absorbed dose by established dosimetry techniques, whereby appropriate changes in management may be made. As with other types of radiation therapy, applying these therapies under a dosimetry-guided paradigm allows clinicians to minimize the risk of long-term toxic side effects, as well as assess for potential benefit in a particular patient.

**DOSIMETRY FOR RPT**

The process of obtaining patient-specific dosimetry for RPT involves characterizing the time-ordered distribution of radiopharmaceutical in the body, especially those tissues that are receiving the greatest radiation dose, or those that are naturally most sensitive to radiation. Techniques currently available for obtaining data regarding the distribution of radiopharmaceutical in a patient include the following: whole-body (WB) emission counting (1-dimensional projection of γ-emitting activity in a patient); planar γ-imaging (2-dimensional projection of γ-emitting activity in a patient); SPECT imaging (3-dimensional [3D] reconstruction of γ-emitting activity in a patient); PET imaging (3D reconstruction of β+-emitting activity in a patient); and blood or urine sampling (average activity concentration in compartment).

Generally, it is not optimal to rely on the use of only one of these technologies independently for patient-specific dosimetry, as each has weaknesses. With that said, the dosimetric accuracy and precision that can be obtained by use of only one of these data-collection techniques may be appropriate depending on

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particular RPT and specific patient management needs. Additionally, the number of data-collection time points can influence the accuracy of dosimetric calculations, with increased data collection being associated with improved dosimetric precision (11,14–19). Considerations needed when developing a dosimetry plan for a given RPT and patient should include: tissues of interest for dosimetry, potential impact of dosimetry on patient management (thus necessitating a certain level of accuracy and precision), and the ability of a patient to undergo dosimetric data collection. Even within a particular RPT, these factors vary on a per-patient basis, thus necessitating flexibility in dosimetry methods and associated reimbursement mechanisms.

Normal organs receiving the highest levels of absorbed dose in the body tend to be organs that are involved in concentrating and excreting the radiopharmaceutical, such as the liver, kidneys, bladder, and gastrointestinal tract. Significant radiation dose is also commonly observed in the spleen and secretory tissues (salivary glands, adrenal glands, pituitary gland). Although not typically receiving the highest absorbed dose, the bone marrow is a particularly radiosensitive tissue, and one that is of importance in RPT dosimetry. For a given RPT, usually only 1 or 2 of these organs will limit the quantity of radiopharmaceutical that can be administered without exceeding toxicity thresholds. A summary of approved and late-stage investigational agents, and their most commonly limiting normal organ tissues (20–29), is listed in Table 1.

In addition to consideration of dose-limiting normal organs, tumor dosimetry provides valuable information regarding potential patient benefit or the need for modifications to administered activity to reach a certain probability of benefit. Although an extensive review of tumor and normal organ dose–response relationships is beyond the scope of this document, typically solid tumor doses (from low–linear energy transfer sources) in excess of 100 Gy are needed to achieve high rates of response, whereas doses of less than approximately 50 Gy often do not provide therapeutic benefit from RPTs (7,9,10,30–33). In some cases, potential patient benefit may be minimal, thereby leading to a decision to not proceed with therapy. In this situation, unnecessary radiation exposure to the patient and public can be avoided, as well as an overall reduction in health-care costs. On the other hand, if a patient’s organ dosimetry is favorable, and tumor targets could benefit from dose escalation, it likely makes sense to administer additional radioactivity to achieve optimal therapeutic outcomes.

<table>
<thead>
<tr>
<th>Radiopharmaceutical Indication</th>
<th>Dose-limiting Tissues</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>131I-Nal</strong></td>
<td>Thyroid cancers</td>
</tr>
<tr>
<td><strong>90Y-microspheres</strong></td>
<td>Intrahepatic tumors, including primary and metastatic disease</td>
</tr>
<tr>
<td><strong>177Lu-DOTATATE</strong></td>
<td>Low-grade neuroendocrine tumors</td>
</tr>
<tr>
<td><strong>131I-MIBG</strong></td>
<td>Paraganglioma, pheochromocytoma</td>
</tr>
<tr>
<td><strong>223RaCl2</strong></td>
<td>Metastatic castration-resistant prostate cancer (mCRPC)</td>
</tr>
<tr>
<td><strong>177Lu-PSMA-617 (investigational)</strong></td>
<td>Metastatic castration-resistant prostate cancer (mCRPC)</td>
</tr>
<tr>
<td><strong>177Lu-DOTATOC (investigational)</strong></td>
<td>Low-grade neuroendocrine tumors</td>
</tr>
<tr>
<td><strong>131I-Iomab-B (investigational)</strong></td>
<td>Acute myeloid leukemia (AML)</td>
</tr>
</tbody>
</table>

## PRIMARY STAKEHOLDERS

Like any medical service and procedure, dosimetry for RPT needs to meet the requirements and expectations of an array of health-care stakeholders that span the entire billing process. Stakeholder interests should be contextualized in terms of the marginal increase in the cost of care, which overall tends to be dominated by the radiopharmaceutical cost in these procedures. Indeed, Centers for Medicare and Medicaid Services (CMS) reimbursement for 177Lu-DOTATATE or 131I-metaiodobenzylguanidine (131I-MIBG) often exceeds $200,000 for a course of therapy. By comparison, CMS reimbursement for services relating to dosimetry and treatment planning is unlikely to exceed $10,000–$15,000 for a course of therapy, depending on the workflow (see “Specific Coding Examples” for details). This represents, at most, a 5%–7% increase in the total cost of care. With this as context, stakeholder interests relating to dosimetry are described below.

- The most important group of stakeholders, patients, benefits from improved quality of care. In a given patient, dosimetry-guided RPT has significant potential for toxicity prevention, tumor control improvement, or total avoidance of futile medical intervention.
- Despite the expense of performing dosimetry, medical payers are expected to see a reduction in long-term costs due to avoidance of unnecessary (and typically vastly expensive) cancer therapies in a subset of patients, as well as potential for improved patient outcomes, which further reduces expense liabilities.
- Clinicians stand to benefit from cost-recovery on existing dosimetry practices, reduced liability from adoption of dosimetric guidance (avoiding over-, under-, and futile administration of RPT), and by remaining competitive in offering the highest level of care possible for patients.
- Radiopharmaceutical development and manufacturing entities can benefit from increases in administered activity to patients who stand to benefit most from doing so, and potentially from improved therapeutic windows in late-stage trials (thus reducing the number of patients needed to conduct trials).
- Technical providers of imaging services stand to benefit from increased use of imaging services, in particular existing γ-camera imaging infrastructure.
- The general public stands to benefit from improved control over the release of radioactive patients, which results in...
DOSIMETRY TECHNIQUES

General Workflows
As mentioned in the section “Dosimetry for RPT,” multiple data-collection methods are available for dosimetry. Additionally, different radiopharmaceuticals have workflows that are conducive to their typical administration schedule. For example, high-specific-activity 131I-MIBG (Azeda; Progenics Pharmaceuticals Inc.) is nominally administered as 2 treatments separated by at least 90 d, whereas 177Lu-DOTATATE (Lutathera; Novartis) is administered over 4 therapeutic administrations, each separated by approximately 60 d. Given the goals of using dosimetry to enhance the safety and efficacy of RPT, it is important to have dosimetry results at a time or times in which treatment decisions can be made. In the case of fractionated therapies (e.g., 177Lu-DOTATATE [Lutathera]), acquiring dosimetry data after the administration of each therapy can allow for adaptation in subsequent administrations to meet specific treatment planning goals. In the case of high-specific-activity 131I-MIBG (Azeda), however, this may or may not be possible, due to potentially reaching or exceeding normal-tissue limits in the first treatment. Likewise, 90Y-microsphere therapies are often administered with a single intraarterial infusion, in which case dosimetry and treatment planning are needed before the first therapeutic administration. Therefore, the 2 main dosimetry/treatment planning workflows are as follows:

- Administration of a small amount of the therapeutic, or a predictive surrogate, for purposes of dosimetry and treatment planning before administration of the primary RPT. This workflow is typically used for 90Y-microspheres, 131I-MIBG, and 131I-NaI.
- Administration of a full RPT administration, followed by dosimetry for modification of subsequent treatments. This workflow is commonly used for 177Lu-DOTATATE and could be used for various agents currently under investigation.

Dosimetric Sampling
Within a given workflow, a dosimetry schedule should be created based on the needs of a particular RPT and patient. The goal of this schedule should be the accurate determination of dose to relevant tissues (dose-limiting organs or tumors); however, the exact imaging and data-collection sequence will vary with situation. Several specific schedule examples are presented in the section “Specific Coding Examples,” however, the following general statements can be made regarding the dosimetry of each agent and tissue type.

131I-NaI. Thyroid uptake should confirmed and quantified with pretreatment imaging. SPECT/CT and planar γ-imaging are appropriate for this when using 124I-NaI or 131I-NaI, and PET/CT is appropriate when using 124I-NaI. Generally, a single imaging time point is adequate for determination of initial tumor uptake fraction; however, quantification of dose to tumor requires anatomic imaging, for example, PET/CT or SPECT/CT, and multiple imaging time points. Quantification of dose to lungs, relevant in cases in which significant lung metastatic disease exists, requires multiple imaging time points and at least 1 anatomic reference scan (e.g., SPECT/CT or PET/CT). Accurate quantification or prediction of marrow dosimetry requires blood sampling at multiple time points and WB planar or SPECT/CT imaging at multiple time points. In summary, a complete and optimal dosimetry workup requires serial blood sampling, serial WB planar imaging, and at least one SPECT/CT that is concordant with one of the planar imaging time points. Some practices have developed population-based biologic clearance models, which may allow for a reduction in the needed data (omission of one or more planar or blood sampling time points); however, these approximations may reduce dosimetric accuracy somewhat (35–37).

131I-MIBG and 177Lu-DOTATATE. For both agents, marrow and kidney dosimetry are normal organs of interest. Optimal renal dosimetric sampling can be achieved by multiple SPECT/CT imaging time points over the first approximately 7 d after administration of the therapeutic or a surrogate. Bone marrow dosimetry for these agents can be performed by addition of WB planar imaging and blood sampling at multiple time points. The blood contribution to total marrow dose is less for 131I-MIBG than for 177Lu-DOTATATE, and therefore fewer collections may be needed. Tumor dosimetry, similar to kidney dosimetry, is best performed with serial SPECT/CT imaging. Some investigators have proposed 177Lu-DOTATATE imaging time-point reduction strategies for kidneys and tumors (11,18,19); however, these approximations may reduce dosimetric accuracy somewhat (14). Similarly, rather than WB imaging to determine marrow dose from 131I-MIBG, some recommendations include the use of WB counting (rather than imaging) in pediatric patients who would otherwise require general anesthesia for imaging (38,39).

222RaCl2. Because of the low administered activity and photon emission abundance, the retention and distribution of 222Ra in a patient is typically assessed by planar imaging only; however, quantitative SPECT/CT has been investigated (40–43).

90Y-Microspheres. 90Y-microspheres are unique among RPTs, due to their nature of maintaining a fixed irradiation geometry after administration. Because of this, only a single imaging time point is needed for dose assessment. For treatment planning purposes, typically 99mTc-macroaggregated albumin (99mTc-MAA) is administered in a way that is consistent with the desired 90Y-microsphere administration method (same catheter position in the hepatic arterial tree, same infusion rate). Dose to tumor, liver, and potentially lung and stomach are of interest after this MAA administration. Liver and tumor dosimetric predictions, as well as evaluation for gastric shunting, are made by way of a single SPECT/CT image after administration of MAA. The axial field of view of a single SPECT/CT acquisition is often not adequate for inclusion of the entire lungs, thus lung shunting should be evaluated by collection of an additional SPECT/CT scan, or by conjugate-view planar imaging. In general, lung shunting has been reported to be overestimated by planar imaging, and therefore SPECT/CT may be preferred (44–46). After administration of 90Y-microspheres, additional 3D imaging at a single time point (SPECT/CT or PET/CT) is needed to confirm microsphere distribution and associated dosimetry. Some discordance is expected when comparing 99mTc-MAA predicted dosimetry and 90Y-SPECT/CT estimated dosimetry due to the difference in image quality; however, comparison of these measurements can confirm general treatment distribution and potential eligibility for subsequent 90Y-microsphere administrations in the case of progression or undercoverage.

Simplified Dosimetry Methods
As mentioned above, simplified dosimetry methods for 131I-NaI, 177Lu-DOTATATE, and 131I-MIBG have been proposed (18,19,35,
Dosimetry Calculations

Two main methods exist for assessing patient-specific dosimetry, regardless of RPT type: absorbed fraction (e.g., MIRD schema) calculations, and 3D voxelwise dosimetry. These 2 methods are not mutually exclusive, meaning that in a single patient dose to 1 tissue (e.g., bone marrow) may be best assessed by an absorbed fraction calculation, whereas dose to another tissue may be best assessed by a 3D voxelwise dose calculation (e.g., liver). Precise methods and considerations regarding these 2 calculations methods are well described elsewhere (49–52) and thus beyond the scope of this document; however, a general diagram of dosimetry calculation steps, including final treatment plan generation, is shown in Figure 1. As described in the previous section, the exact combination of input data required for dosimetry depends on RPT- and patient-specific factors.

Similarly, the radiation dosimetry and treatment planning workflow will vary depending on specific information required by the physician provider for treatment planning purposes. The primary difference between these 2 dose calculation strategies is that absorbed fraction calculations typically result in mean dose to whole organs, whereas voxelwise calculations can provide a 3D dose map within the patient anatomy, including isodose lines and dose volume histograms. This distinction is relevant when considering the appropriateness of existing treatment planning current procedural terminology (CPT; a registered trademark of the American Medical Association) codes.

CPT CODE DESCRIPTIONS

Depending on the specific clinical workflow, personnel effort, documentation, and medical necessity, several existing CPT codes may be applicable to activities relating to radiopharmaceutical dosimetry and treatment planning. A list of existing and potentially pertinent codes and associated relative value units ([RVUs], data obtained from CMS.gov (53)) is provided in Table 2. Detailed

![Diagram of dosimetry calculation steps](image-url)
<table>
<thead>
<tr>
<th>CPT</th>
<th>Short description</th>
<th>Long description</th>
<th>Physician time (min)</th>
<th>Physician RVU</th>
<th>Physicist/technologist time (min)</th>
<th>Non-facility RVU</th>
<th>Facility RVU</th>
</tr>
</thead>
<tbody>
<tr>
<td>78800</td>
<td>Single area planar</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); planar, single area (e.g., head, neck, chest, pelvis), single day imaging</td>
<td>27</td>
<td>0.64</td>
<td>88</td>
<td>7.53</td>
<td>2.06</td>
</tr>
<tr>
<td>78801</td>
<td>Multiple area planar</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); planar, 2 or more areas (e.g., abdomen and pelvis, head and chest), 1 or more days imaging or single area imaging over 2 or more days</td>
<td>30</td>
<td>0.73</td>
<td>99</td>
<td>8.31</td>
<td>3.13</td>
</tr>
<tr>
<td>78802</td>
<td>WB single day</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); planar, WB, single day imaging</td>
<td>30</td>
<td>0.80</td>
<td>109</td>
<td>9.21</td>
<td>4.87</td>
</tr>
<tr>
<td>78804</td>
<td>WB 2 or more days</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); planar, WB, requiring 2 or more days imaging</td>
<td>40</td>
<td>1.01</td>
<td>216</td>
<td>19.42</td>
<td>–</td>
</tr>
<tr>
<td>78803</td>
<td>SPECT single area/single day</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); tomographic (SPECT), single area (e.g., head, neck, chest, pelvis), single day imaging</td>
<td>42</td>
<td>1.09</td>
<td>130</td>
<td>11.38</td>
<td>–</td>
</tr>
<tr>
<td>78830</td>
<td>SPECT/CT single area/single day</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); tomographic (SPECT) with concurrently acquired CT transmission scan for anatomic review, localization and determination/detection of pathology, single area (e.g., head, neck, chest, pelvis), single day imaging</td>
<td>45</td>
<td>1.49</td>
<td>141</td>
<td>14.46</td>
<td>–</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>CPT</th>
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<th>Long description</th>
<th>Physician time (min)</th>
<th>Physician RVU</th>
<th>Physicist/technologist time (min)</th>
<th>Non-facility RVU</th>
<th>Facility RVU</th>
</tr>
</thead>
<tbody>
<tr>
<td>78831</td>
<td>SPECT</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); tomographic (SPECT), minimum 2 areas (e.g., pelvis and knees, abdomen and pelvis), single day imaging, or single area imaging over 2 or more days</td>
<td>55</td>
<td>1.82</td>
<td>224</td>
<td>20.87</td>
<td>–</td>
</tr>
<tr>
<td>78832</td>
<td>SPECT/CT</td>
<td>Radiopharmaceutical localization of tumor, inflammatory process or distribution of radiopharmaceutical agent(s) (includes vascular flow and blood-pool imaging, when performed); tomographic (SPECT) with concurrently acquired CT transmission scan for anatomic review, localization and determination/detection of pathology, minimum 2 areas (e.g., pelvis and knees, abdomen and pelvis), single day imaging, or single area imaging over 2 or more days</td>
<td>60</td>
<td>2.12</td>
<td>264</td>
<td>27.19</td>
<td>–</td>
</tr>
<tr>
<td>78835</td>
<td>Quantification</td>
<td>Radiopharmaceutical quantification measurement(s) single area (list separately in addition to code for primary procedure)</td>
<td>17</td>
<td>0.47</td>
<td>23</td>
<td>3.00</td>
<td>–</td>
</tr>
<tr>
<td>78814</td>
<td>Limited PET/CT</td>
<td>PET with concurrently acquired CT for attenuation correction and anatomic localization imaging; limited area (e.g., chest, head/neck)</td>
<td>60</td>
<td>2.20</td>
<td>Carrier-priced</td>
<td>0.00</td>
<td>–</td>
</tr>
<tr>
<td>78580</td>
<td>—</td>
<td>Pulmonary perfusion imaging (e.g., particulate)</td>
<td>20</td>
<td>0.74</td>
<td>93</td>
<td>6.96</td>
<td>–</td>
</tr>
<tr>
<td>77300</td>
<td>Basic radiation dosimetry calculation</td>
<td>Basic radiation dosimetry calculation, central axis depth dose calculation, time-dose factor, nominal standard dose, gap calculation, off axis factor, tissue inhomogeneity factors, calculation of nonionizing radiation surface and depth dose, as required during course of treatment, only when prescribed by the treating physician</td>
<td>15</td>
<td>0.62</td>
<td>14</td>
<td>1.93</td>
<td>–</td>
</tr>
<tr>
<td>77370^3</td>
<td>—</td>
<td>Special medical radiation physics consultation</td>
<td>0</td>
<td>0.00</td>
<td>65</td>
<td>3.75</td>
<td>–</td>
</tr>
<tr>
<td>77261</td>
<td>—</td>
<td>Treatment planning: (simple, intermediate, complex)</td>
<td>36</td>
<td>1.30</td>
<td>–</td>
<td>2.06</td>
<td>2.06</td>
</tr>
<tr>
<td>77262</td>
<td>—</td>
<td>—</td>
<td>54</td>
<td>2.00</td>
<td>–</td>
<td>3.13</td>
<td>3.13</td>
</tr>
<tr>
<td>77263</td>
<td>—</td>
<td>—</td>
<td>82</td>
<td>3.14</td>
<td>–</td>
<td>4.87</td>
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</table>

(continued)
Stochastic effects, such as secondary hematologic malignancies, have also been shown to result from radiation exposure and chemotherapy. Current models suggest that these effects are not associated with a dose threshold, but rather the effect risk is thought to increase with increasing cumulative treatment. Rather than individualized dosimetry for toxicity avoidance, stochastic effects are better informed by population-level dosimetry data for risk modeling.

### Table 2

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<tr>
<th>CPT</th>
<th>Short description</th>
<th>Long description</th>
<th>Physician time (min)</th>
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<th>Non-facility RVU</th>
<th>Facility RVU</th>
</tr>
</thead>
<tbody>
<tr>
<td>77295</td>
<td>—</td>
<td>3-dimensional radiation treatment plan, including dose-volume histograms</td>
<td>112</td>
<td>4.29</td>
<td>165</td>
<td>14.07</td>
<td>--</td>
</tr>
</tbody>
</table>

**SPECIFIC CODING EXAMPLES**

Below is a series of clinical workflows that may be encountered, including reimbursement coding that is relevant to imaging, dosimetry, and treatment planning. We have intentionally omitted descriptions and codes related to radiopharmaceuticals, radiopharmaceutical administrations, patient consultation, and follow-up. Coding for these related activities are left for other documents.

**Example 1. 90Y-Radioembolization (with Pretreatment 99mTc-MAA Mapping)**

A patient was determined to be a candidate for 90Y radioembolization. Dosimetric planning for treatment began by preparation of a calibrated quantity of 99mTc-MAA. An interventional radiologist localized the catheter to a satisfactory location within the arterial supply of a liver for infusion of the 99mTc-MAA. Catheter tip placement was optimized based on tumor location and the perfused volume indicated by iodine-enhanced digital subtraction fluoroscopy or cone beam CT imaging in the intervention suite. After infusion of the 99mTc-MAA, the patient was relocated to a SPECT/CT scanner for imaging. Acquired were SPECT/CT images centered on the liver (78830) and a conjugate planar image including the extent of the lungs and liver (78830), and a conjugate planar image including the extent of the lungs and liver (bundled with 78830) were acquired. On the basis of institutional policy, this LSF was considered sufficiently low that a more accurate 3D evaluation was not needed. This LSF was transcribed in the patient medical record.

Dosimetry proceeded by use of 510(k)-cleared medical device software for 3D microsphere dosimetry. A physicist, physician, or another qualified individual segmented the whole liver, the perfused portion of the liver, the tumor, and the tumor plus a planning margin to account for breathing motion and potential microinvasion. A 3D dose plan normalized to a nominal administered activity was reviewed by the authorized user, and it was determined that an administered activity of 3.52 GBq was appropriate to maximize tumor dose, without exceeding dose limits to normal liver parenchyma. Three-dimensional dosimetry statistics for this final treatment plan were generated, and a treatment plan report was generated and signed by the physicist and authorized user (77295). An independent qualified individual reviewed this plan for appropriateness and accuracy, including performing a simplified dose calculation via the partition model. This secondary dose verification was documented in the medical record (77300).

Approximately 2 wk after the initial mapping procedure, the patient returned for treatment. The interventional radiologist placed the catheter tip at the same location within the liver arterial vasculature, and 90Y-microspheres were infused according to manufacturer-recommended methods. Because stasis was reached during administration, only 3.24 GBq were administered. The patient was transferred for posttreatment Bremsstrahlung SPECT/CT imaging, whereby a single SPECT/CT view, centered on the liver (78830), and a conjugate planar image including the extent of lungs and liver (bundled with 78830) were acquired. On the basis of these images, dosimetry was performed to assess the delivered dose (77300).

The following is a summary of the procedure and corresponding CPT codes:

- MAA mapping SPECT/CT and planar, 78830;
- Lung shunt quantification, 78835 (2 units);
- 3D radiation treatment planning, 77295;
- Plan check/simple dosimetry, 77300;
- 90Y SPECT/CT and planar, 78830; and
- Treatment verification (simple dosimetry), 77300.

**Comments.** If posttreatment 90Y PET/CT imaging covering the liver and lungs is performed rather than posttreatment SPECT/CT imaging, these codes may also be considered applicable in some situations.

Examples of these codes are included in the section “Specific Coding Examples”; however, the codes can generally be divided into those for γ-imaging (78800, 78801, 78802, 78803, 78804, 78830, 78831, 78832, 78850, 78858, 78814), dosimetry and treatment planning (77300, 77261, 77262, 77263, 77295), and ancillary services (77370, 78835). Notable exclusions from the table below include more general PET imaging codes (i.e., 78811, 78812, 78813, 78815, 78816), which might be applicable in the case where PET or PET/CT is used for pretreatment dosimetry. Codes relating to brachytherapy dosimetry and treatment planning (i.e., 77316, 77317, 77318) may also be considered applicable in some situations.

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and planar, 78814 (limited area PET/CT) would take the place of the posttreatment imaging code 78830. If pretreatment MAA mapping is not performed, the pretreatment imaging, LSF assessment, treatment planning codes, and treatment plan verification codes (78830, 78835, 77295, 77300) would not be applicable. If treatment planning is performed using methods other than a full-3D voxelwise calculation (e.g., partition method or whole liver mean dose determination), simple, intermediate, or complex treatment planning codes (77261, 77262, or 77263) should be used in place of 77295. If SPECT/CT is not acquired after 99mTc-MAA administration, 78800 or 78801 for the planar imaging would be billed in lieu of 78830. Assuming lung shunt quantification is performed from the planar images, 78850 could be used in lieu of 78835.

Secondary dose determination, simple, intermediate, or complex treatment planning codes, and treatment plan verification codes (77261, 77262, or 77263) should be used in place of 77295. If SPECT/CT is not acquired after 99mTc-MAA administration, 78800 or 78801 for the planar imaging would be billed in lieu of 78830. Assuming lung shunt quantification is performed from the planar images, 78850 could be used in lieu of 78835 units. Additionally, 77295 would no longer be applicable (see the earlier text).

Example 2. 177Lu-DOTATATE (with Tumor, Marrow, and Kidney Dosimetry)

After clinical evaluation and results from diagnostic 68Ga-DOTATOC PET/CT imaging, a patient was deemed to be eligible for treatment with 177Lu-DOTATATE RPT. No pretreatment dosimetry was performed; however, eligibility for subsequent 177Lu-DOTATATE administrations (every 8 wk) would be determined on the basis of prior and cumulative radiation doses from treatment.

The patient presented for the first therapeutic administration. The patient had a peripheral intravenous catheter placed, and an infusion of nephroprotective amino acids was started. After approximately 30 min had elapsed (~200 cc of fluid infused), it was confirmed by a nuclear medicine technologist that no signs of extravasation were present. The RPT (7.4 GBq of 177Lu-DOTATATE) was administered through the same intravenous catheter by standard institutional practice, and the amino acid infusion proceeded until completion, approximately 4 h after the start of infusion.

After completion of RPT administration, a blood sample was collected for dosimetric purposes. The patient was monitored and released. In the following days, at 24 h after injection, 72 h after injection, and 120 h after injection, the patient returned for dosimetric sampling. Each dosimetric sampling consisted of blood collection, WB planar imaging, 78804, and abdominal SPECT/CT (78832). On the basis of the 24-h SPECT/CT acquisition, it was determined that extravasation of 177Lu had not occurred.

Dosimetry proceeded by use of 510(k)-cleared medical device software for generalized 3D RPT dosimetry. A physicist, physician, or another qualified individual segmented organs of interest (whole liver, spleen, kidneys) as well as the 3 largest tumor lesions. Marrow dosimetry was performed by absorbed-fraction (MIRD) methods based on WB, blood, and normal organ time-integrated-activity quantification. A 3D dose plan normalized to a nominal administered activity was reviewed by the authorized user, and it was determined that a cumulative administered activity of 25.5 GBq was appropriate to maximize tumor dose, without exceeding dose limits to normal tissues (kidneys, liver, marrows) and within the limits of radiopharmaceutical availability. Three-dimensional dosimetry statistics for this final treatment plan were generated, and a treatment plan report was generated and signed by the physicist and authorized user (77295). An independent qualified individual reviewed this plan for appropriateness and accuracy, including performing a simple dose calculation via established absorbed-fraction (MIRD) methods. This secondary dose verification was documented in the medical record (77300).

After 8 wk, the patient returned for the second (Tx 2). On the basis of the target cumulative administered activity of 25.5 GBq, the patient was deemed eligible for an additional full administration of 7.4 GBq. The treatment was administered, and the patient underwent the same dosimetric sampling regimen as described above. Posttreatment dosimetry was performed to evaluate for deviation from expected tumor and normal organ doses. On the basis of compliance with the original treatment plan, treatment 3 (Tx 3) proceeded in the same manner with a 7.4 GBq administration and posttreatment dosimetry. The final treatment (Tx 4) was delivered in compliance with the initial treatment plan, with an administered activity of 3.3 GBq (cumulative 25.5 GBq). After the terminal treatment, dosimetric sampling was repeated. On the basis of these data, dosimetry was performed and combined with results from all 4 treatments (77300). A final patient-specific dose report was generated and documented in the medical record.

The following is a summary of the procedure and corresponding CPT codes:

- Tx 1 177Lu WB planar imaging (3 d), 78804;
- Tx 1 177Lu abdominal SPECT/CT (3 d), 78832;
- 3D radiation treatment planning, 77295;
- Plan check/simple dosimetry, 77300;
- Tx 2 177Lu WB planar imaging (3 d), 78804;
- Tx 2 177Lu abdominal SPECT/CT (3 d), 78832;
- Dosimetry assessment, 77300;
- Tx 3 177Lu WB planar imaging (3 d), 78804;
- Tx 3 177Lu abdominal SPECT/CT (3 d), 78832;
- Dosimetry assessment, 77300;
- Tx 4 177Lu WB planar imaging (3 d), 78804;
- Tx 4 177Lu abdominal SPECT/CT (3 d), 78832; and
- Dosimetry assessment, 77300

Comments. If tumors are not included within the abdominal SPECT/CT field of view, and tumor dosimetry is needed, additional SPECT fields of view would be required. This would not change the coding unless only a single posttreatment SPECT/CT were planned, in which case 78832 would be submitted in lieu of 78830. If marrow dosimetry is not performed, WB planar imaging (78804) and blood sampling should be omitted unless otherwise deemed medically necessary.

Example 3. 177Lu-DOTATATE (Dialysis Patient, Marrow Dosimetry)

After clinical evaluation and results from diagnostic 68Ga-DOTATOC PET/CT imaging, a patient was deemed to be eligible for treatment with 177Lu-DOTATATE RPT. In addition to having advanced neuroendocrine tumors, this patient had poor kidney function due to obstruction and was therefore receiving hemodialysis 3 d per week. Because of the compromised kidney function, blood clearance of any therapeutic radiopharmaceutical was expected to be significantly inhibited compared with the typical patient presentation. For this reason, the decision was made to initially administer 3.7 GBq of 177Lu-DOTATATE (rather than the standard 7.4 GBq), followed by bone marrow dosimetry to develop a treatment plan for subsequent administrations. For this patient, treatment was deemed to be palliative, and therefore kidney, liver, and tumor dosimetry were considered to be secondarily important to the most likely normal-tissue toxicity (bone marrow).
RPT was administered (methods consistent with what was described in Example 6.2), and the following dosimetric sampling was performed:

- Blood sampling at 4 h after administration (end of AA infusion);
- Hemodialysis performed from 4.5 to 7 h after administration;
- Blood sampling at 7.5 h after administration;
- Blood sampling and WB conjugate planar imaging at 24 h after administration;
- Blood sampling and WB conjugate planar imaging at 46 h after administration;
- Hemodialysis performed from 47 to 49 h after administration;
- Blood sampling performed at 49 h after administration; and
- Blood sampling and WB conjugate planar imaging at 96 h after administration.

After completion of dosimetric sampling, a special medical physics consult (77370) was ordered by the treating physician. The consult request was made to evaluate the effect of hemodialysis and blood retention of $^{177}$Lu-DOTATATE and associated marrow dosimetric effects. Dosimetry calculations were performed by a qualified medical physicist in addition to evaluating the impact of dialysis, and a consultation report was generated and documented. It was determined that minimal blood clearance occurred between dialysis sessions. Absorbed-fraction (MIRD)–based marrow dosimetry indicated significant elevation of population average dose values (more than 4 times the approved label average value). On the basis of this analysis performed by the qualified medical physicist, it was determined that addition of a hemodialysis session at 24 h after administration would be beneficial for marrow dosimetry due to increased peptide removal after the initial tumor uptake phase. A treatment plan was developed (77262) that included this modification while targeting a total administered activity of 14.8 GBq, with the remaining activity (11.1 GBq) to be split between treatments 2 and 3. An independent qualified individual reviewed this plan for appropriateness and accuracy, including performing a simple dose calculation via established absorbed-fraction (MIRD) methods. This secondary dose verification was documented in the medical record (77300).

Treatments 2 and 3 were completed according to the treatment plan, with postadministration dosimetry performed as described above. Bone marrow dosimetry (77300) was performed after each treatment, with dose reports (fraction and cumulative) being documented in the patient medical record.

The following is a summary of the procedure and corresponding CPT codes:

- Tx 1 $^{177}$Lu WB planar imaging (3 d), 78804;
- Med physics special consult: dialysis pharmacokinetics, 77370;
- Treatment planning (intermediate), 77262;
- Tx 1 marrow dosimetry, 77262;
- Tx 2 $^{177}$Lu WB planar imaging (3 d), 78804;
- Tx 2 marrow dosimetry, 77300;
- Tx 3 $^{177}$Lu WB planar imaging (3 d), 78804; and
- Tx 3 marrow dosimetry, 77300.

Comments. The choice of 77262 (intermediate) rather than 77261 (simple) or 77263 (complex) in this example is based on the time-sensitive nature of radiation dose delivery, including appropriate timing of dialysis; however, this treatment plan did not consider many specific treatment areas or organs at risk, and therefore 77263 would likely not be appropriate. A reduction in administered activity in this example led to elimination of 1 treatment administration compared with standard administration workflows – this led to substantial and immediate payer cost savings, and reduced risk of severe toxicity experienced by the patient.

**Example 4. $^{131}$I-MIBG (with Tumor, Marrow, and Kidney Dosimetry)**

After clinical evaluation and results from diagnostic $^{123}$I-MIBG SPECT/CT imaging, a patient was deemed to be eligible for treatment with $^{131}$I-MIBG RPT. Per the Food and Drug Administration–approved label for this RPT, pretreatment dosimetry was performed using a small quantity of the therapeutic radiopharmaceutical.

Radiopharmaceutical was administered (185 MBq) in a manner consistent with manufacturer recommendations and institutional policy. After administration, dosimetric sampling was collected. Dosimetric sampling consisted of blood sample collection at 4, 24, 48, and 96 h after administration and imaging (WB conjugate planar + SPECT/CT of the abdomen) at 24, 48, and 96 h after administration.

Dosimetry proceeded by use of 510(k)-cleared medical device software for generalized 3D RPT dosimetry. A physicist, physician, or another qualified individual segmented organs of interest (whole liver, spleen, kidneys) as well as the 3 largest tumor lesions. Marrow dosimetry was performed by absorbed-fraction (MIRD) methods based on WB, blood, and normal organ time-integrated-activity quantification. A 3D dose plan normalized to a nominal administered activity was reviewed by the authorized user, and it was determined that a cumulative administered activity of 26 GBq was appropriate to maximize tumor dose, without exceeding dose limits to normal tissues (in this case bone marrow). $^{131}$I-MIBG is typically administered over 2 treatments, and therefore a plan of administering 13 GBq in each treatment, separated by at least 90 d. Three-dimensional dosimetry statistics for this final treatment plan were generated, and a treatment plan report was generated and signed by the physicist and authorized user (77295). An independent qualified individual reviewed this plan for appropriateness and accuracy, including performing a simple dose calculation via established absorbed-fraction (MIRD) methods. This secondary dose verification was documented in the medical record (77300).

The patient returned for initial treatment. In accordance with the treatment plan, 13 GBq of $^{131}$I-MIBG was administered. The patient underwent posttreatment dosimetric sampling with the same blood collection and imaging time points as described above. Dosimetry was performed (77300) over the 96 h after administration, and a posttreatment dose report was documented and reviewed by the treating physician (77300). On the basis of exposure rate measurements, the patient was retained with “in-patient” status until the end of day 2, at which time the patient met Nuclear Regulatory Commission and state release criteria. Before and after release, the patient was monitored for treatment-related adverse events.

After 90 d had elapsed, the patient returned for an additional treatment of 13 GBq (26 GBq cumulative) in accordance with the treatment plan. Dosimetry was again performed (77300), with a final cumulative dose report being generated, documented, and reviewed by the treating physician.

The following is a summary of the procedure and corresponding CPT codes:
• Pre-Tx 131I WB planar imaging (3 d), 78804;
• Pre-Tx 131I abdominal SPECT/CT (3 d), 78832;
• 3D radiation treatment planning, 77295;
• Plan check/simple dosimetry, 77300;
• Tx 1 131I WB planar imaging (3 d), 78804;
• Tx 1 131I abdominal SPECT/CT (3 d), 78832;
• Dosimetry assessment, 77300;
• Tx 2 131I WB planar imaging (3 d), 78804;
• Tx 2 131I abdominal SPECT/CT (3 d), 78832; and
• Dosimetry assessment, 77300

Comments. Many patients who can benefit form 131I-MIBG therapy are quite young (below the age of 4), and therefore require general anesthesia for dosimetric imaging. In these cases the treating physician, in collaboration with the multidisciplinary team, may choose to forgo dosimetry after Tx 1 and Tx 2, or develop a nonstandard pretreatment dosimetry workflow in consultation with a qualified medical physicist (77370). An example of a modified dosimetric sampling would be standard blood collections; WB planar imaging at 24 h; and WB counting (nonanesthetized) at 4, 24, 48, and 96 h after administration. In general, these modifications preclude tumor dosimetry; however, dose to the primary limiting organ (bone marrow) can be assessed with reduced precision. Additional details regarding abbreviated dosimetry methods can be found in the EANM procedure guidelines for 131I-MIBG therapy (39).

CURRENT DEFICIENCIES AND FUTURE NEEDS

Although the coding strategies described herein are appropriate to meet the immediate need for baseline support of dosimetry and treatment planning for RPT, the existing CPT code set does not contain a sufficient spectrum of codes to describe the current and anticipated process of care for RPT procedures. Some services fit within the scope of existing codes; however, many services remain unsupported or undersupported by existing codes. New and dedicated codes for theranostics should be developed, with collaboration between relevant stakeholders (Society of Nuclear Medicine and Molecular Imaging [SNMMI], American Society for Radiation Oncology [ASTRO], American College of Radiology [ACR], Society of Interventional Radiology [SIR], American Association of Physicists in Medicine [AAPM], and others). What follows are several notable deficiencies among the current coding structure; however, this list is neither intended to be comprehensive nor authoritative.

Partition and Volume-Based 90Y-Microsphere Treatment Planning

Although 77295 may be appropriate when the clinical case rises to a level of complexity requiring generation and review of 3D isodose volumes relative to normal tissue and tumor targets, a common method of calculation in somewhat simpler cases (e.g., single lesion, well-defined uptake, limited volume of perfusion) involves an approximation of uniform activity distribution in the target tumor and normal liver. Under this approximation, one must determine the volume of treated liver + tumor, the total liver volume, the tumor–to–liver concentration ratio, and the fraction of activity shunting to lungs and other normal tissues. On the basis of these data, calculations can be performed to provide a range of potential treatment plans, from which the authorized user can select the most appropriate. The effort for these activities may exceed what is included in 77261–77263, and thus new codes may need to be developed based on plan complexity.

WB Counting

In some cases, particularly pediatric patients, it may be more appropriate to use serial WB counting in lieu of serial WB planar imaging for the purposes of bone marrow dose assessment. WB counting may involve use of a scintillation spectrometer (i.e., shielded NaI thyroid uptake probe) or use of an ion chamber survey meter. The WB counting procedures, which can allow for data acquisition without general anesthesia in pediatric patients, is not currently supported by any existing code.

More Than 2 SPECT/CT or WB Planar Scans

As indicated by examples provided in the section “Specific Coding Examples” in this paper, it is sometimes necessary to obtain more than 2 imaging fields of view (areas) or imaging time points to adequately characterize the spatial and temporal distribution of radiopharmaceutical in organs and tumors of interest. 78804 and 78832 provide reimbursement for only 2 WB planar images and SPECT/CT areas or imaging timepoints, respectively. Revision of these code, or creation of a modifier to account for additional timepoints, should be undertaken.

RPT “Simulation”

Although not commonly performed (with the exception of 131I-MIBG), administration of a small quantity of the therapeutic radiopharmaceutical or an appropriate surrogate followed by dosimetric sampling may emerge as a useful technique for treatment planning. This process may or may not include supportive compounds, such as infusion of renal-protective amino acids, as these compounds are known to alter the pharmacokinetics of the therapeutic radiopharmaceutical. Dedicated codes for this workflow and associated radiopharmaceutical costs may be needed.

Blood Collection and Counting

Analysis of biologic samples is needed for determination of bone marrow dose in most cases. Analysis may include whole-blood spectroscopic counting, plasma spectroscopic counting, and determination of the patient’s hematocrit. These procedures, typically performed by a nuclear medicine technologist, appear unsupported by the current code set.

Sequential PET/CT for RPT

Interest is growing in the use of positron-emitting surrogates for RPTs, such as 64Cu-DOTATATE as a surrogate for 177Lu-DOTATATE. Currently there are no codes for multiple-time-point PET/CT imaging, such as what is available for SPECT/CT.

Consensus Regarding “Simplified” Dosimetry

As discussed in the section “Primary Stakeholders,” there are some emerging data suggesting that adequate dosimetry can be performed from a limited number of imaging time points based on population pharmacokinetic data. These emerging techniques should be examined by experts in the field, to develop consensus recommendations regarding which clinical scenarios are well-suited to simplification or time-point reduction.

DISCLAIMER

The opinions provided in this paper are those of members of the SNMMI Dosimetry Task Force based on their coding experience. Always check with your local insurance carriers, as policies vary...
by region. The billing strategies described in this paper are unlikely to be accepted universally, and so the final decision for coding for any procedure must be made by the physician, considering regulations of insurance carriers and any local, state or federal laws that apply to the physician’s practice. Neither SNMMI nor any of its officers, directors, agents, employees, committee members, or other representatives shall have any liability for any claim, whether founded or unfounded, of any kind whatsoever, including but not limited to any claim for costs and legal fees, arising from the use of these opinions.

DISCLOSURE

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