Validation of Monte Carlo $^{131}$I radiopharmaceutical dosimetry workflow using a 3D-printed anthropomorphic head and neck phantom

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Abstract

**Purpose:** Approximately 50% of head and neck cancer (HNC) patients will experience loco-regional disease recurrence following initial courses of therapy. Retreatment with external beam radiotherapy (EBRT) is technically challenging and may be associated with a significant risk of irreversible damage to normal tissues. Radiopharmaceutical therapy (RPT) is a potential method to treat recurrent HNC in conjunction with EBRT. Phantoms are used to calibrate and add quantification to nuclear medicine images, and anthropomorphic phantoms can account for both the geometrical and material composition of the head and neck. In this study, we present the creation of an anthropomorphic, head and neck, nuclear medicine phantom, and its characterization for the validation of a Monte Carlo, SPECT image-based, $^{131}$I RPT dosimetry workflow.

**Methods:** 3D-printing techniques were used to create the anthropomorphic phantom from a patient CT dataset. Three $^{131}$I SPECT/CT imaging studies were performed using a homogeneous, Jaszczak, and an anthropomorphic phantom to quantify the SPECT images using a GE Optima NM/CT 640 with a high energy general purpose collimator. The impact of collimator detector response (CDR) modeling and volume-based partial volume corrections (PVCs) upon the absorbed dose was calculated using an image-based, Geant4 Monte Carlo RPT dosimetry workflow and compared against a ground truth scenario. Finally, uncertainties were quantified in accordance with recent EANM guidelines.

**Results:** The 3D-printed anthropomorphic phantom was an accurate recreation of patient anatomy including bone. The extrapolated Jaszczak recovery coefficients were greater than that of the 3D-printed insert ($\sim 22.8$ ml) for both the CDR and non-CDR cases (with CDR: 0.536 vs. 0.493, non-CDR: 0.445 vs. 0.426, respectively). Utilizing Jaszczak phantom PVCs, the absorbed dose was underpredicted by 0.7% and 4.9% without and with CDR, respectively. Utilizing anthropomorphic phantom recovery coefficient overpredicted the absorbed dose by 3% both with and without CDR. All dosimetry scenarios that incorporated PVC were within the calculated uncertainty of the activity. The uncertainties in the cumulative activity ranged from 23.6% to 106.4% for Jaszczak spheres ranging in volume from 0.5 to 16 ml.

**Conclusion:** The accuracy of Monte Carlo-based dosimetry for $^{131}$I RPT in HNC was validated with an anthropomorphic phantom. In this study, it was found...
INTRODUCTION

Approximately 50% of head and neck cancer (HNC) patients will experience loco-regional disease recurrence following initial courses of therapy.1–6 Although many patients can be cured from HNC, the 5-year overall survival rate of patients with locally advanced HNC is approximately 50%,7 and cure rates have improved only marginally over the last 30 years.8 Retreatment of HNC is technically challenging and accompanied by a significant risk of irreversible damage to normal tissues that can translate into profound adverse effects on patient health-related quality of life.9–22 Several studies have ascertained that despite improvement in long term survival (2 years) in approximately 10%–20% of recurrent HNC patients following external beam radiotherapy (EBRT), there is considerable risk for high-grade toxicities after treatment.5–13,15–23 Despite the ability of EBRT to provide disease control, efforts to reduce the potential of acute and long-term tissue injury in locoregionally recurrent HNC patients following EBRT are clearly warranted.

Radiopharmaceutical therapy (RPT) is one potential treatment that can be combined with EBRT to maintain disease control of recurrent HNC and mitigate high-grade toxicities. RPT involves a biological delivery vector labeled with a radioactive isotope that is designed to preferentially attach to or be internalized by cancer cells. The radioactive drug conjugate is infused directly into a patient’s bloodstream allowing it to reach tumor sites located throughout the body. Due to this unique delivery route, the dose-limiting organs are different than EBRT; and, when combined with EBRT, the therapeutic window in the patient can be expanded, potentially allowing physicians to treat more aggressively.

CLR 131,24 an alkyl phosphocholine (APC) analog with broad cancer targeting abilities, is a promising RPT agent paired to the radioisotote. 131I emits both therapeutic beta particles (mean energy of 606 keV) and diagnostic gamma rays (364 keV), which can be localized using SPECT. APCs enter cells through specialized plasma membrane microdomains called lipid rafts,24 which are expressed 6–10 times more in cancer cells in comparison to normal tissues. The increased specificity of CLR 131 for malignant cells shows promise for increased therapeutic efficacy in both preclinical and early clinical studies. Preclinical studies confirm preferential uptake and retention of CLR 131 in murine HNC xenografts and tumor growth inhibition.25 Tumor targeting has been observed in patients in on-going and completed clinical trials.26–28 Additional tumor growth inhibition has been demonstrated when CLR 131 is combined with EBRT.25

Given these results, a clinical trial (NCT04105543) is underway at UW-Madison to investigate the efficacy of combining CLR 131 with EBRT with curative intent in treating relapsed HNC.29 The premise of the trial is to evaluate the combination of CLR 131 and EBRT in the treatment of recurrent HNC, wherein the radiation dose delivered by the CLR 131 permits a dose reduction delivered by EBRT (e.g., standard of care 60–70 Gy will be targeted irrespective of radiation modality). The overall hypothesis of the clinical trial is that the addition of CLR 131 will enable a dose reduction of EBRT, which is safe and tolerable while maintaining favorable tumor response rates and diminishing the adverse impact of radiation treatment on subject-specific symptoms, such as salivary flow, swallowing function, and overall quality of life.

To achieve the objectives of the trial,30 patient-specific dosimetry is needed to accurately characterize the tumor absorbed dose from CLR 131 at the voxel level. Longitudinal single photon emission computed tomography/computed tomography (SPECT/CT) scans are used to map the time-dependent biodistribution of CLR 131 in the patient and serve as input for voxel-level dosimetry calculations performed by an in-house Monte Carlo-based software called RAPID.31 The accuracy of the CLR 131 dose calculation depends on the quantitative accuracy of 131I SPECT images. Nuclear medicine phantom scans are used both for quantitative calibration (i.e., convert counts to units of activity) and correction of partial volume effects (PVEs). PVEs, which are dependent on both the size and shape of the structure being imaged,32–34 are caused by inherent resolution limitations of imaging systems, and can significantly degrade SPECT/CT image quality and quantitative accuracy.

Collimator detector response (CDR) modeling has been recommended by medical internal radiation dosimetry (MIRD) pamphlet 2435 for SPECT image reconstruction due to septal penetration of high energy 131I photons through the collimator. Without CDR modeling, noise can be quite apparent on reconstructed images, and small structures can be difficult to resolve; however, some have observed Gibbs ringing artifacts...
on images reconstructed with CDR enabled, which can affect the resultant activity distribution.\textsuperscript{36}

Anthropomorphic nuclear medicine phantoms have been investigated for a variety of organs (kidney,\textsuperscript{37–40} liver,\textsuperscript{38,40–42} spleen,\textsuperscript{38,40} and pancreas\textsuperscript{38,39}), but few groups have investigated the use of nuclear medicine head and neck phantoms. 3D-printed head and neck phantoms have been created for CT and MR applications but few for nuclear medicine applications. A prototype head and neck phantom was described by Alqahtani et al. in which the performance of gamma imaging systems was evaluated, but the study did not extend to dosimetry.\textsuperscript{43} In these studies, results have generally indicated that anthropomorphic phantoms account for heterogeneities that are not present in traditional nuclear medicine phantoms and are often more accurate than non-anthropomorphic phantoms. 3D-printed anthropomorphic head and neck phantoms that account for the changing contour of the head and neck region and the heterogeneous material composition of the patient have not been investigated. Recently published review articles succinctly demonstrate the state of 3D-printed phantoms.\textsuperscript{44,45}

In this work, we aim to validate our image-based RPT dosimetry workflow for the HNC clinical trial with the introduction and characterization of a novel, 3D-printed, anthropomorphic, head and neck medical phantom that mimics the geometry and material composition of the head and neck region. Imaging studies will be conducted with both traditional Jaszczak nuclear medicine phantoms and the anthropomorphic phantom to investigate \textsuperscript{131}I SPECT/CT calibration, partial volume corrections (PVCs), and the role of CDR modeling on dosimetry. The resultant data from the imaging studies will then be used as input for the \textsuperscript{131}I RPT dosimetry workflow and compared to ground truth scenarios. Finally, the uncertainties from the imaging studies will be investigated, and the impact of uncertainties on the dosimetry will be discussed.

2 METHODS

2.1 Phantom creation

The original CT data for the anthropomorphic phantom were accessed from a publicly available database provided by radiation therapy oncology group (RTOG) 0522 study (Figure 1a).\textsuperscript{46} The CT data had a voxel resolution of 0.98 × 0.98 × 2.4 mm\textsuperscript{3}. Important structures such as the tumor, thyroid, lacrimal glands (both left and right), parotid glands (both left and right), and bone were contoured manually using Amira v.5.3.3 (Figure 1b). The labels were then converted to .stl files, which is a file format native to the stereolithography (SL) CAD software created by 3D systems and required for 3D printing. The .stl files were then modified for final assembly using Magics (Materialise NV, Leuven, Belgium), which is a .stl editing software package (Figure 1c,d). Modifications included the addition of flanges to seal the main volume, the incorporation of mounting points for the assembly of bones and other anatomical features, and ports for filling and sealing the tumor and gland voids.

The final phantom design consisted of nine custom parts, along with commercial hardware for assembly. This included seven 3D-printed pieces, made with two different materials, and two laser-cut acrylic plates. The main chamber (skin) was printed using SL out of Accura60, a proprietary photopolymer resin (3D Systems, Rock Hill, SC, USA), and then clear-coated for water resistance. The thyroid gland was fused to the skin and can be filled independently. The skull and spine were split into three separate parts and printed using SL out of Somos PerFORM, a proprietary photopolymer resin (DSM, Heerlen, Netherlands). The lacrimal glands were fused to the front piece of the skull and can be filled independently. Lastly, the parotid voids and tumor void were printed using SL out of Accura60 (3D Systems) with approximately 2–3-mm wall thickness. The top and bottom of the chamber were formed with quarter inch acrylic plates that were laser-cut to their final size with a PLS6.75 Laser Cutting System (Universal Laser Systems, Scottsdale, AZ, USA). Commercial hardware included o-rings for sealing and nylon bolts for assembly of separate components. The final 3D-printed head-and-neck anthropomorphic phantom is shown in Figure 1e, and a sagittal slice of a CT image of the phantom is depicted in Figure 1f. Table 1 shows the physical material properties taken from material datasheets of the Accura60 and Somos PerFORM comprising the phantom.

2.2 Phantom studies and SPECT/CT image acquisition

Three phantom studies were conducted, the parameters of which are summarized in Table 2. The first was conducted using a uniform 20-cm cylindrical phantom of volume 5.64 L to calculate the calibration factor to convert the reconstructed SPECT images to units of activity concentration. The second study was with a Jaszczak phantom containing 0.5–, 1–, 2–, 4–, 8–, and 16-ml hollow sphere inserts, with corresponding diameters of 9.9, 12.4, 15.4, 19.8, 24.8, and 31.3 mm, injected with 9.9 insert to background activity concentration and was used to calculate spherical-based recovery coefficients (RCs) for PVC determination. The third study was with the anthropomorphic phantom containing a 22.8-ml tumor insert and injected with a 9.9:1 insert to background activity concentration. The resultant images were used to calculate an anthropomorphic-based recovery coefficient and served as the basis for the image-based RPT dosimetry calculation. SPECT/CT
Figure 1: Representation of the head and neck (H&N) anthropomorphic phantom development process. (a) Original CT image obtained from a radiation therapy oncology group (RTOG) managed publicly available image repository and (b) relevant contoured structures. (c and d) 3D rendering of patient’s skin, bony anatomy, and fillable organ anatomies. (e) Final 3D-printed H&N anthropomorphic phantom. (f) CT image of H&N anthropomorphic phantom.

Table 1: Physical properties of the bulk materials

| Material         | Tensile strength (MPa) | Elongation at break (%) | Tensile modulus (MPa) | Flexural strength (MPa) | Water absorption (%) | Hardness (shore D) |
|------------------|------------------------|-------------------------|-----------------------|------------------------|---------------------|--------------------|
| Accura60         | 58–68                  | 5–13                    | 2690–3100             | 87–101                 | N/A                 | 86                 |
| Somos Perform    | 68–80                  | 1.1–1.2                 | 9800–10500            | 120–146                | 0.1–0.2             | 93–94              |

Abbreviation: N/A, not applicable.

Table 2: Parameters used for SPECT/CT phantom studies

| Study             | Initial $^{131}$I activity concentration (kBq/ml) | Insert to background ratio | Number of frames | Seconds per frame |
|-------------------|--------------------------------------------------|---------------------------|-----------------|------------------|
| 20-cm cylindrical phantom | 105                                               | N/A                       | 120             | 20               |
| Hot sphere Jaszczak   | 259                                               | 9.9:1                     | 120             | 60               |
| Anthropomorphic      | 257                                               | 9.9:1                     | 120             | 60               |

Abbreviation: N/A, not applicable.
acquisitions were conducted using a GE Optima NM/CT 640 with a high energy general purpose collimator. All images were acquired over 360 degrees into 128 × 128 pixel matrices per angle. Body contouring was enabled. A photopeak energy window centered at 364 keV and 20% in width (327.6–400.4 keV) and a triple energy window scatter correction with two 20% windows adjacent (267.3–326.7 keV and 401.4–490.6 keV) to the photopeak window were used in the acquisition. A CT was acquired after the SPECT acquisition for CT-derived attenuation corrections (Hawkeye CT, 120 kVp, 20 mA, 487 mAs) and for accurate determination for the location of hot spheres and tumor insert. Raw image data were reconstructed using the GE Xeleris 4.0 ordered subset expectation maximum algorithm with 10 iterations and 10 subsets with CT-based attenuation correction and both with and without GE’s CDR modeling named “resolution recovery” (without: non-RR, with: RR). No postreconstruction filtering was applied. The reconstructed SPECT image matrices were 128 × 128 × 128 voxels with a voxel size of 4.42 × 4.42 × 4.42 mm³, and the reconstructed CT images had a voxel size of 0.98 × 0.98 × 5.0 mm³. High-resolution CT images of the anthropomorphic phantom were also acquired using a Siemens SOMATOM Definition Edge (120 kVp, 452 mA, 226 mAs) with a voxel size of 0.977 × 0.977 × 2.0 mm³. As this high-resolution CT scanner is utilized for photon EBRT treatment planning simulation scans, the scanner is highly characterized, and the electron density CT calibration curve was created utilizing a Gammex 467 Tissue Characterization Phantom.

2.3 Calculation of calibration factor and recovery coefficients

To convert the recorded SPECT image data into units of activity, the calibration factor, Q, was calculated according to Equation (1),

\[ Q = \frac{R}{V \cdot C} \cdot e^{\lambda \Delta t}, \]  

(1)

where \( R \) is the mean count rate of a voxel in a volume of interest (VOI), \( V \) is the volume a voxel, \( C \) is the activity concentration at the time of syringe measurement, and the last term accounts for physical decay between source preparation time and scan time (\( \Delta t \)).

If \( R_{\text{voxel}} \) is the count rate in a voxel, then the activity in the voxel, \( A_{\text{voxel}} \), is given by Equation (2).

\[ A_{\text{voxel}} = \frac{R_{\text{voxel}}}{Q}. \]  

(2)

To account for PVE, recovery coefficients were calculated according to Equation (3), where the measured activity concentration in the object VOI was determined by quantifying the mean value of each hot insert. Con-

\[ RC = \frac{\text{Measured activity concentration in object VOI}}{\text{True activity concentration in object VOI}}. \]  

(3)

To correct the activity in a voxel, the RC is then used in Equation (4), where \( R_{\text{voxel}} \) is the count rate in the voxel.

\[ A_{\text{voxel}} = \frac{R_{\text{voxel}}}{Q} \cdot RC. \]  

(4)

To account for PVE, recovery coefficients were calculated using Equation (3), where the measured activity concentration in the object VOI was determined by quantifying the mean value of each hot insert. Con-

\[ RC_{\text{fit}} = R_{\text{plateau}} - \frac{R_{\text{plateau}}}{1 + e^{\left(\frac{-V}{\beta}\right)^\gamma}}. \]  

(5)

2.4 Monte Carlo dose calculation

An in-house radiopharmaceutical dosimetry platform called RAPID was used to calculate the mean absorbed dose to the tumor structure in the anthropomorphic phantom. RAPID utilizes nuclear medicine images to define the radionuclide activity in each voxel, and the absorbed dose rate distribution is calculated on the CT, which defines the material composition and mass density of the simulation geometry. The acquired SPECT/CT data from the anthropomorphic phantom study were used to calculate a voxel-level dose rate distribution in the phantom. The dose rate in each voxel was then integrated assuming single timepoint exponential decay to produce a total absorbed dose distribution in the phantom.

Three scenarios were considered: (1) ground truth using a simulated, idealized, SPECT dataset, which was created based upon a segmented CT such that the known activity concentrations were assigned at the voxel level, (2) SPECT-based dosimetry without RCs, and (3) SPECT-based dosimetry with phantom informed RCs. The calibration factor, \( Q \), was applied to the recorded SPECT/CT image of the anthropomorphic phantom. If RCs were utilized, the measured RCs from the Jaszczak phantom and the anthropomorphic phantom were then applied to the anthropomorphic phantom tumor volume to correct the activity concentrations. The resultant dose distributions from the SPECT image-based dosimetry were compared against the separate ground truth scenario, which was run by creating an artificial activity distribution assuming a uniform concentration of 26.2 kBq/cc and 259 kBq/cc in the background and tumor insert, respectively. The SPECT data were
interpolated to the same resolution as the CT. The dose calculation grid was of dimensions $278 \times 280 \times 155$ voxels with a $0.977 \times 0.977 \times 2.00$ mm$^3$ voxel size. The simulation was run with enough particles (8000 decays per voxel) to ensure Monte Carlo statistical uncertainty below 1.0% in the tumor region. Simulations were run on the University of Wisconsin-Madison Center for High Throughput Computing.

### 2.5 Uncertainty analysis

Uncertainty analysis for the activity determination of the Jaszczak phantom study was performed according to the recommended European Association of nuclear medicine (EANM) guidelines for determining error propagation in radiopharmaceutical dosimetry.$^{48}$ The fractional uncertainty of the volume determination is given by Equation (6), where $a$ is the voxel size, and $D$ is the equivalent sphere diameter of the contoured structure.

$$\frac{u(v)}{v} = 3 \left( \frac{a^2}{\pi D} \right).$$  \tag{6}$$

The uncertainty of the calibration factor (calculated using Equation (1)) is given by Equation (7) and combines in quadrature two components: the assumed 5% error of the radionuclide calibrator activity measurement ($u(A_{cal})$), and the error of the count rate for the scans to determine the calibration factor $u(C_{ref})$, taken to be the standard deviation of the mean count rate for two timepoints.

$$\left[ \frac{u(q)}{q} \right]^2 = \left[ \frac{u(A_{cal})}{A_{cal}} \right]^2 + \left[ \frac{u(C_{ref})}{C_{ref}} \right]^2.$$  \tag{7}$$

The uncertainty of the expression used to fit the recovery coefficient curve (Equation (5)) is given by Equation (8). $g_c$ is a matrix of dimensions $4 \times 1$ containing both the partial derivatives of first order of $RC$ with respect to $R_{plateau}$, $\beta$, $\gamma$, and $v$, and $V_C$ is a $4 \times 4$ matrix containing the covariance matrix from the least-squares fitting process and the uncertainty of the volumes, $u^2(v)$.

$$u^2(RC) = g_c^T V_C g_c.$$

The uncertainty in the measured mean counts $C$ associated with the accuracy of the VOI definition is given by Equations (9) and (10),

$$\frac{u(C)}{C} = \frac{\varphi \cdot u(v)}{2RC \cdot v}.$$  \tag{9}$$

$$\varphi = \text{erf} \left( \frac{2r^2}{\sigma \sqrt{2}} \right) - \frac{2\sigma}{r \sqrt{2\pi}} \left[ 1 - e^{-\frac{2r^2}{\sigma^2}} \right],$$  \tag{10}$$

where $r$ is $D/2$ (the equivalent sphere diameter of the structure), and $\sigma$ is the standard deviation of the Gaussian point spread function derived from the system spatial resolution, assumed to be 3 mm.$^{49}$

The cumulative activity uncertainties were calculated by adding the system sensitivity, recovery coefficient, and count rate errors in quadrature and subtracting out the covariance of the count rate and recovery coefficient on volume.$^{35}$ The uncertainty of the activity was estimated using Equation (11).

$$\left[ \frac{u(A)}{A} \right]^2 = \left[ \frac{u(q)}{q} \right]^2 + \left[ \frac{u(RC)}{RC} \right]^2 + \left[ \frac{u(C)}{C} \right]^2$$

$$- \frac{\varphi}{RC^2 v} \frac{\partial RC}{\partial v} u^2(v).$$  \tag{11}$$

Estimates of the uncertainty for the anthropomorphic phantom tumor volume were extrapolated from the results of the Jaszczak uncertainties assuming the tumor volume to be spherical.

### 3 RESULTS

#### 3.1 Phantom geometry

Figure 2a–c shows photographs of the head and neck phantom and labels for the fillable and removable inserts. Figure 2a shows an image of the phantom assembled and filled with water. Figure 2b shows the anterior half of the skull insert with the 3D-printed tumor volume and parotid glands attached to it by simple fasteners. Figure 2c shows a side profile of the phantom depicting the lacrimal gland (integrated into the skull) and the thyroid gland (integrated into the shell).

#### 3.2 Material properties of phantom

Figure 3 shows axial, coronal, and sagittal CT slices of the phantom. Figure 3a shows an axial slice in which the jaw, spine, parotid glands, and tumor volume are present. Figure 3b shows a coronal slice of the phantom in which the skull and the C1-C5 vertebrae are present. Figure 3c depicts a sagittal view of the phantom in which the thyroid gland and the parotid gland are present.

Table 3 provides the radiological properties of the bulk materials. Densities were provided by the manufacturer. The measured hounsfield unit (HU) corresponds to regions of interest (ROIs) drawn on bulk portions
FIGURE 2  Photographs showing the (a) main chamber (skin), (b) lower jawbone assembly, and (c) compiled head and neck phantom. Labels for the fillable inserts including the tumor volume (TV), parotid gland (PG), lacrimal gland (LG), and thyroid (Th) are superimposed on the photographs.

FIGURE 3  CT scan of phantom showing (a) axial, (b) coronal, and (c) sagittal views. Labels for the fillable inserts including the tumor volume (TV), parotid gland (PG), and thyroid (Th) are superimposed on the photographs.

TABLE 3  Radiological properties of the bulk materials

| Material       | Density (g/cc) | Measured HU  | $\mu_{177\text{Lu}}$ (cm$^{-1}$) | $\mu_{131\text{I}}$ (364 keV) (cm$^{-1}$) | $\mu_{18\text{F}}$ (511 keV) (cm$^{-1}$) |
|----------------|---------------|--------------|-------------------------------|------------------------------------------|------------------------------------------|
| Accura60       | 1.21          | 292.48 +/- 8.57 | 0.1579                        | 0.1063                                   | 0.1068                                   |
| Somos PerFORM  | 1.61          | 876.36 +/- 14.77 | 0.2137                        | 0.1390                                   | 0.1402                                   |
| Water          | 1.00          | -4.05 +/- 7.95 | 0.1295                        | 0.0896                                   | 0.0896                                   |

of the material by the Siemens high-resolution CT scanner. The tabulated attenuation coefficients were computed by utilizing formalisms from the literature, which have calculated a bilinear relationship between measured narrow beam linear attenuation coefficients and Hounsfield units at the photon energies of interest for each isotope. The bilinear relationships were utilized from Brown et al. for $^{131\text{I}}$, Burger et al. for $^{18\text{F}}$, and Kabasakal et al. for $^{177\text{Lu}}$.

Figure 4 depicts a portion of the CT calibration curve for the high-resolution CT scanner that the anthropomorphic phantom was scanned in. It also plots the two 3D-printed materials alongside the curve in which the clinical calibration curve would slightly overpredict the mass density of the material.

Table 4 shows a comparison between the measured volume of fillable chambers in the phantom and the vol-
TABLE 4  Comparison of fillable volumes in anthropomorphic phantom (ml)

| Fillable void | Measured | DICOM contour | Absolute difference | Percent difference |
|---------------|----------|---------------|---------------------|-------------------|
| ParotidL      | 8.75     | 8.44          | −0.31               | −3.54             |
| ParotidR      | 8.39     | 8.39          | 0.00                | 0.01              |
| LacrimalL     | 1.42     | 1.51          | 0.09                | 6.34              |
| LacrimalR     | 1.49     | 1.34          | −0.15               | −10.07            |
| Thyroid       | 14.12    | 15.47         | 1.35                | 9.56              |
| GTV           | 22.77    | 22.50         | −0.27               | −1.19             |

Abbreviations: DICOM, digital imaging and communications in medicine; GTV, gross tumor volume.

3.3 Phantom imaging studies

Using Equation (1), the calibration factor \( Q \) was calculated to be 104.2 cps/MBq using RR and 21.4 cps/MBq for non-RR. For the RR case, the average measured activity in the background and tumor compartments were 24.4 kBq/cc and 127.5 kBq/cc, respectively, where the background compartment was a large ROI in the shoulder region containing only water. There was a −6.76% difference between the actual background (26.2 kBq/cc) and measured background. For the non-RR case, the average measured activity in the background and tumor compartments were 24.7 kBq/cc and 110.1 kBq/cc. The difference between the actual background and measured background was −5.81%.

Figure 5 depicts an axial SPECT/CT slice of the Jaszczak phantom after image acquisition and shows the difference in activity recovered with and without resolution recovery. For the RR case, the average measured activity in the background compartment and 16-ml hot sphere compartments were 26.5 kBq/cc and 133.2 kBq/cc, respectively, and there was a 2.35% difference between the actual background (25.9 kBq/cc) and measured background. For the non-RR case, the average measured activity in the background compartment and 16-ml hot sphere compartments were 26.9 kBq/cc and 109.0 kBq/cc, respectively, and there was a 3.66% difference between the actual background (25.9 kBq/cc) and measured background.

Figure 6 shows the recovery coefficient for the Jaszczak phantom and 3D-printed insert as a function of sphere insert size for both resolution recovery (RR) and non-RR. As the size of the Jaszczak sphere increased, the recovery coefficients increased. Consistent with theory, the curve fit of the Jaszczak spheres in both cases with the non-RR and RR applied demonstrated that the spheres recovered more activity than the aspherical 3D-printed volume. The Jaszczak phantom results are fitted to the curvefit as in Equation (5), and the 95% confidence bounds for each fit are presented in Figure 6.
The extrapolated recovery coefficient was greater than that of the 3D-printed insert for both the non-RR and RR cases (with RR: 0.536 vs. 0.493, non-RR: 0.445 vs. 0.426, respectively).

The calculated uncertainties for the RR case of the Jaszczak phantom imaging study are given in Figure 7. The largest uncertainties were the mean counts and RC, which are dependent on the volume determination and the RC fitting function. The range of uncertainties for volume (17.3%–54.7%), RC (19.2%–86.0%), mean counts (14.3%–107.2%), calibration factor (7.47%), and cumulative activity (23.6%–106.4%) were calculated.

3.4 Monte Carlo dosimetry

Figure 8 shows the difference in the calculated mean tumor dose in comparison to the ground truth scenario with an absorbed dose calculated to be 62.15 Gy. For the RR (non-RR) case, the mean dose to the tumor was underestimated by 46.8% (53.1%) before applying any RC, underestimated by 4.9% (0.7%) after applying a Jaszczak informed RC, and overestimated by 3.0% (3.0%) after applying the 3D-printed insert RC.

The extrapolated activity uncertainty estimate from the Jaszczak phantom study presented in Figure 7 for the volume of the tumor was 19.4%; this uncertainty effectively places all PVC dosimetry results within the error estimates.

Figures 9 and 10 shows (a) axial, (b) coronal, and (c) sagittal slices of the absolute absorbed dose rate for the ground truth scenario and the resolution recovery image-based scenario, respectively. Figure 11 depicts the respective relative percent differences calculated between the ground truth and the SPECT-derived activity distributions for the resolution recovery case.

Large dose gradients were present in the RR dosimetry. This is consistent with both spill out due to imaging system limitations and Gibbs ringing artifacts that have been documented when using CDR modeling. As depicted in relative difference maps in Figure 11, there are stark regions of red, indicating that the SPECT image has much higher dose than the corresponding ground truth. This is attributable to those certain structures like bone material and 3D-printed material were assumed to be impermeable to water, and thus activity was not assigned to them in the ground truth scenario.

4 DISCUSSION

It was found that, in this study, the Jaszczak-based PVCs were sufficient to correct the activity distributions. While the Jaszczak-derived RC for the non-RR reconstruction was the most accurate scenario, the anthropomorphic phantom derived RC was most accurate for the RR reconstruction, within 3% of the ground truth. The tumor volume studied was quite spherical in shape; it is possible that the disease presentation of patients enrolled in the trial could deviate from spherical behavior in which case the Jaszczak-based recovery coefficients may be less accurate. In the context of uncertainty analysis, the uncertainty in the activity determination was found to be around 23.6% for the 16 ml Jaszczak sphere, and the extrapolated uncertainty estimate for the anthropomorphic tumor volume was 19.4%, placing all the recovered activity dosimetric results within the calculated error and shows that the RCs determined from the anthropomorphic phantom were dosimetrically similar to those derived from the Jaszczak hot sphere phantom. The uncertainties were even higher (up to >100%) for smaller Jaszczak volumes. Uncertainties of this magnitude must be considered and conveyed to physicians for clinical cases. This is consistent with previous work in which VOI delineations have a substantial impact on the activity quantification and dose, which are especially...
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**FIGURE 9** Absolute absorbed dose for the ground truth scenario: (a) axial, (b) coronal, and (c) sagittal views

**FIGURE 10** Absolute absorbed dose for the resolution recovery scenario: (a) axial, (b) coronal, and (c) sagittal views

**FIGURE 11** Relative dose differences between the ground truth scenario and SPECT-derived activity distribution with resolution recovery for (a) axial, (b) coronal, and (c) sagittal views

important for small structures. Additionally, uncertainties have been reported to be up to 102% for small targets for $^{177}$Lu in human clinical trials, placing our results in line with other RPT uncertainties.

Because the calibration factor depends on the accuracy of scatter and attenuation correction, and they vary with phantom geometry, the shape of the phantom can directly affect the accuracy of quantifying activity in the background compartment. This may be the reason for the calibration factor slightly underpredicting the activity concentration in the background compartment (an ROI in the shoulder containing only water) of the anthropomorphic phantom. These types of shape-dependent calibration factors have been previously reported.

The calibration factor derived from the homogeneous Jaszczak phantom study was used to keep the anthropomorphic phantom study in line with the imaging parameters and recommended guidelines for upcoming clinical trial patient studies.

In addition to the difference in shape of the phantom’s background region, another important distinction of the anthropomorphic phantom is the inclusion of 3D-printed bone-mimicking material, which more resembles the density and corresponding HU of bones in the head and neck to afford a more realistic dosimetric scenario. The bone-type material was measured to have a bulk HU value of 876.36 and density of 1.61 g/cc, which is representative of cortical bone, oftentimes found in the skull. Uncertainties and differences of 3D-printed materials have been demonstrated in previous work.
by Craft et al. In that work, slight differences in density and composition of printed materials resulted in dosimetric differences for clinical photon and electron beams. In the context of \(^{131}\text{I}\) RPT dosimetry, not only are dose computations affected because 3D-printed materials may not lie on the CT calibration curve (see Figure 4), but CT-based attenuation corrections for image reconstruction may be affected as well. These material property uncertainties could affect the dosimetry, and the further characterizing of 3D-printed materials should be carefully considered.

RR, the CDR compensation technique included in GE's reconstruction software, Xeleris 4.0\(^{57,58}\) was investigated because, in theory, high energy photon emissions of \(^{131}\text{I}\) requires modeling of the high energy collimators that inherently reduce spatial resolution due to the collimator geometry. We found that there was only a small difference in the dosimetric results between non-RR and RR; however, utilizing RR increased the spatial accuracy of the activity quantification and has been recommended to be used.\(^{35}\) Additionally, the spatial distribution of activity was less clear and difficult to discern the presence of activity in smaller structures (see Figure 5c) thus limiting its applicability. The CDR model in Xeleris only models the intrinsic and geometric components of the collimator and neglects the collimator scatter and penetration, which can lead to significant higher count recovery for \(^{131}\text{I}\).\(^{35}\) In addition to using the manufacturer's RR, Monte Carlo CDR modeling has been studied in previous work to model the CDR explicitly to account for septal penetration and scatter\(^{59,60}\) and should be considered to improve the accuracy of the image reconstruction.

5 | CONCLUSIONS

The 3D-printed anthropomorphic phantom was created to more accurately represent the geometry and material composition of the head and neck region in comparison to typical non-anthropomorphic Jaszczyk-type nuclear medicine phantoms. This phantom was used to evaluate the accuracy of a Monte Carlo, image-based, RPT dosimetry workflow in comparison to a ground truth scenario. We investigated the determination of RCs from different phantoms, the impact of CDR modeling, and assessed the uncertainty in the activity determination. After correcting the reconstructed SPECT images using volume-based RCs, the dosimetry workflow was determined to be accurate within the calculated uncertainty. The Jaszczyk-based PVCs were sufficient to correct the activity distributions in this study; however, aspherical volumes may warrant the utilization of anthropomorphic phantom PVC. Further work should characterize the 3D-printed materials rigorously and include Monte Carlo-based reconstructions, which include more accurate CDR modeling.

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CONFLICT OF INTEREST

Benjamin L. Cox is a co-founder of Phantech LLC, a manufacturer of imaging phantoms. Bryan Bednarz and Joseph Grudzinski are co-founders and have ownership interest in Voximetry, Inc., a nuclear medicine dosimetry company in Madison, WI. Other authors have no conflict of interest.

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