Patient-specific Extravasation Dosimetry Using Uptake Probe Measurements

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Abstract—Extravasation is a common problem in radiopharmaceutical administration and can result in significant radiation dose to underlying tissue and skin. The resulting radiation effects are rarely studied and should be more fully evaluated to guide patient care and meet regulatory obligations. The purpose of this work was to show that a dedicated radiopharmaceutical injection monitoring system can help clinicians characterize extravasations for calculating tissue and skin doses. We employed a commercially available radiopharmaceutical injection monitoring system to identify suspected extravasation of $^{18}$F-fluorodeoxyglucose and $^{99m}$Tc-methylene diphosphonate in 26 patients and to characterize their rates of biological clearance. We calculated the self-dose to infiltrated tissue using Monte Carlo simulation and standard MIRD dosimetry methods, and we used VARSkin software to calculate the shallow dose equivalent to the epithelial basal-cell layer of overlying skin. For 26 patients, injection-site count rate data were used to characterize extravasation clearance. For each, the absorbed dose was calculated using representative tissue geometries. Resulting tissue-absorbed doses ranged from 0.6 to 11.2 Gy, and the shallow dose equivalent to a 10 cm$^2$ area of adjacent skin in these patients ranged from about 0.1 to 5.4 Sv. Extravasated injections of radiopharmaceuticals can result in unintentional doses that exceed well-established radiation protection and regulatory limits; they should be identified and characterized. An external injection monitoring system may help to promptly identify and characterize extravasations and improve dosimetry calculations. Patient-specific characterization can help clinicians determine extravasation severity and whether the patient should be followed for adverse tissue reactions that may present later in time.

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Key words: extravasation; internal dosimetry; radiation dose; biokinetics; extremity dose

INTRODUCTION

Most diagnostic nuclear medicine exams and therapeutic infusions are accomplished by administering radiopharmaceuticals intravenously (Boellaard et al. 2015). An extravasation, also known as an infiltration, occurs when a radiopharmaceutical is inadvertently injected into tissue surrounding the injection site instead of into the vasculature. Extravasations can result from improper initial placement of the intravenous (IV) access device or by failure of the vessel wall (Hadaway 2007). Extravasations occur relatively frequently (mean 10.4%, N = 5418, 20 nuclear medicine centers), as previously described (Hall et al. 2006; Bains et al. 2009; Krumrey et al. 2009; Osman et al. 2011; Silva-Rodriguez et al. 2014; McIntosh and Abele 2016; Muzzafar et al. 2017; Wong et al. 2019; Currie and Sanchez 2020) and can result in significant dose to underlying tissues and skin (Patton and Millar 1950; Shapiro et al. 1987; Rhymer et al. 2010; Bonta et al. 2011; Kawabe et al. 2013; Goodman and Smith 2015; van der Pol et al. 2017; Tylski et al. 2018). However, because radiation effects on patients may take years to manifest and are rarely studied (van der Pol et al. 2017), dose resulting from extravasations should be more fully evaluated.

Factors that influence tissue absorbed dose from extravasation include infiltrated tissue volume as well as radioactivity distribution, retention, absorption, and clearance. Extravasation clearance rate has been estimated to be 2 to 10 h (Esser 2017). Serial imaging with positron emission tomography (PET) or single-photon-emission computed tomography (SPECT) can provide more accurate estimates of radioactivity and clearance (Breen and Dreidger 1991; Williams et al. 2006; Bonta et al. 2011; Terwinghe et al. 2012; Kawabe et al. 2013; Tylski et al. 2018). However, clinicians must promptly recognize that a tissue infiltration has occurred, imaging systems must be available, and staff must know how to evaluate the resulting extravasation image data. In lieu of
imaging, manual serial measurements of the injection site can be made using a scintillation counter or other radiation detection system to determine retention and clearance parameters (Esser 2017). This manuscript describes an efficient, automated serial measurement system used to identify and characterize radiopharmaceutical extravasations.

Radiation dose estimates guide decision-making with respect to follow-up actions that may be appropriate. The purpose of this work was to show that a dedicated radiopharmaceutical injection monitoring system can help clinicians and technologists characterize extravasations for calculating tissue and skin doses.

**MATERIALS AND METHODS**

**Radiation detector**

We employed a commercially available detector (Lara® System, Lucerno Dynamics, Cary, NC) to characterize 26 extravasations of 18F-fluorodeoxyglucose (18F-FDG) and 99mTc-methylene diphosphonate (99mTc-MDP). The Lara radiopharmaceutical injection monitoring system comprises one scintillation detector placed on the patient’s skin proximal to the injection site and another on the opposite arm as a reference (Fig. 1). Each detector incorporates a single bismuth germanate (BGO) crystal and a silicon photomultiplier (SiPM). The detectors are neither shielded nor collimated, so their response is omnidirectional. Photon energy response is variable, depending on radionuclide, as previously described (Knowland et al. 2018). Each Lara detector records photon counts per second (cps) and generates a plot of counts vs. time. Reference detector output may be subtracted from injection-site detector output to correct for background photon counts such as from photons originating in the patient’s torso.

**Radiation dosimetry**

Using mathematical methods (Bolch et al. 2009) recommended by the special committee on Medical Internal Radiation Dose (MIRD) of the Society of Nuclear Medicine and Molecular Imaging (SNMMI), we calculated radiation absorbed doses (Gy) to representative volumes (cm³) of subdermal tissue containing infiltrated radiopharmaceutical.

Using a slightly modified version of VARSKI 6.1 (Hamby and Mangini 2018), a computer code for skin dosimetry, we also calculated the shallow dose equivalent (Sv) to the highest relevant area of the skin (10 cm²). In the MIRD formalism, the absorbed dose 

\[ D(r_T \rightarrow r_S) \]

from activity in a source region that irradiates a target region is

\[ D(r_T \rightarrow r_S) = \bar{A}(r_S, \tau) \sum_i \varphi_i(r_T \rightarrow r_S) \rho_T, \]

where \( \bar{A}(r_S, \tau) \) is the time-integrated activity in the source region, and \( \varphi_i(r_T \rightarrow r_S) \) is the absorbed fraction (fraction of energy emitted from a source region that deposits in a target region), and where \( \rho_T \) is the mass of the target region (Bolch et al. 2009).

**Count-rate curve**

To determine the time-dependent number of radioactive decays in the source region from an extravasation, we used the Lara detector count-rate curve (one measurement per second), which reflects the “effective” disappearance of infiltrated activity (combined effects of radioactive decay and biological clearance) following 26 extravasations of 18F-FDG or 99mTc-MDP. We then identified an appropriate mathematical function for the curve and best-fit parameters by least-squares regression analysis using commercially available curve-fitting software (Curve Expert Professional, Hyams Development, Huntsville, AL). We integrated analytically to yield area under the fitted curve representing total counts from injection through complete disappearance.

**Converting counts to activity present**

Detector photon count rate can be converted to absolute activity (MBq) using a three-dimensional region of interest (ROI) within the patient’s nuclear medicine image. We determined an activity calibration factor by dividing the fitted curve at imaging time by the ROI activity. We then converted the fitted curve to units of activity by multiplying it by the calibration factor. In the absence of quantifiable injection-site image data (e.g., injection site outside of the imaging field-of-view), extravasated activity was estimated based on overall image quality relative to a non-extravasated infusion.

**Absorbed energy fraction**

In the MIRD schema, the absorbed fraction \( \varphi_i(r_T \rightarrow r_S) \) can be determined experimentally using calibration sources and phantoms, or it may be calculated using Monte Carlo track simulations. Infiltrated tissue may present in many different shapes and sizes. To assess and compare potential tissue doses, we used Monte Carlo simulations and modeled the infiltrated tissue as one of three representative geometries of unit-density tissue: (a) a thin, right circular cylinder having a radius (r, cm) and height (h, cm) lying beneath the dermis where the tissue volume = \( \pi r^2 h \) (cm³), (b) as a sphere

Fig. 1. Photo of the injection monitoring system used on a nuclear medicine patient.
where the tissue volume = \((4 \pi r^3)/3\), and (c) as an ellipsoid where the tissue volume = \((4 \pi a b c)/3\) where a, b, and c were the radii of the ellipsoid. We calculated absorbed fractions for each representative geometry using the GEANT4 Application for Tomographic Emission (GATE)\(^6\) Monte Carlo simulation code. Each simulation consisted of 1 MBq distributed uniformly within water.

**Subdermal tissue self-dose**

The mass of infiltrated tissue depends on the volume of extravasated radiopharmaceutical and penetration into the subdermal fascia. We calculated the absorbed doses (Gy) to infiltrated tissues by taking into account the tissue mass, total energy emitted in the source region, and the energy absorbed fraction according to the MIRD schema (Bolch et al. 2009).

**Relevant skin dose**

The National Council on Radiation Protection and Measurements recommends (NCRP 2018) for occupational exposure that the absorbed dose in skin at a depth of 70 \(\mu m\) be limited to 0.5 Gy averaged over the most highly exposed 10 \(cm^2\) of skin. Skin dose assessments in units of shallow dose equivalent (Sv) are required by the Code of Federal Regulations in 10 CFR 20.1201(c) for a contiguous 10 \(cm^2\) area of skin at a tissue depth of 0.007 cm (7 mg \(cm^{-2}\)). For regulatory compliance with recommended skin dose limits, the software code VARSkin, version 6.1 (Hamby and Mangini 2018), was written to calculate occupational dose from radioactive contamination on or near the skin. We applied it to patient radiopharmaceutical infiltrations. For cases involving low-LET radiations, dose expressed in units of Gy and Sv are numerically (approximately) equivalent.

Because infiltrated tissue lies beneath and adjacent to the skin epidermis, we defined the relevant target for calculating dose to overlying skin as a thin layer comprising the sensitive epithelial basal cells with an area of 10 \(cm^2\) and at a tissue depth beneath the skin surface of 0.007 cm (7 \(\mu m\) or 7 mg \(cm^{-2}\)). We assumed that the dose limits to patient skin should be the same or less than those for occupational exposures. We modeled infiltrated subdermal tissue as a three-dimensional thin cylinder, and calculated the relevant skin dose using a modified VARSkin 6.1 computer code by setting the distance between the infiltrated source tissue and the sensitive basal cell layer to 10 \(\mu m\) (1 mg \(cm^{-2}\)) and removing backscatter correction.

**RESULTS**

Fig. 2 shows one example of the 26 actual recorded count rate data and the corresponding curve fit to a single-exponential function for an extravasated patient.

Details of the tissue geometries modeled for each infiltration, together with calculated energy absorbed fractions for \(^{18}\text{F}\) and \(^{99m}\text{Tc}\) are shown in Table 1. Infiltrated tissue self-dose (Gy) and the skin shallow dose equivalent (Sv) for 26 infiltrated patients are show in Table 2. In each case, either (or both) the infiltrated tissue dose or the adjacent skin shallow dose equivalent exceeded the limiting value 0.5 Gy or 0.5 Sv.

**DISCUSSION**

In this work, we investigated extravasations of \(^{18}\text{F}\)-FDG and \(^{99m}\text{Tc}\)-MDP, but the methods described herein also apply to all other radiopharmaceuticals and amounts administered. The positron energy of \(^{18}\text{F}\) resulted in significant tissue self-dose and significant dose to overlying skin. Despite the relatively low absorbed fractions for \(^{99m}\text{Tc}\), we found that \(^{99m}\text{Tc}\)-labeled agents can produce significant tissue absorbed doses. Our results in 26 cases exceeded radiation protection (NCRP 2018) and regulatory\(^7\) limits for extremity tissue (0.5 Gy) and skin (0.5 Sv).

The literature contains several examples of adverse tissue reactions following extravasation of diagnostic and therapeutic radioisotopes such as \(^{201}\text{TI}\) (van der Pol et al. 2017), \(^{90}\text{Y}\) (Williams et al. 2006; Siebeneck 2008), \(^{89}\text{Sr}\) (Kawabe et al. 2013), \(^{131}\text{I}\) (Breen and Dreidger 1991; Bonta et al. 2011; van der Pol et al. 2017), and \(^{32}\text{P}\) (Minsky et al. 1987). We found one published example of radiopharmaceutical extravasation leading directly to a highly localized cancerous lesion (Benjegerdes et al. 2017): following extravasation of \(^{223}\text{Ra}\)-dichloride, the patient developed aggressive squamous cell carcinoma at the injection site.

Because extravasations are common (Wong et al. 2019) and can lead to adverse tissue reaction, prompt identification and mitigation are important factors. In our review, none of the technologists or patients reported immediate...
pain or edema during or following the injection—even in cases of extravasation—emphasizing the difficulty in prompt extravasation identification. Mitigation steps such as elevation of the arm, application of heat (Yucha et al. 1994; Goolsby and Lombardo 2006), and flushing with saline can accelerate clearance and decrease radiation doses.

Once an extravasation has been identified, accurate dose calculation enables clinicians to identify patients who should be followed for adverse tissue reactions or late stochastic effects. Absence of immediate visible skin reactions is a common explanation for not reporting and following up after extravasation events. However, given the expected time for presentation of symptoms, it is unlikely that extravasation-related injury would be discovered. van der Pol et al. (2017) reported that, despite an extensive literature review, only 3,016 published cases of diagnostic radiopharmaceutical extravasation were found. Of those, only three cases included dosimetry calculation and patient follow-up. All three patients who were followed were found to suffer adverse tissue reactions. In one case, a radiation ulcer was diagnosed after 2 y. In a second case, the radiation ulcer diagnosis was made after 3 y. Of the remaining 3,013 cases, none described dosimetric parameters or follow-up (van der Pol et al. 2017).

In cases of 99mTc-MDP extravasation, immediate skin reactions are not likely. Our data review suggests that the shallow dose equivalent to the skin may be low even in cases where the absorbed dose to infiltrated tissue is high. Absence of prompt skin reactions should not dissuade clinicians from considering delayed detrimental effects to tissue and skin. Proper documentation and patient follow-up may protect medical institutions from frivolous litigation and unwarranted regulatory review.

CONCLUSION

Extravasation events in nuclear medicine are rarely fully characterized—including accurate dosimetry and appropriate clinical follow-up. Accurate dosimetry should include the determination of infiltrated fraction of administered activity, clearance half-times, and resulting radiation doses to infiltrated tissue and overlaying skin. We investigated injection-site count-rate data for 26 cases of extravasation of 18F-FDG and 99mTc-MDP, assuming three source-tissue geometries. For cases reported in this paper, radiation absorbed doses to infiltrated tissue ranged from 0.6 Gy to 11.2 Gy, and the shallow dose equivalent to a 10 cm² area of adjacent skin ranged from about 0.1 Sv to 5.4 Sv.

With patient radiation safety in mind, we maintain that both diagnostic and therapeutic extravasation events should be identified and characterized. Severe extravasations affect the diagnostic or therapeutic quality of nuclear medicine procedures, and the unintended dose to tissue and skin may eventually be clinically significant. A dedicated radiopharmaceutical injection monitoring system can be used to improve the accuracy of dosimetry and assist in determining the need for patient follow-up.

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Table 1. Representative tissue geometry details and energy absorbed fractions.

| Geometry | Dimensions (cm) | Absorbed fraction for 18F | Absorbed fraction for 99mTc |
|----------|----------------|--------------------------|---------------------------|
| Cylinder | h = 0.1, r = 4 | 73%                      | 11%                       |
| Ellipsoid| a = 2.13, b = 1.07, c = 0.53 | 95% | 13% |
| Sphere  | r = 1.07        | 97%                      | 13%                       |

Table 2. Detailed dosimetry results.

| Case # | Radiopharmaceutical | Effective clearance half-time (min) | Mean absorbed dose to infiltrated fascia (Gy) | Shallow dose equivalent to skin (Sv) |
|--------|----------------------|-----------------------------------|------------------------------------------|-----------------------------------|
| 1      | 18F-FDG              | 9                                 | 0.6                                      | 0.3                               |
| 2      | 18F-FDG              | 43                                | 7.6                                      | 3.7                               |
| 3      | 18F-FDG              | 93                                | 2.7                                      | 1.3                               |
| 4      | 18F-FDG              | 24                                | 8.4                                      | 4.1                               |
| 5      | 18F-FDG              | 13                                | 0.8                                      | 0.4                               |
| 6      | 18F-FDG              | 22                                | 0.7                                      | 0.3                               |
| 7      | 18F-FDG              | 44                                | 0.9                                      | 0.4                               |
| 8      | 18F-FDG              | 39                                | 11.2                                     | 5.4                               |
| 9      | 18F-FDG              | 70                                | 1.0                                      | 0.5                               |
| 10     | 18F-FDG              | 38                                | 8.7                                      | 4.2                               |
| 11     | 18F-FDG              | 22                                | 3.8                                      | 1.9                               |
| 12     | 18F-FDG              | 41                                | 0.6                                      | 0.3                               |
| 13     | 99mTc-MDP            | 360                               | 8.4                                      | < 0.1                             |
| 14     | 18F-FDG              | 46                                | 1.0                                      | 0.5                               |
| 15     | 99mTc-MDP            | 64                                | 1.5                                      | < 0.1                             |
| 16     | 99mTc-MDP            | 218                               | 5.3                                      | < 0.1                             |
| 17     | 99mTc-MDP            | 38                                | 0.9                                      | < 0.1                             |
| 18     | 99mTc-MDP            | 49                                | 1.2                                      | < 0.1                             |
| 19     | 99mTc-MDP            | 64                                | 1.5                                      | < 0.1                             |
| 20     | 18F-FDG              | 18                                | 1.1                                      | 0.5                               |
| 21     | 18F-FDG              | 22                                | 5.1                                      | 2.5                               |
| 22     | 99mTc-MDP            | 36                                | 0.9                                      | < 0.1                             |
| 23     | 18F-FDG              | 24                                | 6.8                                      | 3.3                               |
| 24     | 18F-FDG              | 79                                | 2.9                                      | 1.4                               |
| 25     | 18F-FDG              | 26                                | 0.8                                      | 0.4                               |
| 26     | 18F-FDG              | 22                                | 3.6                                      | 1.8                               |

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1Official Transcript of Proceedings, NRC ACMU1. Available at https://www.nrc.gov/docs/ML0903/ML090340745.pdf. Accessed 25 August 2020
helpful advice concerning modification and implementation of VARSkin 6.1 for deep-tissue sources.

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