Accurate Dose Calibrator Activity Measurement of $^{90}$Y-Ibritumomab Tiuxetan

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This investigation examined the accuracy of dose calibrator activity measurement of the $\beta$-emitting radiopharmaceutical $^{90}$Y-ibritumomab tiuxetan. **Methods:** Five different facilities independently measured $^{90}$Y in a 10-mL syringe geometry with 30 dose calibrator models from 3 different manufacturers. The activities ranged from 81.4 MBq (2.2 mCi) to 1,406 MBq (38 mCi) over the volume range of 3–9 mL. **Results:** The mean dial settings for $^{90}$Y measurement were 375, 51 $\times$ 10, and 897 $\times$ 100 for Atomlab, CRC, and Mark V dose calibrators, respectively. The maximum volume dependence was 0.28%/mL. **Conclusion:** This study demonstrated that when measuring all volumes of $^{90}$Y-ibritumomab tiuxetan activity prescriptions, only a single dial setting for a given manufacturer’s dose calibrator is required for accurate measurements. Volume corrections are not necessary. For best accuracy, an individually determined dial value should be used.

**Key Words:** $^{90}$Y measurement; dose calibrator; ibritumomab tiuxetan

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The U.S. Food and Drug Administration has approved for commercial use the radioimmunotherapeutic agent $^{90}$Y-ibritumomab tiuxetan ($^{90}$Y-Zevalin; IDEC Pharmaceuticals Corp.) for the treatment of non-Hodgkin’s lymphoma (1). This radiopharmaceutical is generally prepared at a commercial radiopharmacy and then supplied to medical facilities as a unit dosage in a 10-mL syringe with volumes ranging from 3 to 9 mL, dependent on the prescribed activity for an individual patient.

All Nuclear Regulatory Commission and Agreement State licensees must determine and record the activity of unsealed by-product material before medical use. Except in certain Agreement States, this activity determination does not require the use of a dose calibrator, pursuant to 10 CFR part 35.63 (2), provided unit dosages are obtained from an appropriately licensed manufacturer or preparer. However, because the package insert for $^{90}$Y-ibritumomab tiuxetan (1) states that patient dosages should be measured immediately before administration, licensees may prefer to directly measure activity with a dose calibrator.

Commercial reentrant ionization chambers (dose calibrators) are the de facto standard instrument to measure radioactivity in nuclear medicine. The dose calibrator measurement of $\beta$-emitting radionuclides depends on the bremsstrahlung radiation produced from the $\beta$-interaction with the source matrix, its container, and the calibrator chamber wall. The use of different volumes or containers may result in measurement errors, as is the case for low-energy photon emitters.

The purpose of this study was to involve the National Institute of Standards and Technology (NIST), dose calibrator manufacturers, and a commercial radiopharmacy in a common effort to investigate the applicability of a single calibrator dial setting for a particular manufacturer’s dose calibrator model and determine the significance of volume corrections for accurate measurement of the $\beta$-emitting radiopharmaceutical $^{90}$Y-ibritumomab tiuxetan in a syringe geometry.

**MATERIALS AND METHODS**

Dose calibrator measurements of $^{90}$Y were performed independently at 5 different sites: Capintec, Inc., NIST, Cardinal Health Nuclear Pharmacy Services, Cardinal Health Radiation Management Services (Nuclear Associates), and Sun Nuclear Corp. (sites 1, 2, 3, 4, and 5, respectively). Thirty dose calibrators of the pressurized argon well reentrant design were used, including CRC (Capintec), Mark V (Cardinal Health Radiation Management Services), and Atomlab (Sun Nuclear; distributed by Biodex Medical Systems Inc.). Table 1 summarizes the various dose calibrators and procedures used at each site.

The $^{90}$Y was delivered to site 3 by MDS Nordion in a 2-mL closed-septum vial containing a $^{90}$Y-chloride solution with a product data sheet that indicated the NIST-traceable activity concentration, volume, and total activity. The radioactive solution was transferred from the vial to a 10-mL syringe (Becton Dickinson & Co.). Dose calibrator measurements for $^{90}$Y after transfer to the syringe were based on vial measurements and an activity difference method. This procedure specifies measurement of the activity...
in the vial, both before and after removal of source material, but with the volume in the vial restored to its initial value with saline before remeasurement. The difference between these 2 vial measurements is the activity drawn into the syringe, which is still NIST traceable. Site 3 established traceability for the $^{90}$Y vial measurements through prior proficiency testing in a measurement assurance program with NIST. Calibrated dial settings for $^{90}$Y measurement for each calibrator were determined by adjusting the dial settings to read the correct activity; the standard uncertainty on the activity value in the present study was based primarily on the standard uncertainty on the activity provided by Nordion, which was $\pm 5\%$. Sites 4 and 5 received a calibrated activity in a syringe from site 3; site 1 received a calibrated activity in an MDS Nordion vial and performed a nontraceable volumetric transfer of activity into the syringe.

Activity measurements were made either at a start volume of 9 mL and after sequential 1-mL volume withdrawals to a final volume of 3 mL or at a start volume of 3 mL and after sequential 1-mL volume additions to a final volume of 9 mL. The latter procedure was different in that only the volume was varied; the activity remained constant.

The correction factor for each volume was obtained by comparing the measured activity with the calculated activity for each volume. The volume correction factor is given by:

\[
\text{Correction factor} = \frac{\text{calculated } ^{90}\text{Y activity}}{\text{measured } ^{90}\text{Y activity}}
\]

where the calculated $^{90}$Y activity for each volume is a constant either for the 3-mL start volume or for the 9-mL start volume; it is the original calibrated activity multiplied by the respective volume divided by 9.

### TABLE 1
Dose Calibrators Used and $^{90}$Y Activity Measurement Procedures at Each Site

<table>
<thead>
<tr>
<th>Site</th>
<th>Calibrator</th>
<th>$^{90}$Y Activity measurement</th>
<th>Procedure</th>
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| 1    | 5 CRC-15R  | $^{90}$Y in saline*            | Calibrated dial setting determination:  
Start activity and volume of 1,406 MBq (38 mCi) in 3 mL  
Volume dependence:  
Start activity and volume of 81.4 MBq (2.2 mCi) in 3 mL  
Sequential addition of 1 mL saline up to 9 mL |
| 2    | CRC-12, CRC-15R, 35R, and Atomlab 100 | $^{90}$Y in saline | A. Start activity and volume of 1,184 MBq (32 mCi) in 8 mL  
B. Dilute with YCl$_3$/HCl carrier to give enough volume for preparation of syringes and liquid scintillation master  
C. Accurately (to within 0.1%) dispense large volume master into syringes whose needles had been previously sealed to prevent leaks—total of 14 syringes prepared over the volume range, with several repeated preparations at 3, 5, 7, and 9 mL to study syringe variability and repeatability  
D. Prepare nominally 200-fold dilution of large volume master for liquid scintillation measurements  
E. Determine activity concentration of diluted solution and master by liquid scintillation counting using 2 different, independent techniques |
| 3    | 3 CRC-15R  | $^{90}$Y ibritumomab tiuxetan in formulation buffer | Start activity and volume of 1,184 MBq (32 mCi) in 9 mL  
Sequential withdrawal of 1 mL of solution volume down to 3 mL  
Sequential addition of 1 mL ibritumomab tiuxetan in formulation buffer up to 9 mL |
| 4    | 3 Mark V   | $^{90}$Y in saline            | Start activity and volume of 509 MBq (13.8 mCi) in 3 mL  
Sequential addition of 1 mL saline up to 9 mL |
| 5    | 15 Atomlab†| $^{90}$Y in saline‡           | Start activity and volume of 481 MBq (13.0 mCi) in 3 mL  
Sequential addition of 1 mL saline up to 9 mL |

*Activity for calibrated dial setting determination and volume dependence study are different because of use of different syringes.  
†Ten new and 5 repaired units as old as 11 y; all 4 models of this type of dose calibrator use the same chamber assembly, so there is no need to distinguish between models.  
‡During initial volume test, on the 8 mL step, a backfill into the saline vial resulted in source material loss, but initial calibration for activity measurement was unaffected. Volume test was repeated, with a start activity of 92.5 MBq (2.5 mCi) in 3 mL. Repeated results are reported; no backfill was detected.
The measurements performed by site 2 used a different approach involving a direct determination of the solution activity by liquid scintillation counting (3–5) and determination of calibration settings for a set of syringes, each independently prepared with different volumes covering the 3- to 9-mL range. The amount of \(^{90}\text{Y}\) solution added to each of the syringes was carefully controlled using an automated dispenser (Hamilton Co.) having an accuracy of 0.1%. In addition to the measurements made with \(^{90}\text{Y}\)-ibritumomab tiuxetan, the same procedure was repeated by site 2 for syringes containing \(^{90}\text{Y}\) in a carrier solution containing additional \(\text{YCl}_3\) and 1 \(\text{mol} \cdot \text{L}^{-1}\) \(\text{HCl}\) to ascertain what effect the different source matrix may have on dose calibrator measurement because of differences in self-absorption or bremsstrahlung production.

The calibrated dial settings were also converted to response values for the ion chamber of the dose calibrator to compare the different manufacturers’ values. The relationship between chamber response, \(C_R\), normalized to \(^{60}\text{Co}\) and dial setting, \(\text{DS}\), can be expressed as:

\[
C_R = \frac{\text{DS} \cdot 0.0855}{M}
\]

where \(M\) is the display multiplier of 10; and

\[
C_R = \frac{120(1.009 - \text{DS})}{M}
\]

for Mark V, where \(M\) is the display multiplier of 10.

**RESULTS**

The calibrated dial settings for \(^{90}\text{Y}\) measurement for each dose calibrator at each site are given in Table 2. For the Atomlab, the mean calibrated dial setting measured at site 5 was 375, with a range of 363–394 for 15 calibrators. Of the 15 calibrators, 10 were new and their dial settings exhibited a narrower range: 372–378. The mean calibrated dial setting at site 2 for 1 Atomlab was 393, with a range of 387–399.

For the 5 new CRC calibrators, the mean calibrated dial setting measured by site 1 was 50 \(\pm\) 10, with a range of 47 \(\times\) 10 to 53 \(\times\) 10. The mean calibrated dial setting at site 2 for 3 CRCs was 55.7 \(\times\) 10, with a range of 54 \(\times\) 10 to 58 \(\times\) 10. The mean calibrated dial setting at site 3 for 3 CRCs was 47.3 \(\times\) 10, with a range of 47 \(\times\) 10 to 48 \(\times\) 10. There was no overlap in the ranges between sites 1 and 2 or between sites 3 and 2. For the Mark V calibrators, the mean calibrated dial setting measured was 897 \(\times\) 100, with a range of 896 \(\times\) 100 to 897 \(\times\) 100; no other sites studied these calibrators.

Based on the calibrated dial settings, the Atomlab chamber response value at site 2 was 0.01272 and the mean value at site 5 was 0.01333, a factor of 4.8% higher. The mean CRC response was 0.01369 at site 2 and 0.01317 at site 1, a factor of 3.8% lower. The mean CRC response at site 3 was 0.01292, a factor of 5.6% lower than the site 2 value. The mean Mark V response at site 4 was 0.01071.

For the measurements at site 2, the expanded \((k = 2)\) uncertainty on the calibrated dial settings based on the liquid scintillation activity calibration was determined to be 1.6% and was calculated from the quadratic combination of the average SD and the mean deviate estimate calculated from range statistics (6). Similar uncertainty analysis was not performed at the other sites.

Volume correction factors determined by each site, normalized to 6 mL, are shown in Figures 1–4. The equation and correlation coefficient \((r)\) for each line, resulting from linear regression analysis, are given in each figure. The largest volume dependence was determined to be 0.28%/mL. Using this maximum observed volume variation and the applicable volume range of 6 mL, the volume effect for all dose calibrators included in this study should be limited to 1.7%.

A comparison of \(^{90}\text{Y}\) measurements made with ibritumomab tiuxetan and the NIST standard solution in the identical measurement geometry indicated no difference in results to within the expanded measurement uncertainty of 1.6%. Moreover, variability in measurement results for both solutions due to variability in syringe manufacture was found to be less than 0.26% (SD), inclusive of 0.1% vari-

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<td><strong>Calibrated Dial Setting for Each Dose Calibrator at Each Site</strong></td>
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<td><strong>Site</strong></td>
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*Uncertainties on the dial settings are expanded uncertainties determined from the combined standard uncertainty on the \(^{90}\text{Y}\) activity calibration. The uncertainties at the other sites were not evaluated; thus, only ranges of values are given.

For the CRC dose calibrators, all dial settings are \(\times\) 10; that is, the instrument readout must be multiplied by a factor of 10 to obtain the correct activity value. For the Mark V dose calibrators, all dial settings are \(\times\) 100.
ability due to uncertainty in filling volume, at a volume of 5 mL for 8 syringes.

DISCUSSION

Proper dose calibrator measurement of pure β-emitting radionuclides is important for the safe and accurate dosing of various radionuclide therapies in nuclear medicine. According to NUREG-1556, volume 13 (7), and Nuclear Regulatory Commission Information Notice 2002-19 (8), accurate measurement of pure β-emitters is a potential problem. This study was performed to explore the pitfalls and examine the possible solutions when using dose calibrators for accurate measurement of 90Y-ibritumomab tiuxetan.

This study indicated that use of a single calibrated dial setting for a given manufacturer’s dose calibrator resulted in accurate measurements of 90Y in a syringe geometry regardless of volume in a 3- to 9-mL range. Although different model dose calibrators have very different dial settings, their chamber response values are very similar. The 90Y measurement variation based on the calibrator response values for the range of observed dial settings was determined to be within ±5%, which is the level of the standard

FIGURE 1. Correction factor (CF) as function of volume normalized to 6 mL, determined by site 1 based on measurements obtained on 1 CRC dose calibrator. Volume dependence is 0.28%/mL.

FIGURE 2. Correction factor (CF) as function of volume normalized to 6 mL, determined by site 2 based on measurements obtained on 3 CRC and 1 Atomlab dose calibrators. Volume dependence is 0.21%/mL and 0.16%/mL, respectively.

FIGURE 3. Correction factor (CF) as function of volume normalized to 6 mL, determined by site 3 based on average of measurements obtained on 3 CRC dose calibrators. Bold line (●) represents results for sequential 1-mL volume withdrawal from 9 to 3 mL, and dashed line (○) represents results for sequential 1-mL volume addition from 3 to 9 mL. Volume dependence is 0.20%/mL and 0.11%/mL, respectively.

FIGURE 4. Correction factor (CF) as function of volume normalized to 6 mL, determined by sites 4 (●) and 5 (■) based on average of measurements obtained on 3 Mark V and 15 Atomlab dose calibrators, respectively. Volume dependence is 0.12%/mL and 0.13%/mL, respectively.
uncertainty on the activity value as provided by the supplier, MDS Nordion. This uncertainty was propagated to the activity values provided to all participants in the study by the commercial radiopharmacy, with the exception of NIST, which performed its own independent activity calibration, and site 1, which used nominal techniques.

For best accuracy, it is recommended that the single calibrated dial setting be an individually determined value, using the reported range for the appropriate manufacturer’s dose calibrator as a guide. (The limited range of reported Mark V values does not allow users to check their calibrated dial setting for reasonableness in this manner.) Specifically, we recommend, first, that each radiopharmacy establish a $^{90}\text{Y}$-calibrated dial setting based on NIST-supplied or NIST-traceable activity sources so that each activity source supplied to a medical facility can be used as a secondary reference standard and, second, that each medical facility determine its own calibrated dial setting based on the initial $^{90}\text{Y}$ activity received from a commercial radiopharmacy or, alternatively, based on measurement of a NIST-traceable activity source in the same syringe geometry.

Using this approach, volume correction factors should not be necessary when measuring the activity of $^{90}\text{Y}$-ibritumomab tiuxetan at any volume in the range of 3–9 mL using the same type of syringe used in this study.

The type of syringe holder (e.g., T-handle or hook style dipper) used and inconsistent use of the protective well liner may cause further measurement variations. It is recommended that facilities always have the well liner installed during dose calibrator measurement of $^{90}\text{Y}$ and not interchange syringe holders once they have established their calibrated $^{90}\text{Y}$ dial setting.

**CONCLUSION**

This study demonstrated that, for accurate measurements, no adjustment is necessary for a dose calibrator dial setting when measuring different volumes of $^{90}\text{Y}$-ibritumomab tiuxetan activity prescriptions. Medical facilities need only establish their own calibrated dial setting for $^{90}\text{Y}$ using their first prescription measurement based on the stated activity of the radiopharmacy.

**ACKNOWLEDGMENT**

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**REFERENCES**