

The MIRD Schema for Radiopharmaceutical Dosimetry: A Review

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Internal dosimetry evaluates the amount and spatial and temporal distributions of radiation energy deposited in tissue from radionuclides within the body. Historically, nuclear medicine had been largely a diagnostic specialty, and the implicitly performed risk-benefit analyses have been straightforward, with relatively low administered activities yielding important diagnostic information whose benefit far outweighs any potential risk associated with the attendant normal-tissue radiation doses. Although dose estimates based on anatomic models and population-average kinetics in this setting may deviate rather significantly from the actual normal-organ doses for individual patients, the large benefit-to-risk ratios are very forgiving of any such inaccuracies. It is in this context that the MIRD schema was originally developed and has been largely applied. The MIRD schema, created and maintained by the MIRD committee of the Society of Nuclear Medicine and Molecular Imaging, comprises the notation, terminology, mathematic formulas, and reference data for calculating tissue radiation doses from radiopharmaceuticals administered to patients. However, with the ongoing development of new radiopharmaceuticals and the increasing therapeutic application of such agents, internal dosimetry in nuclear medicine and the MIRD schema continue to evolve—from population-average and organ-level to patient-specific and suborgan to voxel-level to cell-level dose estimation. This article will review the basic MIRD schema, relevant quantities and units, reference anatomic models, and its adaptation to small-scale and patient-specific dosimetry.

Key Words: radiobiology/dosimetry; radionuclide therapy; radiopharmaceuticals; dosimetry; MIRD; radiopharmaceutical therapy

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Internal dosimetry evaluates the amount and spatial and temporal distributions of radiation energy deposited in tissue from radionuclides within the body. It has been applied to the determination of tissue-absorbed doses and related

quantities for occupational exposures in radiation protection, environmental exposures in radiation epidemiology, and diagnostic and therapeutic radiopharmaceuticals in nuclear medicine. For diagnostic radiopharmaceuticals, dosimetry is generally based on reference, or standard, anatomic models and average human or, oftentimes, animal kinetic (i.e., time-activity) data. Historically, nuclear medicine had been largely a diagnostic specialty, and the associated risk-benefit analyses implicitly performed by the clinician have been straightforward: relatively low administered activities yield important diagnostic information whose benefit far outweighs any potential risk associated with the attendant normal-tissue radiation doses. Although the dose estimates based on anatomic models and population-average kinetics may deviate rather significantly from the actual normal-organ doses for individual patients, the large benefit-to-risk ratios for diagnostic nuclear medicine are very forgiving of any such inaccuracies. Such dose estimates are nonetheless useful for first-order assessment of the relatively low stochastic risk (probability of harm) associated with diagnostic agents as well as for dosimetric intercomparison of different radiopharmaceuticals and imaging procedures and methodologic refinement of such procedures to minimize patient doses. It is in this context that the MIRD schema was originally developed and has been largely applied. The MIRD schema refers, of course, to the formalism created and maintained by the MIRD committee of the Society of Nuclear Medicine and Molecular Imaging; it comprises the notation, terminology, mathematic formulas, and reference data for calculating tissue radiation doses from radiopharmaceuticals administered to patients (1–3).

By incorporation of appropriate radionuclides and appropriately large activities into target tissue-avid radiopharmaceuticals, a sufficiently high radiation dose may be delivered to produce a therapeutic response in tumor or other target tissue. With escalating administered activities and associated normal-tissue doses, serious radiation injury can ensue, however. To optimize radiopharmaceutical therapy, it is critical to establish patient-specific target-tissue and at-risk normal-tissue radiation doses with reasonable accuracy and precision

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and to develop and apply reliable dose–response relationships for target tissues and dose–toxicity relationships for normal tissues. With the ongoing development of new radiopharmaceuticals and the increasing therapeutic application of such agents, internal dosimetry in nuclear medicine and the MIRD schema continue to evolve—from population-average and organ-level to patient-specific and localized (or voxel-level) dose estimation (3,4).

This article will review the basic MIRD schema, relevant quantities and units, reference anatomic models, and, briefly, its adaptation to patient-specific dosimetry in the setting of radiopharmaceutical therapy. Multiple-choice questions and answers, with detailed explanations of the correct answers, are provided. Radiopharmaceutical absorbed doses and related dose metrics are determined largely by calculation, rather than by direct measurement, and internal dosimetry is, to some extent, unavoidably mathematic. However, the mathematics used are essentially arithmetic, and knowledge of calculus and other advanced topics is not required to read and understand this article.

QUANTITIES AND UNITS

Before addressing the computational engine of the MIRD schema—the actual dose-calculation formulas—one should first be conversant with its language, that is, the quantities, parameters, and symbols that populate the schema. These are detailed in Table 1. For completeness, additional important quantities encountered in radiation dosimetry (5–8) are presented next.

Absorbed Dose

Perhaps the most widely used and biologically meaningful quantity for expressing radiation dose, the absorbed dose, D , is defined as follows:

$$D \equiv \frac{d\bar{E}}{dm}, \quad \text{Eq. 1}$$

where $d\bar{E}$ is the mean energy imparted by ionizing radiation to matter and dm is the mass of matter to which the energy is imparted. The SI unit of absorbed dose is the gray (1 Gy = 1 J kg⁻¹), and the older unit is the rad (1 rad = 100 erg g⁻¹); 1 Gy equals 100 rad and 1 rad equals 1 cGy (or 10 mGy).

Linear Energy Transfer

The quality as well as the quantity of radiation are important determinants of the frequency or severity of radiogenic biological effects. The quality of a radiation is related to the characteristics of the microscopic spatial distribution of energy-deposition events. Sparsely ionizing radiations such as x- and γ -rays and intermediate- to high-energy electrons and β -particles are characterized as low-quality radiations, whereas densely ionizing radiations such as low-energy electrons (e.g., Auger electrons), protons, neutrons, and α -particles are typically characterized as high-quality radiations. For the same absorbed dose, the probability or

severity of biological effects is generally less for sparsely ionizing than for densely ionizing radiations.

The quality of radiation is characterized by the linear energy transfer:

$$\text{Linear energy transfer} \equiv \frac{dE}{dl}, \quad \text{Eq. 2}$$

where dE is the energy deposited by a charged particle (or the secondary charged particle produced by the primary radiation) over a pathlength dl traversed in matter. The SI unit of linear energy transfer is the J m⁻¹, and the older unit is the keV μm^{-1} ; 1 J m⁻¹ equals 6.25×10^9 keV μm^{-1} and 1 keV μm^{-1} equals 1.60×10^{-10} J m⁻¹. In practice, linear-energy-transfer values are most commonly expressed in the older unit of keV μm^{-1} .

Relative Biological Effectiveness

The influence of linear energy transfer and other radiation properties on the probability or severity of biological effects is quantified by the relative biological effectiveness:

$$\text{Relative biological effectiveness (A)} \equiv \frac{D_{\text{reference}}}{D_A}, \quad \text{Eq. 3}$$

where $D_{\text{reference}}$ is the absorbed dose of reference radiation (typically a widely available sparsely ionizing radiation such as ⁶⁰Co γ -rays) required to produce a specific, quantitatively expressed biological effect under specific irradiation conditions and D_A is the absorbed dose of radiation A (i.e., the radiation for which the relative biological effectiveness is being determined) required to produce the same probability or severity of the same specific biological effect under the same conditions as those for the reference-radiation irradiation.

Equivalent Dose

An important dosimetric quantity, defined by the International Commission on Radiological Protection (ICRP) and used in radiation protection, is the equivalent dose, which is derived from the absorbed dose. Specifically, equivalent dose is the product of an organ- or tissue-absorbed dose and a radiation weighting factor (w_R), which ranges from unity for sparsely ionizing radiations such as x- and γ -rays and β -particles to 20 for densely ionizing radiations such as α -particles (9). The high weighting-factor value for α -particles reflects the greater potential per unit absorbed dose of high-linear-energy-transfer, densely ionizing radiations to induce stochastic effects in an organ or tissue (such as cancer and germ-cell, or hereditary, mutations) relative to that of a unit absorbed dose of sparsely ionizing radiations. The equivalent dose is thus defined as follows:

$$H(r_T, \tau) \equiv \sum_R w_R D_R(r_T, \tau), \quad \text{Eq. 4}$$

where w_R is the radiation weighting factor for radiation R , and $D_R(r_T, \tau)$ is the contribution of radiation R to the mean absorbed dose to tissue r_T over a time interval τ . The SI unit for equivalent dose is the J kg⁻¹, with the special

TABLE 1
Terminology and Nomenclature Used in MIRD Schema (1–3)

Parameter	Symbol	Definition	Unit*			Comment
			SI	Older		
Administered activity	A_0	Activity of radiopharmaceutical administered to patient	MBq	mCi		Base SI unit of activity is becquerel ($\equiv 1$ disintegration per second [dps]); older unit of activity, curie, corresponds to 3×10^{10} dps; 37 MBq equals 1 mCi and 37 kBq equals 1 μ Ci; it is important to distinguish activity administered (often mistakenly referred to as dose) from actual radiation dose
Source region	r_s	Region (such as tissue or organ) containing radionuclide and within which radiation is emitted as radionuclide undergoes decay	Not applicable	Not applicable		
Target region	r_T	Region (such as tissue or organ) within which energy of emitted radiations is deposited	Not applicable	Not applicable		For systemically administered and distributed radiopharmaceutical, every tissue or organ is both source and target region
Time	t	Time after administration of radiopharmaceutical	h	h		Time of administration of radiopharmaceutical is designated as time zero, that is, as $t = 0$
Activity	$A(r_s, t)$	Activity in source region r_s at time t	MBq	mCi		Plot of activity in source region r_s vs. time t is source-region time-activity curve
Time-integrated activity	$\bar{A}(r_s, \tau)$	Number of radioactive decays in source region r_s from time $t = 0$ to time $t = \tau$	MBq-s	μ Ci-h		$\bar{A}(r_s, \tau) = \int_0^\tau A(r_s, t) dt$; integration time τ is commonly set to infinity, yielding total number of decays in source region r_s for complete decay of administered radionuclide; time-integrated activity was originally called cumulated activity
Time-integrated activity coefficient†	$\bar{a}(r_s, \tau)$	Number of radioactive decays in source region r_s from time $t = 0$ to time $t = \tau$ per unit administered activity	MBq-s MBq ⁻¹	μ Ci-h mCi ⁻¹		$\bar{a}(r_s, \tau) = \bar{A}(r_s, \tau)/A_0$; time-integrated activity coefficient was originally called residence time
Absorbed dose	$D(r_T, \tau)$	Absorbed dose to target region r_T from time $t = 0$ to time $t = \tau$	Gy	rad		Integration time τ is commonly set to infinity, yielding total absorbed dose to target region r_T for complete decay of administered radionuclide
Absorbed dose coefficient	$d(r_T, \tau)$	Absorbed dose to target region r_T from time $t = 0$ to time $t = \tau$ per unit administered activity	Gy MBq ⁻¹	rad mCi ⁻¹		$d(r_T, \tau) = \frac{D(r_T, \tau)}{A_0}$
Mass	$M(r_s), M(r_T)$	Mass of source region r_s and target region r_T , respectively	g	g		
Energy	E_i	Energy (or mean energy) per particle or photon of i th radiation emitted by administered radionuclide	MeV	MeV		
Number of radiations per decay	Y_i	Number of i th radiation emitted per radioactive decay for administered radionuclide	Not applicable	Not applicable		
Energy per decay	Δ_i	Energy (or mean energy) of i th radiation emitted by administered radionuclide per radioactive decay	MeV	MeV		$\Delta_i = E_i Y_i$
Absorbed fraction	$\phi(r_T \leftarrow r_s, E_i)$	Absorbed fraction, that is, fraction of energy E_i of i th radiation emitted within source region r_s that is absorbed in target region r_T	Not applicable	Not applicable		
Specific absorbed fraction	$\Phi(r_T \leftarrow r_s, E_i)$	Specific absorbed fraction, that is, fraction of energy E_i of i th radiation emitted within source region r_s that is absorbed in target region r_T per unit mass of target region r_T	g ⁻¹	g ⁻¹		$\Phi(r_T \leftarrow r_s, E_i) = \frac{\phi(r_T \leftarrow r_s, E_i)}{M(r_T)}$
S value	$S(r_T \leftarrow r_s)$	Absorbed dose to target region r_T per radioactive decay in target region r	mGy MBq ⁻¹ s ⁻¹	rad μ Ci ⁻¹ h ⁻¹		$S(r_T \leftarrow r_s) = \frac{1}{M(r_T)} \sum_i E_i Y_i \phi(r_T \leftarrow r_s, E_i) = \frac{1}{M(r_T)} \sum_i \Delta_i \Phi(r_T \leftarrow r_s, E_i)$

*Although older units are presented for completeness, SI units should be used exclusively.

†In modern MIRD-schema nomenclature (1), dosimetric quantity normalized to administered activity is identified by term coefficient appended to quantity name.

named unit of sievert. The older unit is the erg g⁻¹ or the special named unit of rem; 1 Sv equals 100 rem, and 1 rem equals 1 cSv (or 10 mSv). Current ICRP-recommended values of w_R , which is a dimensionless quantity, are 1.0 for photons, electrons, positrons, and β -particles and 20 for α -particles (9).

Effective Dose

The effective dose is another radiation-protection quantity derived from the absorbed dose that accounts for both the radiation weighting factor w_R and the relative stochastic risk for given organ equivalent doses, accounting for the differences among tissues of the age- and sex-averaged radiogenic risks of cancer induction (applicable to somatic tissues) and germ-cell mutation (applicable to gonadal tissues). Effective dose thus is not a dose metric for a specific individual but rather a population-averaged quantity that may be compared with a radiation protection standard or among different radiation exposures. Numerically, effective dose is the sum of all the weighted equivalent doses for all organs and tissues of the body of a reference anatomic model, whether the body is irradiated uniformly or nonuniformly:

$$E(\tau) \equiv \sum_T w_T \left[\frac{H^M(T, \tau) + H^F(T, \tau)}{2} \right], \quad \text{Eq. 5}$$

where $H^M(T, \tau)$ and $H^F(T, \tau)$ are the dose equivalents to target tissue T in the reference male and reference female phantom, respectively, over the dose-integration time interval τ , and w_T is a tissue weighting factor defined as the fraction of total stochastic risk associated with individual target tissue T . Values for the tissue weighting factor w_T , a dimensionless quantity, are recommended in ICRP publication 103 (9).

It must be emphasized that assessment of an individual patient's excess cancer risk associated with a nuclear medicine or other medical imaging procedure based on the effective dose for that procedure in combination with any available cancer risk factors is not appropriate, as the tissue weighting factors w_T , as well as the risk factors, are average values not applicable to any individual.

THE COMPUTATIONAL ENGINE—THE DOSE-CALCULATION FORMULAS—OF THE MIRD SCHEMA

The fundamental equations of the MIRD schema, yielding the mean absorbed dose $D(r_T, \tau)$ to a target region r_T over an integration period τ (i.e., from time $t = 0$ to time $t = \tau$) after administration of a radiopharmaceutical for a particular radionuclide in a particular anatomic model (I), are as follows (Table 1 defines the parameters in Eqs. 6–11):

$$D(r_T, \tau) = \sum_{r_S} \int_0^\tau A(r_S, t) \frac{1}{M(r_T)} \sum_i E_i Y_i \varphi(r_T \leftarrow r_S, E_i) dt, \quad \text{Eq. 6}$$

$$= \sum_{r_S} \int_0^\tau A(r_S, t) \sum_i \Delta_i \Phi(r_T \leftarrow r_S, E_i) dt, \quad \text{Eq. 7}$$

$$= \sum_{r_S} \int_0^\tau A(r_S, t) S(r_T \leftarrow r_S) dt, \quad \text{Eq. 8}$$

where

$$S(r_T \leftarrow r_S) = \frac{1}{M(r_T)} \sum_i E_i Y_i \varphi(r_T \leftarrow r_S, E_i), \quad \text{Eq. 9}$$

$$= \frac{1}{M(r_T)} \sum_i \Delta_i \varphi(r_T \leftarrow r_S, E_i), \quad \text{Eq. 10}$$

$$= \sum_i \Delta_i \Phi(r_T \leftarrow r_S, E_i). \quad \text{Eq. 11}$$

As noted in Table 1, the integration time τ is commonly set to infinity (i.e., $\tau = \infty$), yielding the total absorbed dose to target region r_T for complete decay of the administered radiopharmaceutical. As applied to diagnostic radiopharmaceuticals, its traditional application, the MIRD schema implicitly assumes that activity and time-integrated activities are uniformly distributed within source organs and that radiation energy is uniformly deposited within target organs.

Although identical in terms of the calculation performed, Equations 6–8 are made progressively simpler in appearance by combining multiple parameters into various single parameters, particularly with the introduction of the S value (Eqs. 9–11). The ingenuity of the MIRD schema lies in reducing a challenging, seemingly intractable computational task—calculation of the absorbed dose to several irregular target regions (i.e., organs) from the nonuniform and time-varying distribution of a radionuclide among several irregular source regions (i.e., also organs)—to a series of more manageable components. More specifically, the MIRD schema separates the relevant biology from the relevant physics, as most clearly shown in Equation 8: all the biology is combined in the time-integrated activity, $\tilde{A}(r_S, \tau)$, and all the physics in the S value, $S(r_T \leftarrow r_S)$. The S values are, of course, radionuclide- and anatomic model-specific (10), as the energies and frequencies per decay (E_i and Y_i , respectively) of emitted radiations (11–13) depend on the radionuclide, and the absorbed fractions ($\varphi(r_T \leftarrow r_S, E_i)$) depend on the anatomic model (i.e., the sizes, shapes, and separations of the organs) as well as the radionuclide and its emitted radiations (14). Conceptually, the S value $S(r_T \leftarrow r_S)$ is equivalent to the absorbed dose to target region r_T per decay of the radionuclide in source region r_S .

Although perhaps daunting in appearance, Equations 6–8 are conceptually quite simple, as shown when expressed in literal terms:

$$\begin{array}{c}
 \text{Absorbed dose} \\
 \text{to an organ,} \\
 \text{target organ } r_T, \\
 \text{from the administered} \\
 \text{radionuclide in itself or} \\
 \text{in another organ,} \\
 \text{a source organ } r_S
 \end{array}
 = \frac{
 \begin{array}{c}
 \tilde{A}(r_S, \tau) \\
 \downarrow \\
 \text{Number of radioactive} \\
 \text{decays of the administered} \\
 \text{radionuclide in source} \\
 \text{organ } r_S
 \end{array}
 \cdot
 \begin{array}{c}
 \sum_i \Delta_i = \\
 \sum_i E_i Y_i \\
 \downarrow \\
 \text{Radiation energy} \\
 \text{emitted per decay of} \\
 \text{the administered} \\
 \text{radionuclide}
 \end{array}
 \cdot
 \begin{array}{c}
 f(r_T \leftarrow r_S, E_i) \\
 \downarrow \\
 \text{Fraction of radiation} \\
 \text{energy emitted in} \\
 \text{source organ } r_S \text{ that} \\
 \text{is deposited in} \\
 \text{target organ } r_T
 \end{array}
 }{
 \begin{array}{c}
 \text{Mass of the target} \\
 \text{organ } r_T \text{ in which the} \\
 \text{radiation energy} \\
 \text{is deposited} \\
 \uparrow \\
 M(r_T)
 \end{array}
 }$$

Summing the dose contributions above from each source organ r_s to the target organ r_T over all the source organs of the body then yields the total absorbed dose to target organ r_T .

For any given dose calculation, time-activity data and time-integrated activities are typically available for only a limited number of source regions (15). It is important that these include the total body, so that the rest-of-body (also known as the remainder of body) time-integrated activity $\tilde{A}(\text{rest of body}, \infty)$ —and the potentially large rest-of-body dose contribution—may be calculated:

$$\tilde{A}(\text{rest of body}, \infty) = \tilde{A}(\text{total body}, \infty) - \sum_{r_S} \tilde{A}(r_S, \infty), \quad \text{Eq. 12}$$

where the total-body time-integrated activity, $\tilde{A}(\text{total body}, \infty)$, is not included in the summation term, $\sum_{r_S} \tilde{A}(r_S, \infty)$. The rest-of-body S values may be calculated using the formula provided by Coffey and Watson (16):

$$\begin{aligned}
 S(r_T \leftarrow \text{rest of body}) &= \frac{M(\text{total body})}{M(\text{rest of body})} \\
 S(r_T \leftarrow \text{total body}) - \sum_{r_S} \frac{M(r_S)}{M(\text{rest of body})} S(r_T \leftarrow r_S), & \quad \text{Eq. 13}
 \end{aligned}$$

where $S(r_T \leftarrow \text{rest of body})$ and $S(r_T \leftarrow \text{total body})$ are the rest-of-body-to- r_T and total-body-to- r_T S values, respectively; $M(\text{total body})$ and $M(\text{rest of body})$ are the total-body and rest-of-body masses, respectively; and

$$M(\text{rest of body}) = M(\text{total body}) - \sum_{r_S} M(r_S). \quad \text{Eq. 14}$$

S VALUES AND ANATOMIC MODELS

As noted in the previous section, S values and related quantities (Eqs. 9–11) depend on the particular radionuclide and anatomic model. The relevant radionuclide decay data are available in any number of authoritative sources, such as the MIRDO radionuclide data and decay schemes (11,12) and ICRP publication 107 (13); these publications are noteworthy in that they provide tabulations of decay data in a form specifically designed for internal dosimetry. Derivation of S values, on the other hand, is a challenging computational task. Ideally, S values would be computed by Monte Carlo

radiation-transport simulations or other computational means using segmented organs in whole-body CT or MR images of each individual patient. Only then would one have truly patient-specific S values. Although this is generally impractical and not done for diagnostic radiopharmaceuticals, this approach is actually being pursued for radiopharmaceutical therapies using certain commercially available software packages (3,17). Even then, however, only a limited number of organs are imaged in a typical partial-body CT or MRI study and incorporated into a patient-specific dose calculation. More commonly, organ-level dosimetry is performed using precomputed tables of radionuclide S values for source-organ/target-organ pairs selected from a series of anatomic computerized models (or phantoms) of age-dependent average individuals. For illustrative purposes, a partial tabulation of S values for ^{131}I in the ICRP voxel-based phantom of the reference adult male is presented in Table 2 (18,19). Such models, or phantoms, may be classified by their respective format type (i.e., mathematic representation) and morphometric fidelity (i.e., patient anatomic specificity), as outlined in Table 3.

Average individuals are typically taken to be reference persons defined, for example, by the ICRP (20,21); these include males and females at roughly the 50th percentile for heights and weights at specified ages: the newborn, 1-y-old, 5-y-old, 10-y-old, 15-y-old, and adult. Historically, the MIRDO schema has used S values computed using the age-specific stylized computational phantoms of reference individuals developed at the Oak Ridge National Laboratory (ORNL) (Fig. 1A) (22). These computational phantoms include models of organ and body contours based on regular 3-dimensional mathematic topologies (spheres, ellipsoids, cones, and tapered cylinders, for example). The ICRP subsequently developed a family of reference voxel-based phantoms (Fig. 1B) (18,23). These phantoms are ultimately derived from CT image sets of cadavers of actual individuals, and thus, the anatomy is more realistic than that of the ORNL stylized phantoms. Besides having greater anatomic fidelity, the voxel-based ICRP phantoms include a larger number of source and target regions (55 and 42, respectively) than the ORNL phantoms (20 and 20, respectively). A comparison of ^{131}I S values—for selected source and target regions—for the ORNL stylized phantoms and the ICRP voxel-based phantoms of the reference adult male and female is presented graphically in Figure 2 (24). Although yielding reasonably accurate results overall,

TABLE 2
S Values for ¹³¹I-Derived from ICRP Voxel-Based Computational Phantom of Reference Adult Male (18)

Source region	Target region													
	Adrenals	Brain	Gallbladder wall	Heart wall	Kidneys	Liver	Lung	Muscle	Pancreas	Spleen	Stomach wall	Thymus	Thyroid	Urinary bladder wall
Adrenals	1.81E-03	8.14E-09	2.05E-06	6.72E-07	6.95E-06	2.02E-06	5.76E-07	2.05E-07	2.18E-06	3.63E-06	1.66E-06	1.62E-07	9.69E-08	5.21E-08
Brain	8.52E-09	2.70E-05	8.18E-09	2.84E-08	5.55E-09	1.26E-08	4.27E-08	6.17E-08	6.16E-09	1.26E-08	9.22E-09	8.29E-08	1.60E-07	2.97E-10
Gallbladder content	1.88E-06	7.38E-09	6.79E-06	5.59E-07	1.56E-06	4.56E-06	4.14E-07	1.47E-07	6.14E-06	4.00E-07	1.39E-06	1.48E-07	7.78E-08	7.19E-08
Heart content	6.76E-07	2.46E-08	5.68E-07	9.22E-06	3.50E-07	9.19E-07	1.85E-07	1.57E-07	6.12E-07	9.65E-07	1.60E-06	1.69E-06	5.35E-07	1.48E-08
Kidneys	6.97E-06	5.02E-09	1.60E-06	3.49E-07	8.60E-05	1.21E-06	2.92E-07	2.24E-07	2.11E-06	1.74E-06	1.01E-06	8.97E-08	5.42E-08	9.79E-08
Liver	2.02E-06	1.19E-08	5.32E-06	9.77E-07	1.21E-06	1.82E-05	8.13E-07	1.65E-07	1.95E-06	4.18E-07	1.38E-06	2.49E-07	1.35E-07	4.52E-08
Lungs	5.52E-07	4.33E-08	4.23E-07	2.11E-06	2.84E-07	1.18E-07	3.04E-05	2.23E-07	3.56E-07	9.65E-07	6.67E-07	2.07E-06	1.05E-06	1.15E-08
Muscle	2.06E-07	8.49E-08	1.51E-07	1.71E-07	2.27E-07	1.69E-07	2.30E-07	1.41E-06	1.61E-07	2.06E-07	1.61E-07	2.06E-07	2.65E-07	3.12E-07
Pancreas	2.17E-06	5.40E-09	6.95E-06	6.15E-07	2.11E-06	1.96E-06	3.55E-07	1.56E-07	2.07E-04	9.80E-07	3.77E-06	1.20E-07	6.47E-08	9.54E-08
Small intestine content	1.21E-06	3.04E-09	1.35E-06	2.99E-07	1.73E-06	6.75E-07	1.87E-07	1.99E-07	3.38E-06	8.30E-07	1.64E-06	6.01E-08	3.46E-08	4.92E-07
Spleen	3.66E-06	1.17E-08	4.25E-07	9.56E-07	1.74E-06	4.20E-07	1.03E-06	2.00E-07	9.77E-07	1.56E-04	2.54E-06	1.97E-07	1.40E-07	3.26E-08
Stomach content	1.57E-06	8.29E-09	1.19E-06	1.79E-06	1.02E-06	1.18E-06	6.58E-07	1.51E-07	3.00E-06	2.35E-06	5.89E-05	1.99E-07	1.05E-07	5.05E-08
Thymus	1.64E-07	8.24E-08	1.55E-07	2.33E-06	9.23E-08	2.59E-07	1.83E-06	1.93E-07	1.29E-07	2.04E-07	2.13E-07	1.14E-03	8.41E-06	4.48E-09
Thyroid	9.72E-08	1.62E-07	8.07E-08	6.49E-07	5.60E-08	1.38E-07	9.30E-07	2.40E-07	6.75E-08	1.39E-07	1.11E-07	8.25E-06	1.44E-03	2.49E-09
Urinary bladder content	5.64E-08	2.83E-10	7.43E-08	1.58E-08	1.05E-07	4.69E-08	1.21E-08	3.14E-07	1.05E-07	3.36E-08	5.53E-08	4.93E-09	2.74E-09	2.19E-05

Data are mGy/MBq⁻¹ s⁻¹. ICRP voxel-based phantoms actually include many more source and target regions than those tabulated here, but for readability only partial list of S values is presented. (Adapted from reference (19).)

the modeling of anatomy with simplistic shapes in the ORNL stylized phantoms introduces some inaccuracies in S values as compared with the ICRP voxel-based phantoms. For several source–target pairs, in fact, the differences in S values are up to 10-fold. Modern implementations of the MIRDSchema (including the MIRDCalc software (19,25)) generally use the S values for the family of ICRP voxel-based phantoms (18).

Among the different format types, polygon mesh-based phantoms perhaps provide the greatest flexibility and anatomic fidelity. The ICRP recently developed mesh-type reference phantoms that address certain shortcomings of the voxel-based reference phantoms (26). Such phantoms can be used, for example, to create posture (e.g., standing, walking, kneeling)-specific anatomic models to elucidate the impact of posture on internal dose estimates (27).

Differences in body size and contour and internal organ anatomy between a reference phantom (regardless of how realistic such a phantom may appear) and an individual patient can potentially introduce sizable errors in estimates of mean organ doses to the patient, mainly because of errors in estimation of the self-dose. The self-dose typically accounts for most of the dose to an organ. To a first approximation, self-dose S values scale inversely with organ mass, consistent with the short range (typically about 1 mm or less in tissue) and the assumption of complete local (i.e., intraorgan) absorption of particulate radiations such as α-particles, β-particles (including positrons), and Auger and conversion electrons. Therefore, an individual's self-dose S value for a particular organ may be obtained from the respective reference-phantom S value by scaling by the ratio of the phantom-to-individual organ mass (assuming the individual's organ masses have been measured by, for example, CT or MRI). Recently, more accurate scaling approaches for the photon- as well as the particulate-radiation dose contributions have been developed (28).

TIME-ACTIVITY DATA AND TIME-INTEGRATED ACTIVITIES

Each radiopharmaceutical is, of course, characterized by its own time-dependent biodistribution (expressed, for example, as the activity or activity concentration in the different tissues of the body at different times after administration), and this varies not only across different species but also among different subjects of the same species. The biology-related aspects of internal dosimetry—measurement of time–activity data and their reduction to time-integrated activities—are thus particularly challenging.

Initial human absorbed dose estimates for new radiopharmaceuticals—to satisfy regulatory requirements as well as for ethical considerations—are derived from animal biodistribution studies, typically in mice or rats. Such small-animal models are relatively inexpensive, easy to maintain and handle, and available in sufficiently large numbers to achieve reasonable statistical reliability of measured time–activity data (the methodology for measuring biodistribution

TABLE 3
Classification of Computational Anatomic Phantoms by Format and Morphometric Category (3)

Parameter	Class 1	Class 2	Class 3	Class 4
Format*	Stylized: organs and total body represented by regular 3-dimensional shapes such as spheres and ellipsoids.	Voxel: CT- or MRI-based 3-dimensional digital representation of segmented organ as assemblies of cubic volume elements (or voxels)	NURBS (nonuniform rational B-spline): anatomic volumes defined by NURBS equations characterized by set of control points; shape and volume of NURBS surface vary with coordinates of control points	Polygon mesh-based: Polygonal mesh is composed of set of vertices, edges, and faces that specify shape of polyhedral object in 3 dimensions; surfaces of phantom are defined by large number of polygonal meshes such as tetrahedrons
Category†	Reference: average human anatomy corresponding, for example, to men and women at roughly 50th percentile for heights and weights at discrete ages (newborn, 1-y-old, 5-y-old, 10-y-old, 15-y-old, and adult); there is no match to anatomy of any specific patient; S values can be precomputed for all source/target organ combinations in reference phantom series	Patient-dependent: use of computational phantom library covering various combinations of height and weight percentiles in modeled patient population; internal organ anatomy is volumetrically rescaled to outer body size (contour) but is not patient-specific; S values can be precomputed for all source/target organ combinations for each member of patient-dependent phantom library	Patient-sculpted: use of computational methods to change body contour of selected patient-dependent phantom from phantom library; resulting phantom would then appear to have same body size and shape as that of patient, but again, internal organ anatomy is not patient-specific; as patient-sculpted phantom does not exist until it is created (using patient's CT/MR-based body contour), S values cannot be precomputed but must be generated by Monte Carlo computer simulation once that phantom is available	Patient-specific: matching of phantom internal and external anatomy to CT- or MRI-measured anatomy of specific patient; as with patient-sculpted phantoms, S values cannot be precomputed until patient-specific phantom is developed

*From left to right, classes require increasing computer memory.

†From left to right, classes involve improving anatomic fidelity.

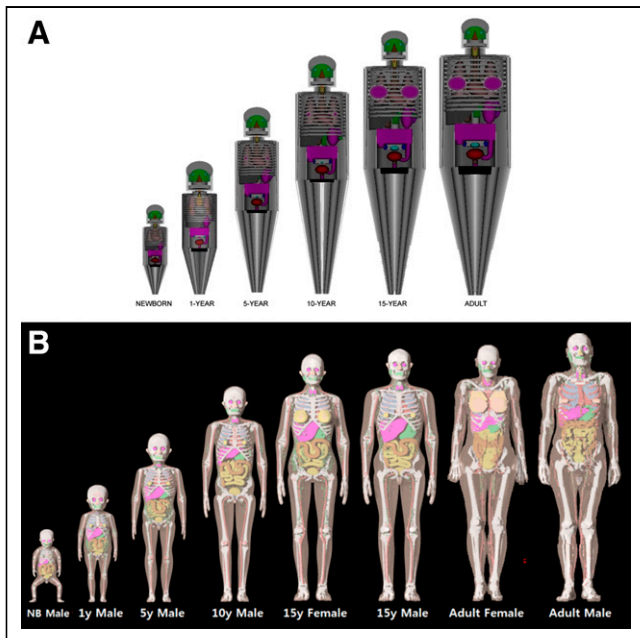


FIGURE 1. (A) Family of stylized reference phantoms from ORNL (22). Each phantom has both male and female sex organs. (B) Family of voxel-based reference phantoms from ICRP (18,23). Total of 12 phantoms are in ICRP reference series. In this figure, only male versions are shown for ages below 15 y. (Reprinted from (3).)

data for radiopharmaceuticals in small-animal models is beyond scope of this article). The translation of such data to human anatomic models is problematic, however, and no entirely satisfactory method for doing so has been developed thus far (29,30). The most widely used approach is based on the assumption that a radiopharmaceutical's body mass-normalized activity concentration (equivalent to the SUV, routinely used to parameterize clinical PET scans) in a particular organ at a particular time after administration is the same among subjects of different body masses, whether subjects of the same species (e.g., patients) or subjects across species (e.g., mice vs. humans) This, in turn, is based on the implicit assumption that the biology of the radiopharmaceutical is identical (except, of course, for statistical variations) across subjects and species and that differences in organ activity concentrations are therefore related exclusively to differences in the radiopharmaceutical's volume of distribution and therefore, in first order, the total-body mass. These considerations yield the following equation relating the organ activity concentration of a radiopharmaceutical in the human (*h*) anatomic model to that measured in the animal (*a*) model (in Eq. 15 and subsequent equations, square brackets are, as usual, used to indicate concentrations):

$$[A(r_S, t)]_h = [A(r_S, t)]_a \frac{BM_a}{BM_h}, \quad \text{Eq. 15}$$

where $[A(r_S, t)]_h$ and $[A(r_S, t)]_a$ are the activity concentrations (in percentage of the injected activity per gram, for example) in source region (i.e., organ) r_S at time t after administration in the animal model and the human anatomic model, respectively,

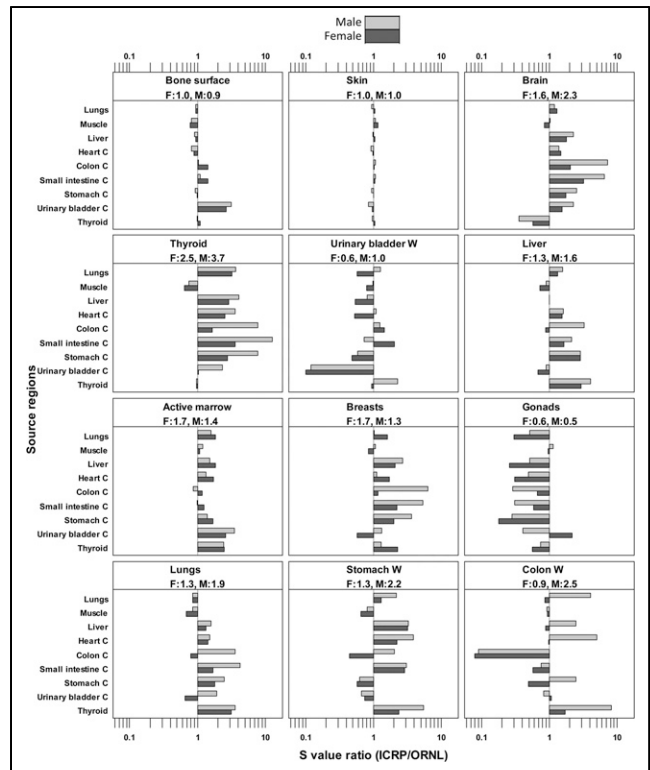


FIGURE 2. Ratio (median) of ^{131}I S values for ICRP voxel-based phantoms to values for ORNL stylized phantoms for selected source-region/target-region pairs. (Reprinted from (24).)

and BM_a and BM_h are the total-body masses of the animal model and the human anatomic model, respectively (3).

After administration of a radiopharmaceutical, the radioactivity is distributed among the tissues of the body and eliminated from the body—simultaneously by a combination of physical decay and biologic processes. Physical decay corresponds to the radioactive decay of the radionuclide in the radiopharmaceutical and, of course, occurs at the same rate regardless of its chemical form and whether the radionuclide is in vivo or ex vivo. Biologic processes reflect the behavior of the radiopharmaceutical in vivo independent of radioactive decay. The combination of the simultaneous processes of physical (i.e., radioactive) decay and biologic clearance yields the so-called effective clearance, which is faster than either physical clearance or biologic clearance alone, as illustrated in Figure 3A. Biologic clearance data are often referred to as decay-corrected, since the time-activity data have been adjusted, or corrected, to eliminate the effect of radioactive decay on the clearance of the radionuclide in the radiopharmaceutical. Effective clearance data are sometimes referred to as non-decay-corrected, since these data include the effect of radioactive decay and thus correspond to the overall clearance of the radionuclide in the radiopharmaceutical. It is effective time-activity data that are ultimately required for radiation dose calculations.

Once organ and total-body time-activity data have been measured, either preclinically in an animal model or

clinically for an individual patient, these data must be reduced to time-integrated activities. This is generally accomplished by fitting a mathematic function to these data—most often, an exponential function—once the measured data have been corrected for radioactive decay to the time of administration of the radiopharmaceutical:

$$[A(r_S, t)]_{\text{corrected}} = \sum_{j=1}^n [A(r_S, 0)]_j e^{-(\lambda_j)_b t}, \quad \text{Eq. 16}$$

where $[A(r_S, t)]_{\text{corrected}}$ is the activity concentration in source region (i.e., organ) r_S at time t after administration corrected for radioactive decay to the time of administration of the radiopharmaceutical; $[A(r_S, 0)]_j$ and $(\lambda_j)_b$ are the activity concentration in source region (i.e., organ) r_S at time $t = 0$ (i.e., the time of administration of the radiopharmaceutical) and the biologic (b) clearance constant, respectively, of the j th exponential term; and n is the number of exponential terms (or components) in the fitted function. The corresponding version of this function for the data not corrected for radioactive decay is as follows:

$$[A(r_S, t)]_{\text{not corrected}} = \sum_{j=1}^n [A(r_S, 0)]_j e^{-(\lambda_j)_e t}, \quad \text{Eq. 17}$$

where $A(r_S, t)_{\text{not corrected}}$ is the activity concentration in source region (i.e., organ) r_S at time t after administration not corrected for radioactive decay to the time of administration of the radiopharmaceutical, and $(\lambda_j)_e$ is the effective (e) clearance constant, reflecting the processes of both physical decay and biologic clearance such that

$$(\lambda_j)_e = (\lambda_j)_b + \lambda_p, \quad \text{Eq. 18}$$

where λ_p is the physical (p) decay constant of the radionuclide. Integration of Equation 17 to infinite time (i.e., for complete decay of the administered radionuclide) then yields the total time-integrated activity, $\tilde{A}(r_S, \infty)$, in source region r_S with mass $M(r_S)$:

$$\tilde{A}(r_S, \infty) = M(r_S) \sum_{j=1}^n \frac{[A(r_S, 0)]_j}{(\lambda_j)_e}. \quad \text{Eq. 19}$$

The physical decay constant λ_p is related to the physical half-life $(T_{1/2})_p$ of the radionuclide by the following equation:

$$\lambda_p = \frac{\ln(2)}{(T_{1/2})_p} = \frac{0.693}{(T_{1/2})_p}. \quad \text{Eq. 20}$$

For the j th exponential component, the biologic clearance constant $(\lambda_j)_b$ and effective clearance constant $(\lambda_j)_e$ are likewise related to the biologic half-life $\{(T_{1/2})_b\}_j$ and the effective half-life $\{(T_{1/2})_e\}_j$, respectively:

$$(\lambda_j)_b = \frac{\ln(2)}{\{(T_{1/2})_b\}_j} = \frac{0.693}{\{(T_{1/2})_b\}_j}, \quad \text{Eq. 21}$$

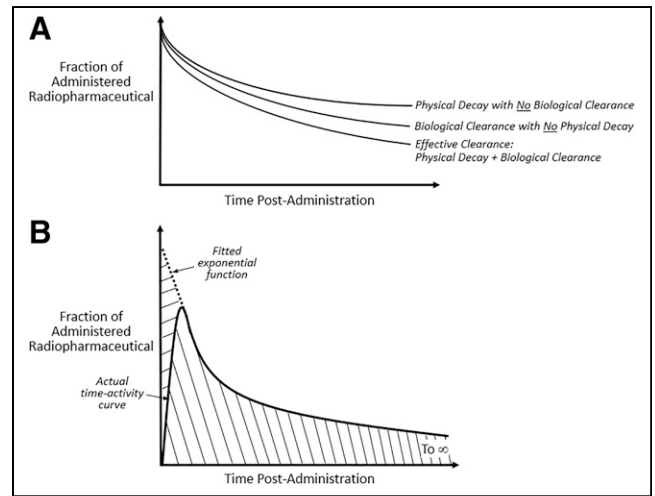


FIGURE 3. (A) Idealized exponentially decreasing time-activity curves illustrating relationship among physical, biologic, and effective half-times in tissue or organ for administered radiopharmaceutical. Each curve is characterized by clearance constant and corresponding half-time, with effective clearance constant being greater than either physical or biologic clearance constant (Eq. 18) and effective half-time being shorter than either physical or biologic half-time (Eq. 23). (Reprinted from (3).) (B) Actual time-activity curve will be more complex, including rising component at early times after administration as radiopharmaceutical exits blood and enters tissue parenchyma of organ. In many cases, this rising portion of time-activity curve is brief compared with remainder of curve and is therefore ignored in fitting mathematic function to curve for purpose of estimating time-integrated activity. More specifically, although fitting of monotonically decreasing exponential function to organ's time-activity data will overestimate total time-integrated activity to infinite time (i.e., corresponding to complete decay of administered radionuclide), magnitude of this overestimate will be small for small-molecule radiopharmaceuticals. However, for slowly localizing, large-molecule radiopharmaceuticals such as radiolabeled antibodies, this overestimate of time-integrated activity may be significant. Graphically, total time-integrated activity (i.e., area under curve) for actual time-activity curve (represented by upper left-to-lower right slant-line hatching) will be overestimated by amount corresponding to lower left-to-upper right slant-line hatching.

$$(\lambda_j)_e = \frac{\ln(2)}{\{(T_{1/2})_e\}_j} = \frac{0.693}{\{(T_{1/2})_e\}_j}, \quad \text{Eq. 22}$$

and the effective half-time $\{(T_{1/2})_e\}_j$ of the j th exponential component is related to the physical half-time $(T_{1/2})_p$ and the biologic half-time $\{(T_{1/2})_b\}_j$ of the j th exponential component as follows:

$$\{(T_{1/2})_e\}_j = \frac{(T_{1/2})_p \{(T_{1/2})_b\}_j}{(T_{1/2})_p + \{(T_{1/2})_b\}_j}. \quad \text{Eq. 23}$$

Rearranging Equation 22 to solve for $(\lambda_j)_e$ in terms of $\{(T_{1/2})_e\}_j$ and substituting that expression into Equation 19

yields a formula for the total time-integrated activity, $\tilde{A}(r_S, \infty)$, in source region r_S in terms of $\left\{ \left(T_{1/2} \right)_e \right\}_j$:

$$\tilde{A}(r_S, \infty) = 1.44 M(r_S) \sum_{j=1}^n [A(r_S, 0)]_j \left\{ \left(T_{1/2} \right)_e \right\}_j. \quad \text{Eq. 24}$$

Actual organ time-activity curves include a rising component at early times after administration as the radiopharmaceutical exits the blood and enters the parenchyma of the organ (as illustrated in Fig. 3B). A mathematic function more complex than a sum of exponential terms (Eqs. 16 and 17) is required for reliable fitting to such data, and the reader is referred to section 5.7.1. (analytic curve fitting) in the 2022 MIRD primer (3) for a review of curve fitting for radiopharmaceutical time-activity data. An alternative approach to curve fitting and analytic integration of complex time-activity data is a numeric approach: connecting consecutive data points by straight-line segments and integrating piecewise by the trapezoidal rule. However, integration by the trapezoidal rule extends only to the final data point. Beyond that point to infinity, the integral is indeterminate by the trapezoidal rule and must be estimated by other means (e.g., assuming, conservatively, elimination of activity beyond the final data point by physical decay only).

Curve-fitting computer programs are widely available. These include commercially available software packages such as Excel (part of the Office suite (Microsoft)), OLINDA/EXM (31) (part of the Hermes Hybrid Viewer Dosimetry Module (Hermes)), PMOD (PMOD Technologies Ltd.), and Prism (GraphPad Software), among many others. Recently, the MIRD committee has developed and will release MIRDfit, an Excel-based fitting and integration program that will be part of and freely downloadable from the MIRDsoft website.

BEYOND ORGAN DOSIMETRY: SUBORGAN, VOXEL-LEVEL, AND CELL-LEVEL DOSIMETRY

As noted, as it is applied to diagnostic radiopharmaceuticals, its traditional application, the MIRD schema implicitly assumes that activity and time-integrated activities are uniformly distributed within source organs and that radiation energy is uniformly deposited within target organs. However, a remarkable strength of the MIRD schema is its generality: by judicious selection of source and target regions, it can be used to calculate the dose to virtually any target region from virtually any source region, from microscopic to macroscopic to whole organs and whole body (32). For several organs, including brain and kidney, suborgan MIRD-schema dosimetry has been developed (33,34). MIRD pamphlet no. 17 (35) introduced voxel S values and extended the MIRD schema to arbitrary macroscopic activity distributions in 3 dimensions for calculation of the resulting macroscopic dose distribution; in this context, the term *macroscopic* refers to volume elements, or voxels, 3 mm or greater in dimension. And, with the publication of

MIRD cellular S factors (36) and the release of the MIRD-cell computer program (37–39), the MIRD schema has been extended to cellular and subcellular source and target regions. This applet, freely downloadable from the MIRDsoft website, models the cell-level distribution of radiopharmaceuticals, calculates the distribution of the radiation dose at the cellular and subcellular levels for both isolated cells and clusters of cells using cellular S values, and mathematically models the biologic responses of the radiolabeled and unlabeled cell populations.

ADAPTATION OF THE MIRD SCHEMA TO PATIENT-SPECIFIC AND TUMOR DOSIMETRY

With the ongoing emergence of new and effective radiopharmaceutical therapies such as ^{177}Lu -DOTATATE (Lutathera; Novartis) and ^{177}Lu -vipivotide tetraxetan (Pluvicto; Novartis), there is widespread, and growing, interest in the development of individualized (i.e., patient-specific) radiation dosimetry, including both normal-organ and tumor dosimetry. The tissue doses associated with radiopharmaceutical therapy are much higher than those for diagnostic applications of radiopharmaceuticals, and severe, potentially life-threatening deterministic effects may therefore result. Consequently, optimally balancing tumor (or other target-tissue) and normal-organ radiation doses requires patient-specific dosimetry. The practical implementation of patient-specific dosimetry, as outlined diagrammatically in Figure 4 (40), remains challenging, however. The patient-independent fixed-administered-activity approach (e.g., MBq, MBq/kg of body mass, and MBq/m² of body surface area), which does not require any kinetic or other patient measurements, remains widely used, therefore. The fixed administered activity for a particular therapeutic radiopharmaceutical is typically established by chemotherapy-like dose-escalation phase 1 or 2 clinical trials. However, several commercial software packages are now available for patient-specific normal-organ and tumor dosimetry in the setting of radiopharmaceutical therapy (17).

Given its generality, the MIRD schema is adaptable to specific patients, including patients with tumors. As previously noted, patient-specific S values can be computed by Monte Carlo radiation-transport simulations or other computational means using segmented organs in whole-body CT or MR images of each individual patient to yield patient-specific S values. This is being pursued in the setting of radiopharmaceutical therapy. Alternatively, various mathematic formulas may be applied to estimate, approximately, patient-specific organ S values from those of reference anatomic models. A first-order approach for such estimation has been implemented in MIRDcalc (19,25). The OLINDA/EXM computer program also includes an adjustment for organ size for organ-level dosimetry (31). S values for organ sizes that deviate from those of the relevant reference anatomic model (i.e., the closest in total-body mass to that of the reference model) may be adjusted by scaling each organ's self-dose by

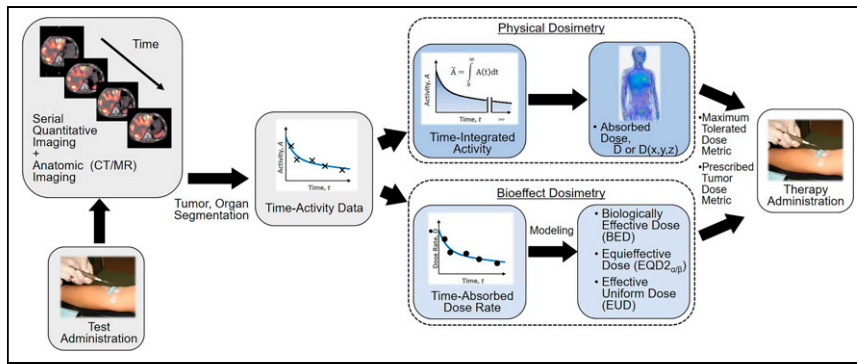


FIGURE 4. General workflow for radiopharmaceutical therapy dosimetry. Test administration may be either pretherapy tracer administration or first cycle of multidose therapy regimen. Calculation of absorbed doses (in terms of either tumor and organ mean doses or dose distributions) or of dose rates D is performed using dose factors (e.g., S values), dose point kernels, or Monte Carlo analysis. Physical dosimetry yields absorbed dose, and bioeffect dosimetry yields dose metrics such as biologically effective dose, equieffective dose, or effective uniform dose (for bioeffect dosimetry). (Reprinted from (40).)

the ratio of the patient-to-reference phantom organ mass raised to a constant power. The value of the power was set to $-2/3$ for photon self-dose scaling and -1 for particulate-radiation self-dose scaling. Estimations of cross-dose (i.e., nonself-dose) contributions are unchanged when using adjusted organ masses. MIRDcalc also supports approximate tumor dosimetry by calculating the self-dose to a sphere from time-integrated activity uniformly distributed within the sphere. MIRDcalc allows user selection not only of the spheric and nonspheric volumes but also of composition (i.e., the relative amounts of bone and soft-tissue composing the tumor). Calculation of tumor self-doses is based on S values provided by Olguin et al. (41), with interpolation (i.e., log-log interpolation) or extrapolation of published data points for user-specified volumes.

CONCLUDING REMARKS

The MIRD schema, which originated in the 1960s, has withstood the test of time, and established itself as a worldwide standard for radiopharmaceutical dosimetry. The schema has evolved, and it continues to do so as technology and the field of nuclear medicine have advanced. Stylized reference anatomic models (22) have been replaced by more anatomically realistic voxel-based models (18,23), with a greater number of source and target regions (Fig. 1). Even more anatomically realistic models are becoming available as well (Table 2). Three-dimensional suborgan dosimetry is doable using voxel S values (35), as is cellular and subcellular dosimetry using cell S values and the MIRDcell software (38,39). With ongoing advances in radiopharmaceutical therapy, adaptation of the MIRD schema to patient-specific dosimetry, including tumor dosimetry (41), is well under way, with scaling of reference anatomic-model organ S values to individualized S values and first-order calculation of tumor self-doses (19,25). Further, the MIRD committee is now addressing the radiation biology

of radiopharmaceutical therapy (3) (including α -particle therapy (42)) and the methodologies for acquisition of the relevant time-activity data for patient-specific dosimetry (43-45).

Importantly, in addition to its traditional hard-copy resources—the peer-reviewed MIRD pamphlets and associated publications—the MIRD committee has created and is expanding the MIRDsoft website for providing a suite of freely downloadable software tools for radiopharmaceutical dosimetry. MIRDsoft currently hosts MIRDcalc (for organ-level and tumor dosimetry, including such novel features as error propagation) (19,25) and MIRDcell (for not only cellular and subcellular dosimetry but also bioeffect modeling) (38,39). Near-term additions to MIRDsoft will include MIRDfit (for curve fitting, with error propagation, of radiopharmaceutical time-activity data), MIRDy90 (for ^{90}Y -microsphere dosimetry for radioembolic therapy of liver tumors), MIRDrelease (for assessment of the releasability of radiopharmaceutical therapy patients and of the duration of postrelease precautions), and MIRDct (for individualized CT dosimetry).

Like fine wine, the MIRD schema has aged well indeed!

DISCLOSURE

Pat Zanzonico receives, on behalf of the MIRD committee, royalties on sales of the 2022 MIRD primer. No other potential conflict of interest relevant to this article was reported.

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