MIRD Pamphlet No. 22 (Abridged): Radiobiology and Dosimetry of α-Particle Emitters for Targeted Radionuclide Therapy*

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The potential of α -particle emitters to treat cancer has been recognized since the early 1900s. Advances in the targeted delivery of radionuclides and radionuclide conjugation chemistry, and the increased availability of α -emitters appropriate for clinical use, have recently led to patient trials of radiopharmaceuticals labeled with α -particle emitters. Although α -emitters have been studied for many decades, their current use in humans for targeted therapy is an important milestone. The objective of this work is to review those aspects of the field that are pertinent to targeted α -particle emitter therapy and to provide guidance and recommendations for human α -particle emitter dosimetry.

Key Words: α -particle emitters; human α -particle emitter dosimetry; targeted α -particle emitter therapy

J Nucl Med 2010; 51:311–328 DOI: 10.2967/jnumed.108.058651

Several reviews have been published on the topic of α -particle–emitting radionuclides, which have been the subject of considerable investigation as cancer therapeutics (1–8). In the context of targeted therapy, α -particle emitters have the advantages of high potency and specificity. These advantages arise from the densely ionizing track and short path length of the emitted positively charged helium nucleus in tissue. The practical implication of these features, as well as the distinction between α -particles and the more widely

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used β -particle emitters for targeted radionuclide therapy, is that it is possible to sterilize individual tumor cells solely from self-irradiation with α -particle emitters. This is generally not possible with β -particle emitters given achievable antibody specific activity, tumor-cell antigen expression levels, and the need to avoid prohibitive toxicity (5). These attributes combine to provide the fundamental strength and rationale for using α -particle–emitting radionuclides for cancer therapy. Current approaches to cancer treatment are largely ineffective once the tumor has metastasized and tumor cells are disseminated throughout the body. There is also increasing evidence that not all tumor cells are relevant targets for effective tumor eradication and that sterilization of a putative subpopulation of a small number of tumor stem cells may be critical to treatment efficacy (9). The eradication of such disseminated tumor cells, or of a subpopulation of tumor stem cells, requires a systemic targeted therapy that is minimally susceptible to chemo- or radioresistance, is potent enough to sterilize individual tumor cells and microscopic tumor cell clusters (even at a low dose-rate and low oxygen tension), and exhibits an acceptable toxicity profile (10). α -Particle–emitting radionuclides address this critical need. To accomplish these goals, a reliable, cost-effective source of α -particle emitters is needed for research and development and for routine use in clinical practice. Improved chemical labeling and stability will be needed to achieve the desired biodistribution and associated dose distribution necessary for successful therapy with acceptable acute and long-term toxicities. These limitations have slowed the development and clinical use of α -emitter targeted therapy relative to the use of β - and Auger-electron-emitting radionuclides.

Received Sep. 29, 2008; revision accepted Jun. 29, 2009.

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The first clinical trial of an α -particle emitter in radiolabeled antibody therapy used ²¹³Bi conjugated to the antileukemia antibody HuM195 and was reported in 1997 (*11,12*), 4 years after ²¹³Bi was first suggested for therapeutic use (*13*). This trial was followed by a human trial of the antitenascin antibody 81C6 labeled with the α -emitter ²¹¹At in patients with recurrent malignant glioma (*14*). In addition to these 2 antibody-based trials, a clinical trial of unconjugated ²²³Ra against skeletal metastases in patients with breast and prostate cancer was recently completed (*15*). More recently, a patient trial of ²¹¹At targeting ovarian carcinoma has been initiated (*16*). Future trials of α -emitters are anticipated using antibodies labeled with ²¹¹At or ²¹³Bi and directed against tumor neovasculature (*17–19*). A conjugation methodology for ²²⁵Ac was recently described (20), and a phase I trial of this radionuclide with the antileukemia antibody HuM195 in leukemia patients has recently been initiated (21). Table 1 summarizes clinical trials involving α -particle–emitting radiopharmaceuticals.

This report focuses on α -emitter dosimetry as it relates to human use in targeted therapy. A review of α -particle radiobiologic studies is provided with a focus on the radiobiology of α -emitters that are relevant to targeted therapy in humans. Closely related to the radiobiology of α -emitters is the concept of relative biological effectiveness (RBE), which is also reviewed. The dosimetry of α -emitters has been addressed in a large number of publications. The criteria for microdosimetry, the different approaches for performing such calculations, and selected studies in which such calculations have been performed are briefly described.

TABLE 1. Summary of Recently Reported Clinical Trials Using α -Particle Emitters				
Radionuclide	Delivery vehicle	Type of cancer	Comments	Reference
²¹¹ At	Antitenascin IgG	Glioblastoma multiforme	Ongoing phase I trial using surgical cavity injection of labeled antitenascin lgG; median survival of 60 wk; 2 patients with recurrent glioblastoma multiforme survived nearly 3 y	14
	MX35 F(ab') ₂	Ovarian carcinoma	Ongoing phase I trial using MX35 F(ab') ₂ ; bone marrow and peritoneal absorbed doses of 0.08 and 8 mGy/MBq, respectively	16
²¹³ Bi	Anti-CD33 IgG	Myelogenous leukemia (acute or chronic)	Phase I completed with no toxicity, substantial reduction in circulating and bone marrow blasts; phase I/II in cytoreduced patients, 4 of 23 patients at very high risk showed lasting complete response (up to 12 mo)	11,21
	Antineurokinin receptor peptide	Glioblastoma	Two patients treated with ²¹³ Bi; 1 with oligodendroglioma treated by distillation in resection cavity alive more than 67 mo	148
	Anti-CD20 IgG (rituximab)	Relapsed or refractory non-Hodgkin lymphoma	Phase I study with 9 patients treated to date	149
	9.2.27 lgG	Melanoma	Sixteen patients; intralesional administration led to massive tumor cell kill and resolution of lesions; significant decline in serum marker melanoma-inhibitory-activity protein at 2 wk after treatment	150
²²³ Ra	RaCl ₂	Skeletal breast and prostate cancer metastases	Phase IA dose-escalation studies completed involving single-dose infusion of 46–250 kBq/kg in 25 patients with no dose-limiting hematologic toxicity; phase IB study in 6 patients to evaluate repeated injections (2 or 5 fractions) totaling up to 250 kBq/kg; phase II randomized trial in 33 patients with metastatic breast or prostate cancer treated with external beam plus saline or 4 times 50 kBq/kg dose of ²²³ Ra at 4-wk intervals; shows significant (-66%) decrease in bone alkaline phosphatase compared with placebo and 15 of 31 patients with prostate-specific antigen decrease > 50% from baseline vs. 5 of 28 in control group	151,152
²²⁵ Ac	Anti-CD33 IgG	Acute myelogenous leukemia	Phase I trial, ongoing, at first dose-level of 18.5 kBq/kg (0.5 μCi/kg); 1 of 2 patients had elimination of peripheral blasts and reduction in marrow blasts	21



Therapeutic nuclear medicine is already a highly multidisciplinary field. Therapy with α -particle emitters is easily one of the more multidisciplinary endeavors within this enterprise. This review is intended to provide the necessary background including the physics and dosimetry perspective to aid in the design, conduct, and analysis of clinical trials using α -emitting radiotherapeutics.

α -PARTICLE RADIOBIOLOGY

Interest in α -particle–emitting radionuclides for cancer therapy is driven by the physical and radiobiologic properties of α -particles as compared with those of photons and electrons (Fig. 1). The energy deposited along the path of an α -particle per unit path length is shown in Figure 2. As shown in the figure, the energy deposition along the path, or linear energy transfer (LET), of an α -particle can be 2 or 3 orders of magnitude greater than the LET of β -particles emitted by radionuclides such as ¹³¹I and ⁹⁰Y.

One of the first studies demonstrating the biologic effects of heavy charged particles was by Raymond Zirkle in 1932 (22). He examined the effect of polonium α -particles on cell division in fern spores and showed a much greater biologic effect when the spore nucleus was placed in the Bragg peak of the α -particle track, compared with the plateau region of the track (23). Much of the subsequent radiobiology of α -particles was established in a series of seminal studies performed by Barendsen et al. in the 1960s (24–32). These studies first demonstrated the now familiar and accepted features of α -particle irradiation. A subsequent series of studies on the mutation and inactivation





of 3 different mammalian cell types exposed to helium, boron, or nitrogen ions spanning LET values in the range of 20-470 keV· μ m⁻¹ was key in evaluating the various biophysical models that had been posited to explain lowversus high-LET effects (33-36). The work was also instrumental in providing both the experimental results and the biophysical analysis to help understand the RBEversus-LET relationship established by Barendsen. The biophysical analysis in the last paper of the series (33)provided compelling theoretic support for the concept of 2 types of radiation-induced cellular inactivation. The first type is that due to the accumulation of multiple events that can be repaired at low doses (i.e., sublethal damage) but that saturate the cellular repair mechanisms at higher doses. This type of inactivation yields the characteristic linearquadratic dose-response curve for low-LET radiation, corresponding to a small number, approximately 3-9 (i.e., \sim 100–300 eV) ionizations in a distance of about 3 nm associated with a low probability of producing lethal lesions. The second type of inactivation arises from a single lethal event for high-LET radiation. In this case, a larger number of ionizations, more than 10, over the 3-nm distance depositing more than 300 eV produces lethal lesions with a high probability. It is important, however, to remember that these studies were performed using external beams of α -particles in which the incident α -particles were generally orthogonal to an α -permeable surface on which the cells were cultured as a monolayer of adherent cells.

As initially demonstrated experimentally by Fisher et al. (37), and then theoretically by Humm et al. (38), and most recently by Kvinnsland et al. (39), the spatial distribution of α -particle emitters has an important impact on the absorbed dose distribution and, correspondingly, on the slope of the cell-survival curve. Neti and Howell recently provided experimental evidence of a lognormal cellular uptake of ²¹⁰Po citrate among a cell population uniformly exposed to the radiochemical and showed that this distribution can substantially alter the cell survival curve (40). Although many of the results obtained from the external-beam studies (summarized in Table 2) are generally applicable regardless of the α -particle distribution, specific parameters such as the average number of α -particle traversals to induce a lethal event or the D₀ value (i.e., the absorbed dose required to reduce cell survival to 0.37) are highly sensitive to experimental factors such as the geometry of the cells, the thickness or diameter of the cell nucleus, the distribution of DNA within the nucleus (i.e., the phase of the cell cycle), and the number and spatial distribution of the α -particle sources relative to the target nuclei.

The distinction between DNA double-strand breaks (DSBs) caused by a single high-LET track versus DNA damage caused by multiple low-LET tracks is illustrated in Figure 3. This basic observation underpins almost all the radiobiology of α -particles.

TABLE 2. α -Particle Beam Findings That Are Also Applicable to Internally Administered α -Particle Emitters					
No.	Finding				
1	RBE > 1 for cell sterilization, chromosomal damage/				
	cancer induction relative to low-LET radiation				
2	Reduced susceptibility to modulation by				
	radiosensitizers and radioprotectors				
3	Reduced capacity to repair sublethal damage				
4	Higher induction of DNA DSBs at low total absorbed doses				
5	Monoexponential surviving fraction curve after				
	uniform irradiation (absence of a shoulder)				

Traversals Required for Cell Kill

The average number of α -particle nuclear traversals required to kill a cell, as measured by loss of the subsequent ability to form a colony, ranges from as low as 1 (41) to as high as 20 (42). If bystander effects are included, the lower end of the range would include 0. The variability in this value when bystander effects are not considered arises because of the high sensitivity of this determination to the geometry of the cell and the nucleus during irradiation and also the LET of the incident α -particles and the LET distribution within the nucleus.

Quoting from a publication of Raju et al. (43), "The notion that a cell will be inactivated by the passage of a single α particle through a cell nucleus prevailed until Lloyd and her associates (42) demonstrated that 10 to 20 5.6 MeV α particles were required to induce one lethal lesion in flattened C3H 10T1/2 cells. Studies by Bird, et al. (44) showed that approximately four ³He ions were required to pass through the cell nucleus to induce one lethal lesion in V79 cells at the G₁/S-phase border, cells in late S phase required five to eight ³He ions. Todd, et al. (45) investigated the effect of 3.5 MeV α particles on synchronized T-1 cells, and observed that approximately one α particle out of four to five traversing a cell nucleus is effective in inducing one lethal lesion. Roberts and Goodhead (46) estimated that one out of six 3.2 MeV α -particle traversals through a C3H 10T1/2 cell nucleus is lethal. Barendsen (47) concluded that the probability of inactivation per unit track length of high-LET α particles is approximately 0.08 μ m⁻¹ for both T-1 and C3H 10T1/ 2 cells consistent with the results of Roberts and Goodhead for C3H10T1/2 cells (46)." In a study comparing high-LET effects of Auger versus α-particle emitters, Howell et al. found that about 9 decays of ²¹⁰Po were required to reduce cell survival to 37% (D₀) when it was distributed between the cytoplasm and nucleus of V79 cells; the energy deposited in the cell nucleus corresponds to about 2 complete (maximum chord length) traversals of the cell nucleus (48). In a murine lymphoma cell line, approximately 25 cell-bound α -particle–emitting ²¹²B immunoconjugates were required to reduce clonogenic



FIGURE 3. Single high-LET track has high probability of yielding DNA DSB, whereas probability of DSB induction with low-LET tracks becomes comparable only at higher absorbed doses. (Reprinted with permission of (154).)

survival by 90% (49). The theoretic efficiency of DSB production when an α -particle passes through DNA was examined by Charlton et al. (50) and was found to be surprisingly low; approximately one eighth of 10-MeV α -particles passing through a 54-nucleotide section of DNA produce a DSB. One passage in 4 of 1.2-MeV α -particles produces a DSB.

Barendsen's estimate of the inactivation probability per unit track length and Goodhead et al.'s determination of the number of lethal lesions per micrometer track through the nucleus (33) suggest another approach for estimating inactivation probability. Along these lines, Charlton and Turner introduced the total α -particle path length (or chord length) through the nucleus as a useful parameter (51). This was used to derive λ , the mean free path between lethal events for α -particles traveling through nuclei. Drawing from an extensive compilation of experimental data, the investigators found that this parameter ranged from 1.5 to 64.4 μ m. As expected, λ was found to be dependent on the LET (Fig. 4). An inactivation probability per unit track length through the nucleus has also been used in a model describing radiation-induced cellular inactivation and transformation. By incorporating aspects of a state vector model for carcinogenesis (52) into the inactivation/transformation model, Crawford-Brown and Hofmann (53) have described a model that successfully predicts both cell survival and transformation after irradiation by α -particles of different LETs at absorbed doses below 1 Gy. This model was used to examine the impact on model predictions of including a correlation between initiation of cellular transformation and cellular inactivation. At absorbed doses greater than 1 Gy, a significant difference was observed in the predicted probability that a cell is transformed and survives.

Cell Survival Curve

Cell survival curves (i.e., surviving fraction, SF, vs. absorbed dose, D) for low-LET radiation such as x-rays exhibit an initial "shoulder" that is thought to reflect the repair of radiation damage. This type of cell survival curve can be represented by the linear-quadratic equation

$$SF = e^{-\alpha D - \beta D^2}.$$
 Eq. 1

The parameters α and β are, respectively, sensitivity per unit dose (D) and per unit dose squared (D²). As the absorbed dose exceeds a certain threshold level, presumably



FIGURE 4. Dependence of mean free path on LET. LET is plotted (e.g., from 200 to 100 keV/ μ m after 250 keV/ μ m) so that stopping powers on low-energy side of Bragg peak can be identified. (Reprinted with permission of (*51*).)





the dose at which the radiation damage repair rate is reduced relative to the rate of induced damage, the relationship between surviving fraction and absorbed dose approaches log-linearity. As shown in Figure 5, the cell survival curve for α -particle radiation is log-linear at low as well as high absorbed doses; that is, it does not exhibit a shoulder region, reflecting the reduced capability of cells to repair α -particle damage. The equation describing this is

$$SF = e^{-D/D_0}, \qquad Eq. 2$$

with the parameter D_0 equal to the absorbed dose required to yield a surviving fraction of 37%. The log-linear aspect of cell survival curves after α -particle irradiation reflects a reduced repair capacity, not the absence of repair. That α -particle damage is repaired has been demonstrated by several studies, as described in the "Radiomodulation" section. Repair of damage is not inconsistent with singleevent lethality and a log-linear survival curve. The key distinction is whether death is a result of accumulated damage or of a single event. Cell survival curves that exhibit an initial shoulder reflect cell death that results from the accumulation of damage, whereas log-linear cell survival curves reflect cell death arising from a single event, without the need to accumulate damage. In both situations, repair is possible.

Oxygen Effect

In addition to dose rate, the influence of oxygen concentration has long been recognized as an important factor in the response of cells to radiation (54,55). Figure 6 demonstrates that this effect is strongly influenced by the LET of the radiation. The oxygen enhancement ratio (OER), or relative radiosensitivity of cells to oxygen concentration, is 1 for charged particles with an LET greater than 140 keV/ μ m (24). The initial LET of 4- to 8-MeV α -particles typical of the α -emitters of interest in

targeted α -emitter therapy ranges from 110 to 61 keV/ μ m. The OER values in this LET range are 1.3 to 2.1. Because the LET of the emitted α -particles increases well beyond the 140 keV/ μ m threshold for OER = 1 as the Bragg peak is approached, the ability of α -particles to overcome radioresistance due to hypoxia will depend on the spatial distribution of the α -emitters relative to the hypoxic region. The ability to overcome hypoxia, noted above, is strictly radiobiologic. There are studies suggesting that hypoxia may alter the phenotype of the cell via cell signaling pathways associated with increased concentrations of hypoxia-inducible factor 1α , leading to a cell phenotype that is inherently more resistant to radiation and other cytototoxic agents, including chemotherapeutics (56). The classic OER effect has been explained as a free radicalmediated effect in which the presence of oxygen "fixes" free radical-induced damage, thereby making repair of the damage more difficult (57). In this case, the reduced OER effect with α -particle radiation may be explained by the



FIGURE 6. OER as function of LET. OER was measured using cultured human kidney-derived cells incubated in air or nitrogen. α -Particles of different energies generated by cyclotron or 250-kVp x-rays (average LET \approx 1.3 keV/ μ m) were used. (Data replotted from (24).)

preponderance of oxygen-independent direct DNA damage (vs. oxygen-dependent indirect, i.e., free radical-mediated, DNA damage) characteristic of α -particles.

Dose Rate

The influence of absorbed dose rate on cell survival for low-LET emissions is well established. As the dose rate is lowered and the exposure time extended, the biologic effect of a given dose is generally reduced (58). The primary explanation for this effect is that lower dose rates provide a greater time interval for DNA damage repair. Because high-LET damage is not easily repaired, dose rate or even dose fractionation should not impact cellular survival. Barendsen examined changes in survival after α -particle irradiation over a dose-rate range of 0.5–100 rad/min, and no dose-rate effect was observed (26).

Oncogenesis

Although not of prime concern in cancer therapy, a much higher incidence of cancer induction is associated with α -particle irradiation (59). Accordingly, the radiation weighting factor for α -particles is 20, meaning that a committee review of the relevant experimental and human data has determined that per unit absorbed dose, α -particles are associated with a 20-fold greater risk of cancer induction than is a similar absorbed dose of photons or β -particles (60). A review of human and animal data related to cancer risk estimates has called the value of 20 into question for bone cancer and leukemia risk, particularly at low absorbed doses (61). Consideration of dose to target cells on bone surfaces as opposed to the average bone dose gives an RBE for bone cancer risk of 3-12. The authors (61) noted that these estimates may also change since there is evidence that bone cancer risk may be best assessed by calculating dose to a 50-µm layer of marrow adjacent to the endosteal bone surface as opposed to a single 10-µm layer as currently assumed. Likewise, a factor of 2 to 3 is more consistent with the experimental data for leukemia induction. All these estimates are based on α -particle emitters not projected for use in targeted α -emitter therapy. The few studies that have been performed to examine carcinogenesis of the short-lived α -emitters of interest in targeted α -emitter therapy have used ²¹¹At. Neoplastic changes, predominately papillary carcinomas in various organs, were seen in a few animals but not more than what was expected for untreated mice. Brown and Mitchell (62) reported a 13% incidence of plasmocytoma in tumor-bearing mice of the same strain 13-21 mo after treatment with 200-750 kBq of 6-211At-astato-2-methyl-1,4-naphthoquinol bis(diphosphate salt). The frequency of low-grade B-cell non-Hodgkin lymphoma was high but similar to that of the control population. A high incidence of pituitary adenomas and mammary tumors has been seen in rats treated with ²¹¹At (63, 64). These tumors, however, were partially attributed to secondary effects associated with a hormonal imbalance resulting from thyroid or ovarian tissue compromise.

Fractionation

The fundamental rationale for fractionation in externalbeam radiotherapy is based on the differential repair capacity of most normal organs compared with most tumors. This is expressed in terms of early versus late responding tissues, corresponding to high versus low α/β ratios (65). Fractionation tends to spare normal organs without a reduced efficacy against tumors. As shown in Figure 7, this is not the case with high-LET radiation (26). Cultured cells derived from human kidneys showed the same surviving fraction for a single total absorbed dose of α -particle radiation or the same total dose delivered in 2 equal fractions, separated by 12 h. For the same cell line, similar results have been observed when the total dose was delivered in 3 equal fractions at 4, 8, and 12 h after cell plating (25). Extension of the biologically effective dose formalism to account for RBE effects has also demonstrated that fractionation is theoretically not likely to confer a normal tissue-sparing effect for high-LET radiation (66). Similar conclusions may be drawn for the chronic, exponentially decreasing dose rates delivered by internally administered α -particle emitters.

Radiomodulation

Few examples of agents that can modulate α -particle radiation–induced damage have been reported. In the early 1960s, Barendsen et al. compared the radioprotective effects of cysteamine and glycerol (25). The surviving fraction of T₁ (human kidney–derived) cells increased by a factor of 3.7 for 250-kVp x-irradiation and only 1.2 for



FIGURE 7. Effect of fractionation on cell survival: cell survival curve obtained with single doses of 200-kV x-rays (a), curve obtained when 200-kV x-ray doses are separated by 12 h (4.5 Gy, then 2.5 or 4.5 Gy) (b), and curve obtained with 3.4-MeV α -particles (c). In (c), circles correspond to single exposure, and squares to 2 equal exposures separated by 12 h. (Adapted from (26).)

²¹⁰Po α -particle radiation. Similar results were observed with glycerol; cell survival was increased by 2.0 and 1.2 for 250-kVp x-rays and ²¹⁰Po α-particles, respectively. Qualitatively consistent but quantitatively different results have been obtained with the radiosensitizer Wortmannin. This irreversible and potent inhibitor of DNA-dependent protein kinase is involved in the nonhomologous end-joining DNA repair pathway invoked in the repair of DNA DSBs (67). In V79 Chinese hamster cells, Wortmannin led to a 3- to 4-fold increase in genotoxic damage, as measured by the induction of micronuclei. High-LET irradiation, as delivered by a boron neutron-capture reaction, leading to the release of α -particles with an average energy of 2.3 MeV, vielded an increase in micronucleus induction of approximately 2-fold. This finding suggests that the more complex double-strand damage induced by high-LET radiation is a substrate of the nonhomologous end-joining pathway (68,69). In vivo studies in mouse testes have shown that soybean oil, S-(2-aminoethyl)isothiuronium bromide hydrobromide, and cysteamine afford some protection against the cytotoxic effects of 5.3-MeV α -particles emitted by ²¹⁰Po (70–73). When spermatogonial cell survival was used as the biologic endpoint, dose modification factors of 2.2, 2.4, and 2.6, respectively, were obtained. No modification of the spermatogonial response to α -particles was observed when dimethyl sulfoxide or vitamin C was used (74,75).

That DNA damage and its repair are at the core of α -emitter radiobiologic effects is supported by many years of experimental and theoretic work. It is important, however, to keep in mind that all the foundation work regarding the radiobiology of α -emitters was performed well before modern molecular biology came into existence. In light of the remarkable and far-reaching gains in our understanding of the molecular mechanisms involved in cancer genesis, the cellular response to radiation, and DNA single and DSB repair, a reexamination of α -particle radiobiology using modern tools is warranted.

RBE

The biologic effect of ionizing radiation is influenced by the absorbed dose, the dose rate, and the quality of radiation. Radiation quality is characterized by the spatial distribution of the energy imparted and by the density of ionizations per unit path length, referred to as the LET or stopping power of a charged particle (22,60). Depending on the effect considered, greater ionization density along a track will increase the probability of inducing a biologic effect. Compared with electrons and β -particles, α -particles exhibit a high density of ionization events along their track (76). Electrons and β -particles that are emitted by radionuclides generally range in energy from several megaelectron volts to as low as several kiloelectron volts, with corresponding LET values ranging from about 0.1 to 1 keV/µm (β-particles actually are characterized by a spectrum of energies; the bottom end of the spectrum is zero). The exception to these is Auger electrons, which have energies

as low as several electron volts and corresponding LET values as high as 25 keV/ μ m. α -Particles emitted by radionuclides range in energy from 2 to 10 MeV, with initial LET values ranging from 60 to 110 keV/µm. A given tissue-absorbed dose resulting from α -particles, therefore, is likely to yield considerably greater biologic effects (again depending on the effect being considered) than the same absorbed dose delivered by typical electrons or β -particles. To account for differences in energy deposition pattern exhibited by different quality radiations, the concept of RBE has been established. An authoritative review of this concept, its derivation, and appropriate application has been published by the International Commission on Radiological Protection (ICRP) (60,77), and the reader is encouraged to consult this source for additional information. In radiobiology, RBE equals the ratio of absorbed doses of 2 types of radiation that produce the same specified biologic effect.

RBE Defined

RBE is calculated as the absorbed dose of a reference radiation (e.g., x-rays, γ -rays, β -particles), $D_r(x)$, required to produce a biologic effect, x, divided by the absorbed dose of the test radiation, $D_t(x)$, required to produce the same biologic effect:

$$RBE(x) = \frac{D_r(x)}{D_t(x)}.$$
 Eq. 3

RBE is thus an experimentally determined value defined for a particular biologic effect and therefore for a particular biologic system.

The experimentally determined value can be influenced by the variability of the biologic system across different laboratories. This issue has been examined for studies in vitro (78). The methodology used for calculating the absorbed dose of the 2 radiation types will also impact the result. Ideally, this should not be the case. The methodology used should provide the true absorbed dose value or specific energy distribution ("Case for Microdosimetry" section) to the relevant biologic target for both the test and the reference radiations. In practice, however, this is a challenge even for studies in vitro (79). In the setting of human α -particle emitter dosimetry, consistency and reproducibility will be as important as accuracy. This issue is discussed in greater detail in "Recommendations for Dosimetry of Deterministic Effects" section.

The fact that the RBE is related to the pattern of ionizing energy deposition along a particle track leads to a third factor that will impact the results. The RBE for a particular radiation type will also depend on the initial emission energy of the particle (i.e., how close the particle is to the end of its track [the Bragg peak]). This factor has been examined by Charlton et al. (80) and Howell et al. (81). In the studies by Howell et al., a uniform distribution of decays was assumed to calculate the D₀ for 7 α -emitting isotopes covering a wide range of initial energies. Using the D_0 obtained for x-rays for the cell line used in the α -emitter calculations, a linear relationship between RBE and initial α -particle energy was obtained over an initial α -energy ranging from 5 to 8.5 MeV. The straight line was given by RBE = 2.9–0.167E_i, where E_i is the initial α -particle energy in megaelectron volts. This is an approximate scaling of the equation derived from in vivo experimental data by Howell et al. (*81*). In addition to effects related to the Bragg peak, nonuniform biodistribution of the α -emitters also leads to microdosimetric effects that impact RBE and the slope of the cell-survival curve (*37–39*).

If the reference radiation yields a dose-response relationship that is not log-linear for the biologic system examined, the RBE value will depend on the specific biologic quantitative endpoint selected (e.g., D_{50} , D_{37} (= D_0), D_{10} , etc., which determines whether the comparison falls in the shoulder or in the log-linear region of a doseresponse or survival curve). The type of biologic endpoint (e.g., survival, mutation) and the dose rates of the test and reference radiations will also influence the RBE value. Strictly speaking, the test radiation should be delivered in a manner identical to that of the reference radiation (e.g., chronic or acute). However, acute externally administered x- and γ -rays are often used as the reference radiation when RBE values are determined for internally administered radionuclides. Given the often-sizeable difference in biologic responses to acute-versus-chronic low-LET radiation, the dose rate at which the reference radiation is delivered can impact the resulting RBE (48). The dose-rate pattern delivered by radiopharmaceuticals is generally well represented by multicomponent exponential functions. Howell et al. have delivered such patterns with external beams of 137 Cs γ -rays (82). This approach was used to study the bone marrow response to exponentially decreasing dose rates of 137 Cs γ -rays (83). The response of granulocyte-macrophage colony-forming cells in the marrow to decreasing dose rates with half-times ranging from 62 h to ∞ (i.e., constant dose rate) were studied and compared with the response to acute exposures. Mean lethal doses for chronic irradiation were up to 40% higher than those for acute exposures. Thus, care must be taken when comparing RBE values based on different reference radiations.

Based on a review of experimental literature, an RBE value of between 3 and 5 was recommended for cell killing by a panel convened by the Department of Energy in 1996 (84). Because human studies using α -particle emitters have yet to be analyzed for deterministic effects, an RBE of 5 was recommended for projecting the possible deterministic biologic effects associated with an estimated α -particle absorbed dose.

RBE, Q, and w_R

The discussion thus far has focused on RBE. RBE is occasionally confused with quality factors, Q, and radiation

weighting factors, w_R. This confusion reflects the historical evolution of RBE which was originally defined as relative biological efficiency and intended to apply to both radiobiology (deterministic effects) and protection (stochastic effects). As currently recommended by the ICRP, however, RBE is not to be used directly in radiation protection but only as a starting quantity to derive the radiation weighting factor $w_{\rm R}$, which replaced the quality factor Q in the most recent ICRP recommendations (85,86). The RBE values used to arrive at $w_{\rm R}$ relate to stochastic endpoints such as cancer induction, rather than deterministic endpoints such as normal-tissue toxicity and tumor cell sterilization in cancer therapy patients. The ICRP radiation weighting factor for α -particles is 20. This value, intended only for stochastic effects caused by α -particle irradiation, is based on animal experiments and from analysis of historical α -emitter exposures. In contrast to RBE values, weighting factors are not directly measured values but rather are consensus recommendations of the ICRP (60).

The radiation weighting factor $w_{\rm R}$ is a unitless factor that converts average absorbed dose (in units of grays) to equivalent dose in an organ or tissue. The SI unit for equivalent dose is referred to by the special name *sievert*. The sievert is not a unit in the conventional sense but is intended to indicate that the absorbed dose value has been adjusted to reflect a biologic risk that is associated with stochastic effects. Although the sievert is often used in the context of deterministic effects, this use is not strictly correct because the ICRP has stipulated that the sievert should be used only to designate the risk of incurring stochastic biologic effects such as cancer. The ICRP has reported on RBE for deterministic effects (RBE_M), but no special name has been chosen by the ICRP for the product of absorbed dose and a factor such as RBE that specifically reflects similar scaling for a deterministic effect (77).

α -PARTICLE DOSIMETRY

Radiation dosimetry offers a means for standardizing and comparing the efficacy of different radiation-based treatments. It provides a logical basis for understanding the effects that various radiation qualities have on biologic matter. For α -particle emitters, accurate dosimetry calculations require knowledge of the activity distribution as a function of time at the cellular and subcellular levels (87). Furthermore, an accurate representation of the geometry at this level is also required. For in vitro experiments (i.e., cell survival studies), the activity distribution is straightforward, consisting of uptake on the surface or within the cell, along with a known fraction in the surrounding solution. In these experiments, the cell and nucleus can be approximated as concentric spheres, the dimensions of which can easily be measured. However, for clinical applications, these idealizations give way to complex activity and tissue geometries. In these cases, modeling the 3-dimensional geometry of a spheroid (88,89) or using microscopic data from tissue biopsy samples (90) can provide information on the target geometry. Determining the activity distribution, however, remains difficult. Autoradiography (91) may provide a snapshot of the activity distribution at a single instance in time. However, the determination of the activity as a function of time may require mathematic modeling (92-94) of the carrier molecules as they diffuse through tissue and bind to markers on cell surfaces. Ideally, such modeling should be validated using animal model measurements in vivo.

Case for Microdosimetry

There are 2 methods for calculating the energy deposited by individual α -particles. One method uses the MIRD formalism to calculate the average dose to the target (cell nucleus) for a variety of source compartments (cell surface, cytoplasm, and nucleus). Extensive tables have been produced for various combinations of α -particle–emitting radionuclides and cellular geometries (95,96). The basis for using mean absorbed dose is related to the biologic properties of low-LET radiations such that a large number, often several thousands, of statistically independent radiation deposition events in a single cell nucleus is required to induce a demonstrable biologic effect. In such a case, the statistical variation of the energy imparted to different cell nuclei is minimal. In contrast, for high-LET irradiation, such as α -particles, the effect of even a single event in the cell nucleus is so great that the mean absorbed dose can be a misleading index of biologic effect. This is due to several reasons. Foremost is that the number of α -particles that traverse a cell nucleus is often few, and therefore stochastic variations become important. In addition, the path of the α -particle through the cell nucleus is also critical. An α -particle that crosses directly through a cell nucleus will deposit a large amount of energy, whereas one that merely grazes the surface will deposit little or no energy. Thus, a second method for *a*-particle dosimetry-microdosimetry-takes into account the stochastic nature of energy deposited in small targets. The fundamental quantities in classic microdosimetry are specific energy (energy per unit mass) and lineal energy (energy per unit path length through the target) (97). Microdosimetry was originally proposed by Rossi (98) to explain the stochastic nature of energy deposited in matter by external ionizing radiation. It has subsequently been adapted to the case of internally deposited α -particle emitters (99–101).

Microdosimetric Techniques

Microdosimetric spectra may be calculated using either analytic or Monte Carlo methods (102). Analytic methods use convolutions (via Fourier transforms) of the singleevent spectrum to calculate multievent distributions (98). The single-event spectrum represents the pattern of specific energy depositions for exactly 1 α -particle hit. Kellerer developed a method to efficiently determine the multipleevent spectrum through the use of Fourier transforms (103). Although analytic codes are computationally efficient, they are often limited to simple source-target geometries because the single-event spectrum must be known for each source-target configuration. Monte Carlo codes offer greater flexibility than analytic methods and can simulate a wide variety of geometries and source configurations. Idealizations are often made to simplify the coding and reduce calculation time. In nearly all Monte Carlo codes, α -particles are assumed to travel in straight lines. This approximation is valid for α -particles having energies less than 10 MeV (97). In addition, the range of δ -rays (energetic electrons originating from the α -particle track that cause secondary ionizations in the vicinity of the track) and the width of the α -particle track (~100 nm) are often ignored because the targets that are studied (i.e., cell nucleus) are much larger than these dimensions (104). The rate of α -particle energy loss is characterized by the stopping power. These data for a variety of media can be obtained from the literature (105-108). Inherent in the stopping-power formulation is the continuous slowingdown approximation. As the name implies, this approximation assumes that α -particles lose energy continuously as they traverse matter. Thus, the calculated specific energy imparted depends on the choice of stopping powers used.

Criterion for Adopting Microdosimetry

The rationale for microdosimetry was outlined by Kellerer and Chmelevsky (109). They suggested that the stochastic variations of energy deposited within the target must be considered when the relative deviation of the local dose exceeds 20%. For example, a small cell nucleus with a diameter of 5 μ m irradiated by α -particles would require an average dose of at least 100 Gy for the relative deviations to be less than the 20% threshold. Thus, the necessity for microdosimetric methods will depend on the source distribution, the target size and shape, and the expected mean dose. For small average doses (such as those expected by nontargeted tissues) microdosimetry may be important in characterizing the pattern of energy deposition and in understanding how this pattern relates to clinical outcomes. However, in tumor, where the mean dose may be large, a microdosimetric treatment may not be necessary.

Microdosimetry Implementation Techniques

Although microdosimetry has increased our understanding of stochastic patterns of energy deposition by α -particles in both simple and complex geometries and has made it possible to explain in vitro observations, application to clinical practice has been limited because time-dependent activity distributions at the subcellular level are complex and not well characterized in vivo. Roeske and Stinchcomb (110) described a technique for determining dosimetric parameters that are important in α -particle dosimetry. These parameters consist of the average dose, SD of specific energy, and the fraction of cells receiving zero hits. The individual values are determined using tables of the "S" value (111), and the first and second moments of the single-event spectra. The average dose is determined by multiplying the S value by the cumulated activity within the source compartment. Dividing the average dose by the first moment of the single-event spectrum yields the average number of hits. Subsequently, the fraction of cells receiving zero hits (or any number of hits) can be determined using the average number of hits and the Poisson distribution. The SD is the product of the average number of hits and the second moment of the single-event spectrum. Individual moments may be determined using either analytic methods or Monte Carlo calculations. Stinchcomb and Roeske (*112*) have produced tables of the S value and the individual moments for several geometries and source configurations appropriate for α -particle therapy. These tables were also used in the analysis of cell survival after α -particle irradiation (*112*).

Applications of Microdosimetry

Early applications of microdosimetry were performed to assess the probability of cancer induction after exposure to α -emitters. These exposures were generally not intended for therapeutic purposes, and carcinogenesis was of concern. In one such application, the specific energy distributions for plutonium oxide in dog lung were calculated. The calculations accounted for the size distribution of the inhaled aerosol and the associated deposition probabilities in the lung for various particle sizes. The distribution of target sites; the probability of an α -particle intersecting a target site; and the range, energy loss, straggling characteristics, and δ -ray production of α -particle tracks were also considered. The analysis provided an improved understanding of the relationship between dose, as described by microdosimetric specific energy spectra, and response, as measured by the incidence of lung tumors in beagle dogs (113).

In radioimmunotherapy, microdosimetry has been used in several α -particle applications. These applications can be broadly characterized as theoretic studies of simple cellular geometries, experimental analysis of cell survival after α -particle irradiation, and the microdosimetry of realistic geometries such as multicellular spheroids and bone marrow. The work in each of these categories will be discussed separately.

Roesch (99) described an approach for calculating microdosimetric spectra. Fisher et al. (37) subsequently applied this approach to several geometries that have therapeutic application, including sources distributed on and within individual cells, sources distributed within spheric clusters of cells, and sources located in cylinders (i.e., blood vessels) that deposited energy within spheric cell nuclei a short distance away. These calculations showed the number of α -particle emissions originating from cell surfaces that would be needed to inactivate cancer cells with high efficiency. The basic geometries that described the spatial distribution of α -emitters relative to the spatial distribution of target spheres have served as the basis of those used in several theoretic studies. In one such study, Humm (114) used a Monte Carlo method with

a model of cell survival to estimate the surviving fraction of cells located outside a capillary and cells located within a tumor with uniformly distributed ²¹¹At. Although the mean dose was similar for these 2 types of geometries, there was a significant variation in the expected cell survival due to the differences in the specific energy spectra. In particular, the fraction of cells receiving no α -particle hits increased with distance from the capillary (due to the short range of the α -particles). The surviving fraction versus mean specific energy was biexponential. That is, for low doses, the slope of this curve was similar to that of a uniformly irradiated tumor. However, with increasing doses, the curve was less steep and asymptotically approached a value corresponding to the fraction of nonhit cells. Building on the previous analysis, Humm and Chin (38) analyzed how specific energy spectra are affected by cell nucleus size, binding fraction, cell volume fraction, and nonuniform binding. Their results indicated that nonuniform distributions of α -particle emitters can result in expected survival curves that deviate significantly from the classic monoexponential curves produced by a uniform, external source of α -particles. In these studies, although the inherent cell sensitivity (z_0) was held constant, the slope of the cell survival curve as a function of absorbed dose to the medium was highly dependent on the source configuration. Furthermore, simulations in which cells were more uniformly irradiated resulted in steeper cell survival curves than those in which the distribution of α -emitters was highly heterogeneous. The effects of cell size and shape on expected cell survival were further studied by Stinchcomb and Roeske (115). In their analysis, the cell and nucleus were assigned various shapes ranging from spheres to ellipsoids where the ratio of the major-to-minor axis was varied from 1 to 5 while the volume of the nucleus was held constant. Separately, the dimensions of the nucleus were varied while the shape was held constant. Calculations of specific energy spectra and resulting cell survival demonstrated that the expected surviving fraction was not a strong function of the target shape, provided the volume was fixed. However, significant variations in cell survival were observed as the volume of the nucleus was varied. More recently, Aubineau-Laniece et al. developed a Monte Carlo code to simulate cylindric geometries as a model for bronchial airway bifurcations (116). In a series of reports on α -particles from radon progeny, Fakir et al. (117–119) demonstrated that for uniform surface emissions, there were significant variations in cellular energy deposition. Larger variations in the hit frequencies and energy deposited were observed when a nonuniform distribution of activity was also considered. Palm et al. (120) examined the microdosimetric effects of daughter products from ²¹¹At. Separate simulations were performed assuming the daughter products decayed at the site of ²¹¹At emission or that they diffused away from the site. Based on an analysis of experimental data, the ²¹⁰Po daughter product seemed to diffuse from the decay site, decreasing the energy deposited in the cell nucleus by a factor of 2. All these studies illustrate the need to accurately model the source-target geometry. Moreover, approximations, such as using mean values, may impact both the specific-energy spectrum and subsequent calculation of cell survival (39).

Application to Cellular Clusters

Single-cell survival analyses after α -particle irradiation has also been extended to multicellular clusters. Charlton (89) described a multicellular spheroid model and simulated α -particle energy deposition events within individual cell nuclei. A cell survival model that takes into account the effects of varying LET (51) was combined with the distribution of α -particle tracks throughout cells within the spheroid. Simulating a uniform source distribution (average 1 decay per cell, 50% cell packing), this analysis demonstrated that cell survival decreased significantly (from 57% to 37%) as the spheroid diameter increased from 75 to 225 µm. The number of hits per cell also increased in larger spheroids when longer-ranged α-particle emitters were considered. Cell survival subsequently decreased from 46% to 26% in 200-µm-diameter spheroids as the packing fraction was increased from 40% to 70% (also with 1 decay per cell). The decrease in cell survival was due to the increased crossfire dose as the packing fraction was increased. In a separate simulation, the total number of decays per spheroid was kept constant while a small fraction of cells (20%) was assumed not to take up any activity. This process simulated the effects of cells that lacked a specific targeting moiety. It is interesting to note that the unlabeled fraction did not significantly alter the expected cell survival. In these studies, the specific energy distribution is highly nonuniform and varies with depth below the spheroid surface. Thus, a single dose or specificenergy distribution is not representative of that through the entire tumor. By combining the specific-energy distribution with cell survival models, it is possible to gain insight into those factors that will influence the therapeutic efficacy of a particular targeting approach. However, most of these cell survival models do not take into account second-order processes such as the bystander effect that may play an important role in modeling cellular clusters and micrometastases. Refinement of these models is currently an active area of research (121,122).

Application to Bone Marrow

Bone marrow is often the dose-limiting organ in radioimmunotherapy. The dosimetry of bone marrow is difficult because of its complex geometry and the presence of tissue inhomogeneities. Thus, idealized models, as have been used in the previous studies, must be replaced by more realistic geometries. The work to date on estimating specific energy spectra for bone marrow has focused largely on using histologic samples obtained from humans or animal models. Akabani and Zalutsky (90) obtained histologic samples of beagle bone marrow and manually measured chord length distributions. Using a Monte Carlo program, they calculated the single-event specific energy distribution for sources both in the extracellular fluid and on the surface of red marrow cells. These single-event distributions were combined with a model of cell survival. This analysis demonstrated that activity concentrated on the cell surface resulted in significantly greater cell killing than did activity in the extracellular fluid. The effect of LET on the survival of human hematopoietic stem cells in various geometries was studied by Charlton et al. (80). These geometries were determined from human marrow samples obtained from cadavers. Microdosimetric spectra and cell survival were calculated for 3 different source-target geometries: isolated cells labeled on their surfaces, a nontargeted distribution of decays in an extended volume, and nontargeted decays in marrow with 36% of the marrow volume occupied by fat. Two different radionuclides, ¹⁴⁹Tb and ²¹¹At, were considered. These simulations indicated that for targeted decays ¹⁴⁹Tb was 5 times more effective than ²¹¹At when compared on a hit-by-hit basis. This enhancement was due to the lower energy of ¹⁴⁹Tb resulting in a higher LET of the incident α -particles. Those authors also concluded that cell survival was a function of the position of the decay relative to the cell nucleus. Using a model similar to that of Charlton et al. (80), Utteridge et al. (123) considered the risk of the development of secondary malignancies (i.e., leukemia) from α -particles. This risk may be important in evaluating the future therapeutic application of α -particles in patients who have an excellent prognosis. Three α -emitting radionuclides were considered on the basis of the relative range (short, medium, and long) of the particle. In this analysis, the authors calculated the fraction of cells that are hit and would survive (as these would potentially cause secondary malignancies). They determined that the lowest fraction occurred for low energies and the highest fraction occurred for the highest-energy α -particle emitter.

RECOMMENDATIONS FOR DOSIMETRY OF DETERMINISTIC EFFECTS

Beyond providing a rational basis for a starting administered activity value for a phase I study, dosimetry has an important role in guiding clinical trial design to help maximize the likelihood of a successful, minimally toxic implementation. This is particularly important because α -emitter targeted therapy has the potential to be both highly effective and also quite toxic. Which of these 2 aspects emerges in a therapeutic trial will depend on having an understanding of the physical and biologic factors that impact response and toxicity. It is essential that clinical trials investigating targeted α -particle therapy be rationally designed; otherwise, there is the risk that α -emitters may be abandoned before they have been properly tested in the clinic.

This increased importance of dosimetric analysis is coupled with a greater difficulty in obtaining the human data necessary to perform dosimetry. In contrast to most targeted therapy trials to date, collection of biodistribution data for dosimetry from pretreatment imaging studies will not be possible for most α -particle–emitting radionuclides with therapeutic potential. This places a greater emphasis on preclinical studies and extrapolation of results obtained from such studies to the human. Several of the α -emitting radiotherapeutics decay to α -emitting daughters whose distribution may not be that of the carrier. Aside from understanding the biodistribution and dosimetry of the α -emitter–labeled carrier, therefore, the biodistribution and dosimetry of the daughter must also be considered (124–131).

In this section, the focus of the discussion and the recommendations that are made are specific to deterministic effects.

Recommendations

After stability and radiochemical purity of the radiopharmaceutical have been established and an appropriate target identified, the following progression of studies is proposed. Elements of these recommendations have also been described elsewhere (132-134).

- 1. Determine cellular targeting kinetics and properties.
 - A. Determine number of sites per cell and fraction of cells expressing target.
 - B. Determine distribution of binding sites per cell among the targeted cells.
 - C. Determine binding and dissociation constants for cell targeting (e.g., antibody affinity).
 - D. Determine internalization rate and fraction internalized.
 - E. Determine fate of internalized radionuclide.
 - F. Determine median lethal dose in targeted versus nontargeted cells.
 - G. Determine cell-level dosimetry for targeted and nontargeted cells.
- 2. Perform animal (xenograft or transgenic) model studies.
 - A. Evaluate maximum tolerated administered activity.
 - B. Identify likely dose-limiting organs.
 - C. Collect macroscopic (whole-organ) pharmacokinetics.
 - D. Collect microscopic (e.g., by autoradiography or optical imaging) biodistribution in dose-limiting organs.
 - E. Evaluate stability of the radiopharmaceutical in vivo.
 - F. Evaluate efficacy at maximum tolerated administered activity.
 - G. Perform cell- and organ-level dosimetry for the animal model.
- 3. Extrapolate data obtained in steps 1 and 2 to the human to arrive at initial activity for a phase I study.
 - A. Develop and fit a pharmacokinetic model to data obtained in steps 1 and 2.
 - B. Replace model parameter values with estimated human values; simulate biodistribution in humans.
 - C. Use model-derived biodistribution to estimate absorbed dose to dose-limiting organs identified in step 2B.
- 4. Assess radiopharmaceutical distribution during the phase I study.

- A. Image (if possible).
- B. Collect and count blood samples.
- C. Collect, count, and autoradiograph biopsy samples (if practical).

If there are concerns (not addressed by animal studies) about possible renal, urinary bladder wall, or gastrointestinal toxicity related to the localization of activity in luminal contents versus the organ wall:

- D. Collect and count urine samples.
- E. Collect and count fecal samples.

Steps 1-3 are general guidelines. The primary objective is to collect adequate preclinical data so as to have an understanding of the α -emitters' likely biodistribution and kinetics in humans. This objective is particularly important because pretherapy patient imaging will not be possible. It is essential that this approach not be seen as mandatory for moving α -emitter–labeled radiopharmaceuticals to the clinic; in particular, step 3 may be replaced by a projected conservative (worst-case) scenario analysis or by a direct translation of small-animal pharmacokinetics to the human using standard methods to adjust for differences in body size and organ mass (135). The autoradiography proposed in steps 2D and 4C will clearly be subject to the practical constraint of α -emitter half-life. For short-lived α -emitters, microscopic imaging of fluorescently tagged agents may be a viable alternative to autoradiography in animal models.

Conventional Versus Cell-Level Dosimetry. In most cases, a microdosimetric analysis will not be necessary for targeted therapy applications because the activity level administered and mean absorbed doses to targeted cells are larger than in the cases described here and the resulting stochastic deviation is expected to be substantially less than 20%. In such cases, standard dosimetry methods may be applied (111,136). The standard approach to dosimetry calculations has been described by the MIRD Committee (111). In this formalism, the absorbed dose to a target volume from a source region is given as the total number of disintegrations in the source region multiplied by a factor (the S value) that provides the absorbed dose to a target volume per disintegration in the source region. The sum of these products across all source regions gives the total absorbed dose to the target. MIRD cellular S values have been published for cell level dosimetry calculations for situations in which the number of disintegrations in different cellular compartments can be measured or modeled (95). With these S values, the absorbed dose to the nucleus may be calculated from α -particle emissions uniformly distributed on the cell surface, in the cytoplasm, or in the nucleus.

Conventional Dosimetry for Organs and Tumors. Estimation of the average absorbed dose to a particular normal organ or tumor volume is based on the assumption that the radioactivity is uniformly distributed in the organ and that the energy deposited by the emitted α -particles is also distributed uniformly within the organ. With some exceptions (137–141), the cross-organ dose from α -particle and electron emissions can be assumed negligible for human organ and tumor dosimetry. Care is required in applying S values for α -emitters because α -emitters may have multiple decay pathways and multiple radioactive daughters that should be considered. For example, S values for ²¹³Bi will not include the emissions from the ²¹³Po daughter, which has a 4- μ s half-life and contributes 98% of the α -particles emitted by ²¹³Bi decay (the remaining 2% come from decay of ²¹³Bi itself). This consideration and also the importance of separately accounting for absorbed dose due to electron and photon emissions from that due to α -particles requires that the dosimetry calculations be based on absorbed fraction calculations rather than on S values. The methodology is described by the following equations (presented using the recently published updated MIRD schema) (142):

$$D_{\alpha}(r_T, T_D) = \tilde{A}(r_S, T_D) \cdot \frac{\sum_i \Delta_i^{\alpha} \phi(r_T \leftarrow r_S; E_i^{\alpha})}{M(r_T)}, \quad \text{Eq. 4}$$

$$D_{e}(r_{T}, T_{D}) = \tilde{A}(r_{S}, T_{D}) \cdot \frac{\sum_{i} \Delta_{i}^{e} \phi(r_{T} \leftarrow r_{S}; E_{i}^{e})}{M(r_{T})}, \quad \text{Eq. 5}$$

$$D_{ph}(r_T, T_D) = \frac{\sum\limits_{r_S} \left(\tilde{A}(r_S, T_D) \cdot \sum\limits_i \Delta_i^{ph} \phi(r_T \leftarrow r_S; E_i^{ph}) \right)}{M(r_T)},$$
Eq. 6

$$D_{RBE}(r_T, T_D) = RBE_{\alpha} \cdot D_{\alpha}(r_T, T_D) + RBE_e$$

$$\cdot D_e(r_T, T_D)$$

$$+ RBE_{ph} \cdot D_{ph}(r_T, T_D), \qquad \text{Eq. 7}$$



FIGURE 8. Decay scheme for ²¹³Bi.

where $D_x(r_T, T_D)$ is absorbed dose to the target region, r_F , from emission type x, over the dose integration period, T_D ; $D_{RBE}(r_T, T_D)$ is RBE-weighted dose to the target region, r_T ; r_T and r_S are the target and source region (or tissue), respectively; $\tilde{A}(r_T, T_D)$ is time-integrated activity or total number of nuclear transitions in the target region, r_T ; $M(r_T)$ is the mass of the target region; Δ_i^x is mean energy emitted per nuclear transition for the *i*th emission of particle type x(alpha, electron, or photon); $\phi(r_T \leftarrow r_S; E_i^X)$ is the fraction of energy emitted per nuclear transition in the source region, r_S , that is absorbed in the target region, r_T by the *i*th emission of particle type x that is emitted with initial energy E; and RBE_{α} , RBE_e , and RBE_{ph} are RBEs for α -particles (α), electrons (e), and photons (ph), respectively $(RBE_e = RBE_{ph} = 1)$.

The total number of nuclear transitions in a particular tissue or region is typically obtained by longitudinal imaging, or counting tissue samples for radioactivity. Values for the Δ_i 's are obtained from decay-scheme tabulations published for each radionuclide (143). The absorbed fraction for each decay type, ϕ , must be calculated from tabulations of absorbed fractions for the particular tissue geometry. In almost all cases, non-cell-level dose calculations, the absorbed fractions for α -particles,

TABLE 3. Electron Emissions Considered in Absorbed Dose Calculations						
	Electrons					
	Energy	Isotope % per	Effective % per	Mean energy		Electron
Isotope	(keV)	disintegration	disintegration	(keV/disintegration)	$\Delta^{\rm e}$ (Gy-kg/Bq-s)	range (mm)
²¹³ Bi	200	0.20	0.20	0.40	6.41E-17	0.5
²¹³ Bi	347	2.55	2.55	8.85	1.42E-5	1.4
²¹³ Bi	423	0.40	0.40	1.69	2.71E-16	1.9
²¹³ Βi (β)	444	97.80	97.80	434.23	6.96E-14	2.1
²⁰⁹ ΤΙ (β)	659	100.00	2.20	14.50	2.32E-15	4.2
²⁰⁹ Pb (β)	198	100.00	100.00	198.00	3.17E-14	0.5
Sum				657.67	1.05E-13	
Mean energy and range values are listed for β-emissions. Dominant contributors to electron absorbed dose are shown in bold.						

TABLE 4. α-Particle Emissions Considered in Absorbed Dose Calculations						
	α-Particles					
	Energy	Isotope % per	Effective % per	Mean energy		α-range
Isotope	(keV)	disintegration	disintegration	(keV/disintegration)	Δ^{lpha} (Gy-kg/Bq-s)	(μm)
²¹³ Bi	5,549	0.16	0.16	8.88	1.42E-15	42.0
²¹³ Bi	5,869	2.01	2.01	117.97	1.89E-14	45.5
²¹³ Po	7,614	0.003	0.003	0.22	3.58E-17	66.0
²¹³ Po	8,375	100.00	97.80	8,190.75	1.31E-12	75.6
Sum				8,317.82	1.33E-12	

can be assumed equal to 1; the absorbed fractions for electrons are likewise usually assumed equal to 1. A description of the methods used to calculate these values is beyond the scope of this review but are provided in the references (141,144,145), one of which (141), in particular, describes absorbed fractions that are tabulated by α -particle energy for bone marrow trabeculae. For α -emitters that decay via a branched decay scheme, as in ²¹³Bi, for example (Fig. 8), it is important to account for the relative yield of each branch in determining the total energy emitted by each type of emission (i.e., the Δ_i 's). In the case of ²¹³Bi, Tables 3 and 4 summarize the electron and α -particle emissions. The tables illustrate how to tally the total electron and α -particle energy. As shown, 2.2% of ²¹³Bi decays results in ²⁰⁹Tl with the emission of an α -particle; the initial energy of the emitted α -particle is either 5.5 or 5.8 MeV, with the probability of each given by the yields shown in Table 2. In the remaining 97.8% of decays, ²¹³Bi decays to ²¹³Po with the emission of a β -particle. ²¹³Po itself decays rapidly via the emission of an 8.4-MeV α -particle to ²⁰⁹Pb, which in turn decays to ²⁰⁹Bi with the emission of a 198-keV β-particle. The exercise illustrates that a careful accounting of emissions is required in tallying the energy emitted per disintegration of the administered α -emitter, even when the decay scheme is relatively simple as for ²¹³Bi. Although outside the scope of this review, the photon S values (Table 5) can be calculated on the basis of tabulations of photon absorbed fractions to different source-target organ combinations and photon energies (146).

Units

The issue of identifying the most appropriate dosimetry quantities and units is particularly important for α -emitters because, as noted earlier, there can be confusion regarding the calculation of dosimetry quantities that relate to stochastic versus deterministic effects. It is incorrect to assign the unit sievert to the quantity defined by Equation 7. The sievert is not a unit in the conventional sense but, rather, is intended to indicate that the absorbed dose value has been scaled to reflect a biologic risk that is associated with stochastic effects. Although the product of deterministic RBEs and absorbed dose in grays has been referred to as a sievert, this is not strictly correct because sievert should be used only to designate the risk of incurring stochastic

biologic effects such as cancer. No special named unit has been widely adopted to reflect a dose value that has been multiplied by an RBE and that specifically reflects the magnitude of deterministic effects. The MIRD Committee has proposed that the barendsen (Bd) be defined as the special named unit for the product of deterministic RBE and absorbed dose and has published a commentary to this effect (*147*). To avoid confusion during the transition period, the MIRD Committee recommends that the 3 absorbed dose values, for α -, electron, and photon emissions, be provided separately and reported in the absorbed dose unit, gray. This removes any ambiguity as to interpretation of reported absorbed doses for α -emitter therapy applications.

Daughters

The example provided above is for an α -emitter with a relatively simple decay scheme. Each disintegration of the parent ²¹³Bi leads to a single α -particle emission; there are no long-lived α -emitting daughters. This is not the case for the longer-lived α -emitters ²²³Ra, ²²⁵Ac, and ²²⁷Th, which decay via α -emitting daughters. Because emission of an α -particle by the parent atom leads to a 50- to 100-nm recoil of the resulting daughter, daughter atoms may not remain conjugated to the molecular carrier. In the most complex scenario, the biologic distribution of the daughter will depend on the site of parent decay (124). In practice, the biologic distribution of long-lived daughters tends to be dominated by the chemical fate of the daughter atom. For example, ²¹³Bi, the longest-lived daughter of ²²⁵Ac, concentrates in the kidneys. Likewise, ²²³Ra, the daughter of ²²⁷Th, localizes to bone. Dosimetry calculations for such radionuclides must, therefore, account for the biodistribution of both the parent and all daughters.

TABLE 5. Individual Photon S Factors and SummedPhoton S Factor Used for ²¹³ Bi Photon Dosimetry (25)				
	Photon			
Isotope	energy (keV)	S factor (Gy/MBq-s)		
²¹³ Bi	440	5.78E-11		
²¹³ Bi	79	9.84E-13		
²⁰⁹ TI	117	1.60E-12		
²⁰⁹ TI	467	6.71E-12		
²⁰⁹ TI	1,566	2.37E-11		
$Sum = S_{WB \leftarrow WB}$		9.08E-11		

ACKNOWLEDGMENT

We thank David E. Charlton for providing guidance, vigorous discussions, and for his critical reading of the manuscript.

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