Ex Vivo Activity Quantification in Micrometastases at the Cellular Scale Using the α-Camera Technique

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Targeted α-therapy (TAT) appears to be an ideal therapeutic technique for eliminating malignant circulating, minimal residual, or micrometastatic cells. These types of malignancies are typically infraclinical, complicating the evaluation of potential treatments. This study presents a method of ex vivo activity quantification with an α-camera device, allowing measurement of the activity taken up by tumor cells in biologic structures a few tens of microns. Methods: We examined micrometastases from a murine model of ovarian carcinoma after injection of a radioimmunoconjugate labeled with 211At for TAT. At different time points, biologic samples were excised and cryosectioned. The activity level and the number of tumor cells were determined by combined information from 2 adjacent sections: one exposed to the α-camera and the other stained with hematoxylin and eosin. The time-activity curves for tumor cell clusters, comprising fewer than 10 cells, were derived for 2 different injected activities (6 and 1 MBq). Results: High uptake and good retention of the radioimmunoconjugate were observed at the surface of tumor cells. Dosimetric calculations based on the measured time-integrated activity indicated that for an injected activity of 1 MBq, isolated tumor cells received at least 12 Gy. In larger micrometastases (≤100 μm in diameter), the activity uptake per cell was lower, possibly because of hindered penetration of radiolabeled antibodies; however, the mean absorbed dose delivered to tumor cells was above 30 Gy, due to cross-fire irradiation. Conclusion: Using the α-camera, we developed a method of ex vivo activity quantification at the cellular scale, which was further applied to characterize the behavior of a radiolabeled antibody administered in vivo against ovarian carcinoma. This study demonstrated a reliable measurement of activity. This method of activity quantification, based on experimentally measured data, is expected to improve the relevance of small-scale dosimetry studies and thus to accelerate the optimization of TAT.

Key Words: 211At; radioimmunotherapy; quantification; small-scale dosimetry

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On the basis of the characteristics of the α-particle, targeted α-therapy (TAT) has been acknowledged as a potentially efficient therapeutic technique for decades. α-particles have a short range in tissue, making the irradiation specific, and a high linear energy transfer, making them highly cytotoxic. These properties make them suitable for the treatment of hematologic malignancies or minimal residual or micrometastatic diseases. The first clinical trials using TAT have been started for the treatment of leukemia (5,6), ovarian carcinoma (7), and residual tumor cells after surgery of malignant brain tumors (8).

Micrometastatic diseases are infraclinical and hardly or not localized with in vivo imaging techniques. Thus, targeted radionuclide therapies of tumor cells or micrometastases require complex evaluations. Activity quantification of micrometric lesions by imaging is difficult in patients because of the limited spatial resolution. Only rough estimations of the activity of isolated tumor cells are feasible, and they are based on assumptions of tumor burden size (5,6).

Consequently, preclinical experiments are required to assess the targeting efficiency of the radiolabeled vector and to identify the optimal conditions for treatment delivery. The first method for selecting the most promising vector for delivering α-emitters to tumors is based on in vitro measurements of the activity taken up by isolated cells (9–11). However, in vivo evaluations are complex, even in animal studies, for determining tumor uptake of radioimmunoconjugates in the context of circulating cells, residual cells, or minimal diseases. To perform accurate dosimetric studies and estimate the therapeutic window of a potential treatment, it is essential to measure the temporal distribution of activity within the micrometric lesions. To that purpose, small-scale ex vivo quantifications (i.e., assessment of the temporal distribution of the radionuclide, presumably presenting the number of radiolabeled vectors per tumor cell) can provide invaluable information.

Recently, a device based on autoradiography, the α-camera, was developed and evaluated by our group (12). This device provides images of activity distribution (of α-emitters) with good spatial resolution (around 35 μm). In the present study, we used the α-camera to develop a method to quantify the activity taken up by small micrometastases excised from a murine model of ovarian carcinoma after TAT. We hypothesized that this method would allow estimations of relevant, small-scale, mean absorbed doses and thus provide a means to confirm the delivery of potentially therapeutic absorbed doses to tumor cells.
MATERIALS AND METHODS

Overview

The general method of quantification is described in Figure 1. The samples comprised small micrometastases excised from a nude mouse model of ovarian carcinoma (OVCAR-3) after TAT. The method was developed to measure the tumor cell uptake of a monoclonal antibody (mAb), MX35, radiolabeled with the α-emitter 211At.

The excised biologic samples were snap-frozen and cryosectioned as previously described (12). Ten to 20 pairs of consecutive sections were analyzed for each sample: one section for α-camera imaging and the other section for staining with hematoxylin and eosin (H&E) (in some cases, 3 consecutive sections were analyzed [Fig. 2]). The α-camera image of radioactivity and the digitalized image of the H&E section were manually coregistered with Adobe Photoshop CS5 software. A region of interest (ROI) was manually drawn on the α-camera image with the ImageJ software (version 1.6.0_29; National Institutes of Health), as shown in Figure 1. The total pixel intensity within the ROI was quantified. A calibration curve was used to derive a coefficient that related the pixel intensity value to the activity of the α-emitter, and this was used to estimate the activity in each ROI. This value was corrected for decay as discussed below (Eq. 4).

Similarly, an ROI was drawn on the digitalized H&E section to evaluate the number of cells that corresponded to the measured activity. The number of cells in an ROI was estimated with the following equation:

\[
Nb_{Cell,ROI} = S_{ROI} \times \Delta e \times P / V_{Cell} \quad \text{(Eq. 1)}
\]

where \(S_{ROI}\) is the ROI area, \(\Delta e\) represents the tissue section thickness (12 μm), \(P\) represents the cell packing within micrometastases (\(P = 1\) was assumed), and \(V_{Cell}\) is the cell volume calculated as:

\[
V_{Cell} = \frac{4}{3} \pi R_{Cell}^3. \quad \text{(Eq. 2)}
\]

where \(R_{Cell}\) stands for the cell radius. The distribution of apparent radii of isolated OVCAR-3 cells on the H&E images was used to determine mean radius. The maximal value in this distribution was used as the mean radius (\(R_{Cell}\)).

The activity per cell (\(A_{cell}\)) at a given time after injection was derived by dividing the activity measured in the ROI of the α-camera image (\(A_{ROI}\)) by the number of cells within this part of the micrometastasis, \(Nb_{Cell,ROI}\):

\[
A_{cell} = \frac{A_{ROI}}{Nb_{Cell,ROI}} = \frac{PL_{ROI} \times Coeff_{Calib}/\frac{1}{d} - e^{-d}}{S_{ROI} \times \Delta e \times P / V_{Cell}} \quad \text{where} \ d = \lambda \times \Delta t. \quad \text{(Eq. 3)}
\]

where \(PL_{ROI}\) stands for the total pixel intensity in the defined ROI on the α-camera image, \(Coeff_{Calib}\) is the coefficient that transforms the pixel intensity into activity, \(\Delta t\) is the acquisition time, and \(\lambda\) is the decay constant of the α-emitter.

α-Camera Images

Three α-camera systems were set up in parallel. The α-camera system consisted of a cooled, high-sensitivity charge-coupled device (SXV-H9; Starlight Xpress Ltd.), coupled to a photographic lens (60 mm f/2.8 Micro-Nikkor; Nikon Corp.) (12). Specimens to be imaged (i.e., tissue sections on a scintillating sheet) were placed on an adjustable stand below the photographic lens. The distance of the specimen to the lens was adjusted to capture the entire specimen within the field of view.

A series of 600- or 1,200-s images (~10) was acquired for each group of specimens. Postprocessing was performed as described previously (12).

The coefficient for converting the pixel intensity in an α-camera image into activity was determined during the experiments. This coefficient depended on the specimen-to-lens distance. To establish the calibration curve for each camera, each sample was imaged with the α-camera at a given pixel resolution (proportional to the specimen-to-lens distance), then measured with a γ-counter (Wizard 1480; Wallac Oy). At each distance, and for numerous samples, the calibration coefficients were calculated by dividing the γ-counter–measured activity (Bq) by the total pixel intensity per second of acquisition (pixel intensity unit/s) for the whole tissue section. The calibration curves comprised the coefficients of calibration plotted as a function of the pixel resolution. Experimental points were fitted with a second-order polynomial curve.

Because the exposure times were long (~2 h), a correction was applied (Eq. 3) to the measured activity (\(A_{ROI,Meas}\)) equal to \(PL_{ROI} \times Coeff_{Calib}\) to correct for radioactive decay during the time of acquisition; that is, the activity was not constant during the acquisition period.

FIGURE 1. Illustration of different steps involved in activity quantification process.

FIGURE 2. Description of sectioning scheme performed on biologic samples. (A) Method for all biologic samples: first section was used for α-camera imaging and consecutive section for H&E staining. In some cases, 3 consecutive sections were taken (B) to evaluate accuracy of cell number determination.
was determined in this third section (Fig. 2B), was evaluated. The number of cells in each identified cluster (Fig. 2A). The size of this introduced sampling error depended on the certainty, are detailed in the supplemental materials (available online only at http://jnm.snmjournals.org).

Astatine

The α-emitter $^{211}$At was produced in a cyclotron (Scanditronix MC32; Cyclotron and PET Unit, Rigshospitalet) via the $^{209}$Bi($α,2n$)$^{211}$At reaction and isolated with a dry distillation procedure (13). The half-life of $^{211}$At is 7.2 h, and the emitted α-particles (7.45 and 5.87 MeV, with ranges in tissue of 70 and 48 μm) were generated through 2 branches of decay (58% and 42%, respectively).

mAb

The murine IgG1 class mAb MX35 (F(ab')$_2$ fragments) targets the NaPi2b cell surface glycoprotein expressed on more than 90% of human epithelial ovarian cancers (14,15).

Cells

The human ovarian cancer cell line NIH:OVCAR-3 (16) was obtained from the American Type Culture Collection. Cell cultures were maintained at the Department of Oncology at Sahlgrenska University Hospital.

Antibody Labeling

The mAb MX35 was labeled with $^{211}$At using the reagent m-MeATE (N-succinimidyl 3-(trimethylstannyl)benzoate), as previously described (17). The specific activity achieved was 650 MBq/mg (i.e., 1 of 250 mAbs labeled with an $^{211}$At atom) for all the experiments. The radiotoxicity purity was more than 99%, determined by methanol precipitation, and the immunoreactivity was about 85%.

Animals and Study Groups

Females, nude BALB/c nu/nu mice (n = 44; age range, 5–7 wk [Charles River Laboratories]) were inoculated intraperitoneally with 1 $\times$ 10$^7$ NIH:OVCAR-3 cells in 0.2 mL of cell medium. After 3–8 wk (to obtain micrometastases of variable sizes), the mice were intraperitoneally treated with 6 MBq (5.8–7 MBq) or 1 MBq (0.8–1.1 MBq) of $^{211}$At-MX35 F(ab')$_2$ in 0.75 mL of phosphate-buffered saline. For each activity (1 and 6 MBq), animals were sacrificed at 1.5 h (n = 6 + 6), 4 h (n = 6 + 6), 8 h (n = 6 + 6), and 24 h (n = 4 + 4). The spleen, stomach, liver, peritoneal wall samples, and suspected lymph nodes were excised and snap-frozen. All animal experiments were approved by the ethics committee of the University of Gothenburg.

Time–Activity Curve

The activity per cell ($A_{cell}$; Eq. 3) was derived by quantification of activity in hundreds of tumor cell clusters at 4 different time points after injection. $A_{cell}$ values were sorted into groups according to the size of the cluster, as measured on the digitalized H&E sections. Two cluster sizes were considered: those with radii less than 13.5 μm (corresponding to <5 cells in a 12-μm-thick section) and those with radii between 23.4 and 42.8 μm (between 15 and 50 cells in a 12-μm-thick section). The $A_{cell}$ versus time point curve (i.e., time–activity plot) for each group was fitted using the equation $y = (1 - \exp(-a \times t)) \times \exp(-b \times t)$. The time-integrated activity per cell was derived as the area under the fitting curve.

Dosimetry

Dosimetric calculations at the cellular level were performed to estimate the mean absorbed dose delivered to tumor cells for each injected activity (1 or 6 MBq). We used a validated code, based on analytic calculations, that is able to determine the mean absorbed doses to isolated cells or cells within spheroids, exposed to α-particles emitted from variable compartments (cell surface, cytoplasm, nucleus, extracellular domain) (18). In the case of spheroids, the contributions both from radiolabeled antibodies on the cell and from cross-fire are used to calculate the mean absorbed dose. The time-integrated activity derived above was used as an input for the algorithm. Cells and nuclei were modeled as 2 concentric homogeneous spheres of density 1 g/cm$^3$ with radii of 6.9 and 5.5 μm, respectively. We calculated the mean absorbed dose delivered to isolated cells, based on the time-integrated activity determined for clusters with a 13.5-μm radius, and the mean absorbed dose delivered to cells within a micrometastasis with a radius of 42.8 μm (modeled by a spheroid containing 200 cells), based on the number of decays per cell (which was measured experimentally for this cluster size). The spatial resolution (≈35 μm) of the α-camera images in this study did not allow for discrimination of differences in cell uptake of the radiolabeled antibody according to position in the cluster, but no cold spots were observed within identified tumor cell clusters, therefore a uniform activity distribution was assumed.

Statistical Analysis

Mean values of activity per cell were compared using a 2-sample $t$ test with the PRISM 5 software (GraphPad Software, Inc.). P values of less than 0.05 were considered significant. Data are expressed as mean ± SD.

RESULTS

α-Camera Calibration Coefficient

The calibration coefficient for the α-camera (Fig. 3) was derived by plotting the measured activity (Bq) divided by the pixel intensity per second of acquisition as a function of the relative to-lens distance. The experimental data were satisfactorily fitted to a second-order polynomial curve: $y = b_0 + b_1 \times x + b_2 \times x^2$ ($b_0 = 3.973 \times 10^{-3} ± 9.07 \times 10^{-4}$; $b_1 = -3.32 \times 10^{-4} ± 1.14 \times 10^{-4}$; $b_2 = 1.93 \times 10^{-5} ± 3.3 \times 10^{-6}$).

Accuracy of Number of Cells Determination

Figure 4 shows the distributions of $\frac{Nb_{Cells,ROI}}{Nb_{Cells,true}}$ for 2 sizes of clusters (<13.5-μm radius or 23.4- to 42.8-μm radius). The clusters with large radii showed a gaussian distribution, with a mean value of 0.96 and an SD of 0.26, indicating a relatively good
Figure 6A represents the time–activity plots (decay-corrected) of tumor cell clusters by the peritoneal surface of the spleen. The activity distribution from the α-camera images closely matched the anatomic features. α-camera images were acquired from samples of different animals, and one can notice micrometastases of various sizes.

Figure 6B represents the time–activity plots (decay-corrected) of tumor cell clusters by the peritoneal surface of the spleen. The activity distribution from the α-camera images closely matched the anatomic features. α-camera images were acquired from samples of different animals, and one can notice micrometastases of various sizes.

Here, $B_{\text{max}}$ represents the total number of antigens on the tumor cell surface, $SA$ is the specific activity of the $^{211}$At-MX35 antibody (650 MBq/mg), $M$ is the molar mass of MX35 F(ab’)$_2$ (100,000 g/mol), $N_A$ is the Avogadro constant, $r$ is the immunoreactive fraction (0.82), $A_{\text{inj}}$ is the injected activity (6 or 1 MBq), and $V_{\text{inj}}$ is the injected solution volume (0.75 mL). The fits of both curves provided a $B_{\text{max}}$ of $7 \times 10^5 \pm 1.8 \times 10^5$, the $K_{\text{on}}$ was 44,000 L mol$^{-1}$ s$^{-1}$, and the $K_{\text{off}}$ was $2.2 \times 10^{-3}$ s$^{-1}$ ± 0.67 × 10$^{-3}$. The value of $K_{\text{on}}$ was fixed as a constant in the fitting equation, based on previous in vitro measurements (19).

Figure 6B represents the time–activity plots (decay-corrected) for tumor clusters with 23.4- to 42.8-μm radii for both injected activities (1 and 6 MBq). The curve shapes were similar for large
DISCUSSION

Because of the characteristics of $\alpha$-particles, TAT is expected to be a potent treatment for circulating, minimal, and residual malignancies. The development of radioimmunoconjugates for those indications involves complex evaluations. Consequently, many evaluations rely on in vitro measurements of radiolabeled vector binding and tumor cell survival (20) or on the estimation of in vivo efficacy for different kinds of malignant lesions (e.g., subcutaneous tumors grafted on mice) (21). For ovarian carcinoma, TATs were evaluated with a model mimicking the human minimal disease (i.e., peritoneal spread of micrometastases) to determine the efficacy of $^{211}$At-MX35 and compare it with other radiolabeled specific (19) and nonspecific antibodies (22). The endpoint was the fraction of animals that presented no visible signs of tumor at 8 wk after therapy. These studies contributed to the initiation of a clinical trial (7); however, numerous questions remained unanswered and prevented proper optimization. In general, it has been difficult to measure pre- and post-therapy tumor statuses; thus, the efficacy evaluation can be misleading (23).

We demonstrate here a method using ex vivo measurements to assess activity uptake in micrometastases and even in isolated tumor cells. We showed that the time-integrated activity could be measured directly in targeted tumor clusters or tumor cells, enabling comparisons of the in vivo action of different radioimmunoconjugate regimens. Previous attempts to measure activity uptake on micrometric tumor structures with autoradiography have been confined to snapshot measurements that provided the activity distribution at only 1 time point after treatment injection (24,25). The present method showed that the $\alpha$-camera could be used to determine both spatial and temporal activity distributions, despite the need to analyze samples from different sacrificed animals.

Uncertainties

To validate this technique and evaluate its accuracy, we detailed and estimated the uncertainties associated with each step of the calculation (supplemental materials). The global relative uncertainty on the value of activity per cell was estimated to be approximately 75% for a case example of a tumor cluster of 42.8 $\mu$m in radius. We expected that an analysis of 10–15 clusters would be sufficient to achieve an acceptable confidence level (i.e., a relative uncertainty close to 20%). This process highlighted difficulties in determining the number of cells in very small tumor cell clusters (<13.5 $\mu$m in radius)—that is, close to 1 cell. The distribution of the ratio (measured value:true estimation of the value) was non-gaussian, indicating a potential systematic error. This error was not observed for large clusters. To investigate the impact of this error, we created another time–activity curve based on 2 H&E-stained sections to determine the number of cells. We analyzed several clusters at different time points ($n = 20 + 25$ at 1.5 h, $n = 30$ at 4 h, and $n = 9$ at 8 h). As expected, the activity per cell determined by this method was slightly higher when based on 3, rather than 2, adjacent sections. This difference was not significant, given the error bars. For example, from the 2 animals

FIGURE 5. Coregistration of $\alpha$-camera images and their consecutive H&E section. Bright areas represent areas of high activity. (A) Whole section of spleen with numerous tumor cell clusters for which activity uptake is significant 4 h after injection. (B) Blow up of previous image showing 4 clusters and corresponding activity spots on $\alpha$-camera image. (C) Nonvascularized micrometastasis (400-$\mu$m thick and 800-$\mu$m wide) attached to spleen. Eight hours after injection, activity distribution appeared slightly uniform. (D) Detail of section of drop of ascitic fluid containing tumor cell clusters.

and small clusters, but the uptake per cell was significantly lower for large clusters (Fig. 6B). For 6-MBq injections, the activity per cell was constant from 1.5 h ($\approx 0.027 \pm 0.010$ Bq/cell) to 4 h ($\approx 0.034 \pm 0.012$ Bq/cell), then decreased slightly at 8 h ($\approx 0.015 \pm 0.010$ Bq/cell). For 1-MBq injections, the maximum uptake was observed around 1.5 h ($\approx 0.005 \pm 0.002$ Bq/cell) and decreased thereafter.

The time-integrated activity for small tumor clusters was over 1,700 decays per cell for a 6-MBq injection, as derived from the lines shown in Figure 6A. This value decreased to 500 decays per cell for a 1-MBq injection. Using these values, we estimated the mean absorbed dose delivered to isolated cells, and they were approximately 40 and 12 Gy for injections of 6 and 1 MBq, respectively. For the large tumor clusters, around 750 and 180 decays per cell were detected for the 6- and 1-MBq injections, respectively. Assuming a uniform activity distribution in those clusters, the calculations showed that cells received between 125 and 245 Gy for the 6-MBq injections and between 30 and 60 Gy for the 1-MBq injections (with $\approx 85\%$ of the mean absorbed dose originating from cross-fire irradiation).
have performed in vivo assessments of tumor uptake. With the present method, it was possible to make experimentally based measurements and create a time–activity curve for micrometastases as small as a few tumor cells. Using this data, we derived relevant values of the mean absorbed dose delivered to tumor cells. We found good radiolabel stability for up to 8 h, with a constant number of antibodies on the surfaces of cells. At 24 h, and to a lesser degree at 8 h, the measured activity per cell declined. It was not possible to ascertain whether this decline resulted from instability of the antigen–antibody association, from conjugate instability (less probable according to in vitro results demonstrating good stability of the labeling method (17)), or from a measurement artifact (at late time points, the signal-to-noise ratio had declined on the α-camera images; thus, data were less reliable). Additionally, cell damage after 8–24 h of acute irradiation could induce measurement errors. We assume that ultrastructural changes in irradiated cells, receiving more than 10 Gy of α-particle irradiation, could affect the results derived from our method.

**Small-Scale Dosimetry**

When time–activity curves for 6- and 1-MBq injections were fitted, we found in vivo values of $B_{\text{max}}$ ($7 \times 10^5$) and antibody dissociation constant, $K_d$ ($5 \times 10^{-8}$ M). Previously (19), in vitro measurements indicated that OVCAR-3 cells possessed approximately $1 \times 10^6$ antigens, and the antibody $K_{\text{off}}$ was negligible. In the present work, the maximal uptake on tumor cells depended on the injected activity. This could argue for the possibility of antibody dissociation. From previous in vitro measurements, theoretic modeling estimated that the number of decays per cell would be more than 800 for an injected activity of 1 MBq, taking into account a specific activity of 134 MBq/mg, which was used at that time (19). Those values were previously used for small-scale dosimetry and the establishment of dose–effect curves. However, the 2 dose–effect curves reported at that time for injections of nonspecific and specific radiolabeled antibodies were different, possibly indicating an overestimation of the mean absorbed dose delivered in the case of specific antibodies (22). In the present study, the measured time-integrated activity was around 500 decays per cell for a 1-MBq injection (specific activity, 650 MBq/mg, or 4.8 times the specific activity used in previous studies). This value was 7.7 times less than the theoretically modeled value (scaling for the specific activity). On the basis of our experimental time-integrated activity, we calculated that the mean self-absorbed dose delivered to OVCAR-3 cells was around 12 Gy, a highly therapeutic dose, because a $D_0$ (the mean absorbed dose required to reduce cell survival to 37%) of 0.56 Gy was determined in vitro for OVCAR-3 cells (9).

Interestingly, clusters with radii greater than 20 μm had significantly lower uptake than smaller clusters, regardless of the injected activity. At 1.5 h, after a 1-MBq injection, the activity was less than 0.025 Bq/cell for large micrometastases and around 0.05 Bq/cell for an isolated tumor cell ($P = 0.018$). For large tumor clusters (>100 μm in diameter), slow penetration of radiolabeled antibodies through the tumors was reported (28–30). We therefore expected that the number of radiolabeled antibodies per tumor cell (after decay-correction) would increase with time. For both injected activities, this was not observed in the first 8 h after the injection. Within large micrometastases (>500 μm) containing more than 5,000 cells, the activity distribution showed no ring-effect—that is, presence of activity only at the edge of the micrometastasis—at early time points (Fig. 5C). In contrast, the distribution appeared uniform.
Despite the lower uptake per cell in the large clusters, the time-integrated activity was still high; an average of 500 decays per cell was measured within micrometastases of 23.4-42.8 μm in radii (i.e., between 15 and 50 cells in a 12-μm-thick section) for 1-MBq injections. Although activity distribution was unknown in micrometastases, the range of possible mean absorbed doses could be estimated by considering 2 opposing scenarios: a uniform distribution of activity or a skewed distribution. In the latter scenario, all the labeled antibodies (500 decays per cell × number of cells in the micrometastasis) were assumed to be attached to tumor cells at the edge of the lesion. When a uniform distribution of decays was assumed within a large micrometastasis (42.8-μm radius), the mean absorbed dose delivered to cells was between 30 (cells on the edge) and 60 Gy (cells in the core). When a skewed distribution of decays was assumed for the micrometastasis, the lowest mean absorbed dose in the tumor was 26 Gy. Again, those values are expected to be therapeutic, and they may explain the good therapeutic outcomes previously observed (22). Further studies on penetration of MX35 within intraperitoneal larger metastases are ongoing.

CONCLUSION

The present work demonstrated a method for measuring time–activity curves at the cellular scale from experimental data. Our results provided information on the binding of a specific radio-labeled antibody to ovarian carcinoma micrometastases. We measured a high activity uptake, particularly for micrometastases of only a few tumor cells. This quantitative method represents a step forward in the development of small-scale dosimetry, because further modeling was not required to derive the spatial and temporal activity distributions in micrometric biologic structures. This method is not limited to preclinical experiments; in future, it could be applied to the analysis of tumor material excised from patients undergoing surgery.

DISCLOSURE

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