Intratumoral Distribution of Fluorine-18-Fluorodeoxyglucose In Vivo: High Accumulation in Macrophages and Granulation Tissues Studied by Microautoradiography

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While 2-deoxy-2-[18F]fluoro-D-glucose ([18F]FDG) is a useful tumor imaging agent, its intratumoral distribution has not been described well at the cellular level. In order to demonstrate cellular localization of [18F]FDG and 2-deoxy-D-[3H]glucose (³H-DG) uptake by the tumor in vivo. C3H/He mice transplanted subcutaneously with FM3A tumors were studied 1 hr after intravenous injection of [18F]FDG or 3H-DG using microand macro-autoradiography. Fluorine-18-FDG and ³H-DG showed the same distribution pattern in the tumor with both autoradiographic methods. The newly formed granulation tissue around the tumor and macrophages, which were massively infiltrating the marginal areas surrounding necrotic area of the tumor showed a higher uptake of [18F]FDG than the viable tumor cells. A maximum of 29% of the glucose utilization was derived from nontumor tissue in this tumor. The comparison of double-tracer autoradiographic distribution patterns of [18F]FDG and [6-3H]-thymidine showed the differences and the similarities between glucose utilization and the DNA synthesis. Whole proliferating tissue metabolizes [¹⁸F] FDG but not vice versa. High accumulation of [¹⁸F]FDG in the tumor is believed to represent high metabolic activity of the viable tumor cells. Our results showed that one should consider not only the tumor cells proper but also the nonneoplastic cellular elements, which appear in association with growth or necrosis of the tumor cells, for precise analysis of [¹⁸F]FDG uptake in tumor-bearing subjects, especially after anti-neoplastic treatment.

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Luorine-18-2-deoxy-2-glucose ($[^{18}F]FDG$) has been shown to be a useful tumor-detecting agent (1-4). The mechanism of accumulation of this tracer into the malignant tissue is due to the enhanced rate of glucose utilization by neoplastic cells (5-7). Due to increased metabolic demand for glucose, the hexokinase (a key enzyme for glycolysis) activity is increased (8). 2-Deoxyglucose has been shown to be a substrate for hexokinase (9). Therefore, the 2-deoxyglucose analog [18F]FDG is a particularly appropriate imaging agent for tumors. Clinical applications using positron emission tomography (PET) have been reported on the correlation of the [18F]FDG uptake to: the differentiation of malignant tumor from benign (10); the grade of malignancy (11); and the proliferative activity of tumor cells (12). An autoradiographic study using x-ray films has also reported that ¹⁸F-FDG accumulates into areas of malignant growth (13). High [18F]FDG accumulation in abscesses (14,15) have also been observed. The influence of radiotherapy on [18F]FDG uptake in tumor was reported to be slower than the response of metabolic damage of amino acid and DNA incorporation/synthesis (16). In this study, we report a unique uptake pattern of [¹⁸F]FDG and 2-deoxy-D-[³H]glucose (³H-DG) by the complex of heterogeneous cell elements in a malignant tumor model in mice.

MATERIALS AND METHODS

Animal Model

Male 10-wk-old C3H/He mice were subcutaneously injected with a 0.1-ml suspension of 7×10^6 syngeneic FM3A tumor cells on their left thighs. Solid tumors produced on their thighs were measured with vernier calipers and the product of three principal diameters of the tumor was designated as "tumor volume" as described previously (17). A tumor growth curve until death was obtained as the average of data from six mice. Tracer experiments were performed 10 days after transplantation.

Double-tracer Macro-autoradiography with [¹⁸F]FDG and ³H-Thd

A C3H/He mouse with FM3A tumors was injected with a mixture of 200 μ Ci (7.40 MBq) of [¹⁸F]FDG and 30 μ Ci (1.11

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MBq) of $[6^{-3}H]$ thymidine (³H-Thd, Amersham Japan Ltd., Japan) and killed 1 hr later. The tumor was quickly dissected and prepared for frozen sectioning as reported previously (18). In brief, the trimmed tumor sample was embedded in O.C.T. compound (Miles Inc.) and frozen in isopentane cooled with liquid nitrogen. The frozen sample block was sectioned on a cryostat at -25° C. The 5 μ m-thick section was mounted on a clean glass slide, air-dried and directly contacted with ARG film (MARG ³H-type, Konica, Japan) for 2 hr in order to produce the [¹⁸F] FDG image. Three days (about 39 half-lives of ¹⁸F) after the initial autoradiography when the ¹⁸F radioactivity had decayed, the same section was contacted with another film for 2 mo to produce the ³H-Thd image.

Micro-autoradiography

Sixteen C3H/He male mice were transplanted subcutaneously with FM3A tumor cells. Approximately 10 days following transplantation, four mice were injected intravenously with 1 mCi (37 MBq) of [18F]FDG, four with a mixture of 1 mCi (37 MBq) of [¹⁸F]FDG and 30 μ Ci (1.11 MBq) of ³H-Thd, and eight with 30 μ Ci (1.11 MBq) of ³H-DG (Amersham Japan Ltd., Japan). They were killed 1 hr later. The tumors were quickly dissected and prepared for frozen sectioning as described above. Under a safety light, the 5 µm-thick sections were directly mounted on slides coated with NTB2 nuclear emulsion (Kodak) which was then cooled to -15° C. The slides were immediately deep-frozen on a dry-ice block and placed in exposure boxes cooled with dry-ice. After a 4-hr exposure for [18F]FDG and a 3-wk exposure for ³H-DG, they were transferred to ethanol with 5% acetic acid at -70°C and 18.5°C for 1 min each. After washing off the excess acid ethanol in water for 2 min, autoradiograms were developed in Konidol-X (Konica, Japan) for 4 min, fixed in Fuji general purpose fixer (Fuji, Japan) for 8 min and washed in gently running water for 30 min at 18.5°C. The slides were stained with hematoxylin and eosin. Nonradioactive tumor sections were subjected to the same process for chemographic control. Silver grain numbers were counted in various tumor regions under a transmitted light brightfield microscope using a micrometer. The contamination of [18F]FDG microautoradiogram from simultaneously injected ³H-Thd was a maximum of 0.8% during the 4 hr of exposure in a separate study, and thus was neglected.

The contiguous 5 μ m-thick sections from the same frozen blocks used in the micro-autoradiographic study were mounted on clean glass slides, air-dried and directly coated with ARG films for 30 min to 1 hr in order to image [¹⁸F]FDG distribution, and for 2 mo to image ³H-DG distribution. The exposure duration for the [¹⁸F]FDG autoradiogram was adjusted, depending on the starting time of exposure in consideration of [¹⁸F]FDG decay. After the [¹⁸F]FDG radioactivity had decayed, the sections labeled with simultaneously injected ³H-Thd were coated with ARG films for 2 mo to image ³H-Thd distribution. After this exposure, the sections were stained with hematoxylin and eosin. The micrographs were illustrated and the areas of major cell elements in each tumor were measured.

Light Microscopic Examination

The FM3A tumor tissue was fixed in a conventional 10% neutral-buffered formalin, dehydrated in graded alcohol and finally embedded in paraffin. Thin histologic sections were stained with hematoxylin and eosin using the routine staining procedure.

Quantitative Analysis

To determine the relationship between grain numbers and radioactivity, normal male C3H/He mice were injected intravenously with [¹⁸F]FDG in different doses of 0.05 to 2.3 mCi (1.85 to 85.1 MBq) and were killed 1 min later. The 5 μ m liver sections were then processed for ARG as uniform step-wise standards of radioactive sample, counterstained with eosin, and the grain numbers per unit area were counted under the transmitted light brightfield microscope using micrometer. Other 5 μ m sections of the same samples were attached to thin polyethylene films, air-dried and cut into 6-mm diameter circles with a punch. The radioactivity in the punched out section was then measured with a gamma-counter. Cross calibration between the gamma counter and a well-type dose meter was performed for ¹⁸F. The radioactivity per unit area of the section then was calculated and corrected for decay (18).

The animals used in this study were maintained in the animal care facility of our institution and the study protocol was approved by the laboratory animal care and use committee of Tohoku University.

RESULTS

Tumor Growth Curve

The tumor growth curve of subcutaneously transplanted FM3A tumor cells on the thighs of 6 mice is shown in Figure 1. The in vivo doubling time of the tumor was about 4.4 days at the tracer experiment (10 days after transplantation). The mean survival time of the mice after transplantation was 41.8 ± 3.2 days.

Standard Curve

A standard curve relating grain numbers (grains/100 μ m² in 5 μ m-thick sections) to [¹⁸F]FDG radioactivity (fCi/100 μ m² in 5 μ m-sections) is shown in Figure 2. A linear relationship between the number of grains (y, grains/



FIGURE 1. A FM3A tumor growth curve represented by daily average of every tumor measurement along the survival time of six mice. Day 0 is the day when the tumor cells were transplanted. Tumor volume of less than 100 mm³ was technically unmeasureable in vivo. The tracer study was performed on Day 10 when the mean tumor volume was 506 mm³ and the mean volume doubling time was 4.4 days.



FIGURE 2. A standard curve of the relationship between silver grain numbers and ¹⁸F radioactivity in micro-autoradiography. In the experimental dose range, the grain numbers were linearly related to the ¹⁸F radioactivity. About 8000 grains over 1200 microgrid areas were counted for this study.

100 μ m²) and the corresponding [¹⁸F]FDG radioactivity (x, fCi/100 μ m²) was observed (y = 0.4171x + 0.3538, r = 0.9996, p < 0.001). Background grain numbers (1.24 ± 0.41 grains/100 μ m²) were constant for all the experimental doses. Most of the micro-autoradiographic studies were performed within the confirmed range of linearity.

Double-tracer Autoradiograms with [¹⁸F]FDG and ³H-Thd

Figure 3 shows the autoradiograms of a FM3A tumor section 1 hr after the injection of a mixture of 18 F and 3 H. There were different heterogeneous distribution patterns of silver grains within the tumor which could not be explained by the resolution differences of [18F]FDG and ³H. The [¹⁸F]FDG autoradiogram shows the high density of the tumor rim, and the accentuated dense region within the tumor corresponds to the translucent area on the ³H-Thd autoradiogram. The dense area on the ³H-Thd autoradiogram is consistent with the proliferating cell region by the 1-hr labeling index of DNA synthesis during mitotic cycle, but the dense area on the [¹⁸F]FDG autoradiogram is distributed on the whole section by glucose utilization. Both autoradiograms showed the scarce density on the normal host tissue. The differential uptake ratio of this tumor for $[{}^{18}F]FDG$ was 1.368 ± 0.286 (n = 8), and the tumor-to-blood and tumor-to-muscle ratios were 10,523 and 4.035. The whole area with the grain density of ³H-Thd has the grain density of [18F]FDG but not vice versa, which represents the consistency and the difference be-



FIGURE 3. A combination of double-tracer macro-autoradiograms and microscopy. Images of [¹⁸F]FDG distribution (A) and ³H-Thd (B), a photomicrograph of the specimen (C) which produced the autoradiograms and the illustration of the micrograph (D). The contamination of the [¹⁸F]FDG autoradiogram from ³H-Thd was undetectable and that of ³H-Thd autoradiogram from ¹⁸F-FDG was zero. The mean 1-hr labeling index of the proliferating tumor cells by ³H-Thd was 27.9% in a separate micro-ARG study. T = tumor cells; G = granulation tissue; N = necrosis; H = host normal tissue. Scale bar: 2 mm.

tween glucose utilization and the DNA synthesis in the tumor tissue. For histological confirmation, the following microautoradiography procedure was performed.

Macro- and Micro-autoradiography

The macro-autoradiograms of [18F]FDG and 3H-DG by contiguous sections for micro-autoradiography showed a similar pattern of a heterogeneous grain density in comparison with portions of the tumor mass (Figs. 4 and 5). In the autoradiograms of both [¹⁸F]FDG and ³H-DG, a markedly dense area surrounded the tumor itself and the central translucent area. The other dense spots were located in an area of relatively lower density. The central grain-free area was microscopically confirmed to be extensive necrotic tissue in the tumor. The autoradiograms of ³H-Thd showed a different pattern from those of [¹⁸F]FDG and ³H-DG. The ³H-Thd autoradiograms showed another heterogeneous grain density pattern in the tumor mass. The dense areas on the ³H-Thd autoradiograms were observed in the tumor cell region and were not always consistent with those on the [18F]FDG autoradiograms.

A combination of micro-autoradiography of [¹⁸F]FDG and ³H-DG and light microscopy revealed that the both tracers showed biologically the same distribution pattern, although the grain sizes and numbers were different due to the physical nature of ¹⁸F and ³H. The highest grain concentration was observed on macrophages which were massively infiltrating the marginal areas surrounding extensive tumor necrosis (Fig. 6A–C). Spotty dense areas within the tumor also corresponded to areas of focal necrosis where macrophages were massively pooled (Fig. 7A–C). Whether necrosis was diffuse or focal, the grains concentrated where macrophages were present.

Young granulation tissue consisting of capillary vessels, fibroblasts and macrophages surrounded the tumor mass (Fig. 6C). The second highest grain level was found in this tissue. Within the granulation tissue, the grains were densely concentrated where macrophages were present (Fig. 8). The grains were also localized diffusely on viable tumor cells, but grain concentrations were relatively lower than those on the macrophages or the granulation tissue (Fig. 7).

Tables 1 and 2 show the results of grain counting for [¹⁸F]FDG and ³H-DG micro-autoradiographs. The tumor numbers in the tables correspond to Figures 4 and 5. The macrophage layer surrounding extensive tumor necrosis showed the highest value, which was about 3.5–3.6 times that of the tumor cells. The second highest was the young granulation tissue with capillary vessels, fibroblasts and macrophages surrounding the tumor mass, which showed



FIGURE 4. Macro-autoradiograms of [¹⁸F]FDG distribution in contiguous sections for micro-autoradiography. The ³H-Thd autoradiograms of four tumors (No. 1–4), which were labeled with [¹⁸F]FDG and ³H-Thd simultaneously, are also shown. For a simple and more informative comparison, the micrographs were added. See Figure 3 for definitions. Scale bar = 5 mm.



FIGURE 5. Macro-autoradiograms of ³H-DG distribution in the contiguous sections for micro-autoradiography and the micrographs. See Figure 3 for definitions. The pattern of ³H-DG autoradiograms was similar to that of ¹⁸F-FDG autoradiograms, but was different from that of ³H-Thd autoradiograms in comparison with Figure 4. Scale bar = 5 mm.

about a 2.4–2.5 times greater value than tumor cells followed by the necrotic area with macrophages, which showed about a 2.0–2.3 times greater value than tumor cells. The necrotic area with neutrophils, which were one of major phagocytes, as well as macrophages showed the lowest value. Grain counts for both [¹⁸F]FDG and the ³H-DG autoradiography showed the same pattern. Semiquantitative morphometry of the micrograph is included in Tables 1 and 2. The macrophage layers were about 4.14%-4.38% and the granulation tissues were 3.55%-5.47% of the total area. The macrophage pooled areas were only a part of necrosis. It shows that about 23.85% ($4.38\% \times 3.5$ + $3.55\% \times 2.4$) of radioactivity for [¹⁸F]FDG and 28.59\% ($4.14\% \times 3.6$ + $5.47\% \times 2.4$) of that for ³H-DG (no



FIGURE 6. Micro-autoradiography of [¹⁸F]FDG (A) and ³H-DG (B). This region is the marginal areas surrounding extensive tumor necrosis near the outer surface of tumor. It consists of three layers of different cell elements: granulation tissue (G), macrophage layer (M) and necrosis (N) in order from left to right. Macrophages are observed between necrosis and granulation tissue, and the highest grain density is found in this region. Macrophages are also pooled in the necrosis and grains were distributed in this zone. Histology of the periphery of tumor tissue (C) clearly demonstrated that the macrophages, neutrophils and granulation tissue are the major components between the necrosis and the surrounding intact host tissues. Scale bar = 50 μ m.



FIGURE 7. Micro-autoradiography of [¹⁸F]FDG (A) and ³H-DG (B). This region has a focal necrotic area with a relatively high grain density level within the tumor. Macrophages are observed among the necrotic tissue. The tumor cells around this focal necrosis showed low grain levels. Histology of the focal necrotic area (C) demonstrated that the necrosis is composed of a mixture of macrophages, neutrophils and necrotic degenerative tumor cells. The tumor cells around this area are viable. T = tumor cells and N = necrosis. Scale bar = 50 μ m.

significant difference) in the whole section were derived from macrophages and granulation tissues.

Micro-autoradiographic grains were seen in all tumor tissue sections of [¹⁸F]FDG injected mice, and no grain was observed in the control sections, thus ruling out the possibility of a positive chemogram.

DISCUSSION

The results can be summarized as follows: [¹⁸F]FDG accumulates not only in the tumor cells but also in the inflammatory cell elements which appear in association with growth or necrosis of tumor. Accumulation of [¹⁸F] FDG is relatively higher in macrophages and young granulation tissue than in the tumor cells. Macrophages are generally concentrated in the outer zones of necrosis, and granulation tissue is formed around the tumor, demarcating it from the surrounding intact host tissue. This demarcation seems to give a suitable explanation for the unique macro-autoradiographic pattern of tumor.

However, these findings do not mean that the inflammatory cells are the major source of radioactivity of [¹⁸F] FDG in tumor tissue. Despite more active uptake of [¹⁸F] FDG by inflammatory cells within or around the tumor, it is reasonable to consider that most of the radioactivity of [¹⁸F]FDG in the whole tumor originates from viable tumor cells because they consist of the major component of the tumor mass.

Clinical studies with PET report high $[^{18}F]FDG$ uptake in various malignant tumors, such as gliomas (3), hepatoma (19), lymphoma (20) and lung cancer (21). Minn et al. (12) have assessed [¹⁸F]FDG accumulation in malignant tumors with PET and correlated proliferative activity of the biopsied samples with DNA flow cytometry. These authors observed a clear correlation between the proportion of the cells in S + G₂/M phases of cell cycle or the percentage of S-phase cells and [¹⁸F]FDG accumulation.

Fluorine-18-FDG also accumulates in abscesses (14,15). This phenomenon can be explained by the active uptake of [¹⁸F]FDG by phagocytes within the abscess or by granulation tissue surrounding the abscess. Interestingly, chemically produced sterile abscesses do not accumulate [¹⁸F]FDG (1). This may be due to a different inflammatory response from that of infectious abscesses.

In the post-radiotherapy study of rat AH109A tumor model, [¹⁸F]FDG accumulation by the tumor showed a slower decrease than ¹⁴C-methionine and ³H-thymidine uptakes (*16*), and the reduction patterns of [¹⁸F]FDG uptake have been similar to that of ⁶⁷Ga-citrate, which is also considered to accumulate in phagocytes. In one PET study, the high [¹⁸F]FDG uptake by colon cancer was reported to remain for a few months after radiotherapy and has been explained by the inflammatory reaction in the region (*22*).

The tumor model used in this study was subcutaneously transplanted malignant tumors. The experimental findings may not be directly applicable to human tumors because tissue response to the invading neoplasms is considered to be different in various organs, especially in the brain (23). However, this study suggests that one should bear in mind



FIGURE 8. Micro-autoradiography of [¹⁸F]FDG (A) and ³H-DG (B). This region has young granulation tissue formed around the tumor. Macrophages were present where the grains were densely concentrated. The second highest grain density is found in this region. Scale bar = 50 μ m.

Tumor		Ν	mber of grai	ns / 100 Jm ²			Ratio	s relati	ive to T	umor Ce	lls	Area r whole	atio† tissue	to (\$)
*.0N	ħ	G(fb)	G(fb,mp)	Σ	(dm) N	(du)N	G(fb)	G(fb,mF	W (0	(dw)N	(du)N	U	Σ	z
1	7.4±4.6	17.3±8.2	17.4± 6.9	24.0± 6.7	20.8± 6.4	3.5±2.9	2.34	2.35	3.24	2.81	0.47	4.75	5.23	26.96
2	12.5±3.4	14.1±4.6	30.9± 9.8	43.3±16.2	26.6± 8. 1	6.1±3.7	1.13	2.47	3.46	2.13	0.49	0.89	3.75	33.69
e	10.5±3.9	16.1±6.9	28.5± 8.4	35.7±11.7	20.9± 4.5	7.3±1.9	1.53	2.71	3.40	1.99	0.70	6.53	4.49	25.80
4	9.8±4.9	14.2±5.9	ı	32.7±22.0	26.4±16.0	9.2±3.8	1.45	ı	3.34	2.69	0.94	0.08	4.01	12.52
2	11.7±5.8	20.6±8.3	28.7±12.2	42.4±13.2	23.8±13.5	6.0±3.1	1.76	2.45	3.20	2.03	0.51	2.61	4.78	18.16
9	10.9±9.5	17.5±5.6	25.9± 8.1	42.4±10.5	24.1±13.5	4.7±2.5	1.61	2.38	3.89	2.21	0.43	8.96	4.14	22.31
7	6.9±3.2	20.2±7.0	19.4± 9.9	23.2± 4.2	19.0±14.4	6.4±3.3	2.93	2.81	3.36	2.75	0.93	2.29	2.89	7.61
80	12.1±6.7	19.3±4.0	19.9± 4.0	45.6±10.3	23.3± 5.0	8.4±2.4	1.60	1.64	3.77	1.93	0.69	2.37	5.72	26.42
Mean	10.2±2.1	17.4±2.5 [†]	24.4± 5.4 [‡] '	¶ 36.2± 8.8 [‡]	23.1± 2.7 [‡]	6.5±1.9 [§]	1.79	2.40	3.48	2.32	0.65	3.55	4.38	21.68
s.D.							0.57	0.38	0.26	0.37	0.20	3.00	0.88	8.51
T: ti	umor cells,	, G(fb):	granulation	tissue with :	fibroblasts,	G(fb,mp):	granul	lation t	issue w	ith fib	roblasts	and	nacroph	ages,
ŭ W:	acrophage lé	ayer, N(mp): macrophage	<pre>pooled zone</pre>	in necrosis,	N(np): ne	utrophil	l pooled	l zone i	n necro:	sis, G:	granula	tion ti	ssue,
N: N	ecrosis. ł	Each grain	count data w	∕as mean ± S.I	D. of 8 to 1	12 various 1	points i	in the t	issue.	Five 1	to 15 mi	crogrid	areas	were
counte	ed and ave	eraged for ϵ	each point.	*Tumor numbei	r corresponds	to Figure	4.	The pla	nimetry	was pei	rformed	on the	conti	snonɓ
sectic	on used for	macro-auto	radiography.											

TABLE 1 Results of Microautoradiographic Distribution of [¹⁸F]FDG in FM3A Tumor Tissues

 $\ddagger:$ p<0.001, §: p<0.005 compared to tumor cells.

 \P : p<0.01 compared to macrophages.

TABLE 2 Results of Microautoradiographic Distribution of ³H-DG in FM3A Turnor Tissues

Tumo	ų	Nu	mber of grai	ns / 100 Jm ²			Ratio	s relati	ve to 1	'umor Ce	lls	Area I whole	atio† tissue	to (\$)
No.*	F	G(fb)	G(fb,mp)	Σ	N (mp.)	N(np)	G(fb)	G(fb,mp	W	(dm) N	(du)N	υ	Σ	z
-	48.3±13.4	ł	114.7±41.2	159.0±16.0	63.2±6.5	1	1	2.37	3.29	1.31	I	1.35	5.22	18.52
7	35.6±21.4	33.4±9.4	105.2±27.8	138.3±32.8	64.3±16.7	23.8±12.5	0.94	2.96	3.88	1.81	0.67	6.70	5.39	27.61
٣	38.5±19.2	I	26.6±6.9	139.0±60.4	139.5±55.6	25.7± 4.6	ı	0.69	3.61	3.62	0.67	3.88	6.93	21.30
4	41.3±21.8	71.0±30.6	119.4±13.6	131.5±65.5	109.7±20.2	16.1± 3.2	1.72	2.89	3.18	2.66	0.39	5.70	3.25	20.09
ŝ	42.5±17.8	ı	99.0±16.5	165.7±57.4	66.2±16.4	27.4± 5.7	ı	2.33	3.90	1.56	0.64	1.76	3.49	17.97
و	35.0± 9.4	46.0±20.2	63.0±10.0	137.4±28.6	58.0±17.2	19.0± 8.3	1.31	1.80	3.93	1.66	0.54	3.53	2.73	30.40
٦	41.5±17.9	70.5±22.0	122.0±32.8	143.4±26.7	73.0±12.2	30.2±16.3	1.70	2.94	3.46	1.76	0.73	9.22	3.23	15.23
80	41.7±11.5	75.2±32.7	176.7±44.3	161.4±42.7	67.7±26.2	22.4± 1.5	1.80	4.24	3.87	1.62	0.54	11.60	2.91	31.67
Mean	40.6± 4.2	59.2±18.5†	[:] 103.3±44.2 ⁵	¶147.0±13.0 ^{#:}	* 80.2±28.8 ⁵	23.5±4.9 ^{**}	1.50	2.53	3.64	2.00	0.60	5.47	4.14	22.85
s.D.							0.36	1.03	0.30	0.76	0.12	3.59	1.51	6.19
ц.	tumor cells	s, G(fb):	granulation	tissue with	fibroblasts,	G(fb,mp):	granu	lation t	issue w	vith fit	roblast	s and	macrop	hages,
Μ:	macrophage 1	layer, N(mp): macrophag	e pooled zone	e in necrosis	, N(np): neu	itrophi	l pooled	zone i	in necro	sis, G:	granul	ition t	issue,
 N	necrosis.	Each grain	l count data	was mean ± S.	.D. of 8 to	12 various p	oints	in the t	issue.	Three	e to 7 m	icrogric	l areas	were
coun	ted and av	veraged for	each point.	*Tumor numb	oer correspon	ds to Figure	e 5.	[†] The pl	animetr	y was p	erforme	d on the	e cont	iguous

 $\frac{1}{2}; \ p.0.02; \ \ \frac{9}{5}; \ p<0.002, \ \ \text{**:} \ p<0.001 \ \ \text{compared to tumor cells.}$

section used for macro-autoradiography.

 $\P\colon p{<}0.02$ compared to macrophages.

that higher tumor uptake may reflect not only tumor cell viability and proliferation but also the contribution of secondary inflammatory reaction elements. Further micro-autoradiographic studies with [¹⁸F]FDG will be of help in understanding PET's role in various clinical problems.

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