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In vivo $^{31}$P magnetic resonance spectroscopy and morphometric analysis of the perfused vascular architecture of human glioma xenografts in nude mice

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Summary The relationship between the bioenergetic status of human glioma xenografts in nude mice and morphometric parameters of the perfused vascular architecture was studied using $^{31}$P magnetic resonance spectroscopy (MRS), fluorescence microscopy and two-dimensional digital image analysis. Two tumour lines with a different vascular architecture were used for this study. Intervascular distances and non-perfused area fractions varied greatly between tumours of the same line and tumours of different lines. The inorganic phosphate–nucleoside triphosphate (P/NTP) ratio increased rapidly as mean intervascular distances increased from 100 μm to 300 μm. Two morphometric parameters – the percentage of intervascular distances larger than 200 μm (ivd$_{200}$) and the non-perfused area fraction at a distance larger than 100 μm from a nearest perfused vessel (area$_{100}$) – were deduced from these experiments and related to the P/NTP ratio of the whole tumour. It is assumed that an aerobic to anaerobic transition influences the bioenergetic status, i.e. the P/NTP ratio increased linearly with the percentage of ivd$_{200}$ and the area$_{100}$.

Keywords: $^{31}$P magnetic resonance spectroscopy; fluorescence microscopy; tumour bioenergetic status; vascular morphology

Phosphorus magnetic resonance spectroscopy ($^{31}$P-MRS) studies of the bioenergetic status of tumours have been compared directly with tumour tissue $p_{O_2}$ (Vaupel et al, 1989a) and with several physiological parameters that affect $p_{O_2}$ in tumour tissue, such as the intravascular concentration of oxyhaemoglobin (Rofstad et al, 1988) and tumour blood perfusion (Evelhoch et al, 1986). $^{31}$P-MRS has been demonstrated to be sensitive to tumour size (Okunieff et al, 1986; Wendland et al, 1992), with an increase in tumour size being supposed to have a negative influence on perfusion- and diffusion-limited oxygen delivery and nutrient delivery (Vaupel, 1996). The global bioenergetic status of a tumour, expressed as the P/NTP ratio, depends on the balance between the oxygen and nutrient supply, and the consumption rates of the tumour cells, which is related to the type of energy metabolism. This balance between supply and consumption determines the critical diffusion distances for oxygen and nutrients in tumour tissue. The total oxygen supply and nutrient supply depends on the tumour microcirculation and on the diffusion geometry.

The purpose of this study was to investigate the existence of a possible relationship between the global bioenergetic status and the diffusion geometry of human glioma xenografts in nude mice. A two-dimensional morphometric analysis of the perfused vascular architecture of complete transverse tumour sections was performed. Morphometric parameters, such as the percentage of large intervascular distances and the fraction of non-perfused areas in a tumour section area, are related to the diffusion-limited oxygen delivery and nutrient delivery. Critical diffusion distances for oxygen and nutrients per perfused vessel can be estimated using a Krogh model (Vaupel, 1974; Kreuzer, 1982; Kallinowski et al, 1987; Groebe et al, 1988; Vaupel et al, 1989b; Dewhirst et al, 1994). For example, the mean critical oxygen diffusion distance in gliomas using a Krogh model is approximately 100 μm. When tumour cells have an aerobic energy metabolism, changes in the percentage of intervascular distances larger than approximately 200 μm are expected to affect the global bioenergetic status (P/NTP ratio). This indeed was observed. In addition, the mean fraction of the non-perfused tumour area at a distance larger than approximately 100 μm from the edge of the nearest perfused vessel was also found to be related to the global bioenergetic status of a tumour.

MATERIALS AND METHODS

Animal model
Two tumour lines (E49, n = 14; E98, n = 10), derived from two different primary human gliomas, were grown subcutaneously in the hind limb of athymic mice (Balb/c nu/nu, BonholdGard, Denmark) after several passages in the flank of nude mice. During the $^{31}$P-MRS experiments, motion artefacts were less important for tumours on the hind limb than for tumours on the flank. The experimental procedures were approved by the local ethics committee for animal use.

In vivo $^{31}$P magnetic resonance spectroscopy
MRS measurements were performed on a vertical-bore Bruker spectrometer (4.7 T) using a home-built $^1$H/$^{31}$P double-tunable three-turn solenoid coil with an inner diameter of 13 mm. The solenoid coil was fitted with a Faraday shield to eliminate spurious...
signals from normal tissue adjacent to the tumour. Mice with human glioma xenografts were excluded if host tissue was partially present in the volume sampled by the solenoid (tumour weight < 0.3 g) and when tumours were too large to fit completely in the solenoid (tumour weight > 0.9 g).

The mice were anaesthetized with a flow of 1.5% enflurane in an oxygen-nitrous oxide (3:7) mixture applied through a nose cone. Body temperature was monitored by a rectal probe (36-gauge wire, Hewlett Packard) and maintained at 36.5-37°C by a warm-water blanket with a feedback system. 31P-MR spectra were obtained with a one-pulse sequence with a hard pulse of 12 μs (optimized for maximum signal intensity) and a pulse repetition time of 5 s. The number of scans was 320.

As the 31P-MRS experiments were carried out with an interpulse delay shorter than three times the T1 of the 31P spins of the P (i.e. approximately 4 s), the area of the P peak is not strictly proportional to its concentration (Certaines et al, 1993). However, all in vivo spectra were run with the same acquisition parameters and in this group of tumours little effect is expected on the calculations of the P1/NTP ratio at a pulse repetition time of 5 s.

Fluorescence microscopy

After the MRS experiments, 0.05 ml of phosphate-buffered saline (PBS, pH 7.4) containing a fluorescent perfusion marker, Hoechst 33342 (15 mg kg−1, Sigma, St. Louis, MO, USA), was injected i.v. via a lateral tail vein. One minute after injection, the mice were killed and tumours were quickly removed and frozen in liquid nitrogen, preventing the dye from diffusing too far into tissue. The tumour was cut in two halves: one half was used for the analysis of the perfused vascular architecture and the other half was used for classical histological staining with eosin (cytoplasm) and haematoxylin (nuclei).

Fifteen frozen tissue sections (5 μm) at random locations were made using a freeze microtome. Sections were processed at room temperature by a 15-min incubation with collagen type IV polyclonal antibody (rabbit serum, Euro Diagnostica, Oss, The Netherlands), a marker for the basal lamina of the tumour tissue. Next, the sections were incubated with a second antibody, goat anti-rabbit immunoglobulin labelled with TRITC (Tago, Burlingame, CA, USA).

In this paper any vascular structure, including arterioles, venules and capillaries, in a tumour tissue section stained by collagen type IV antibody is described as a vessel. The whole-tumour sections were analysed in the fluorescence microscope using a digital-image processing system. A detailed description of this method is given by Rijken et al (1995). Briefly, each section was scanned twice on the computer-controlled motorized stage of a fluorescence microscope using two different excitation–emission filters. After processing all fields of each scan, a composite image.
was reconstructed from the individual processed fields, revealing the perfused vessels (Hoechst image) and the total vascular bed (collagen image) in separate scans. When both images were combined, the new matched image showed the perfused and non-perfused vessels. In the next step, the fluorescent rim of Hoechst dye around perfused vessels, due to Hoechst diffusion into adjacent tissue, was deleted by image processing. In Hoechst images only, vascular areas are slightly overestimated.

Data analysis

$^3P$-MR spectra

Zero filling and the convolution difference technique with line broadenings of 30 and 1000 Hz were applied to the free induction decay (FID). The peaks of the $\alpha$, $\beta$, $\gamma$-NTP, $P$ and, when present, phosphocreatine (PCr) were fitted to Lorentzian line shapes with NMR1 software (New Methods Research, Syracuse, NY, USA). The integral of the $P$ peak and the sum of the integral of the $\alpha$, $\beta$, $\gamma$-NTP peaks were used in the calculation of the $P/NTP$ ratio.

Calculation of the $pH_{v_{\text{org}}}$

The pH was deduced from the chemical shift of the $P$ signal with respect to the chemical shift of the PCr signal, or the $\alpha$-NTP signal in the absence of a PCr resonance. A modified Henderson-Hasselbach equation was used, with the following parameters: pK$_{\alpha}$ 6.75, a (acid shift) = 3.29, b (base shift) = 5.7 (Moon et al, 1973; Seo et al, 1983).

Vascular morphology of perfused vessels

The percentage of intervascular distances $> 200 \mu m$ (ivd$^{200}$)

For each perfused vessel, a domain (Yoshii et al, 1988), i.e. the area of tumour tissue that is supposed to be supplied by the nearest perfused vessel, was determined in matched images. As a consequence, one domain contains one perfused vascular structure, and may contain non-perfused vascular structures and avascular regions. With the help of an image analysis system, contours of these domains were represented by line networks in Figure 1. The shortest distance between neighbouring perfused vascular structures was used as an estimation of the ivd. Note that neighbouring perfused vascular structures have adjacent domains. Thus, intervascular distances were not determined between perfused vascular structures that did not have adjacent domains. Calculations of distances were started from perfused vessel walls. Ivds obtained by a domain analysis are always larger than ivds between perfused vessels, measured in perfused regions only (Less et al, 1991). For each tumour, the frequency distribution of the ivds was determined for all values calculated in the 15 tumour sections (Statistica, StatSoft, Tulsa, OK, USA). The percentage of intervascular distances $> 200 \mu m$ (ivd$^{200}$) was obtained from the cumulative frequency distribution for the whole tumour.

The fraction of the non-perfused tumour area at a distance larger than 100 $\mu m$ from the nearest perfused vessel (area$^{100}$)

In matched images, a circle with a radius of 100 $\mu m$ was drawn around every perfused vessel. For each tumour section, the tumour area outside the circles was determined and divided by the total tumour section area. Next, a mean non-perfused area fraction at a distance $> 100 \mu m$ from the nearest perfused vessel was calculated for all 15 tumour sections.
The morphological parameter – the percentage of ivd\textsubscript{200} – is probably less sensitive to the total non-perfused area than the parameter area\textsubscript{100}. Only a few long ivds (> 200 \textmu m) may be responsible for the determination of a large non-perfused area. In other words, the weight of a few long ivds in comparison with all ivds is of less importance than the weight of the non-perfused areas in relation to the total tumour area.

Analysis of the relationship between the P/NTP ratio and the morphometrical parameters

Linear regression analysis was performed between the morphometrical parameters, mentioned above, and the P/NTP ratios using Graphpad (Graphpad PRISM version 2.0, San Diego, USA). The goodness of the fit ($R^2$), the 95% confidence intervals and the $P$-value of the slope are given, i.e. test result of the significant difference of the slope from zero. Values presented in the text are means ± s.d.

RESULTS

Comparison of vascular morphometrical analysis of perfused vessels between the tumour lines (E49, E98) and host tissue

In Figure 2, a frequency histogram of the ivds is given for a single tumour and host tissue, i.e. skeletal muscle of the hind limb. Tumour tissue showed an important tail of long ivds in comparison with host tissue. Ivds larger than 200 \textmu m were not observed in host tissue; thus, the percentage of ivd\textsubscript{200} = 0. The mean ivd of host tissue was 35 ± 21 \textmu m, which was much smaller than the mean ivds per tumour for both lines, which varied between 102 ± 255 \textmu m (E98, no. 6) and 526 ± 613 \textmu m (E49, no. 23) (Figure 3). In skeletal muscle of the hind limb, the mean area\textsubscript{100} was 0.07 ± 0.02 (−) and is smaller than the mean area\textsubscript{100} values found for the tumour lines E49 and E98, which varied between 0.09 ± 0.02 (E98, no. 1 and no. 6) and 0.84 ± 0.02 (E49, no. 23) (Figure 5).

Vascular morphometrical analysis of perfused vessels and $^{31}$P-MRS

The integral of the peaks in the $^{31}$P-MR spectra reflects the quantity of phosphorylated metabolites (α, β, γ-NTP, P, PCT, PME, PDE) in viable tumour cells (Tozer and Griffiths, 1992), in the volume sampled by the solenoid (approximate tumour volume). The P/NTP ratio is accepted as an indication of the energy status of cells (Roelfstad et al, 1988; Vaupel et al, 1989a), where NTP is broken down to NDP and P by the action of NTPases during cellular activities.

There was little or no contamination by PCR and NTP signals from muscle tissue. $^{31}$P-MR spectra of tumours with similar weights showed PCR peaks smaller than NTP peaks, except some well-perfused tumours, e.g. E98 no. 15 (Figure 1). An example of the domain analysis for two different tumour sections of E49 no. 4 and E98 no. 15 is shown in Figure 1A and C with the corresponding $^{31}$P-MRS spectra of the whole tumours (Figure 1B and 2D). Tumour E49 no. 4 showed a large non-perfused area in the centre, whereas tumour E98 no. 15 showed a homogeneously perfused vessel distribution. The tumours had the following values for the morphometrical parameters: no. 4, percentage of ivd\textsubscript{200} = 22, mean area\textsubscript{100} = 0.52 ± 0.07; no. 15, percentage of ivd\textsubscript{200} = 7 and mean area\textsubscript{100} = 0.12 ± 0.04. The P/NTP ratio of no. 4, i.e. 0.45, was higher than the ratio of no. 15, i.e. 0.17.

P/NTP ratio and the mean intervascular distance

Figure 2 shows that a frequency distribution of ivds in tumour tissue is not a normal distribution. An important tail of large ivds (> 200 \textmu m) was found in all tumours (results not shown here).
Standard deviations were mostly larger than mean ivds, with the exception of most E98 tumours, which showed a more homogeneously perfused vessel distribution. Therefore, the mean ivd can only be used as a rough indication for differences in the ivd distribution between tumours. In Figure 3, the mean ivd per tumour for both lines is related to the P/NTP ratio. The P/NTP ratio showed the largest changes between a mean ivd of approximately 100 μm and approximately 300 μm; the P/NTP ratio increased around a mean ivd of approximately 200 μm.

**P/NTP ratio and the percentage intervascular distance greater than 200 μm**

In Figure 4, the relationship between the P/NTP ratio and the percentage of ivd>200 is shown for the tumour lines E49 and E98. A linear relationship was found with a goodness of fit (R^2) = 0.70. The slope of the regression lines from both tumour lines were not significantly different (P > 0.01). The pooled slope was significantly different from zero (P < 0.0001). No relationship was found between the P/NTP ratio and the percentage of ivd > 100 μm and the percentage of ivd > 300 μm (results not shown here). The P/NTP ratio for the host tissue (hind limb skeletal muscle) was 0.09 ± 0.01 and the percentage of ivd=0. The datapoint of the host tissue is depicted in Figure 4, but was not used in the regression analysis. The energy metabolism in muscle tissue cells probably differs from the energy metabolism in glioma tumour cells, so a direct comparison is not allowed.

**P/NTP ratio and the non-perfused area fraction at a distance larger than 100 μm from the nearest perfused vessel (area_100)**

In Figure 5, the relationship between the P/NTP ratio and the mean area_100 is shown. For both tumour lines, there was a linear relation between the P/NTP ratio and the area_100: R^2 = 0.76. The slopes of the linear regression lines of the tumour lines E49 and E98 were not significantly different (P > 0.01), but the pooled slope was significantly different from zero (P < 0.0001). No relationship was found between the P/NTP ratio and the mean area_50 and the mean area_100, as expected from the results obtained for ivds as shown in the previous paragraph. The goodness of fit in Figure 5 was slightly better than in Figure 4, i.e. the morphometrical parameter mean area_100 showed a better correlation with the P/NTP ratio than the percentage of ivd>200. The datapoint of the host tissue was lower than the datapoints of the different tumours and was not used in the regression analysis (see previous paragraph).

**pH_mrs and morphometrical analysis of perfused vessels**

pH_mrs (approximate pH) was independent of the percentage of ivd>200 and the area_100 over a large range of values (results not shown). All tumours showed a single P, peak corresponding to pH_mrs values from about neutral (pH approximately 7.0 ± 0.1) to basic (pH approximately 7.3 ± 0.1). There were two exceptions in the E49 line: tumour no. 16 showed a split P, peak, corresponding to pH values of 7.3 ± 0.2 and 6.8 ± 0.1; tumour no. 23 had pH_mrs values of 7.12 ± 0.1 and 6.7 ± 0.1. The tumours had the following values for the morphometric parameters: no. 16, percentage of ivd>200 = 37, mean area_100 = 0.63 ± 0.19; no. 23, percentage of ivd>200 = 43 and mean area_100 = 0.84 ± 0.1.

**DISCUSSION**

The relationship between the P/NTP ratio and the morphometrical parameters

In this study, the perfused vascular architecture of complete transverse tumour sections was analysed by two-dimensional morphometric analysis. Two morphometric parameters were evaluated: (a) the percentage of large intervascular distances and (b) the non-perfused area fraction. These were compared with the bioenergetic status of the whole tumour. To our knowledge this is the first study that has related morphometric analysis of the perfused vascular architecture directly to the global bioenergetic status of the same tumour, measured by 31P-MRS.

A domain analysis was used for the calculation of intervascular distances. A domain included non-perfused vascular structures and avascular regions. Polymer infusion techniques can provide similar morphometric information about ivds, but only of perfused regions (Less et al, 1991). Morphometric analysis of only well-perfused regions will probably fail to correlate with the bioenergetic status of the whole tumour, because avascular and non-perfused vascular regions will have a negative impact on the global bioenergetic status if these regions are hypoxic and are lacking nutrients.

The total bioenergetic status (P/NTP ratio) of a tumour depends on the balance between the oxygen supply and nutrient supply and the consumption rates of the tumour cells, which is related to the type of energy metabolism in the different tumour cells. This balance between supply and consumption determines the critical diffusion distances for oxygen and nutrients in tumour tissue. The total oxygen supply and nutrient supply depends on the tumour microcirculation and on the diffusion geometry. This study was restricted to the analysis of the relationship between the diffusion geometry of a tumour and its bioenergetic status. The P/NTP ratio was found to increase rapidly between a mean ivd of approximately 100 μm and approximately 300 μm (Figure 3). This is in agreement with preliminary histological analysis of critical oxygen diffusion distances in tumour sections, using a bioreductive chemical probe for hypoxic cells (Hodgkiss et al, 1991). With use of the bioreductive chemical probe NITP (N-imidazoloto-thio­phylline, a generous gift from Dr R Hodgkiss, Gray Laboratory, England) and the perfusion marker Hoechst, hypoxic areas and perfused vessels were stained simultaneously in the same tumour section. The distances between perfused vessels and hypoxic areas varied from approximately 50 μm to approximately 150 μm, with a mean distance of 113 ± 46 μm. This corresponds to a mean ivd of approximately 200 μm. The complete study will be published separately. The mean critical oxygen diffusion distance of approximately 100 μm was used as a cut-off value in the definition of the morphometrical parameters, i.e. the percentage of ivds > 200 μm (ivd>200) and the non-perfused area fraction at a distance > 100 μm from the nearest perfused vessel.

At a mean distance of approximately 100 μm from a perfused vessel, the probability of an aerobic to anaerobic transition is high. A main question in this study was whether this transition can influence the global bioenergetic status of a tumour? Are there sufficient tumour cells with an aerobic energy metabolism, depending on oxygen for efficient NTP production or may other substrates such as glucose assure NTP production by an intensified (an)aerobic glycolysis?

In vitro studies on tumour cells (Pianet et al, 1991; Gerweck et al, 1993) and ex vivo studies on perfused tumours (Eskey et al, 1993)
showed the effect of a reduced oxygen supply on the oxidative phosphorylation or the inhibition of oxidative phosphorylation (Loesberg et al, 1990). Pianet et al (1991), Gerweck et al (1993) and Eskey et al (1993) found in vitro and ex vivo, that the reduction of the oxygen tension has little influence on the NTP/P ratio in the presence of high glucose concentrations. They argued that (an) aerobic glycolysis is capable of maintaining the energy status. In vivo (DS sarcoma), Vaupel et al (1994) reported that after reduction of the tumour blood flow the amount of NTP remains nearly constant. These findings were explained by an intensified glycolysis due to the recruitment of glucose from the interstitial reservoir of the tumour. Only for tumours with a median oxygen tension below 10-15 mmHg was NTP depletion observed. These conditions were found for tumour masses larger than 1.5% of the body weight. Okunieff et al (1989) observed with 31P-MRS for large tumours (murine fibrosarcoma FS11, approximately 2.5% of the body weight) that i.p. injection of glucose had a positive effect on tumour energy metabolism, and no significant effect on the energy metabolism of small tumours (approximately 0.7% of the body weight).

What can we expect in our glioma tumour model? For a given consumption rate and diffusion coefficient of oxygen and glucose, critical diffusion distances can be estimated using the Krogh model. The consumption rate of oxygen and glucose is related to the energy metabolism of cells. Histochemical evaluation of this metabolism in rat C6 gliomas (Ikezaki et al, 1992) revealed that the energy production is more dependent on aerobic glycolysis than on oxidative phosphorylation: enzymes of the energy-producing tricarboxylic acid cycle and the electron-transport system were reduced, although still present. Rhodes et al (1983) found for human gliomas in vivo, using positron emission tomography (PET), a metabolic uncoupling between the regional oxygen consumption and glucose consumption. The latter is indicative of an increased aerobic glycolysis. In order to determine the importance of the mitochondrial oxidative phosphorylation vs aerobic glycolysis, the in vivo determination of the oxygen and glucose consumption is important. Mean oxygen and glucose consumption rates for macroscopic tissue volumes of human high-grade gliomas (PET-derived data) were estimated to be approximately 0.5 µmol g⁻¹ min⁻¹ for oxygen and approximately 0.5 µmol g⁻¹ min⁻¹ for glucose (Vaupel et al, 1989b). The mean consumption rate of oxygen and glucose was found to be equal in high-grade gliomas, which means an increased aerobic glycolytic activity and/or an increased activity of the pentose phosphate shunt for DNA synthesis (Ikezaki et al, 1992) in comparison with normal brain tissue.

Rough estimations of the oxygen and glucose Krogh radii is possible, using mean consumption rates, mean concentrations in perfused vessels and constant diffusion coefficients (Figure 6). The Krogh radii are approximately 100 µm for oxygen and approximately 200 µm for glucose (Vaupel, 1974; Kallinowski et al, 1987; Groebe, 1988; Vaupel et al, 1989b; Dewhirst et al, 1994). In vivo, a further distribution of Krogh radii will be found, depending on: (a) the distance in a perfused capillary from the inlet, because the pO₂, and the glucose concentration in a capillary decreases between the inlet and outlet; (b) further, the energy metabolism and the related consumption rates of tumour cells may be heterogeneous, i.e. cell regions with aerobic and anaerobic energy metabolism may exist. However, the mean critical oxygen diffusion distance in Figure 6 corresponds well to the mean distance between perfused vessels and hypoxic areas as determined by the use of the bioreductive chemical probe NITP. In addition, the percentage of ivd₂₀₀ and the mean area₁₀₀ were linearly related to the P/NTP ratio, whereas no relationship was found between the P/NTP ratio and higher or lower values than the cut-off value of approximately 100 µm used in the definition of both morphometrical parameters.

The results lead to the following hypothesis: the linear relationship between the P/NTP ratio and the morphometrical parameters in Figures 4 and 5 is due to a slowly changing metabolic steady state: (aerobic) glycolysis + oxidative phosphorylation → anaerobic glycolysis, in which the glioma cells attempt to maintain NTP synthesis by the anaerobic glycolysis during a progressively decreasing glucose supply. At an intervascular distance of approximately 200 µm, the probability of an aerobic to anaerobic transition is high and consequently will affect the local bioenergetic status of glioma cells, which consume oxygen. The relationships found between the P/NTP ratio and the morphometrical parameters possibly indicate that the diffusion-limited supply of oxygen is a major determinant of tissue oxygenation in our glioma tumour model.

As a next step, in vivo and/or in vitro measurements of the oxygen consumption rates of tumour cells in human glioma xenografts, used in this study, will be performed and related to critical diffusion distances of oxygen in vivo.

**pH<sub>mes</sub>**

Two tumours (E49, nos 16 and 23) with 36% extracellular volume, estimated from the eosin- and haematoxylin-stained tumour sections, showed a split P<sub>2</sub> peak. These tumours had the highest values for the percentage of ivd₂₀₀ and the mean area₁₀₀. For no. 16, the percentage of ivd₂₀₀ = 37 and the mean area₁₀₀ = 0.64. For no. 23, the percentage of ivd₂₀₀ = 43 and the mean area₁₀₀ = 0.84. Stubbs et al (1992) suggested that a split P<sub>2</sub> peak is quite possible when the contribution of extracellular P<sub>2</sub> is 35% or more. The ΔpH across the plasma membrane would have to be > 0.3-0.4 pH units.

Gerweck et al (1991) showed for murine fibrosarcoma cells (FS11) that the pH<sub>i</sub> is relatively resistant to changes in pH<sub>j</sub> above 6.9. Below a pH<sub>j</sub> of 6.9 a pH gradient is maintained with pH<sub>i</sub> being consistently more basic than pH<sub>j</sub> by ± 0.35 pH units. The results of Gerweck et al (1991) and Stubbs et al (1992) agree with our findings: for tumour nos 16 and 23 the mean pH<sub>j</sub> is 6.8 ± 0.1 and 6.7 ± 0.1 and the mean pH<sub>i</sub> is 6.8 + 0.35 = 7.2 and 6.7 + 0.35 = 7.1.

No relationship was found between the pH<sub>mes</sub> and the morphometrical parameters. Spatially resolved bioluminescence and fluoroscopic imaging studies of pH values in tumour tissue showed a relationship with the distribution of perfused and non-perfused areas (Hossmann et al, 1992), but global pH<sub>mes</sub> measurements failed in this study.

**ABBREVIATIONS**

MRS, magnetic resonance spectroscopy; ivd, intervascular distance; ivd<sub>200</sub>, intervascular distance larger than 200 µm; Area<sub>100</sub>, non-perfused area fraction at a distance > 100 µm from a nearest perfused vessel; NTP, nucleoside triphosphate; P<sub>2</sub>, inorganic phosphate; PME, phosphomonoesters; PDI, phosphodiesters; pH<sub>mes</sub>, pH measured with 31P-MRS (~ pH<sub>j</sub>); pH<sub>i</sub>, intracellular pH; pH<sub>j</sub>, extracellular pH; pO<sub>2</sub>, oxygen tension; FID, free induction decay; PCr, phosphocreatine.

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