Fraction of radiobiologically hypoxic cells in human melanoma xenografts measured by using single-cell survival, tumour growth delay and local tumour control as end points

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Summary Four human melanoma xenograft lines (A-07, D-12, R-18, U-25) grown orthotopically in Balb/c nu/nu mice were characterized with respect to the fraction of radiobiologically hypoxic cells. The purpose of the study was to establish a firm radiobiological basis for future use of the lines in the development and evaluation of non-invasive assays of tumour hypoxia. The hypoxic fractions were assessed using three different assays, the single cell survival assay, the tumour growth delay assay and the local tumour control assay, and the means ± s.e. were found to be 6 ± 3%, 3 ± 1% and 5 ± 2% respectively (A-07), 26 ± 5%, 25 ± 6% and 22 ± 6% respectively (D-12), 55 ± 9%, 65 ± 8% and 48 ± 7% respectively (R-18) and 52 ± 8%, 59 ± 7% and 47 ± 7% respectively (U-25). The three assays gave numerical values for the hypoxic fraction that were not significantly different for any of the lines. The hypoxic fraction differed significantly among the lines; the R-18 and U-25 lines showed higher hypoxic fractions than the D-12 line ($P < 0.05$), which in turn showed a higher hypoxic fraction than the A-07 line ($P < 0.05$), regardless of the assay. The wide range of the hypoxic fractions and the significant differences among the lines suggest that A-07, D-12, R-18 and U-25 tumours should be useful models in future studies attempting to develop non-invasive assays of tumour hypoxia.

Keywords: treatment–response assay; hypoxic fraction; melanoma xenograft; radiation biology

Many human tumours develop regions of hypoxic cells during growth (Vaupel et al, 1989; 1991; Höckel et al, 1991). The fraction of hypoxic cells differs substantially among individual tumours of the same histological type (Nordsmark et al, 1994; Brizel et al, 1995; Wong et al, 1997). Hypoxia may cause resistance to radiation therapy (Coleman, 1988; Gatenby et al, 1988; Höckel et al, 1993; Nordsmark et al, 1996; Brizel et al, 1997; Fyles et al, 1997) and some forms of chemotherapy (Teicher, 1994) and may promote the development of metastatic disease (Schwickert et al, 1995; Brizel et al, 1996; Höckel et al, 1996; Walenta et al, 1997). Non-invasive methods for detecting and quantifying hypoxia in tumours are needed for prediction of hypoxia-induced treatment resistance and malignant progression in individual patients (Stone et al, 1993).

Several new strategies based on magnetic resonance imaging and spectroscopy, positron emission tomography, single photon emission tomography, electron paramagnetic resonance spectroscopy or phosphorescence imaging are currently being used in attempts to develop non-invasive assays of tumour hypoxia (Hawkins and Phelps, 1988; Moonen et al, 1990; Coleman, 1991; Wilson and Cerniglia, 1992; Negendank, 1992; Bačić et al, 1993). The ultimate aim of these attempts is to arrive at assays for measuring the fraction of the clonogenic cells of tumours that is hypoxic, i.e. the fraction of radiobiologically hypoxic cells, as only the clonogenic cells are of relevance for hypoxia-induced treatment resistance and malignant progression (Kim et al, 1993; Fenton et al, 1995). Rodent tumour lines or human tumour xenograft lines with ‘known’ fractions of radiobiologically hypoxic cells are usually used as model systems in the studies. Successful studies require that the hypoxic fractions of the tumour models have been determined correctly. Most groups involved in the studies have derived the hypoxic fractions of their tumour models from the literature or at best have measured the hypoxic fractions in their own laboratory using a single radiobiological technique. These hypoxic fractions may be misleading, as the hypoxic fraction of a given tumour line may differ significantly among laboratories and the use of different radiobiological assays may result in substantially different hypoxic fractions (Moulder and Rockwell, 1984; Rockwell and Moulder, 1990).

The fractions of radiobiologically hypoxic cells of four human melanoma xenograft lines measured by using single cell survival, tumour growth delay and local tumour control as end points are reported in the present communication. The purpose of the study was to establish a firm radiobiological basis for future investigations aiming at evaluating the usefulness of promising non-invasive methods for detection and quantification of tumour hypoxia.

MATERIALS AND METHODS

Mice and tumours

Adult female Balb/c nu/nu mice (8–12 weeks old), bred at our research institute, were used as host animals for tumours. The mice were maintained under specific pathogen-free conditions at constant temperature (24–26°C) and humidity (30–60%). Sterilized food and tap water were given ad libitum.
The study was performed using the A-07, D-12, R-18 and U-25 human melanoma xenograft lines (Rofstad, 1994). Tumours were initiated from monolayer cultures in exponential growth. Monolayer cells, cultured in RPMI-1640 medium (25 mM Heps and 1-glutamine) supplemented with 13% fetal calf serum, 250 mg ml−1 penicillin and 50 mg ml−1 streptomycin, were detached by trypsinization (treatment with 0.05% trypsin/0.02% EDTA solution at 37°C for 2 min). Approximately 3.5 × 10^6 cells in 10 μl of Ca^2+ and Mg^2+-free Hank’s balanced salt solution (HBSS) were inoculated intradermally in the flanks of mice using a 100-μl Hamilton syringe. The cells were verified to be free from *Mycoplasma* contamination.

The tumours were deposited above the subcutaneous muscle tissue in the deeper part of the dermis. The dermis and the subcutaneous muscle were infiltrated and gradually replaced by malignant tissue during tumour growth. The infiltration of host cells in the tumours was sparse; leucocytes, macrophages and fibroblasts constituted less than 1% of the total number of cells. The growth and histological appearance of the tumours have been described in detail previously (Rofstad, 1994).

**Irradiation**

A Siemens Stabilipan X-ray unit, operated at 220 kV, 19–20 mA, and with 0.5-mm Cu filtration, was used for irradiation. The tumours were irradiated at a dose rate of 5.1 Gy min−1. A 15 × 15 mm hole through a 2-cm-thick lead block served as beam defining aperture. To ensure uniform doses throughout the tumour volume, the tumours were exposed to radiation by two opposing treatment fields through each of which 50% of the dose was delivered.

Tumour volume (V), calculated as \( V = \pi a b^2 \) (a and b are the longer and the shorter of two perpendicular diameters respectively), was within the range of 200–400 mm^3 at irradiation. Tumour diameters were measured with callipers. Hypoxic conditions were obtained by occluding the tumour blood supply with a clamp 5 min before radiation exposure. The mice were kept under anaesthesia during irradiation. Ketamine (Parke Davis, Barcelona, Spain) and azaperone (Janssen Pharmaceutika, Beerse, Belgium) were diluted in phosphate-buffered saline and administered intramuscularly in doses of 33 mg kg−1 body weight and 25 mg kg−1 body weight respectively. The body core temperature of the mice was kept at 36–38°C using a heating pad.

**Single cell survival assay**

Single cell suspensions were prepared from tumours immediately after irradiation using a standardized mechanical and enzymatic procedure. The tumour tissue was minced with crossed scalpels in cold HBSS before enzymatic treatment at 37°C for 2 h. The enzyme solution consisted of 0.2% collagenase, 0.05% pronase and 0.02% DNAase in HBSS. The resulting suspensions were filtered through 30-μm nylon mesh, centrifuged and resuspended in culture medium. Cell concentrations were determined by counting trypan blue-excluding cells in a haemocytometer. The fraction of cells showing trypan blue exclusion was always higher than 80%. The fraction of doublets was always lower than 3%. Larger cell aggregates were not seen.

Fraction of surviving cells was measured in vitro using a plastic surface colony assay. Aliquots of 1 ml of cell suspension were plated in 25-cm² tissue culture flasks. The flasks contained 1 × 10^6 lethally irradiated (30 GY) feeder cells in 4 ml of culture medium. The feeder cells were derived from monolayer cultures and were plated 24 h before the tumour cells were plated. It was verified experimentally that the use of feeder cells increased the plating efficiency of the tumour cells. Moreover, a linear relationship between the number of colonies and the number of cells plated was ensured by the use of feeder cells. The use of feeder cells also inhibited migration of viable tumour cells, hence causing dense and easily scorable colonies.

The cells were incubated at 37°C for 7–21 days in a humidified atmosphere of 5% carbon dioxide in air. One half of the culture medium (2.5 ml) was removed and replaced with fresh medium every seventh day. The cells were fixed in 100% ethanol and stained with methylene blue. Colonies containing more than 50 cells were counted using a stereomicroscope. Plating efficiency was calculated from the number of colonies counted and the number of trypan blue-excluding cells plated and was corrected mathematically for multiplicity (Gillespie et al., 1975). The cell surviving fraction of an irradiated tumour was calculated from the plating efficiency of the cells of the tumour and the mean plating efficiency of the cells of six unirradiated control tumours. The plating efficiencies of unirradiated control tumours were within the ranges of 30–50% (A-07, D-12, U-25) and 50–70% (R-18).

Survival curves were fitted to the data by linear regression analysis to determine the \( D_0 \) values and extrapolation numbers for clamped and unclamped tumours. Only data points at doses judged to be beyond the shoulder region of the survival curve (clamped tumours) and data points at doses eliminating oxic cells (unclamped tumours) were included in the analysis. Fraction of radiobiologically hypoxic cells (RHF) was determined from the vertical displacement of the curves pertaining to clamped and unclamped tumours as

\[
\text{RHF} = \exp \left\{ \ln \left[ n(\text{unclamped}) \right] - \ln \left[ n(\text{clamped}) \right] \right\} = \frac{n(\text{unclamped})}{n(\text{clamped})}
\]

and the s.e. of the RHF [s.e. (RHF)] as

\[
s.e.(\text{RHF}) = \text{RHF} \times \left[ \text{s.e.} \left[ \ln \left( n(\text{unclamped}) \right) \right] \right]^2 + \text{s.e.} \left[ \ln \left( n(\text{clamped}) \right) \right]^2
\]

where \( n(\text{clamped}) \) and \( n(\text{unclamped}) \) represent the extrapolation numbers for clamped and unclamped tumours respectively.

**Tumour growth delay assay**

Tumour volume was measured twice weekly after irradiation as described above. Tumour growth delay, i.e. the time after irradiation at which an irradiated tumour reached twice its volume at irradiation minus the median time required for unirradiated control tumours to double their volumes, was determined and plotted vs radiation dose in a semilogarithmic plot. Dose–response curves were fitted to the data by linear regression analysis. Fraction of radiobiologically hypoxic cells was determined from the horizontal displacement of the curves pertaining to clamped and unclamped tumours as

\[
\text{RHF} = \exp \left\{ \left[ D(\text{unclamped}) - D(\text{clamped}) \right] / D_0 \right\}
\]

and the s.e. of the RHF as

\[
s.e.(\text{RHF}) = \text{RHF} \times \frac{1}{D_0} \times \left[ \text{s.e.}\left[D(\text{unclamped})\right] + \text{s.e.}\left[D(\text{clamped})\right] \right]^{1/2}
\]

where \( D(\text{clamped}) - D(\text{unclamped}) \) represents the horizontal displacement of the curves. The mean of the \( D_0 \) values for clamped and unclamped tumours determined from the single cell survival
data was used in the calculations of the fraction of radiobiologically hypoxic cells.

**Local tumour control assay**

Tumours were examined twice weekly after irradiation and scored as locally controlled if regrowth was not observed within 180 days after treatment. Mice with recurrent tumours were killed when the tumour diameters reached 10–12 mm. Cured mice were killed at day 180 after irradiation and subjected to necropsy and histological examinations for residual tumour tissue. Mitotic figures, morphologically intact tumour cells or any other signs of viable melanoma tissue were never seen in the histological sections.

The percentage of locally controlled tumours was plotted vs radiation dose, and TCD$_{50}$ ± s.e., i.e. the radiation dose that results in 50% local tumour control, was determined by probability regression analysis. Fraction of radiobiologically hypoxic cells was calculated as

$$RHF = \exp \left( \frac{\text{TCD}_{50}(\text{unclamped}) - \text{TCD}_{50}(\text{clamped})}{D_\text{s}} \right)$$

and the s.e. of the RHF as

$$\text{s.e.}(RHF) = \text{RHF} \times \frac{1}{D_\text{s}} \times \left[ \frac{\text{s.e.}[\text{TCD}_{50}(\text{unclamped})]}{\text{TCD}_{50}(\text{unclamped})} \right] + \left[ \frac{\text{s.e.}[\text{TCD}_{50}(\text{clamped})]}{\text{TCD}_{50}(\text{clamped})} \right]$$

The mean of the $D_\text{s}$ values for clamped and unclamped tumours determined from the single cell survival data was used in the calculations of the fraction of radiobiologically hypoxic cells.

**Statistical analysis**

Statistical comparisons of mean values were performed under conditions of normality and equal variance by using the Student’s $t$-test for single comparisons and one-way analysis of variance and the Student–Newman–Keuls test for multiple comparisons. All $P$-values were determined from two-sided tests. A significance criterion of $P < 0.05$ was used. The statistical analysis was performed using SigmaStat statistical software (Jandel Scientific, Erkrath, Germany).

**RESULTS**

Cell survival curves for clamped and unclamped tumours irradiated in vivo and assayed in vitro are presented in Figure 1. The $D_\text{s}$ values describing the terminal slopes of the curves differed among

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**Table 1** $D_\text{s}$ and TCD$_{50}$ values of human melanoma xenografts

| Melanoma | $D_\text{s}$(unclamped) (Gy) | $D_\text{s}$(clamped) (Gy) | TCD$_{50}$(unclamped) (Gy) | TCD$_{50}$(clamped) (Gy) |
|----------|-----------------------------|---------------------------|---------------------------|---------------------------|
| A-07     | 2.39 ± 0.09                 | 2.46 ± 0.07               | 33.0 ± 0.7                | 40.1 ± 0.4                |
| D-12     | 2.17 ± 0.05                 | 2.14 ± 0.06               | 30.0 ± 0.5                | 33.3 ± 0.3                |
| R-18     | 2.69 ± 0.05                 | 2.77 ± 0.09               | 46.2 ± 0.3                | 48.2 ± 0.3                |
| U-25     | 2.51 ± 0.06                 | 2.49 ± 0.05               | 39.2 ± 0.3                | 41.1 ± 0.2                |

$^\text{a}$Mean ± s.e.

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**Table 2** Fraction of radiobiologically hypoxic cells of human melanoma xenografts

| Melanoma | Single cell survival | Tumour growth delay | Local tumour control |
|----------|----------------------|---------------------|----------------------|
| A-07     | 6 ± 3                | 3 ± 1               | 5 ± 2                |
| D-12     | 26 ± 5               | 25 ± 6              | 22 ± 6               |
| R-18     | 55 ± 9               | 65 ± 8              | 48 ± 7               |
| U-25     | 52 ± 8               | 59 ± 7              | 47 ± 7               |

$^\text{a}$Mean ± s.e.
Figure 2 Tumour growth delay vs radiation dose for human melanoma xenografts irradiated under unclamped (□) or clamped (▲) conditions. The curves were fitted to the data by regression analysis. Points and bars represent geometric mean ± s.d. of 30 tumours.

Figure 3 Local tumour control vs radiation dose for human melanoma xenografts irradiated under unclamped (□) or clamped (▲) conditions. The curves were fitted to the data by regression analysis. Points are based on 40 tumours.

The melanoma lines by a factor of 1.2–1.3, but were not significantly different for clamped and unclamped tumours in any of the lines (P > 0.05) (Table 1). The fractions of radiobiologically hypoxic cells determined to be 6 ± 3% (A-07), 26 ± 5% (D-12), 55 ± 9% (R-18) and 52 ± 8% (U-25) (Table 2).

Figure 2 shows growth delay curves for tumours irradiated under clamped and unclamped conditions. Tumour growth delay increased exponentially with radiation dose, i.e., linear curves gave excellent fits to the data sets when the data were plotted in semi-logarithmic diagrams. The slopes of the curves for clamped and unclamped tumours were not significantly different in any of the melanoma lines (P > 0.05). The fractions of radiobiologically hypoxic cells were determined to be 3 ± 1% (A-07), 25 ± 6% (D-12), 65 ± 8% (R-18) and 59 ± 7% (U-25) (Table 2).

Local control curves for tumours irradiated under clamped and unclamped conditions are presented in Figure 3. The TCD₄₀ values differed among the melanoma lines by factors of 1.4–1.5 (clamped tumours) and 1.5–1.6 (unclamped tumours) (Table 1). The slopes of the curves for clamped and unclamped tumours were not significantly different in any of the lines (P > 0.05). The fractions of radiobiologically hypoxic cells determined to be 5 ± 2% (A-07), 22 ± 6% (D-12), 48 ± 7% (R-18) and 47 ± 7% (U-25) (Table 2).

The fraction of radiobiologically hypoxic cells differed among the melanoma lines (Table 2): R-18 and U-25 showed significantly higher hypoxic fractions than D-12 (P < 0.05), which in turn showed a significantly higher hypoxic fraction than A-07 (P < 0.05), regardless of whether single cell survival, tumour growth delay or local tumour control was used as end point. The hypoxic fractions determined for R-18 and U-25 were not significantly different for any of the end points (P > 0.05). The fractions of radiobiologically hypoxic cells determined from the single cell survival assay, the tumour growth delay assay and the local tumour control assay were not significantly different in any of the melanoma lines (P > 0.05).

Individual tumours of the same melanoma line also differed in the fraction of radiobiologically hypoxic cells: the standard deviations of the cell surviving fractions and the tumour growth delays were usually larger for unclamped than for clamped tumours of the same line, as illustrated by the bars in Figures 1 and 2. The coefficient of variation was calculated at each dose level of each curve and the data were subjected to statistical analysis. The coefficient of variation was significantly larger for unclamped tumours than for clamped tumours in all lines (P < 0.05), regardless of whether single cell survival or tumour growth delay was used as end point.
DISCUSSION

Non-invasive methods to determine the fraction of radiobiologically hypoxic cells in tumours are needed for prediction of hypoxia-induced malignant progression and treatment resistance in individual patients (Stone et al. 1993). The development of such assays requires the use of tumour models that are well characterized with respect to the fraction of radiobiologically hypoxic cells. The hypoxic fractions of the human melanoma xenograft lines A-07, D-12, R-18 and U-25 were measured here using single cell survival, tumour growth delay and local tumour control as end points. The purpose of the work was to characterize the lines with respect to radiobiological hypoxia and hence to establish the radiobiological basis for future use of the lines in the evaluation of potentially useful non-invasive assays of tumour hypoxia. The experimental procedures and the methods of calculation were similar to those used previously by others in studies of the fraction of radiobiologically hypoxic cells in experimental tumours (Moulder and Rockwell, 1984; Grau et al. 1990: Rockwell and Moulder, 1990). The novel feature of the work is that the three major assays of hypoxic fraction were applied to the same tumour lines and that the assays gave similar results.

The assessment of the hypoxic fraction of tumours using radiobiological techniques is based on certain assumptions (Moulder and Rockwell, 1984). Some of these assumptions are common for the three major assays. They assume that (a) the survival curves for naturally and artificially hypoxic cells have the same slope and intercept, (b) the majority of the tumour cells are either fully oxic or fully hypoxic, (c) the method of clamping leaves no oxygenated tumour regions and (d) the tumour cells rendered artificially hypoxic are no less viable than the cells in unclamped tumours. In addition, special assumptions are associated with each of the assays. Thus, the single cell survival assay is based on the assumption that no tumour cell subpopulation is selectively enriched or lost during the preparation of single cell suspensions. The tumour growth delay assay requires that the same level of cell inactivation in clamped and unclamped tumours results in the same growth delay. Finally, the local tumour control assay assumes that the same level of cell inactivation is required to control clamped and unclamped tumours. Some of the assumptions can be tested experimentally as described by Moulder and Rockwell (1984). These assumptions have been found to be valid for the tumour lines studied here. Other assumptions cannot be tested experimentally, and the validity of these assumptions is therefore questionable. However, as the hypoxic fractions determined by the single cell survival assay, the tumour growth delay assay and the local tumour control assay were not significantly different, the assumptions that these assays do not have in common were probably adequately met by the A-07, D-12, R-18 and U-25 tumour lines.

The main conclusion of our work is that the fraction of radio biologically hypoxic cells differs significantly among the human melanoma xenograft lines subjected to investigation: the R-18 and U-25 lines showed a higher hypoxic fraction than the D-12 line, which in turn showed a higher hypoxic fraction than the A-07 line. This conclusion is indisputable as the three major assays of hypoxic fraction not only ranked the lines in the same order but also gave numerical values for the hypoxic fraction that were not significantly different for any of the lines. Another important conclusion is that the fraction of radiobiologically hypoxic cells differs substantially among individual tumours of the same line. This conclusion is based on the observation that the coefficients of variation for cell surviving fraction and tumour growth delay were significantly larger for unclamped than for clamped tumours.

The A-07, D-12, R-18 and U-25 tumour lines should be useful models for developing and evaluating non-invasive assays of tumour hypoxia for several reasons. Firstly, the tumour lines have been well characterized with respect to the fraction of radiobiologically hypoxic cells. Secondly, the hypoxic fractions of the lines cover a broad range, from approximately 5% to more than 50%.

Thirdly, the fact that the hypoxic fraction differs substantially among individual tumours of the same line, particularly in the A-07 line, may be used to assess to what extent a non-invasive assay of tumour hypoxia is influenced by the biochemical properties of the tumour cells in addition to the hypoxic fraction. Moreover, previous studies have shown that several biological characteristics of the donor patients' tumours have been retained in these tumour lines, including angiogenic potential; growth, histopathological and pathophysiological parameters; organ-specific metastatic pattern; and sensitivity to dacarbazine, heat and radiation treatment (Rofstad, 1994).

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