Energy metabolism in human melanoma cells under hypoxic and acidic conditions in vitro

R Skøyum, K Eide, K Berg and EK Rofstad

Institute for Cancer Research, The Norwegian Radium Hospital, Montebello, 0310 Oslo, Norway

Summary The response to treatment and the malignant progression of tumours are influenced by the ability of the tumour cells to withstand severe energy deprivation during prolonged exposure to hypoxia at normal or low extracellular pH (pH_e). The objective of the present work was to demonstrate intertumour heterogeneity under conditions of microenvironment-induced energy deprivation and to investigate whether the heterogeneity can be attributed to differences in the capacity of the tumour cells to generate energy in an oxygen-deficient microenvironment. Cultures of four human melanoma cell lines (BEX-c, COX-c, SAX-c, WIX-c) were exposed to hypoxia in vitro at pH_e 7.4, 7.0 or 6.6 for times up to 31 h by using the steel-chamber method. High-performance liquid chromatography was used to assess adenylate energy charge as a function of exposure time. Cellular rates of glucose uptake and lactate release were determined by using standard enzymatic test kits. The adenylate energy charge decreased with time under hypoxia in all cell lines. The decrease was most pronounced shortly after the treatment had been initiated and then tapered off. BEX-c and SAX-c showed a significantly higher adenylate energy charge under hypoxic conditions than did COX-c and WIX-c whether the pH_e was 7.4, 7.0 or 6.6, showing that tumours can differ in the ability to avoid energy deprivation during microenvironmental stress. There was no correlation between the adenylate energy charge and the rates of glucose uptake and lactate release. Intertumour heterogeneity in the ability to withstand energy deprivation in an oxygen-deficient microenvironment cannot therefore be attributed mainly to differences in the capacity of the tumour cells to generate energy by anaerobic metabolism. The data presented here suggest that the heterogeneity is rather caused by differences in the capacity of the tumour cells to reduce the rate of energy consumption when exposed to hypoxia.

Keywords: adenylate energy charge; energy metabolism; extracellular pH; hypoxia; melanoma

The metabolic microenvironment of tumours differs substantially from that of the corresponding normal tissue (Vaupel et al, 1989). Many tumours develop hypoxic regions and regions with low extracellular pH (pH_e) during growth (Wike-Hooley et al, 1984; Stone et al, 1993). Two types of hypoxia have been recognized in tumours: chronic hypoxia, resulting from limitations in oxygen diffusion; and acute hypoxia, resulting from transient cessations in blood flow (Horsman, 1995). Low pH_e in tumours is mainly a result of increased rate of glycolysis and poor vascularity, leading to inadequate removal of H+ (Vaupel et al, 1989). The rate of glycolysis is up-regulated in hypoxic tumour cells (Robin et al, 1984), and anaerobic glycolysis leads to accumulation of lactic acid (Busa and Nuccitelli, 1984; Wike-Hooley et al, 1984; Vaupel et al, 1989) the pH_e might be particularly low in the hypoxic regions of tumours (Kallinowski and Vaupel, 1988; Tannock and Rotin, 1989; Martin and Jain, 1994).

Tumour treatment response depends on the metabolic microenvironment of the tumour cells (Wike-Hooley et al, 1984; Sutherland et al, 1988). Hypoxic tumour cells are resistant to radiation therapy (Coleman, 1988) and some forms of chemotherapy (Teicher, 1994). The radiation dose required to inactivate tumour cells under hypoxic conditions is 2.5–3.0 times higher than that required under aerobic conditions (Stone et al, 1993). Tumour cells at low pH_e also show increased radiation resistance, although the increase is much less than that due to hypoxia (Tannock and Rotin, 1989; Durand, 1991). The cytotoxic activity of chemotherapeutic drugs is increased, unchanged or decreased at low pH_e, depending on the tumour cells and the mechanism of action of the drug (Tannock and Rotin, 1989; Teicher, 1994).

The malignant progression of tumours is also influenced by the metabolic microenvironment of the tumour cells (Hill, 1990). The expression of several specific genes involved in the malignant progression is increased in hypoxic tumour cells and tumour cells at low pH_e (Brown and Giaccia, 1994; Dachs and Stratford, 1996). Tumour cells can show increased metastatic potential following transient exposure to hypoxia (Young et al, 1988; Young and Hill, 1990) or low pH_e (Schlappack et al, 1991). Moreover, hypoxia followed by reoxygenation can lead to increased resistance to some chemotherapeutic drugs (Rice et al, 1987; Luk et al, 1990; Sanna and Rofstad, 1994) and to the development of new cell subpopulations increased DNA content (Rice et al, 1986; Wilson et al, 1989) or a doubling of the number of chromosomes (Rofstad et al, 1996a).

The probability of microenvironment-induced tumour treatment failure or malignant progression is influenced by the ability of the tumour cells to withstand severe energy deprivation during prolonged exposure to hypoxia at normal or low pH_e (Coleman, 1988; Tannock and Rotin, 1989; Vaupel et al, 1989; Hill, 1990). The energy status of tissues is controlled by the balance between the rates of ATP formation and utilization. The ATP formation is reduced markedly in tumour cells in a hypoxic microenvironment,
mainly because of inhibition of respiration but also because of inhibition of glycolysis by low \( \text{pH}_i \) (Rotin et al, 1986; Gerweck et al, 1993). ATP utilization is subsequently reduced, leading to decreased DNA and protein synthesis and cessation of cell proliferation (Born et al, 1976; Heacock and Sutherland, 1990; Casciani et al, 1992). The rate of energy deprivation during exposure to hypoxia at normal or low \( \text{pH}_i \) might thus differ between tumours because of differences in the capacity of the tumour cells to generate energy and/or to reduce the energy consumption.

The main purpose of the work reported here was to demonstrate intertumour heterogeneity under conditions of microenvironment-induced energy deprivation and to investigate whether the heterogeneity can be attributed to differences in the capacity of the tumour cells to generate energy in an oxygen-deficient microenvironment. Cell cultures of four human melanoma lines were exposed to hypoxia in vitro at \( \text{pH} \), 7.4, 7.0 or 6.6, and adenylate energy charge was measured as a function of exposure time. As glucose is the main energy source of tumour cells under hypoxic conditions (Wike-Hooley et al, 1984; Vaupel et al, 1989), the rates of glucose uptake and lactate release were used as measures of energy generation.

**MATERIALS AND METHODS**

**Cell lines**

Four human melanoma cell lines (BEX-c, COX-c, SAX-c, WIX-c) were included in the study (Rofstad et al, 1991). The cell lines were maintained in monolayer culture in RPMI-1640 medium (25 mM Hepes and L-glutamine) supplemented with 13% fetal calf serum, 250 mg \( \text{L}^{-1} \) penicillin and 50 mg \( \text{L}^{-1} \) streptomycin. The cultures were incubated at 37°C in a humidified atmosphere of 5% carbon dioxide in air and subcultured once a week by trypsinization (treatment with 0.05% trypsin/0.02% EDTA solution at 37°C for 2 min). All solutions were purchased from Life Sciences Technology, UK. The cell lines were verified to be free from *Mycoplasma* contamination by using the Hoechst fluorescence and the mycotrin methods.

**Exposure to hypoxia**

Monolayer cell cultures growing in glass dishes were exposed to hypoxia by using the steel-chamber method (Sanna and Rofstad, 1994). The cultures were incubated at 37°C in a humidified atmosphere of 5% carbon dioxide in air for 24 h before the hypoxia treatment. The culture medium was removed and replaced by fresh medium before the cells were exposed to hypoxia. The medium used during the hypoxia treatment was supplemented with 1.1 mg \( \text{mL}^{-1} \) sodium bicarbonate and adjusted to a \( \text{pH} \) of 7.4, 7.0 or 6.6 using sodium hydroxide or hydrochloric acid. Before use, it was flushed with 5% carbon dioxide in air for 2 h and then the \( \text{pH} \) was readjusted (Boyer et al, 1993). The glass dishes were kept in air-tight steel chambers during the hypoxia treatment. The steel chambers were flushed with a humidified, highly purified gas mixture consisting of 95% nitrogen and 5% carbon dioxide at a flow rate of 5 l min\(^{-1}\). Measurements showed that the concentration of oxygen in the medium was less than 10 p.p.m. after 30 min of flushing. Control cultures were flushed with humidified 5% carbon dioxide in air. Separate dishes were used for measurement of adenylate energy charge and concentrations of glucose and lactate. After medium samples had been collected for measurement of glucose and lactate concentrations, the cells were detached from the dishes by trypsinization and counted by using a haemocytometer and a phase-contrast microscope.

![Figure 1](image1.png)  
*Figure 1* Extracellular \( \text{pH} \) in hypoxic cell cultures vs time under hypoxia. The cells were cultured in medium adjusted to a \( \text{pH} \) of 7.4 (□), 7.0 (■) or 6.6 (△) immediately before the hypoxia exposure. The curves refer to single experiments with BEX-c. Similar curves were achieved in all experiments, irrespective of cell line.

![Figure 2](image2.png)  
*Figure 2* Relative cell number vs time for aerobic (○) and hypoxic (■) cell cultures. The curves refer to single experiments with BEX-c at \( \text{pH}_i \) 7.4. Similar curves were achieved in all experiments, irrespective of cell line and \( \text{pH}_i \).
Measurements of rates of glucose uptake and lactate release

Cellular rates of glucose uptake and lactate release were determined by using standard enzymatic test kits (Boehringer Mannheim, Mannheim, Germany) and spectrophotometric assays (Casciari et al, 1992). The glucose and lactate concentrations in the media of aerobic and hypoxic cell cultures were measured as a function of time. The rates of the concentration changes were divided by the logarithmic mean cell number to obtain rates of glucose uptake and lactate release in terms of mol per cell per second (Rofstad et al, 1996b).

Measurement of adenylate energy charge

The adenylate phosphates were separated by high-performance liquid chromatography and quantified at 254 nm. The cells were lysed in acetonitrile (CH$_3$CN), scraped off the dishes and dried in nitrogen. Low-strength buffer (100 mM potassium dihydrogen phosphate, 1.5% acetonitrile, 0.08% tetrabutylammonium bromide (C$_6$H$_{15}$NBr), pH 5.0) was added before centrifugation and collection of the supernatant. The elution was performed with 80% low-strength buffer and 20% high-strength buffer (150 mM potassium dihydrogen phosphate, 10% acetonitrile, 0.08% tetrabutylammonium bromide, pH 5.0) for 10 min, then with a linear

Figure 3 Medium concentration of glucose (A and B) and lactate (C and D) in hypoxic cell cultures vs time under hypoxia. The curves refer to single experiments with BEX-c at pH 7.4 or 6.6. Similar curves were achieved in all experiments, irrespective of cell line.
gradient to 100% high-strength buffer for 10 min and finally with the high-strength buffer for 10 min. The separation was carried out with a Supelcosil LC-18-T 5-μm cartridge (Supelco, SA, Crans, Switzerland). The flow rate was 1.0 ml min⁻¹. Adenylate energy charge (AEC) was calculated from peak areas:

\[ \text{AEC} = \frac{[\text{ATP}] + \frac{1}{2} [\text{ADP}]}{[\text{ATP}] + [\text{ADP}] + [\text{AMP}]} \]

Statistical analysis

Results are presented as individual values or as arithmetic means ± standard deviation (s.d.). 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. Paired tests were performed where appropriate. 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).

Figure 4 Rates of glucose uptake and lactate release in BEX-c, COX-c, SAX-c and WIX-c under aerobic and hypoxic conditions. The experiments were performed at pH \( \text{pH}_0 \) 7.4. Columns and bars represent mean values ± s.d. of 4–9 independent experiments. ○, Rate of glucose uptake; ■, rate of lactate release

RESULTS

Cell cultures were exposed to hypoxia at pH \( \text{pH}_0 \) 7.4, 7.0 or 6.6 for 3–31 h. In this time interval, the pH decreased by 0.2 pH units at pH \( \text{pH}_0 \) 7.4 and 7.0 and by 0.4 pH units at pH \( \text{pH}_0 \) 6.6 (Figure 1). The cells did not proliferate during the hypoxia treatment. The number of cells per dish increased exponentially with time in aerobic control cultures and stayed unchanged in hypoxic cultures (Figure 2). The cells remained attached to the glass surface during the hypoxia treatment. The fraction of trypan blue excluding cells after 31 h of hypoxia was higher than 95% in all cell lines, independent of the pH.

The glucose concentration in the medium of hypoxic cell cultures decreased linearly with time at pH \( \text{pH}_0 \) 7.4 and 7.0 and, correspondingly, the lactate concentration increased linearly with time (Figure 3). In contrast, the glucose and lactate concentration curves were not linear at pH \( \text{pH}_0 \) 6.6: they showed a break between 10 and 14 h (Figure 3). The rates of glucose and lactate concentration changes were determined by linear regression analysis. Single values were derived from each data set at pH \( \text{pH}_0 \) 7.4 and 7.0. Two values were derived at pH \( \text{pH}_0 \) 6.6: one early value based on analysis
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Figure 5 Rates of glucose uptake and lactate release in BEX-c, COX-c, SAX-c and WIX-c under hypoxic conditions. The experiments were performed at pH₇.4, 7.0 or 6.6. Columns and bars represent mean values ± s.d. of four or five independent experiments. □ Rate of glucose uptake; ■ rate of lactate release.

of the data in the time interval 0–10 h and one late value based on analysis of the data in the time interval 14–31 h. The rates of the glucose and lactate concentration changes in the medium of aerobic cell cultures were determined by linear regression analysis of data pertaining to a 6-h time interval (figure not shown).

The rates of glucose uptake and lactate release in BEX-c, COX-c, SAX-c and WIX-c under hypoxic conditions are compared with those under aerobic conditions in Figure 4. The experiments were performed at pH₇.4. BEX-c showed lower rates of glucose uptake and lactate release than COX-c, SAX-c and WIX-c under aerobic conditions (P < 0.01). The rates of glucose uptake and lactate release were higher under hypoxic conditions than under aerobic conditions in all cell lines (P < 0.05). The magnitude of the hypoxia-induced glycolysis up-regulation was not significantly different in the four cell lines.

Figure 5 shows the rates of glucose uptake and lactate release in BEX-c, COX-c, SAX-c and WIX-c under hypoxic conditions at pH₇.4, 7.0 and 6.6 (early and late). The glucose metabolism differed significantly between some of the cell lines at pH₇.4, 7.0 and 6.6 (early). Thus, the rates of glucose uptake and lactate release were higher in COX-c, SAX-c and WIX-c than in BEX-c at pH₇.4 (P < 0.01) and 7.0 (P < 0.01). SAX-c showed higher rates of glucose uptake and lactate release than BEX-c, COX-c and WIX-c at pH₆.6 (early) (P < 0.05). Significant differences between the cell lines were not detected at pH₆.6 (late).

The glucose metabolism was similar at pH₇.4 and 7.0 and inhibited at pH₆.6 (Figure 5). Thus, the rate of glucose uptake was lower at pH₆.6 (early and late) than at pH₇.4 and 7.0 in all cell lines (P < 0.05). All lines also showed a lower rate of lactate release at pH₆.6 (late) than at pH₇.4 and 7.0 (P < 0.01). However, the rate of lactate release at pH₆.6 (early) was not significantly different from that at pH₇.4 and 7.0 in any of the lines. It should also be noticed that at pH₆.6 (early), the rate of lactate release was higher than the rate of glucose uptake by a factor larger than 2.0.
The adenylate energy charge of the cell lines was 0.94 ± 0.01 (BEX-c), 0.92 ± 0.02 (COX-c), 0.93 ± 0.03 (SAX-c) and 0.93 ± 0.02 (WIX-c) under aerobic conditions at pH<sub>s</sub> 7.4. Chromatograms showing that the adenylate energy charge decreased with time under hypoxia are presented in Figure 6. The ATP peak was reduced and the AMP and ADP peaks were enhanced after protracted hypoxia treatments. The adenylate energy charge followed biphasic curves; the decrease was most pronounced shortly after the hypoxia treatment had been initiated and then tapered off (Figure 7). The magnitude of the hypoxia-induced decrease differed significantly between some of the cell lines. Thus, 6–31 h after the initiation of treatment, the adenylate energy charge was higher in BEX-c and SAX-c than in COX-c and WIX-c at pH<sub>s</sub> 7.4 (P < 0.05), 7.0 (P < 0.05) and 6.6 (P < 0.05). The adenylate energy charge in BEX-c was not significantly different from that in SAX-c and the adenylate energy charge in COX-c was not significantly different from that in WIX-c at any of these pH<sub>s</sub> values. Adenylate energy charge curves were therefore fitted to the combined BEX-c and SAX-c data and to the combined COX-c and WIX-c data in Figure 7. The adenylate energy charge was independent of the pH<sub>s</sub>; the curves at pH<sub>s</sub> 7.4, 7.0 and 6.6 were not significantly different in any of the cell lines.

**DISCUSSION**

The adenylate energy charge of BEX-c, COX-c, SAX-c and WIX-c decreased during exposure to hypoxia. Hypoxia resulted in up-regulated glycolysis in all four lines. However, the up-regulation was not sufficiently large to compensate for the loss in energy generation resulting from the inhibition of the oxidative pathway. The effects of hypoxia on glycolysis and adenylate energy charge reported here for human melanoma cells are thus consistent with those reported previously for rodent tumour cells and other histological types of human tumour cells (Rotin et al, 1986; Heacock and Sutherland, 1990; Casciarli et al, 1992; Gerweck et al, 1992).

The hypoxia-induced decrease in adenylate energy charge differed significantly between the four cell lines. Thus, BEX-c and SAX-c showed a higher adenylate energy charge under hypoxic conditions than did COX-c and WIX-c, whether the pH<sub>s</sub> was 7.4,
7.0 or 6.6. This observation suggests that tumour cells can differ substantially in their ability to avoid energy deprivation during microenvironmental stress. As all cell lines studied here were of melanoma origin, significant intertumour heterogeneity in microenvironment-induced energy deprivation might occur even within single histological types of tumours.

The differences between the cell lines in adenylyl energy charge during exposure to hypoxia cannot be attributed mainly to differences in the capacity of the cells to generate energy by anaerobic metabolism, because there was no correlation between the adenylyl energy charge and the rates of glucose uptake and lactate release. Thus, the adenylyl energy charge was similar for BEX-c and SAX-c, irrespective of the pH₇, whereas BEX-c showed lower rates of glucose uptake and lactate release than SAX-c, except at pH 6.6 (late). SAX-c showed a higher adenylyl energy charge than COX-c and WIX-c at pH 7.4 and 7.0, whereas the rates of glucose uptake and lactate release were similar. Moreover, the adenylyl energy charge at pH 6.6 was similar to that at pH 7.4 and 7.0 in all cell lines, whereas the rates of glucose uptake were down-regulated at pH 6.6.

The adenylyl energy charge of tumour cells subjected to microenvironmental stress is controlled by the balance between the rates of energy generation and energy consumption (Calderwood et al., 1985; Rotin et al., 1986; Gerweck et al., 1992). Exposure of cells to hypoxia leads to continuously decreasing rates of DNA and protein synthesis and cessation of cell proliferation (Born et al., 1976; Shrieve et al., 1983; Pettersen et al., 1986; Heacock and Sutherland, 1990). Cell lines might differ in the capacity to regulate and turn off these energy-requiring processes (Gerweck et al., 1992). Consequently, the higher adenylyl energy charge in BEX-c and SAX-c than in COX-c and WIX-c might have resulted from a faster hypoxia-induced decrease in the energy consumption in BEX-c and SAX-c than in COX-c and WIX-c. This interpretation is consistent with the biphasic shape of the adenylyl energy charge curves. The decrease in adenylyl energy charge was most pronounced shortly after the hypoxia treatment had been initiated and then tapered off with time.

The pH₇ of most tumours in vivo is within the range 6.6–7.4 (Wike-Hooley et al., 1984; Tannock and Rotin, 1989; Vaupel et al., 1989). Experimental studies have suggested that the energy status is particularly low in hypoxic tumour cells at pH 7.0 in the lower part of this range, because of inhibited glycolysis owing to the low pH₇ (Halperin et al., 1969; Rotin et al., 1986; Casiari et al., 1992; Gerweck et al., 1993). However, the adenylyl energy charge at pH 6.6 was similar to that at pH 7.4 and 7.0 in hypoxic cultures of the cell lines studied here, even though the rate of glucose uptake was lower at pH 6.6, than at pH 7.4 and 7.0. It is possible that the melanoma cells used an intracellular energy source at pH 6.6, e.g. a stock of glycogen, to compensate for the reduced rate of glucose uptake. This suggestion is consistent with the observation that the rate of lactate release was higher than the rate of glucose uptake by a factor larger than 2.0 during the first 10 h of the hypoxia treatment at pH 6.6. The rate of lactate release corresponded to the rate of glucose uptake 14–31 h after the treatment had been initiated (Figure 5), indicating that the intracellular energy source had been exhausted. The rate of energy consumption might have been adjusted to the reduced rate of energy generation by then, and hence, the adenylyl energy charge at pH 6.6 was similar to that at pH 7.4 and 7.0 in hypoxic BEX-c, COX-c, SAX-c and WIX-c cultures.

In summary, hypoxia and low pH might cause resistance to treatment and promote the malignant progression of tumours, provided that the tumour cells have the ability to avoid severe energy deprivation during prolonged microenvironmental stress. The present study suggests that tumours differ in this ability and that the differences can be attributed to differences in the capacity to turn off energy-requiring processes in an oxygen-deficient microenvironment rather than to differences in the capacity to generate energy by anaerobic metabolism.

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