Regulation of cell proliferation under extreme and moderate hypoxia: the role of pyrimidine (deoxy)nucleotides

Ø. Åmelllem¹, M. Löffler¹ & E.O. Pettersen¹

¹Department of Tissue Culture, Institute for Cancer Research, The Norwegian Radium Hospital, N-0310 Oslo, Norway; ²Klinikum der Philippus-Universität Marburg, Institut für Physiologische Chemie, Karl-von-Frisch-Straße, D-35033 Marburg, Germany.

Summary. In the present study we have used flow cytometric DNA measurements on synchronised human NHK 3025 cells to measure cell cycle progression under various conditions of reduced oxygenation. Our data indicate that addition of 0.1 mM deoxycytidine or uridine has no effect on the oxygen-dependent arrest in late G1, or on the inhibition of cell proliferation through S-phase under extremely hypoxic conditions. Following reoxygenation of cells exposed to extremely hypoxic conditions in G1, initiation of DNA synthesis in the subsequent cell cycle is delayed by several hours. This G1-induced delay is completely abolished for approximately 60% of the cell population by addition of deoxycytidine to hypoxic G1 cells. This finding supports previous proposals that important steps in the preparation for DNA synthesis occur during G1 of the previous cell cycle, and it indicates that this preparation is connected to the de novo synthesis of pyrimidine deoxy-nucleotide precursors. The results show that cells are able to enter S-phase in the presence of 100-1,300 p.p.m. (0.01-0.13%) oxygen, but they are not able to initiate DNA synthesis under such conditions. However, the cell cycle inhibition induced under moderate hypoxia is partially reversed in the presence of exogenously added deoxycytidine and uridine, while no such reversal is seen in the presence of thymine deoxynucleosides (deoxyadenosine and deoxyguanosine). Thus, both deoxycytidine and uridine could replace reoxygenation under these conditions. These results indicate that the reduction of CDP to dCDP by ribonucleotide reductase, an enzyme which requires oxygen as an essential factor for the formation of tyrosyl radicals for its catalytic activity, does not seem to be the limiting step responsible for the reduced dCDP pool observed under moderate hypoxia. We conclude that the oxygen-dependent catalytic activity of the M2 subunit of ribonucleotide reductase is still intact and functional in NHK 3025 cells even at oxygen concentrations as low as 100 p.p.m. Therefore, the cell cycle inhibition observed is probably due to inhibition of the respiratory chain-dependent UMP synthesis at the stage of dihydroorotate dehydrogenase.

The presence of hypoxic cells in solid tumours has long been considered a problem in cancer treatment, particularly for radiation therapy, but also for treatment with some anticancer drugs. Recent research has indicated that cell growth and cell cycle control are affected in a fundamental manner by reduced oxygenation. The knowledge of how intracellular processes respond to various degrees of hypoxia might therefore give relevant information concerning cell growth regulation as well as suggest ideas for future improvements concerning some cancer therapy regimens.

In general, cells exposed to extremely hypoxic conditions in S-phase immediately arrest, while cells in other phases of the cell cycle proceed to late G1 before they become arrested (Koch et al., 1973a; Bedford & Mitchell, 1974; Löffler et al., 1978; Pettersen & Lindmo, 1981; Probst & Gekeler, 1988). Although it has been reported that some cells arrest in G1 or mitosis during hypoxia (Shrieve et al., 1983; Shrieve & Begg, 1985), it appears that most cells studied so far are able to complete mitosis and divide in the absence of oxygen.

The nature of the oxygen-dependent restriction point in late G1 is still an enigma. This resting state is, not for example, similar to the aerobic non-cycling state G0 (Probst et al., 1988; Åmelllem & Pettersen, 1993). It is probable, though, that the G1 arrest under hypoxia plays a fundamental role in protecting cells from the lethal effect of extreme hypoxia in S-phase (Merz & Schneider, 1983; Spiro et al., 1984; Åmelllem & Pettersen, 1991). Although hypoxia affects the energy status of the cell, it appears that lack of energy is not directly responsible for the G1 arrest observed under such conditions (Löffler, 1985a). Hypoxia has been shown to alter gene expression. Specific proteins expressed under hypoxic conditions have been identified (Anderson & Matovcik, 1977; Scandra et al., 1984; Heacock & Sutherland, 1986). However, the functions of these proteins under and following hypoxic stress seems not to be connected to the specific G1 arrest (Shi et al., 1993). Still, it may well be that the mechanism preventing cells from initiating DNA synthesis under extreme hypoxia is actively regulated in order to protect the cells from damage. Even under moderate conditions of hypoxia, when cellular respiration is not hampered (Froese, 1962; Boag, 1970), such as above 1,300 p.p.m. oxygen, cell cycle progression is inhibited (Koch et al., 1973b; Löffler, 1992).

It appears that reduced de novo synthesis of deoxy-nucleotides might, at least partly, be responsible for this cell cycle inhibition since the intracellular level of pyrimidine precursor pools, dCTP and dTTP, needed for DNA synthesis has been found to be substantially reduced after hypoxia (Löffler et al., 1983). It has been suggested that the size of the dCTP pool may have a regulatory role in the rate of DNA synthesis (Bjursell & Reichard, 1973).

Two enzymes, dihydroorotate dehydrogenase and ribonucleotide reductase, are presumably responsible for the imbalanced deoxynucleotide pools during hypoxia. The first, dihydroorotate dehydrogenase, operates at the stage of the respiratory chain-dependent UMP synthesis. The second, ribonucleotide reductase, requires molecular oxygen as an essential factor for the formation of a tyrosyl radical needed for its catalytic activity (Thelander et al., 1983). Under normoxic growth conditions regulation of the deoxynucleotide triphosphatase (dNTP) pools occurs through de novo synthesis (main route) and secondly through the salvage pathway to ascertain that the proper amount of all four dNTPs are available for DNA replication (Reichard, 1988). The main route includes the two mentioned oxygen-dependent steps, while the salvage pathway, although dependent on oxygen for nucleoside synthesis is independent of oxygen for deoxynucleoside synthesis (see Figure 1). Thus, deoxynucleosides such as deoxyctydine (dC) should be able to support the dNTP pools needed for DNA replication through the salvage pathway during hypoxia.

In the present report we have focused on the possible role of exogenously added deoxynucleosides and uridine in the control of cell proliferation under various hypoxic conditions.
Figure 1. Pathways for the synthesis of pyrimidine deoxyribonucleotide triphosphates. Two enzymes of particular interest to cells exposed to hypoxic conditions are indicated as follows: DHO DH, dihydroorotate dehydrogenase; and RR, ribonucleotide reductase. Urudine or deoxycytidine (dC) added to the medium is rapidly transported into cells by facilitated diffusion. Urudine must pass through the oxygen-dependent main route via RR, while dC can pass through the salvage pathway, which is oxygen independent, to be used for DNA synthesis. Solid arrows indicate one step in the synthesis of pyrimidines and dashed arrows indicate more than one step.

Materials and methods

Cell culture and synchronisation by the method of mitotic selection

NHK 3025 is an established human cell line derived from cervical carcinoma in situ (Nordbye & Oftebro, 1969; Oftebro & Nordbye, 1969). The cells were cultivated as a monolayer in medium E2a (Puck et al., 1957) containing 20% human serum (prepared in the laboratory) and 10% horse serum (Gibco, UK). The cells were in exponential growth by reculturing three times a week. Under these conditions the median durations of the various phases of the cell cycle are: G1, 6.5 h; S, 7.5 h; G2, 1.5 h; and mitosis, 1 h. Under optimal growth conditions, as used here, these cells meet the requirements set up by Anderson et al. (1967) for cells in balanced growth (Pettersen et al., 1977). NHK 3025 cells have a DNA index of 2.5, measured by means of DNA flow cytometry on samples prepared by the method of Vindelov et al. (1983a,b) using chicken and trout red blood cells as internal standards.

Populations of cells with a high degree of synchrony were obtained by collecting detached mitotic cells after a shaking procedure, as described previously (Pettersen et al., 1977). Such synchronised cells are in balanced growth (Running et al., 1981) and have the same cell cycle and phase durations as exponentially growing cells (Pettersen et al., 1977). From one selection the yield was between 4 x 10^4 and 8 x 10^5 cells in 240 ml of medium. After centrifugation (250 g, 5 min), the cells were then seeded in glass Petri dishes and placed in a 37°C incubator with 5% carbon dioxide in air of high humidity for at least 2.5 h to allow attachment to the bottom of the glass dishes.

Hypoxia

The technique of introducing and maintaining extremely hypoxic conditions in cell cultures has been described previously (Pettersen & Lindmo, 1981). Briefly, the cells were seeded in Anumbras glass dishes (7 cm in diameter) and incubated in a carbon dioxide incubator. At the appropriate time the glass dishes were brought from the carbon dioxide incubator into a walk-in incubator room at 37°C. The medium content in each dish was reduced from 10 to 3 ml and placed without lids in a stainless-steel chamber. Deoxygenation took place by continuous flushing of the chamber with a gas mixture (Hydro Gas, Norway) of 97% nitrogen, 3% carbon dioxide and <4, 100 and 1,300 p.p.m. oxygen at 37°C using the set-up described previously (Pettersen et al., 1973; Levhaug et al., 1977). The hypoxic atmosphere in the chamber was established about 12 min after the start of flushing (unpublished results). Untreated control populations were kept in the carbon dioxide incubator all the time after mitotic selection. Medium supplemented with a mixture of deoxycytidine (0.1 mm), deoxyadenosine and deoxyguanosine (0.01 mm each) or with deoxycytidine (0.1 mm) or uridine (0.1 mm) alone was added at the start of the hypoxic exposure, during reoxygenation, or both. All chemicals were obtained from Sigma (USA).

Microscopy

In order to observe cells in a microscope (Zeiss, Germany) under hypoxic conditions we constructed two small steel chambers with a glass lid containing one glass dish each (Belco, USA), which also constitute the bottom of the chambers. First, cells were seeded in glass dishes (3.1 cm in diameter) immediately after mitotic selection and placed in a carbon dioxide incubator. The medium content in each dish was reduced from 2 to 0.6 ml and placed in the steel chambers before deoxygenation took place by continuous flushing as described above. A map of approximately 150 cells was drawn before and after exposure to extremely hypoxic conditions in order to count cells and to calculate the mean multiplicity of doublets (i.e. synchronised cells attached as newly divided sister cells).

Flow cytometry

Cell cycle progression was measured from DNA histograms recorded on a laboratory-built high-resolution flow cytometer using a mercury lamp on a Nikon invertoscope (Lindmo & Steen, 1979; Kærn et al., 1990). Cells were trypanosed into single cells, washed with phosphate-buffered saline (PBS) and fixed in 50% ethanol before the samples were stored at 4°C. The day before flow cytometry analysis the cells were resuspended in PBS containing 60 µg ml⁻¹ RNAse 1A (Sigma, USA) and incubated at 20°C overnight and subsequently stained with 17µg ml⁻¹ propidium iodide (Sigma, USA). The fraction of cells in the various phases of the cell cycle was determined by a commercial computer program (Multicycle, Phoenix Flow Systems, USA) using an algorithm based on the work of Dean and Jett (1974).

Cell survival

Cell survival studies were performed as described previously (Åmellem & Pettersen, 1991). Briefly, 200 cells from either exponentially growing cell cultures or cultures synchronised by mitotic selection were seeded in glass dishes (5 cm diameter). The medium content in each dish was reduced to 1.5 ml prior to the hypoxic treatment. After fixation and staining the colonies containing more than 40 cells were scored as survivors.
**Results**

In the following text the expression 'a balanced mixture of deoxynucleosides' will be used for a solution of E2a medium containing 0.1 mM deoxycytidine, 0.01 mM deoxyadenosine, and 0.01 mM deoxyguanosine. The composition of this solution is similar to that previously used by Löfler (1985).

Addition of deoxynucleosides has no effect on cell proliferation under aerobic conditions (data not shown). A higher concentration of deoxycytidine (0.4 mM) was also tested (data not shown), but the results were equal to the effect of 0.1 mM.

**Cell proliferation under extremely hypoxic conditions**

Our previous investigations on NHIK 3025 cells have shown that under extremely hypoxic conditions no cells enter S-phase and most cells within S-phase remain stationary except cells in the very latest part of S-phase, which are able to complete DNA synthesis (Amellem & Pettersen, 1991). The DNA histograms presented in Figure 2 show NHIK 3025 cells, synchronised by mitotic selection, after 20 h of extreme hypoxia, in the absence (Figure 2b and e) or presence (Figure 2c and f) of a balanced mixture of deoxynucleosides (dN). As shown in this figure, the addition of dN has no effect on the
Figure 3 DNA histograms of synchronised NHIK 3025 cells fixed at various time points after mitotic selection. Histogram a represents untreated control cells 16 h after mitotic selection. Histograms b-f represent cells exposed to extreme hypoxia (<4 p.p.m. oxygen) for 20 h starting 16 h after selection when the majority of the cell population had reached late S/G2 + M (about 90%). Histograms b and c represent cells fixed immediately following hypoxia, while histograms d-f represent cells fixed 6 h after reoxygenation. Cell populations were supplemented with medium containing 0.1 mM deoxycytidine (dC) during hypoxia (histograms c and e), during reoxygenation (histogram d) or during both hypoxia and reoxygenation (histogram f).

cell cycle arrest in G1 during extreme hypoxia (<4 p.p.m. oxygen), i.e. the cells remained stationary. Comparing the DNA histogram in Figure 2d with that in Figure 2e shows that no DNA synthesis (S-phase is indicated by the hatched area in Figure 2d,e and f) takes place when these cells are exposed to extreme hypoxia for 20 h. Even if these cells, during hypoxia, were supplied with dN no cell proliferation was detected by flow cytometry (Figure 2f). In order to make sure that no cell division takes place during 20 h of extreme hypoxia in the presence of dN we performed some additional control experiments. Synchronised cells were exposed to extremely hypoxic conditions in a stainless-steel chamber constructed for observation of cells in a microscope during hypoxia. During a 20 h treatment the multiplicity of doublets remained unchanged, i.e. the synchronised cells attached as newly divided sister cells (doublets), indicating that no cell division took place during hypoxia.

However, unexpected results appeared when NHIK 3025 cells were exposed to extreme hypoxia during cell cycle progression through G1 and mitosis in the presence of deoxycytidine (Figure 3). Recently, we showed that following reoxygenation of cells initially exposed to extremely hypoxic conditions in G1, initiation of DNA synthesis following reoxygenation was delayed for several hours (>6 h) as com-
Figure 4  DNA histograms of synchronised NHIK 3025 cells fixed at various time points after mitotic selection. Histograms a, b and f represent untreated control cells in G1 of the first generation (2.5 or 5 h), in G1 of the second generation (22.5 h) or when most cells are in G1/mid-S of the second generation (25 h) respectively. Histograms c-e and g-i represent cells fixed 20 h after exposure to 100 p.p.m. (0.01%) oxygen starting in early G1 (2.5 h after selection) or late G1 (5 h after selection) respectively. Cell populations were supplemented with medium containing either a balanced mixture of deoxynucleosides (dN) (histogram e), 0.1 mM deoxycytidine (dC) (histogram i), 0.1 mM uridine (U) (histogram b) or 0.01 mM deoxyadenine and deoxyguanine (dA,dG) (histogram d) at the start of the hypoxic exposure.

pared with initiation after reoxygenation of cells initially exposed to extremely hypoxic conditions in G1 (Amellem & Pettersen, 1993). Figure 3 shows DNA histograms of synchronised cell populations preceding and following 20 h exposure to extreme hypoxia and 6 h after reoxygenation. The histogram in Figure 3a represents control cells 16 h after mitotic selection, i.e. at the start of the hypoxic exposure, when about 61% of the cell population has reached G2 and about 33% late S-phase (hatched area). During hypoxia these cells become arrested in G1 (about 93%) in the subsequent cell cycle (Figure 3b) independent of the presence of deoxycytidine following 20 h exposure to extreme hypoxia (Figure 3c). Deoxycytidine had no effect on the delayed entry into S-phase if added during reoxygenation (Figure 3d). However, the unexpected finding was that addition of deoxycytidine during the hypoxic exposure in G2 abolished the delayed entry into S-phase for 60% of the cell population after reoxygenation (Figure 3e and f). In fact, the mean DNA
Figure 5 DNA histograms of synchronised NHIK 3025 cells fixed at various time points after mitotic selection. Histograms a and f represent untreated control cells fixed either 3 h or 23 h (most cells are in G1 of the second generation) after mitotic selection respectively. Histograms b–e represent cells fixed 20 h after exposure to 1,300 p.p.m. (0.13%) oxygen starting in mid-G2 (3 h after selection). Cell populations were supplemented with plain medium or medium containing either a balanced mixture of deoxynucleosides (dN) (histogram e), 0.1 mM deoxycytidine (dC) (histogram d) or 0.1 mM uridine (U) (histogram e) at the start of the hypoxic exposure.
content per cell in the reoxygenated subpopulation in S-phase (Figure 3e and f) was 1.6 times that of G1 cells, indicating that these cells synthesise DNA at approximately the same rate as normoxic cells (Åmellem & Pettersen, 1993).

**Cell proliferation under 100 p.p.m. oxygen**

Figure 4 shows DNA histograms of synchronised cells exposed to 100 p.p.m. oxygen for 20 h starting either early or late in G1 (i.e. 2.5 or 5 h after mitotic selection). Under these hypoxic conditions most of the cells completed half (85%) of the cell population (Figure 4e) or two-thirds (90%) of the cell population, Figure 4g) of S-phase. Addition of (deoxy)nucleosides during hypoxia increased the rate of cell cycle progression significantly. Most of the cells supplied with (deoxy)nucleotides during hypoxia reached either G1 + M or G0 of the subsequent cell cycle (Figure 4e, h and i). However, despite the addition of (deoxy)nucleosides cell proliferation is still slower than under normoxic conditions (compare Figure 4b with 4e). Addition of 0.01 mM deoxycytidine and deoxyguanosine alone had no effect on the hypoxia-induced inhibition of DNA synthesis (compare Figure 4c with 4d). Thus, only the pyrimidines (deoxycytidine and uridine) and not the purines (deoxyadenine and deoxyguanosine) could stimulate cell cycle progression under these hypoxic conditions. More surprising was the effect of uridine (Figure 4h), which has to be converted by ribonucleotide reductase in order to gain deoxynucleotides, i.e. dUDP and dCDP. The histogram in Figure 4h clearly shows that uridine restimulates cell proliferation in cells arrested by hypoxia-induced inhibition of DNA synthesis to the same extent as deoxycytidine (Figure 4i).

The comparison between cells exposed to hypoxia while in early and late G1 revealed a difference in the ability to traverse S-phase in the presence of deoxynucleotides. The fraction of cells that was able to complete DNA synthesis during hypoxia was greater if cells were in late than in early G1 at the onset of hypoxia (compare Figure 4i with 4e).

**Cell proliferation under 1,300 p.p.m. oxygen**

According to the work by Froese (1968) cellular respiration is the same under 1,300 p.p.m. oxygen as under normoxic conditions. In our earlier studies we concluded that there was no inhibition of cell cycle progression for oxygen concentrations above 1,000 p.p.m. (Pettersen & Lindmo, 1983). However, in those experiments, treatment times were short: only 3 h. In the present report we have increased the treatment time up to 20 h.

The DNA histograms presented in Figure 5 show synchronised cells exposed to 1,300 p.p.m. oxygen starting in mid-G1 (3 h after mitotic selection). We found that most of the synchronised cells accumulate in mid-S-phase (about 70%) under these conditions (Figure 5b) and conclude that the rate of cell proliferation was reduced to much the same level when cells were exposed to 1,300 compared with 100 p.p.m. oxygen. In the presence of a balanced mixture of deoxynucleosides (Figure 5c), deoxycytidine (Figure 5d) or uridine (Figure 5e) under these hypoxic conditions the accumulated cells in S-phase were able to complete DNA synthesis. However, cell cycle progression was still reduced in the presence of these (deoxy)nucleosides (Figure 5c, d and e) as compared with normoxic conditions (Figure 5f). Again the positive effect of uridine (Figure 5e) on cell cycle progression under hypoxic conditions was equally effective as deoxycytidine (Figure 5d) or the balanced mixture of deoxynucleosides (Figure 5c). Comparing the effect on cell proliferation under these hypoxic conditions of the balanced mixture of deoxynucleosides (Figure 5c) with the effect of deoxycytidine alone (Figure 5d) revealed that deoxycytidine alone is responsible for the observed effect, indicating that deoxyadenine and deoxyguanosine had no additional effect on cell progression under 1,300 p.p.m. oxygen.

Synchronised NHIK 3025 cells exposed to 5,000 p.p.m. oxygen (data not shown) did not cause any significant delay in cell cycle progression within the time period of 20 h, and thus supplementation of deoxynucleosides did not have any effects under these hypoxic conditions.

**Cell survival**

We have previously shown that the ability of NHIK 3025 cells to survive under conditions of extreme hypoxia depends on the position in the cell cycle, with cells in S-phase being highly sensitive (Åmellem & Pettersen, 1991). Virtually all cells exposed to such low oxygen concentration while in S-phase for 20 h are inactivated, while cell survival for exponentially growing (i.e. asynchronous) cells is about 30% (see Figure 6). In contrast, when exponentially growing cells were exposed to 100 or 1,300 p.p.m. oxygen for 20 h, the ability to form colonies was almost equal to that seen under aerobic conditions, 90 ± 5% and 97 ± 8% respectively (Figure 6). Flow cytometry showed that synchronised cells exposed to hypoxic conditions in G1 had reached mid- to late S-phase in these experiments (see Figures 4 and 5). We observed no differences between exponentially growing and synchronised cells concerning their ability to survive exposure to low oxygen concentrations ranging from 100 to 5,000 p.p.m. This indicates that there are no phase-specific differences in cell cycle sensitivity to cells exposed to 100 p.p.m. oxygen or more. Thus, whereas cells in S-phase are completely inactivated after prolonged exposure to 4 p.p.m. oxygen, only slightly higher oxygen concentrations are without toxicity. Not surprisingly, the addition of deoxynucleotides to either of these hypoxic or aerobic cell cultures had no effect on cell survival (data not shown).

**Discussion**

Our previous investigations have shown that under extremely hypoxic conditions [i.e. <4 p.p.m. oxygen, which according to the work by Froese (1962, 1968) and Boag (1970) reduces respiration essentially to zero] no cells enter S-phase and most cells within S-phase remain stationary except cells in late S, which are able to complete DNA synthesis (Åmellem & Pettersen, 1991). The progression from G1 to G0 is slightly halted, but cell division is completed successfully without the
presence of oxygen (Åmelllem & Pettersen, 1993; Pettersen et al., 1986). The transit time from the completion of mitosis to late G1 is constant, independent of the presence of oxygen, until the cells accumulate in an oxygen-sensitive restriction point near the G1/S border (Åmelllem & Pettersen, 1993).

Cell proliferation during extremely hypoxic conditions

In the present study we find that addition of a balanced mixture of deoxynucleosides has no effect on the arrest maintained in G1 or on the inhibition of cell proliferation in S-phase under extremely hypoxic conditions (Figure 2). Similar results were observed with Ehrlich ascites cells grown in suspension culture (Löffler, 1987). The cell cycle arrest induced by extreme hypoxia in late G1 is presumably not a consequence of ATP depletion, since the addition of deoxycytidine stimulated reprogramming of antimycin-arrested, but not hypoxia-arrested, cells (Löffler, 1985a). Both treatments reduced the ATP pool to 50–60% of the normal value.

In a previous study (Åmelllem & Pettersen, 1993) we showed that cells initially in G2 at the onset of extreme hypoxia completed cell division and were arrested in G1 during hypoxia, but that initiation of DNA synthesis was delayed for more than 6 h following reoxygenation. In contrast, cells initially in G2 at the onset of hypoxia initiated DNA synthesis within 1.5 h after reoxygenation. The delayed initiation of DNA synthesis after reoxygenation of cells initially in G2 at the onset of hypoxia is completely abolished for about half of the cell population if deoxycytidine is present during the extremely hypoxic exposure (Figure 3). Furthermore, the rate of DNA synthesis of the subpopulation that was stimulated after reoxygenation owing to the presence of deoxycytidine during hypoxia is approximately equal to the rate previously observed in normoxic cells (Åmelllem & Pettersen, 1993). This strengthens our previous proposal that it is important for DNA synthesis to be located in G1 in the previous cell cycle, and that they are, at least partly, connected to reduced de novo synthesis of pyrimidine precursors and/or to the regulation of ribonucleotide reductase (see below).

The rate of entry into S-phase was affected by the presence of deoxycytidine during hypoxia, but was unaffected by the presence of deoxycytidine after reoxygenation (Figure 3). A similar observation was noted in Ehrlich ascites cells (Löffler, 1989), indicating that the presence of high amounts of dC during completion of G2 mitosis and early G1 under hypoxic conditions helps the cells to restore (or maintain) their machinery for initiation of DNA synthesis during these phases. The crux of the matter may be that addition of dC helps to restore the dNTP pools even during hypoxia since, as illustrated in Figure 1, further metabolism of dC does not involve any oxygen-dependent steps. Thus, low levels of dCTP and dTTP after prolonged exposure to extreme hypoxia may play an important role in determining the rate with which the cells initiate DNA synthesis after reoxygenation. In this context it is interesting to refer to the early observations by Walters et al. (1973) and Skoog et al. (1973) showing that the pool of dCTP is degraded in G2 and mitosis. As a consequence, this pool must be restored before a new initiation of DNA synthesis can take place. A cell cycle-dependent pattern similar to that obtained for dCTP was also obtained for both mRNA transcripts of ribonucleotide reductase: in each case levels increased as cells progressed into S-phase, and then declined when they progressed into G2 phase and mitosis (Björklund et al., 1990). However, the role that the cell cycle-dependent regulation of both the dCTP pool and initiation of ribonucleotide reductase play in S-phase remains uncertain. It is possible that the reduced dCTP pool may play under hypoxic conditions remains to be elucidated. Several possible mechanisms involved in the cell cycle arrest during extreme hypoxia remains to be elucidated. To our knowledge no evidence points to possible oxygen-dependent steps and disturbances along the salvage pathway that could block the import and synthesis of pyrimidine deoxynucleotides under extremely hypoxic conditions. On the contrary it has been shown that the dCTP pool is three-quarters restored by the addition of deoxycytidine under hypoxic conditions (Löffler et al., 1983). Therefore, the most likely explanation is that some sort of mechanism is activated or inactivated to prevent cells from the lethal effect of initiating DNA synthesis under conditions of extreme hypoxia. In this context it is interesting to note that both 'normal' cell lines and transformed cell lines such as NIH3T3, lacking the normal restriction point controls in G1 (i.e., serum starvation) (Ramengi & Pettersen, 1984, 1985), respond to oxygen depletion in a similar manner. Thus, cancer cells have not been selected to bypass this highly specific arrest point.

Cell proliferation under moderately hypoxic conditions

Under more moderately hypoxic conditions (100–1,300 p.p.m. oxygen), cell cycle arrest in late G1 is less pronounced than under extreme hypoxia, but the rate of DNA synthesis is highly reduced and most cells are unable to complete DNA synthesis under these hypoxic conditions (Figure 4 and 5). However, the presence of deoxycytidine or uridine during moderate hypoxia almost abolishes the inhibition of DNA synthesis but does not establish the normal rate of cell cycle progression. This was previously observed for cells grown in suspension culture, i.e. Ehrlich ascites cells (Löffler, 1987, 1989, 1992; Probst et al., 1989) and indicates that reduced cell proliferation in these and NIH 3T3 cells under conditions of moderate hypoxia (i.e. such as 100 and 1,300 p.p.m. of oxygen) is mainly the result of an insufficient supply of pyrimidine nucleic acid precursors, particularly deoxycytidine (dC can be converted to dTTP in the absence of oxygen). Since cell cycle inhibition was not completely abolished, additional mechanisms regulating the rate of cell proliferation must be present under hypoxic conditions, and there may also be some variation between different cells. In contrast to our results with NIH 3T3 cells, addition of a balanced mixture of deoxycytidine, deoxyadenine and deoxyguanine in 1,300 p.p.m. oxygen was much more efficient than uridine alone in restimulating DNA synthesis in Ehrlich ascites cells (Löffler, 1992). Of particular interest are the data of Figures 4 and 5 showing that the reduced rate of DNA synthesis in the presence of 100 or 1,300 p.p.m. oxygen respectively [where cell respiration is probably not hampered (Froese, 1968; Boag, 1970)] is almost completely counteracted by separate addition of either deoxycytidine or uridine. Both these nucleosides can be used for DNA synthesis through the salvage pathway, while only uridine has to pass through the main route via ribonucleotide reductase (see Figure 1). These results indicate that the reduction of CDP to dCDP by ribonucleotide reductase does not seem to be the limiting step responsible for the reduced dCTP pool observed under moderate hypoxia. Thus, it is more probable that the reduced DNA synthesis observed in NIH 3T3 cells in the presence of 100 or 1,300 p.p.m. oxygen is due to inhibition of the respiratory chain-dependent UMP synthesis at the stage of dihydroorotate dehydrogenase.

Although the dNTP pools are in general sufficient for only a few minutes of DNA synthesis (Reichard, 1988), cells are still able to progress to mid-S-phase (approximately 4 h into normal S-phase) under 100 p.p.m. oxygen. This indicates that the catalytic activity of ribonucleotide reductase in NIH 3T3 cells, although reduced under low levels of oxygen such as 100 p.p.m., is still capable of reducing as much CDP as needed to replicate half the content of DNA in 20 h. As was previously proposed by Probst et al. (1989), the regulation of the enzyme's activity probably occurs through the concentration of the M2-specific tyrosyl radical. This was supported by the observation that de novo synthesis of the M2 protein correlates with DNA synthesis and that the half-life of the protein is only 3 h, whereas the level of the M1 protein is constant throughout the cell cycle (Engström et al., 1985). In mammalian ribonucleotide reductase prepared from calf thymus the half-life of the M2 radical was
found to be of the order of 10 min under anaerobic conditions (Thelander et al., 1983). Although the continuous presence of oxygen is needed during the catalytic reactions, no specific consumption of oxygen in the catalytic cycle was shown (Thelander et al., 1983). Other groups have shown by use of Electron paramagnetic resonance (EPR) spectroscopy that the intracellular concentration of this radical distinctly decreased under hypoxia and reincreased upon reoxygenation in Ehrlich ascites cells, indicating that oxygen is an essential factor for enzyme activity (Lassmann et al., 1989; Probst et al., 1989).

Both the present and previous results (Probst et al., 1989; Löffler, 1992) point to the size of the dCTP pool and not the size of the purine deoxynucleoside pools as important in regulation of cell proliferation under hypoxic conditions. As shown in Figure 4, addition of deoxadenine and deoxyguanine during various oxygen concentrations showed no effect on hypoxia-induced inhibition of cell cycle progression. This is in accordance with the earlier observation that the concentration of dATP remains unchanged during hypoxia in Ehrlich ascites cells (Löffler et al., 1983). Furthermore, in the present study, addition of purine deoxynucleosides in combination with deoxycytidine to cells exposed to 100 or 1,300 p.p.m. oxygen had no additive effect to the stimulating effect of deoxycytidine alone. However, in Ehrlich ascites cells the combination of deoxynucleosides was slightly more effective than dC alone (Löffler, 1985b). Since degradation of purines is dependent on molecular oxygen (xanthine oxidase oxidises hypoxanthine to xanthine and then to urate), such degradation does not take place under hypoxic conditions and consequently addition of purines has no effect on cell cycle progression during hypoxia (Figure 4e).

Cell survival under hypoxic conditions

We have previously shown that the ability of NIHK 3025 cells to survive exposure to extreme hypoxia is dependent on the position in the cell cycle, with cells in S-phase being most sensitive (Amellem & Pettersen, 1991). However, cell survival of exponentially growing cells exposed to 100, 1,300 or 5,000 p.p.m. oxygen for 20 h was almost equal to that seen under aerobic conditions (Figure 6) and not influenced by the addition of deoxynucleotides (data not shown). Similar results were obtained with Chinese hamster cells (CHL-F), when 100% of the cells survived 690 p.p.m. oxygen for 28 h (Bedford & Mitchell, 1974). Probst et al. (1988) furthermore showed that cell damage and loss of reversibility of the shutdown of replicon initiation occurs in Ehrlich ascites cells at oxygen concentrations below 200 p.p.m.

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