Bioenergetic Pattern of Isolated Type II Pneumocytes in Air and during Hypoxia

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ABSTRACT The bioenergetic pattern of a cell clone derived from rat lung with ultrastructural and biochemical characteristics like those of type II pneumocytes (T-II-P), has been studied in a tissue culture system. During air cultivation, these cells have a high rate of aerobic and anaerobic glycolysis associated with high activities of two rate-limiting enzymes in glycolysis (pyruvate kinase [PyKi] and phosphofructokinase [PFK]). This is present despite the rates of oxygen consumption and activities of cytochrome oxidase (CyOx) similar to other lung cells. Presumably the high rate of aerobic glycolysis explains the substantial lactate production previously described in lung slices and in the intact perfused lung.

Hypoxic cultivation results in a decrease in CyOx. Acute re-exposure to air does not restore the oxygen consumption to normal, presumably as a result of decreased mitochondrial O₂ utilization associated with decreased CyOx activity. As a result, hypoxically cultivated T-II-P cells have a decreased capacity for mitochondrial ATP generation in air as compared to air-cultivated cells. During hypoxia, aerobic and anaerobic glycolysis are further increased as well as the activities of PyKi and PFK.

The high rate of glycolysis and high activities of PyKi and PFK in cultivated T-II-P appear to reflect intrinsic genetic regulation. The decreased CyOx activity and increased PyKi and PFK activities in hypoxic T-II-P appear to reflect alterations in enzyme biosynthesis/biodegradation regulated by O₂ availability.

INTRODUCTION

Type II pneumocytes (T-II-P)¹ play a critical role in alveolar function. It is generally accepted that these cells are the site of synthesis and storage of surface active material in the lung (1, 2). In addition, recent work suggests that these cells have an important repair function and may replace type I pneumocytes in several forms of alveolar injury (3, 4).

Cell function is dependent on metabolic energy provision and therefore, the metabolic characteristics of this cell type are of interest. Furthermore, metabolic alterations in T-II-P in response to decreased O₂ availability may be an important determinant of alveolar function in disease states associated with alveolar hypoxia.

Until recently, precise characterization of the metabolic functions of this cell type has not been possible. This has resulted from inability to obtain pure cell preparations which are sufficiently normal to examine metabolic parameters. The use of cell cloning as described by Douglas and Kaign (5) has provided such a preparation. In the present study, a cloned, T-II-P epithelial cell preparation obtained from rats has been used to examine the energy metabolism of these cells during maintenance in air and under conditions of chronic hypoxia.

Hypoxic exposure results in decreased rates of oxygen utilization associated with decreased activities of a key mitochondrial enzyme, cytochrome oxidase (CyOx).

¹Abbreviations used in this paper: CyOx, cytochrome oxidase; F12-10% FCS, Hams F12 tissue culture medium (pH = 7.4) supplemented with 10% fetal calf serum; PFK, phosphofructokinase; PyKi, pyruvate kinase; O₂, oxygen consumption; T-II-P, type II pneumocytes.
The data show that T-II-P possess a high rate of aerobic and anaerobic glycolysis associated with high activities of two rate-limiting glycolytic enzymes, pyruvate kinase (PyKi) and phosphofructokinase (PFK). The exposure of these cells to hypoxic conditions results in the augmentation of the rates of aerobic and anaerobic glycolysis as well as further increases in the activities of the glycolytic enzymes. This bioenergetic adaptation thus provides a possible mechanism for maintenance of cell viability when mitochondrial energy provision is decreased under conditions of alveolar hypoxia.

METHODS

Cell preparation. Cloned type II pneumocytes were prepared as previously described (5). Cell monolayers were initially maintained in T-150 tissue culture flasks (Corning Medical, Corning Glass Works, Medfield, Mass.) at 37°C in 95% air-5% CO₂. The incubating media used initially, and at all subsequent cultivation times, was Hams F12 tissue culture medium (pH = 7.4) supplemented with 10% fetal calf serum (F12-10% FCS). For subculture, the cell monolayers were suspended in F12-10% FCS and 1 ml of cell suspension containing 50–100,000 cells was placed in T-75 culture flasks to which 24 ml of F12-10% FCS was added. The flasks were then incubated in 95% air-5% CO₂ until confluence. In preliminary experiments, it was determined that this population density (50–100,000 cells/T-75 flask) resulted in confluent cell culture flasks after 7 days of in vitro maintenance.

At the time of confluence, two T-75 flasks were harvested (as described below) for either metabolic or enzymatic studies. The T-II-P harvested at confluence were considered the control cells. Enzymatic and metabolic data obtained at this time are referred to as time zero values.

The remaining cell monolayers were refed with 25 ml F12-10% FCS and incubated in stoppered T-75 flasks under either aerobic (PO₂ ~ 140 torr) or hypoxic (PO₂ ~ 15 torr) conditions for an additional 48–96 h. All cultures were refed and regassed at 24 and 72 h.

Hypoxia was achieved by gassing culture flasks with 95% nitrogen-5% CO₂. Multiple measurements of media PO₂ in the hypoxic flasks (Corning 165 pH/Blood Gas Analyzer, Corning Medical, Corning Glassworks) indicated that the PO₂ was maintained at values of ~15 torr throughout the 96-h cultivation period. Aerobic conditions were achieved by gassing with 95% air-5% CO₂. Multiple measurements of media PO₂ in the aerobic flasks confirmed that the PO₂ was maintained at ~140 torr. Media pH was measured (Corning 165 pH/Blood Gas Analyzer) at the different cultivation times and no significant differences between the aerobic and hypoxic systems were noted.

Flasks were harvested at 48 and 96 h for either metabolic or enzymatic studies. In addition, identically prepared culture systems were processed as described (5) for ultrastructural confirmation of cell types.

Enzyme studies

Cell preparation. At the time of harvest (48 and 96 h), the medium was removed and monolayers incubated in 3.5 ml of a dispersal medium (collagenase-trypsin-chicken serum) for 10–15 min. Cell dispersal was verified by direct observation and 6.5 ml of F12-10% FCS added to inactivate proteolytic enzymes. The monolayers were then harvested by a rubber policeman as a 10-ml suspension and an aliquot of this suspension was used for quantitation of cell number. The harvested cells were centrifuged at 700 g for 10 min (2–4°C), washed twice with phosphate-buffered saline (pH = 7.4), suspended in 1.2 ml, 35 mM tris buffer (pH = 7.4), and disrupted by sonication (Heat Systems-Ultrasonics, Inc. Plainview, N. Y.) for 30 s at a power output of 70 W. PyKi, PFK, CyOx, and protein analyses were performed on these cell sonicates.

Enzyme assay

PyKi activity was assayed by a modification of the spectrophotometric method of Valentine and Tanaka (6). A 0.05-ml sample of cell sonicate was added to a 2.95-ml reaction mixture containing: 0.5 ml of 50 mM triethanolamine-HCl buffer (pH = 7.5), 1.45 ml of distilled water, 0.1 ml of 2.25 M KCl, 0.1 ml of 240 mM MgSO₄, 0.2 ml of 6 mM ADP, 0.1 ml of lactate dehydrogenase (18 enzyme units), 0.4 ml of 1.3 mM NADH, and 0.1 ml of 45 mM phosphoenolpyruvic acid (trisodium salt). NADH oxidation at 25°C was followed by measuring the decrease in optical density at 340 nm in a model DB spectrophotometer (Beckman Instruments, Inc., Fullerton, Calif.). Enzyme activity was calculated during a period of zero order kinetics using a molar extinction coefficient of 6.2 × 10⁴ for NADH. Enzyme activity was expressed both as units (micromoles of phosphoenolpyruvate converted to pyruvate per minute) per milligram of cell protein and as units per 10⁶ cells.

CyOx was measured spectrophotometrically by techniques described (7). A 0.1-ml sample of cell sonicate was added to 2.9 ml of reaction mixture containing: 0.3 ml of 10% sodium deoxycholate, 2.5 ml of 13 mM phosphate buffer (pH = 7.4), and 0.1 ml of 3% solution of cytochrome c (reduced by sodium hydrosulfite). Oxidation of cytochrome c at 25°C was followed by measuring the decrease in optical density at 550 nm in a model DB spectrophotometer. Enzyme activity was calculated using a molar extinction coefficient of 19.1 × 10⁴ for cytochrome c and units expressed as micromoles cytochrome c oxidized per minute per milligram of protein and per 10⁶ cells.

PFK was assayed spectrophotometrically by following the conversion of fructose-6-phosphate to fructose-1,6-diphosphate (8). A 0.1-ml sample of cell sonicate was added to a reaction mixture containing 1.735 ml of 0.1 M Tris-HCl, pH 8.2, 0.05 ml of 0.1 M MgCl₂, 0.2 ml of 0.1 M Na₃PO₄, 0.175 ml “enzyme mix”, 0.56 ml of “substrate mix”, and 0.27 ml NADH (1.4 μmol/ml). [Enzyme mix: 4.92 ml of 0.1 M Tris-HCl, pH = 8.2, 0.6 ml aldolase [10 mg/ml]]. [Substrate mix: 0.1 M fructose-6-phosphate; 0.1-M ATP].

NADH oxidation was followed spectrophotometrically at 25°C and enzyme activity calculated during a period of zero order kinetics. Enzyme activity was expressed both as units (micromoles of fructose-6-phosphate converted to fructose-1,6-biphosphate per minute) per milligram of protein and as units per 10⁶ cells.

Aerobic and anaerobic lactate production

Fresh media (F12-30% FCS, 10 mM glucose, pH = 7.4) was added to T-75 flasks at the time of study, and paired stopped flasks were incubated under aerobic (95% air-5% CO₂) or anaerobic (95% N₂-5% CO₂) conditions for 150 min, at 37°C. Aliquots (0.5 ml) of incubating media were removed from the flasks after 30, 90, and 150 min of incubation and immediately added to an equal volume of chilled
3.5% perchloric acid. Lactate production was relatively linear over this time period and the change in lactate concentration from 30 to 150 min was used to calculate the rate of lactate production. The deproteinized samples were centrifuged at 1,000 g and the resultant supernate was used for lactate analyses (9). The cell monolayers were then harvested for cell counting and the final cell pellet was suspended in 1.2 ml of 35 mM Tris. Protein determinations were performed on this cell suspension and lactate production expressed as micromoles lactate produced per hour per milligram protein and per 10^6 cells.

**Oxygen consumption**

T-II-P were harvested and counted, washed twice with F12, and the cell pellets were suspended in 8 ml of F12 (pH = 7.4). 4-ml aliquots of this cell suspension were used to measure O₂ consumption in the YSI 53 O₂ monitor (Yellow Springs Instrument Co., Yellow Springs, Ohio). The cell suspension was equilibrated with room air (PO₂ ~ 150) before measurement and the decrease in O₂ concentration was then determined over a 15-min time period; the O₂ consumption was linear over this time period. Total cell protein of these aliquots were determined and the QO₂ expressed as micromoles oxygen consumed per hour per milligram protein and per 10^6 cells.

**Protein analysis**

Protein content was determined by the method of Lowry et al. (10) using crystalline human albumin as the standard.

**Cell quantitation**

Cell counts were performed in standard fashion using a hemocytometer. Cell viability was assessed using Trypan blue and was >95% at all times.

**Fibroblast studies**

To examine the specificity of the bioenergetic properties demonstrated in T-II-P, a separate series of studies was carried out using cultivated rat lung fibroblasts. Rat lung fibroblasts were obtained using a modification of the method of Balin et al. (11). Whole rat lungs were dissociated (using a collagenase-trypsin-chicken serum medium) into single viable cells. The resulting cell suspension was inoculated into T-75 culture flasks at 5 x 10^5 cells/T-75 flasks (F12-10% FCS, 37°C, 95% air-5% CO₂) and maintained to semi-confluency. Serial passage (three times) of these primary cultures at relatively high density (5 x 10^5 cells/T-75 flask) was then carried out. Under these conditions, fibroblasts replicate rapidly and become the dominant cell type.

The cell populations obtained by this technique have the morphologic characteristics of fibroblasts on light microscopy and demonstrate polymerized collagen between cells on electron microscopy. No lamellar bodies or surfactant associated markers are demonstrable.

In the present studies, the isolated fibroblasts were used to prepare confluent monolayers in T-75 culture flasks (F12-10% FCS, 37°C, 95% air-5% CO₂) and cells processed for metabolic and enzyme studies as described above.

**RESULTS**

Table I summarizes data with respect to O₂ consumption and CyOx activity in control (confluent) cells and in cells cultivated up to 96 h in air and in a hypoxic environment. After confluence (time zero), T-II-P cultivated in air show a significant increase in total QO₂. The QO₂ per 10^6 cells increases by ~75% during the 96 h of aerobic cultivation (from 0.15±0.03 to 0.26 ±0.07 µm [P < 0.01]) [The QO₂ per milligram protein increases by ~50% during this period. The change in QO₂ per milligram protein is less than the change per 10^6 cells because of a relative increase in cell protein during the period of cultivation. As with QO₂, changes in glycolysis and enzyme activity per 10^6 cells were not quantitatively identical with changes per milligram protein because of increases in cell protein during cultivation. To establish that observed alterations do not merely reflect variable changes in cell protein or cell number under the different culture conditions, all data is expressed both per milligram protein and per 10^6 cells.]

The changes in QO₂ are accompanied by significant

| Table I |
|---|---|---|---|---|---|---|---|---|
| | Aerobic cultivation PO₂ ~ 140 | | | Hypoxic cultivation PO₂ ~ 15 | | | |
| | 0 h | 48 h | 96 h | 0 h | 48 h | 96 h |
| O₂ consumption | | | | | | | |
| µm/mg protein | 0.52±0.07 † | 0.69±0.09 | 0.70±0.13 | 0.52±0.07 | 0.32±0.03 § | 0.23±0.08 $ |
| µm/10^6 cells | 0.15±0.03 | 0.23±0.07 | 0.26±0.07 | 0.15±0.03 | 0.11±0.07 ‡ | 0.10±0.07 $ |
| CyOx | | | | | | | |
| U/mg protein | 9.2±1.3 | 9.6±1.7 | 12.4±1.1 | 9.2±1.3 | 7.8±1.2 § | 7.1±1.1 $ |
| U/10^6 cells | 3.0±0.6 | 4.1±0.9 | 5.8±1.7 | 3.0±0.6 | 3.1±0.6 § | 3.7±0.8 $ |

* Micromoles O₂ consumed per hour.
† Mean±SD.
‡ P < 0.01 (compared to air-cultivated cells).
§ P < 0.05 (compared to air-cultivated cells).
$ n = 7 for all studies.

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increases in CyOx activity at 96 h, measured either per milligram of cell protein (from 9.2 ± 1.3 to 12.4 ± 1.1, \( P < 0.01 \)) or per \( 10^6 \) cells (from 3.0 ± 0.6 to 5.8 ± 1.7, \( P < 0.05 \)).

Confluent T-II-P cells appear to undergo increases in oxygen metabolism similar to those noted during in vitro cultivation in other cell systems such as WI-38 cells (12). Increased respiration is accompanied by parallel increases in CyOx as is true in a wide variety of cell and organ systems (7, 13). Assuming that \( \approx 75\% \) of total \( O_2 \) consumption reflects mitochondrial \( O_2 \) uptake, mitochondrial \( O_2 \) utilization at 96 h is similar to that in at least one other lung cell studied under identical conditions, the alveolar macrophage (13).

Because of the temporal changes that occur during in vitro maintenance under standard (aerobic) conditions, an assessment of the effect of any variable on metabolic parameters requires time-matched analysis. During hypoxia, total \( QO_2 \) significantly decreases so that at 96 h, the hypoxic T-II-P reexposed to air used only about one-third of the total \( O_2 \) consumed by the time-matched air-maintained cells (air-0.70 ± 0.13 \( \mu \)mol/mg protein; hypoxia-0.23 ± 0.08 \( \mu \)mol, \( P < 0.01 \)). These differences in \( QO_2 \) are associated with significant differences in CyOx activities between the air-maintained and hypoxic T-II-P.

In addition, there are absolute decreases in \( QO_2 \) after hypoxia (time 0-[confluent]-0.52 ± 0.07 \( \mu \)mol/mg protein; 96 h hypoxia-0.23 ± 0.08 \( \mu \)mol, \( P < 0.01 \)). This absolute decrease in \( QO_2 \) suggests that the changes in \( O_2 \) consumption are not merely an inhibition of development of mitochondrial \( O_2 \) utilization but reflect a true change in bioenergetic pattern during hypoxia.

Table II summarizes the data concerning glycolytic rates. The rate of aerobic glycolysis is high in the control cells (time 0) averaging almost 3 \( \mu \)mol lactate/mg cell protein per h. This is a value which is 10 times as high as aerobic lactate production studied under identical conditions in another lung cell type, the alveolar macrophage, and is three times the value found in peritoneal macrophages (13).

Unlike respiration, there is no significant increase in either aerobic or anaerobic glycolysis (per milligram cell protein) during maturation in air. The ratio of anaerobic to aerobic glycolysis is \( \approx 150\% \) in control cells (time 0) and is unchanged after maturation.

In contrast to the data in air-cultivated cells, there is a striking and progressive increase in both aerobic and anaerobic glycolysis after cultivation under hypoxic conditions. Both aerobic and anaerobic lactate production are significantly greater in T-II-P exposed to hypoxia as compared to those maintained in air. In fact, after 96 h of hypoxic cultivation, the rate of aerobic glycolysis is as high as the rate of anaerobic glycolysis in the air-cultivated cells. The ratio of anaerobic to aerobic glycolysis, however, is similar to that in the air-cultivated cells.

Table III summarizes data with respect to temporal changes in two rate-limiting glycolytic enzymes. The specific activities of both PyKi and PFK are relatively high in control cells, (as compared, for example, to alveolar macrophages [13]), but do not change significantly during air incubation. It appears that one mechanism for the high rates of glycolysis in this cell type are the high activities of rate-limiting glycolytic enzymes.

During hypoxic cultivation, there are dramatic increases in the activities of both PyKi and PFK which parallel the changes in glycolytic capacity. These increases in enzyme activities presumably subserve enhanced glycolytic energy provision under conditions of decreased \( O_2 \) availability.

The results of the metabolic and enzyme studies carried out on confluent fibroblasts are shown in Table IV. The equivalent T-II-P data is shown for comparison. Rates of aerobic and anaerobic glycolysis are significantly less in the cultivated fibroblast. The lower

| TABLE II |
| Changes in Aerobic and Anaerobic Glycolysis* in T-II-P Maintained Under Aerobic and Hypoxic Conditions |

<table>
<thead>
<tr>
<th></th>
<th>Aerobic cultivation ( PO_2 \sim 140 )</th>
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<th>Hypoxic cultivation ( PO_2 \sim 15 )</th>
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<tbody>
<tr>
<td></td>
<td>0 h</td>
<td>48 h</td>
<td>96 h</td>
</tr>
<tr>
<td>Aerobic glycolysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \mu )mol/mg protein</td>
<td>2.89 ± 0.52</td>
<td>2.17 ± 0.55</td>
<td>2.64 ± 0.43</td>
</tr>
<tr>
<td>( \mu )mol/10⁶ cells</td>
<td>1.00 ± 0.26</td>
<td>0.97 ± 0.31</td>
<td>1.40 ± 0.28</td>
</tr>
<tr>
<td>Anaerobic glycolysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \mu )mol/mg protein</td>
<td>3.85 ± 0.53</td>
<td>3.11 ± 0.46</td>
<td>3.71 ± 0.40</td>
</tr>
<tr>
<td>( \mu )mol/10⁶ cells</td>
<td>1.30 ± 0.21</td>
<td>1.40 ± 0.23</td>
<td>2.02 ± 0.52</td>
</tr>
</tbody>
</table>

* Micromoles of lactate produced per hour.
\( \dagger \) \( P < 0.05 \) (compared to air cultivation).
\( \ddagger \) \( P < 0.01 \) (compared to air cultivation).
\( n = 5 \) for all studies.
rates of glycolysis are accompanied by lower activities of PyKi and PFK. O2 consumption and CyOx activities are similar in the two cell types.

DISCUSSION

Studies of isolated lung slices (14) and of the isolated perfused lung (15, 16) have shown what appeared to be an anomaly. Despite exposure in situ to relatively high O2 tensions, (alveolar PO2 ~ 100 torr) lung slices and intact lung have a substantial rate of lactate production. This finding suggests the existence of a significant population of lung cells with a high rate of aerobic glycolysis. In the present studies, T-II-P are shown to possess high rates of glycolysis despite the fact that these cells normally develop and function at PO2 of ~100 torr. The rate of aerobic glycolysis in the T-II-P is 30–40 times the rate of aerobic glycolysis found in the isolated perfused lung (15, 16) or in tissue slices (14). Differences in the experimental conditions could account for some of this difference. However, the high rates of glycolysis observed in T-II-P suggest that this cell type is, at least in part, responsible for the glycolytic characteristics of intact lung and lung slices.

The validity of this interpretation depends on several assumptions. One is that the cloned cell type used is truly a T-II-P. The evidence for this includes: (a) the light and ultrastructural characteristics of these cells are similar to those of type II cells in situ, particularly with respect to the existence of multiple osmiophilic lamellar bodies, (b) lamellar bodies isolated from these cells resemble those isolated from in situ type II cells in morphologic appearance (5). In addition, three key enzymes in the synthesis of surface active material are present in the cultured cells with activities of choline kinase, cholinephosphate cytidylyltransferase, and cholinephosphotransferase increased as high as 10-fold, relative to whole lung homogenates (17). Further, there is significant activity of 1-acyl 2-lyso

<table>
<thead>
<tr>
<th>PyKi</th>
<th>PFK</th>
<th>O2</th>
<th>CyOx</th>
</tr>
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<tbody>
<tr>
<td>0.55±0.10</td>
<td>0.85±0.12</td>
<td>52±15</td>
<td>5.8±1.3</td>
</tr>
<tr>
<td>1.69±0.22</td>
<td>2.69±0.31</td>
<td>176±49</td>
<td>20.4±4.6</td>
</tr>
</tbody>
</table>

* Micromoles of lactate produced per hour.
† Enzyme units.
§ Micromoles O2 consumed per hour.
* P < 0.01 (compared to confluent fibroblasts).
phosphotidylcholine acyl-CoA acyltransferase,\(^2\) an enzyme involved specifically in the biosynthesis of surface-active material.

In addition to these published observations, the morphologic appearance of the cell populations used in the present studies was specifically confirmed and the cells were shown to have the characteristics expected of differentiated T-II-P. The cells cultivated at PO\(_2\) of both \(\sim 140\) and \(\sim 15\) torr demonstrated osmiophilic lamellar bodies. The hypoxic T-II-P did not appear to show significant changes in either size or number of these characteristic inclusion bodies (18).

Precise characterization of most specialized cell types in tissue culture systems are difficult and some reservations are appropriate (19). The use of the cloning technique does, however, insure that the cell population being studied was homogeneous in nature.

The second assumption is that the metabolic activities of these cells are similar to those found in situ. It is well known that cultured cells may transform in vitro and show evidence of dedifferentiation. Unlike differentiated cells, the present cells maintained normal contact properties assuming a confluent pattern with maturation. After confluence, the cells demonstrate a replication rate that is much less than that demonstrated during the log phase growth. Finally, karyotyping indicates that the cells are diploid. It is therefore probable that the metabolic properties of these cells are not the results of tissue culture transformation.

To further evaluate the validity of the assumption that the metabolic activities demonstrated are relatively specific for T-II-P and are not an artifact of tissue culture, a separate series of studies was carried out using cultured lung fibroblasts. The disparity between the rates of glycolysis in the two cultivated lung cell systems in the face of similar rates of O\(_2\) utilization is consistent with the conclusion that the metabolic properties observed in T-II-P reflect a specific property of this cell type rather than a nonspecific property of cultured lung cells.

The rate of aerobic glycolysis in these cells is remarkable. A number of cell types such as cancer cells (Warburg effect), retina, turtle brain, and renal medullary cells (20-22) have been shown to have high rates of aerobic glycolysis. All of these cell types normally exist in a relatively low O\(_2\) environment (tissue PO\(_2\) \(\sim 25\) torr). However, cells in the alveolar wall such as T-II-P, function in an O\(_2\)-rich environment. On the air side, PO\(_2\) values of about 100 torr are present. On the blood side, pulmonary capillary blood is unique (as compared to other capillary beds) in having a rapid increase (rather then decrease) in O\(_2\) tension as blood passes from the (pulmonary) arterial to the (pulmonary) venous end. Thus, a high rate of lactate production is present despite a luxuriant O\(_2\) supply.

One possible explanation for the high rates of aerobic glycolysis is that these cells have an unusually low rate of mitochondrial O\(_2\) utilization in air. This does not appear to be the case. The high rate of aerobic glycolysis in T-II-P is present despite a rate of total oxygen utilization which is comparable to that previously demonstrated in other tissue culture maintained cells (23, 24) and is similar to the rate of O\(_2\) utilization in lung fibroblasts from the same species.

Although it is difficult to interpret measurements of O\(_2\) consumption in isolated cell systems in an absolute sense (the values observed depend heavily on the conditions of measurement, and are influenced by cell density, conditions of incubation, substrate concentrations, etc.), the data suggest that the ability to derive ATP from glycolysis plays some special role in these cells. The lung is often subjected to variable degrees of alveolar hypoxia as the result of disease. It is well known that even after months of atelectasis, alveoli are capable of being reexpanded with preservation of parenchymal integrity. High rates of glycolytic energy provision could presumably play an important role in maintaining cell viability under these conditions.

The finding that the Pasteur effect is only moderate in control (confluent) cells is of particular interest. This finding has been reported in other cell types with high rates of aerobic glycolysis and indicates that such cell types are functioning close to glycolytic capacity despite normal respiration (24).

During hypoxic cultivation, there is a significant decrease in CyOx activity. When these cells are acutely exposed to air (during the measurement of QO\(_2\)), QO\(_2\) is strikingly diminished despite a normal O\(_2\) supply. This observation has two important implications. It indicates that the decrease in CyOx activity most probably reflects a true change in CyOx content rather than merely being a result of changes in low molecular weight regulators or cofactors. Despite reexposure to adequate O\(_2\), QO\(_2\) remains low as compared to control (time zero) cells studied under identical conditions. This must occur because there has been an intrinsic alteration of mitochondrial O\(_2\) utilization. One explanation is that this occurred because of a parallel reduction of CyOx content. In turn, this suggests that molecular O\(_2\) modified the biosynthesis/biodegradation of this enzyme. This type of enzyme regulation has been demonstrated in anaerobic vs. aerobic yeast. In that system, anaerobiosis is associated with the loss of cytochrome oxidase activity (25). Yeast CyOx can be resolved into six or seven polypeptides, the three largest are formed on mitochondrial ribosomes, whereas the others are made on cytoplasmic ribosomes. In the absence of molecular

O₂, two of the mitochondrial peptides are no longer synthesized. Because O₂ is required for the biosynthesis of the complete enzyme, lack of O₂ leads to loss of CyOx activity (26).

The precise mechanism for altered CyOx activity in T-II-P (a mammalian system) has not been studied, but it appears likely that molecular O₂ does play a role in the long-term regulation of CyOx.

The second implication involves the temporal relations of reductions in CyOx activity and reduced O₂. The former is a process which requires hours to days. It therefore seems probable that restoration to control values would also require substantial periods of time. As a result, when the hypoxic cell is restored to a normal O₂ environment it still manifests a profound depression of mitochondrial O₂ utilization with a concomitant reduction in capacity for aerobic ATP generation. Thus, the hypoxically mediated decrease in CyOx, which is presumably part of the bioenergetic adaptations subserving more favorable energetics during hypoxia, limits an optimal energy provision when O₂ availability is increased.

After hypoxic cultivation, there is a substantial increase in the rate of aerobic and anaerobic glycolysis though the relationship between the two remains unaltered. The hypoxically cultivated T-II-P actually has a rate of aerobic glycolysis which is equal to the rate of anaerobic glycolysis in the air-cultivated cell. The augmentation of glycolysis during chronic hypoxia must play a role in the ability of this cell to survive prolonged O₂ depletion.

In considering the basis for the glycolytic properties of these cells, a major finding is the high activities of PyKi and PFK. In the control (time zero) T-II-P, these activities are significantly greater than in the cultivated lung fibroblasts. Glycolytic enzyme activity in these cells is also greater than in both alveolar macrophages cultivated under aerobic conditions (13), and in whole lung homogenates (27). The activities of both PFK and PyKi are important in controlling the rate of glycolysis (28, 29). We have shown in a number of systems that PyKi and PFK activities are quantitative markers for glycolytic capacity (13, 27). A similar relationship has also been demonstrated in tumor cells (30). Presumably, the high activities of these enzymes are intrinsically programmed in T-II-P because all T-II-P cells were only exposed to normal (and not low) O₂ tensions before confluence.

The increase in glycolysis during hypoxia is at least partially mediated by a further increase in the activities of the rate-limiting glycolytic enzymes. This increase in PyKi and PFK, under hypoxic conditions, may in turn be related to another more general mechanism. We have shown that changes in energy metabolism enzyme activities (both glycolytic and those of oxidative phosphorylation) are regulated by O₂ tension in alveolar and peritoneal macrophages (13, 31). The present studies demonstrate the operation of this adaptive mechanism in another cell type.

Enzyme activities rather than enzyme contents were measured. It is possible that the changes in activity with hypoxia could be related to changes in cofactors or low molecular weight regulators of CyOx, PyKi, and PFK. This seems unlikely since (a) enzyme activities were measured under standard conditions by in vitro assay systems and (b) a series of complex changes would be required which both decreased CyOx activity and increase PyKi and PFK activity.

If the content of the various enzymes is changed, the altered enzyme activities may be a result of changes in the rate of biosynthesis or biodegradation of these enzymes as modified by O₂ availability. Thus, the net ability of this cell to withstand hypoxia seems to stem from a combination of (a) intrinsic metabolic characteristics found specifically in T-II-P upon which may be superimposed (b) bioenergetic alterations produced by a general regulatory mechanism involving alterations of enzyme biosynthesis and/or biodegradation evoked by differences in O₂ availability.

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