A Cellular Model of Endothelial Cell Ischemia

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Endothelial cell dysfunction in ischemia may cause increased capillary permeability. We examined the effect of failing ATP synthesis, a major consequence of ischemia, on microfilaments—important structural determinants of the endothelial cell. Glycolytic and mitochondrial ATP synthesis in bovine pulmonary artery endothelial cells was inhibited by glucose depletion and 650 picomole (pmole) oligomycin/μg DNA, respectively. ATP levels were monitored with the luciferase-luciferin assay over a 2-hr time course followed by recovery for 1 hr after removal of the oligomycin and addition of 5.5 mM glucose. ATP levels fell to 83.6 ± 63.8 pmole/μg DNA (n = 11) by 30 min, 26.9 ± 13.8 pmole/μg DNA (n = 11) by 60 min, and 17.2 ± 3.8 pmole/μg DNA (n = 6) by 120 min, whereas control uninjured cells had 541.3 ± 196.8 pmole/μg DNA (n = 6) at 120 min. Fluorescence microscopy of microfilaments stained with rhodamine-phalloidin revealed progressive disassembly and shortening of the microfilaments in >90% of cells over 60 min which correlated with the fall in ATP. Ultrastructural examination revealed that side to side aggregation of microfilaments had occurred over the 120-min time course. Two hours of glucose depletion (305.5 ± 130.8 pmole ATP/μg DNA, n = 6) or oligomycin alone (480.0 ± 90.1 pmole ATP/μg DNA, n = 6) failed to produce the dramatic fall in ATP or the microfilament changes. During cell recovery, there was a rapid reassembly of microfilaments, detected by fluorescence microscopy, which was nearly complete in 85-90% of cells by 45-60 min. ATP levels increased significantly (P = 0.002) to 96.1 ± 36.8 pmole/μg DNA (n = 6) by 30 min. This model should provide insight into the pathogenesis and treatment of the capillary leak seen with ischemia.

INTRODUCTION

Increased capillary permeability is frequently seen in conjunction with acute ischemic tissue injury. Endothelial cell dysfunction as a result of ischemia may be an important factor mediating the changes in permeability. Inhibition of cellular ATP synthesis by depletion of substrate (e.g., glucose) and oxygen is a major biochemical consequence of acute ischemia following interruption of blood flow.

Microfilaments are important structural determinants of cells which are primarily composed of the protein actin, in its polymerized form. ATP is an important regulator of the cycling of actin within the cell between the monomeric (G) and polymeric (F) forms [1, 2]. Cells depleted of ATP have demonstrated progressive disruption of microfilaments which was reversible [2–5].

It is unclear whether the biochemical events (i.e., inhibition of anaerobic and aerobic ATP synthesis) which are major sequelae of acute ischemia following vascular occlusion affect mammalian endothelial cell ATP levels and in turn microfilament organization.

To address this question, we examined the impact of acute glucose depletion (inhibition of glycolysis) and treatment with 650 pmole oligomycin/μg DNA (inhibition of mitochondrial respiration) on the levels of ATP in bovine pulmonary artery endothelial cells as a model of endothelial cell ischemia. The dose of oligomycin, which interferes with electron transport as well as phosphorylation of ADP [6], was derived from previous work...
The effect of this treatment on microfilament organization in endothelial cells was then examined by fluorescence and electron microscopic techniques. The potential reversibility of changes in endothelial cell microfilament organization induced by the metabolic inhibition was also examined and correlated with ATP levels in recovering cells.

**METHODS**

**Cells and culture.** Bovine pulmonary artery endothelial cells (No. AG2791A) obtained from the National Institute of Aging, Aging Cell Culture Repository (Camden, NJ) were cultured at 37°C under 5% CO₂/95% air in medium RPMI 1640 supplemented with 2 mM glutamine (GIBCO), 10% fetal bovine serum (Whittaker, M.A. Bioproducts), 10 mM Heps (Whittaker, M.A. Bioproducts), 100 u/ml penicillin, and 100 μg/ml streptomycin (GIBCO). Cells were grown in 150-cm² flasks (Corning) or on collagen-coated beads (Cytodex 3, Sigma) in a spinner flask (Techne, Cambridge, UK), spinning at 30 rpm. Experiments in which ATP measurements were performed were done with cells on microcarrier beads in suspension or with cells from 150-cm² flasks suspended following a brief exposure (10–15 min) to trypsin (0.05%) and EDTA (0.02%) (Sigma). The cells were suspended at an approximate concentration of 18 μg DNA/ml in a buffer containing 147 mM NaCl, 5 mM KCl, 1.9 mM KH₂PO₄, 1.1 mM Na₂HPO₄, 0.3 mM MgSO₄, 1 mM MgCl₂, 5.5 mM glucose, 10 mM Heps, and 1.5 mM CaCl₂. In some experiments, glucose was absent from the buffer.

Experiments with suspended cells were carried out at 37°C in a shaking water bath.

**DNA measurements.** Endothelial cell DNA content was measured using a fluorometric assay [7, 9]. Briefly, cells were sonicated for 30 sec in a buffer containing 2 M NaCl and then exposed to 1 μg/ml Hoechst dye No. 33258 (Calbiochem). Fluorescence measurements of DNA content were made on a Perkin–Elmer LS5 spectrofluorometer with excitation at 356 nm and emission at 458 nm. Fluorescence measurements were then compared to a standard curve generated using calf thymus DNA (Boehringer–Mannheim). Comparisons between DNA content and cell number obtained from counting cell suspensions in a hemacytometer yielded a relationship of approx 1 μg DNA = 10⁵ cells.

**ATP measurements.** Cellular ATP levels were measured by the luciferase–luciferin method of Stanley and Williams [10] as previously adapted for use with endothelial cells [7] with a few modifications. The luciferase–luciferin (Sigma No. L0633) was reconstituted at 40 μg/ml sterile water (on ice) and diluted 1:50 in a buffer which contained 1% bovine serum albumin, 20 mM glycine, and 2 mM EDTA at pH 8.0. The final working solution was kept on ice and made fresh every 30–45 min. A new standard curve was generated after each batch of diluted enzyme was made. Measurements were made using a Beckman (LS8000) liquid scintillation counter with the photomultiplier tube signals accepted out of coincidence.

**Fluorescence microscopy.** Adherent endothelial cells were grown overnight in six-well plates (Nunc) or on glass coverslips at an approximate density of 1–2 × 10⁵ cells/cm². Metabolic inhibition was carried out after a wash step in which the culture media was removed and replaced with the experimental buffer. At various times throughout the time course of the experiment, adherent cells were stained with rhodamine–phalloidin, a probe specific for the polymerized or F form of actin in microfilaments [11]. The staining procedure [12, 13] was as follows: buffer was removed from the plate and 1 ml per well of a cocktail with final concentrations of 50 μg/ml lysophosphatidyl choline, 1.85% phosphate-buffered formalin, and 165 nM rhodamine–phalloidin in buffer was added to the cells. This was incubated for 10 min at 37°C or 20 min at room temperature. After removal of the excess stain solution, a glass coverslip was sealed to the monolayer in the six-well plates with 5 μl of 90% glycerol, or the glass coverslips with stained cells were...
sealed with the glycerol to microscope slides. The cell monolayers were then examined for their rhodamine fluorescence staining patterns with a Nikon Optiphot fluorescence microscope. Fluorescence micrographs were taken at 400X magnification using Plus X film (Kodak). Fixation of cells with 2% paraformaldehyde prior to permeabilization with 1% Triton X-100 and staining with rhodamine-phalloidin were also used [14] and there was no difference seen in staining patterns or microfilament organization between the one-step and the two-step procedures.

**Transmission electron microscopy.** Adherent endothelial cells, grown in the same manner as for fluorescence microscopy, were fixed in situ with 2% glutaraldehyde at room temperature at various times after metabolic inhibition. The fixed cells were then scraped off of the plastic substrate with a rubber policeman and centrifuged. The cell pellet was embedded in albumin (which gels when exposed to glutaraldehyde). The resultant block was postfixed in 1% OsO4 for 60 min, and then stained en bloc with uranyl acetate for 20 min. Samples underwent dehydration in graded concentrations of ethanol with several final passages in absolute ethanol. They were then transferred through propylene oxide into a monomer mixture (Poly/Bed 812, Polysciences, Inc., Warrington, PA; Araldite; dodecenyl succinic anhydride; and tri(dimethylaminoethyl)phenol) and polymerized at 60°C. One-micrometer thick sections were stained with toluidine blue for review in the light microscope, and selected areas were thin sectioned at 60 nm and mounted on copper grids. Sections were stained with Reynold’s lead citrate and uranyl acetate. Thin sections were examined in a Zeiss 9S-2 electron microscope.

**RESULTS**

**Endothelial cell ATP.** Endothelial cell ATP levels were measured to assess the metabolic impact of acute inhibition of glycolysis and mitochondrial respiration by glucose depletion and exposure to 650 pmole oligomycin/μg DNA, respectively. Endothelial cells were inhibited metabolically for 2 hr and then allowed to recover for 1 hr during which

![Graph](https://via.placeholder.com/150)

**Fig. 1.** Time course of ATP levels in endothelial cells exposed to buffer with glucose (solid circles), glucose plus 650 pmole oligomycin/μg DNA (open circles), no glucose present (solid squares), and no glucose present plus 650 pmole oligomycin/μg DNA (open squares). After 120 min, cells exposed to oligomycin were washed rapidly and 5.5 mM glucose was added for 60 min of recovery. All data points represent the mean ± SD of between 5 and 11 separate observations. See text for details of statistical analysis.
Fig. 2. Time course of changes in microfilament organization in endothelial cells exposed to 650 pmole oligomycin/μg DNA in the absence of glucose over 60 min. Cells were stained with rhodamine–phalloidin and viewed at 400× magnification. (A) Control (+ glucose), uninjured cells; (B) 30 min plus oligomycin in the absence of glucose; (C) 60 min plus oligomycin in the absence of glucose. Note the progressive shortening and apparent aggregation of microfilaments occurring over 60 min of metabolic inhibition.
glucose (5.5 mM) was added to the buffer and oligomycin removed in a rapid wash step. The combined inhibition of glycolytic and mitochondrial ATP synthesis produced a rapid and dramatic fall in endothelial cell ATP levels (Fig. 1). Data from cells exposed to oligomycin in the absence of glucose were compared with those from cells exposed to glucose, glucose plus oligomycin, and glucose depletion over the 3-hr time course. Since there were considerable differences between the variances of the metabolically inhibited cells and the three control groups, Welch’s solution to the Behrens–Fisher problem [15] was used. The three control groups were significantly different ($P < 0.003$) from the metabolically inhibited group (oligomycin in the absence of glucose) with the exception of glucose depletion alone at 180 min, even after adjusting for multiple comparisons with Bonferroni’s inequality. After 120 min when the inhibited cells were washed free of oligomycin and glucose (5.5 mM) was restored, ATP levels increased over the following 30–60 min but not to control levels (Fig. 1). Paired $t$ tests were used to compare the levels of ATP in metabolically inhibited cells (lower curve—Fig. 1) at 120 min of inhibition and following the initiation of recovery at 135, 150, and 180 min. The most significant difference was between 120 and 150 min (30 min of recovery; $P = 0.002$).

**Microfilament Organization—Endothelial Cells**

**Fluorescence microscopy.** Adherent endothelial cells were fixed and stained with rhodamine-phalloidin over the time course of metabolic inhibition depicted in Fig. 1. There was a progressive loss of normal microfilament organization in $>90\%$ of cells over the 2 hr of inhibition with dramatic shortening of F actin into aggregates within the cells (Figs. 2 and 3). When the inhibited endothelial cells were allowed to recover after 2 hr of metabolic inhibition, there was a rapid reassembly of microfilaments within the cells as early as 15 min after addition of
FIG. 3. Time course of changes in microfilament organization during 60 min of recovery after 120 min of metabolic inhibition with 650 pmole oligomycin/μg DNA in the absence of glucose. Cells were stained with rhodamine–phallolidin and viewed at 400X magnification. (A) 120 min exposure to oligomycin in the absence of glucose; (B) 150 min (30 min recovery) oligomycin removed and 5.5 mM glucose added at 120 min; (C) 180 min (60 min recovery) oligomycin removed and 5.5 mM glucose added at 120 min. Note the rapid return to normal microfilament architecture over 30–60 min of recovery.
glucose and removal of oligomycin which was nearly complete in 85–90% of cells by 60 min (Fig. 3). Inhibition of either glycolytic or mitochondrial ATP synthesis alone did not alter the pattern of microfilament organization in the endothelial cells as compared to control cells in buffer containing glucose (data not shown).

**Transmission electron microscopy (EM).** To further characterize ultrastructural changes in endothelial cell microfilament organization associated with inhibition of ATP synthesis, adherent cells were fixed with glutaraldehyde and prepared for transmission EM. Ultrastructural examination of metabolically inhibited endothelial cells showed a change from the diffuse distribution of cytoplasmic microfilaments to aggregation into bundles (Fig. 4). The layer of conspicuous microfilaments normally present parallel to the surface onto which the cells adhere also disappeared. As would be expected, cytoplasmic particulate glycogen disappeared under these conditions. Restoration of ATP synthesis was associated with a gradual restoration of the normal diffuse distribution of microfilaments within the recovering cells (Fig. 5).

**DISCUSSION**

To produce a rapid and sustained fall in endothelial cell ATP levels, it was necessary to inhibit both aerobic and anaerobic ATP synthesis. This kind of situation should be theoretically operative during acute ischemic events *in vivo* where substrate (glucose) and O₂ delivery would be suddenly interrupted. ATP levels did fall in an *in vivo* model of renal ischemia to a range (<10% of controls) which was comparable to that seen in this *in vitro* study [16]. This indicates that *in vivo* ATP levels during ischemia could be affected to the same degree as seen with this *in vitro* model making it theoretically possible for changes in the cell cytoskeleton associated with ATP depletion to occur *in vivo*. The ability of the endothelial cells to maintain ATP synthesis at fairly normal levels via either glycolytic or mitochondrial pathways...
FIG. 4. Transmission electron microscopy of adherent endothelial cells at 120 min of metabolic inhibition. (A) Control-uninjured cell(s) 10,800X magnification. Normal ultrastructural pattern with microfilaments scattered throughout the cytoplasm and a normal basal layer of microfilaments (open arrows); (B) oligomycin in the absence of glucose 10,200X magnification. Note the side to side aggregate (solid arrow) of microfilaments in the metabolically inhibited cell.

alone emphasizes their adaptability in the face of a variety of metabolic stresses. From correlation of microfilament changes seen with fluorescence microscopy to changes in ATP levels occurring over the time course of metabolic inhibition in Fig. 1, it appears that
the approximate threshold level of endothelial cell ATP which correlates with destabilization of microfilaments occurs in the range of 15-20% of control levels (taken from data at 30 min of inhibition and at 60 min of recovery). It is quite interesting that ATP levels in recovering cells do not reach normal control levels over the 60-min recovery period. This may represent a very rapid turnover and utilization of newly synthesized ATP in recovering cells for biosynthetic processes interrupted by the period of metabolic inhibition including reassembly of disrupted microfilaments.

The reversability of the microfilament changes associated with ATP depletion in

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FIG. 5. Transmission electron microscopy of recovering adherent endothelial cells 30 and 60 min after removal of oligomycin and addition of glucose (5.5 mM). Cells were inhibited metabolically for 120 min prior to the recovery period. (A) 30 min recovery 11,500× magnification. Note the presence of normal microfilaments along with the bundled aggregate of microfilaments (solid arrow). (B) 60 min recovery 11,500× magnification. A normal basal layer of microfilaments (open arrow) has returned to the cytoplasm.
endothelial cells (demonstrated by fluorescence microscopy) is further confirmation of earlier observations in other cell types [3–5]. The ultrastructural changes in microfilament organization associated with inhibition of ATP synthesis appear to represent a side to side aggregation of microfilament segments. These changes are very similar (if not identical) to those seen in microfilament organization following hydrogen peroxide (H₂O₂) injury where ATP levels fall secondary to H₂O₂-mediated inhibition of ATP synthesis [7, 8, 13, 17]. Preliminary observations using a fluorometric assay [18] to evaluate the possibility that metabolically inhibited endothelial cells might generate superoxide anion (O₂⁻) or H₂O₂ were negative. Further work will be needed to fully rule out any involvement of oxidants in this type of model, however. The microfilament aggregation seen by electron microscopy may well represent an important part of any cell injury from whatever cause which is characterized by a significant decline in ATP levels. The aggregates may become useful ultrastructural markers of cellular injury in vivo, particularly in animal models of ischemic or inflammatory tissue injury.

There were only subtle changes in endothelial cell morphology accompanying the microfilament disruption associated with metabolic inhibition. It is unlikely that changes in morphology associated with microfilament disruption alone would produce gross changes in capillary permeability during an ischemic event. However, a transmembrane association between microfilaments and fibronectin, an extracellular matrix protein, has been demonstrated [19] and suggests that microfilaments play an important role in cell adherence to the basement membrane. Also in support of this concept is work demonstrating that metabolically inhibited endothelial cells lose their normal adherence properties [20].

Thus, the pathophysiologic significance of these observations may be that ischemic or ATP-depleted endothelial cells may be relatively less adherent and potentially could come off the basement membrane of an ischemic large or small vessel especially during reflow, resulting in exposure of basement membrane which would initiate platelet deposition and the associated inflammatory response [21]. Also, the exposed basement membrane would not be a very effective permeability barrier. It may be very important to develop ways to maintain or restore endothelial cell ATP levels as a means of preventing this type of problem from occurring in vivo during acute ischemia.

In conclusion, ATP depletion which occurs in a cellular model of acute ischemia is correlated with a reversible disruption of endothelial cell microfilament architecture. This may play an important role in endothelial cell dysfunction during acute ischemia in vivo.

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REFERENCES

7. Spragg, R. G., Hinshaw, D. B., Hyslop, P. A.,


