The effects of hyperosmolal coronary perfusion on the haemodynamic,
metabolic and ultrastructural changes of myocardial anoxia*

NORMAN BRACHFELD
M.D.

ROBERT ERLANDSON
Ph.D.

JAMES CHRISTODOULOU
M.D.

CHARLES SMITHEN
M.D.

Department of Medicine, The New York Hospital–Cornell Medical Center and the
Department of Pathology, Memorial Hospital, New York, New York 10021

Summary
Recovery from anoxia has been evaluated in the iso-
volumic non-recirculating paced perfused rat heart.
Seventy studies were performed consisting of 15 min
of aerobic perfusion (AP); AP+15 min anoxic per-
fusion; and AP+15 min anoxic perfusion+15 min re-
oxygenation (recovery). Krebs-Ringer-bicarbonate+
5 mmol glucose (KRB) (290 mmol) was compared to
KRB+mannitol (350 mmol). Mannitol decreased
myocardial water content. It improved recovery of
haemodynamic function after reoxygenation. With
KRB alone left ventricular systolic peak pressure
(LVSP) decreased 32\% and maximum dP/dt by 50\%.
With mannitol added LVSP decreased 18\% and dP/dt
21\% (P<0.01). KRB and mannitol did not differenti-
tially affect total coronary flow, lactate and glucose
extraction, tissue glycogen, creatine phosphate or
adenine nucleotide concentrations.

No difference in submicroscopic appearance was
noted with either perfusate during aerobic perfusion.
Anoxic hearts perfused with isosmolar KRB demon-
strated the most severe ultrastructural alterations
including mitochondrial swelling with disruption of
cristae and extraction of matrix components, myo-
fibrillar fusion and contraction bands, and sub-
sarcolemmal oedema and vacuolization. These changes
were only partially reversed during reoxygenated
perfusion. However, cellular changes were reversed or
markedly improved during both the anoxic and re-
oxygenation perfusion periods with hyperosmolar
solutions. When studied by silicone rubber injection of
the microcirculation, only focal capillary endothelial
swelling was noted and no difference in arteriolar
or capillary filling was observed with either perfusate.
Mannitol appears to improve LV function by direct
myocardial osmotic action unrelated to enhanced
energy production.

Introduction
Both ischaemia and anoxia have been shown to
induce a significant degree of tissue swelling. The
physiological significance of such changes may be
expressed with particularly detrimental effects in
myocardium. Myocardial cellular swelling may dilute
intracellular contents and lead to palpable firmness
of the ventricular wall. Reduction in ventricular
compliance caused by high membrane tensions can
seriously impair haemodynamic performance. Recent
studies by Willerson et al. (1972) demonstrated the
beneficial action of elevated osmolality in protecting
the ischaemic myocardium against the potentially
toxic effects of increased cell volume. Its mechanism
of action, however, is not completely understood.
These studies were designed to evaluate the effects
of a hyperosmolar agent (mannitol) during anoxia
under the controlled conditions provided by the
isolated perfused non-recirculating Langendorff rat
heart preparation. Haemodynamic, metabolic, optical
and electron microscopic and latex vascular injection
studies were performed to explore its mechanism of
action. The results of haemodynamic and metabolic
studies form the subject of this preliminary report.

Methods
Seventy isolated rat hearts were perfused in a
tandem Langendorff apparatus modified to permit
rapid exchange of perfusates varied as to osmolality

* Supported by: USPHS Contract No. N01 HV 71439;
Co.; The Prudential Ins. Co. of America and North American
Reassurance Co.

Address for reprints: Dr Norman Brachfeld, The New
York Hospital, 525 East 68th Street, New York, New York
10021.
and $pO_2$ (Fig. 1). Male, fed, albino rats of the Sherman strain weighing 250–290 g were used. Rats were killed by axial fracture, the heart rapidly removed and mounted on the aortic cannula. The procedure required 40 sec from sacrifice to onset of perfusion. The substrate reservoir was set to deliver a perfusion pressure of 75 mmHg and buffer was maintained at 37°C and a pH of 7.4. After onset of perfusion, a small latex balloon catheter was inserted into the left ventricle. The balloon was inflated (diastolic pressure = atmospheric pressure) and ventricular pressure and its first derivative recorded. A stab wound in the wall of the left ventricle prevented accumulation of thebesian drainage. Heart rate was maintained at a constant of 300 beats/min during aerobic perfusion by bipolar pacing. Coronary flow was measured by collection of fluid ejected via the pulmonary artery. This venous efflux was assayed as described below. Arterial samples were drawn from the perfusion cannula.

**Perfusates**

The control perfusate consisted of modified Krebs-Ringer bicarbonate buffer containing 5 mmol glucose, 0.1 mm Na Lactate and 25 µ of glucagon free insulin/ml solution. Its osmolality was 290 mmol. The experimental perfusate differed by the addition of 20% mannitol sufficient to raise the osmolality of the solution to 350 mmol.

During aerobic studies perfusate $pO_2$ was maintained at 550±10 mmHg by bubbling with 95% O$_2$+5% CO$_2$. For anoxic studies, solutions were bubbled with 95% N$_2$+5% CO$_2$ and $pO_2$ was 50±5 mmHg.

**Experimental design**

The two perfusates were compared during three sequential study periods: (a) 15 min of paced aerobic perfusion (control), (b) 15 min of anoxic perfusion (anoxia), and (c) 15 min of paced aerobic perfusion (recovery). Metabolic, haemodynamic and histological studies were performed on a series of fourteen hearts following each of the three periods. An additional group of seven hearts was perfused with isosmolar buffer during control and anoxic periods and with the hyperosmolal solution during the recovery phase.

**Assay procedures**

The arterio-venous concentration differences of

---

**Fig. 1.** Modified Langendorff rat heart perfusion apparatus. A retrograde, non-circulating tandem system is illustrated allowing for ventricular pressure recording during isovolumic ventricular contraction by inflation of a balloon catheter attached to a pressure transducer, and control of heart rate by an external battery powered pacemaker.
buffer samples were multiplied by coronary flow to express substrate consumption/production. Glucose and lactate assays were performed in triplicate with a coefficient of variation of ±2% by automated techniques previously described (Apstein, Puchner and Brachfeld, 1970). At the termination of each perfusion period, hearts were compressed by Wollenberger clamps taken from liquid nitrogen. Frozen specimens were finely ground and wet/dry weight determined after drying of an aliquot at 105°C for 24 hr. An additional aliquot was weighed in a tared frozen centrifuge tube and homogenized with HClO₄. A sample of the supernatant was extracted with KOH for glycogen determination. The remainder was assayed enzymatically for ATP, creatine phosphate, ADP and AMP.

**Electron microscopic studies**

In a separate series of fourteen hearts the cardiac apex was excised during perfusion and was immediately immersed in s-collidine buffered 2% acrolein and 5% glutaraldehyde fixative for 1·5–2 hr. The tissue was then rinsed in buffer and postfixed for 1 hr in 1% s-collidine buffered osmium tetroxide. After dehydration, specimens were embedded in Maraglas-D.E.R. 732 epoxy resin. Thin sections were stained with uranyl acetate followed by lead citrate and were examined in a Siemens Elmiskop 101 microscope. Thick (1µ) sections for light microscopic examination were stained with toluidine blue.

**Microcirculation studies**

A silicone rubber injection mass was prepared so as to attain a viscosity of 15–25 centipoise after catalysation. The material was introduced at a fixed perfusion pressure identical with that used during buffer perfusion. The mixture was admitted to the retrograde aortic perfusion cannula by a system designed to exclude the interposition of air bubbles.

Hearts continued to beat for 2–3 min and perfusion pressure was maintained until the injection mass appeared in the venous drainage. The heart was then removed from the cannula and allowed to cure overnight at room temperature. Curing took place with non-exothermic cross-linking and minimum volume change. The tissue was then dehydrated by immersion in ethyl alcohol of progressively increasing concentration. It was cleared and rendered transparent by immersion in synthetic methyl salicylate.

Sections of 200–300 µ were prepared for direct microscopic examination and also cut at 20 µ for routine HE staining.

**Results**

**Haemodynamics**

During control aerobic perfusion, hyperosmolarity did not significantly affect peak left ventricular systolic pressure. Active contraction ceased soon after onset of anoxia and pressures could not be recorded. During aerobic recovery neither series of hearts reached pre-anoxic performance levels. Nevertheless, those perfused with the hyperosmolar solution showed a significant improvement over those perfused with KRB alone. Isosmolal perfusate pressures reached 68% of aerobic controls (127±5 → 86±6 mmHg); hyperosmolar pressures reached 77% of control measurements (132±5 mmHg → 102±7 mmHg) P= <0·01. Enhanced performance was more evident when the first derivative of the ventricular pressure curve was analysed (Fig. 2). With KRB buffer alone, mean recovery values of dP/dt were 50% of control (3513±328 mmHg/sec → 1758±172 mmHg/sec). Perfusion with hyperosmolar solution enhanced performance to 79% of control aerobic perfusion (3817±215 → 2998±234 mmHg/sec) P= <0·01. Improved recovery was also demonstrable when hyperosmolality was induced at the termination of the anoxic period. Under these circumstances, peak systolic recovery was 86% and LV dP/dt 81% of aerobic controls.

**Coronary flow**

Coronary flow during perfusion with isosmolal buffer was 11·9±0·7 ml/min, rose markedly at the onset of anoxia to reach 19·2±2·4 ml/min and fell to 8·4±0·9 ml/min with cessation of contractile activity. The mean level during recovery was 7·12±0·6 ml/min, a flow rate that did not differ significantly from aerobic control when flow was corrected for contractile activity. Hyperosmolal perfusion did not significantly change these rates. Coronary flow during
aerobic perfusion was 12·2±1·0 ml/min, rose to 20·9±1·9 ml/min during anoxia and fell to 7·8±0·5 ml/min during recovery.

**Metabolic studies**

**Carbohydrate metabolism**

Variations in osmolality of perfusate solutions did not significantly influence glucose consumption. With isosmolar buffer, consumption was 3·39±0·72 μmol/min during control aerobic perfusion and remained at this level despite almost total cessation of contractile activity and work performance during anoxia. With aerobic recovery, consumption fell to 2·00±1·1 μmol/min. During hyperosmolar perfusion glucose consumption was 3·28±0·61 μmol/min in period (a), 3·00±0·72 μmol/min in period (b), and 2·22±0·56 μmol/min in period (c); *P* = N.S.

Mean lactate production during aerobic isosmolar perfusion with KRB buffer was 0·57±0·16 μmol/min. Production increased to 3·05±0·53 μmol/min during anoxic perfusion. During recovery, production fell markedly and did not differ significantly from aerobic control, 0·78±0·13 μmol/min. The use of the hyperosmolar perfusate did not alter lactate metabolism at any phase of the study. Production was 0·81±0·12 μmol/min in period (a), 3·04±0·26 μmol/min in period (b), and 1·07±0·13 μmol/min in period (c); *P* = N.S.

Tissue glycogen concentration was 103±11 μmol/g dry wt during aerobic perfusion with isosmolar buffer. It fell to 41±5 μmol/g dry wt during anoxic perfusion and remained close to this level during aerobic recovery (43±4 μmol/g dry wt) (Fig. 3). Hyperosmolar tissue glycogen levels were 93±10 μmol/g dry wt during period (a), 37±1 μmol/g dry wt during period (b), and 47·7±7 μmol/g dry wt during period (c); *P* = N.S.

**Tissue high energy phosphate stores**

Hyperosmolality did little to alter the depletion of myocardial high energy phosphate stores during anoxia nor its repletion during recovery (Fig. 4). Creatine phosphate levels of 28·1±1·4 μmol/g dry wt during isosmolar aerobic perfusion fell to negligible levels (0·3 μmol) during anoxia and rose to 20·6±3 μmol/g dry wt with recovery aerobic perfusion. Concentrations obtained with hyperosmolar buffer were (a) 29·4±6·5, (b) 0·2, and (c) 24·6±3 μmol/g dry wt; *P* = N.S. (Fig. 4). Tissue ATP concentration showed a qualitatively similar change. With isosmolar buffer, ATP = (a) 20·1±1·3, (b) 10·1±1·1, (c) 8·5±1·2 μmol/g dry wt. Hyperosmolar perfusion did not significantly improve this balance nor did the presence of the hyperosmolar buffer significantly alter ADP, or AMP concentrations in the recovery phase.

![Fig. 3. Comparison of tissue glycogen levels with and without hyperosmolar mannitol. Glycogen concentration fell during anoxia and remained depressed during recovery. No significant differences between the two perfusates were seen. ■, KRB; ○, KRB + mannitol; *P* = N.S.](image)

![Fig. 4. Summary of tissue studies during recovery from anoxia. No difference in concentrations of glycogen, creatine phosphate, ATP or AMP were observed. Perfusate osmolality did not affect ADP levels (not illustrated). ■, KRB; ○, KRB + mannitol.](image)

**Myocardial tissue water content**

Presence of a hyperosmolar perfusate significantly affected total tissue water content during all three phases of perfusion (Fig. 5). The tissue wet/dry wt ratio with isosmolar perfusion in period (a) was 6·68±0·41 and rose to 7·48±0·56 (*P* = <0·01) with anoxia. It remained elevated throughout the recovery period (7·55±0·17). Hyperosmolar perfusion reduced the ratio during aerobic perfusion (5·80±0·26). Water content during anoxic perfusion was significantly less than that seen with isosmolar buffer and remained fixed during recovery (ratio=6·77±0·19). In studies which utilized the isosmolar buffer during periods (a) and (b) and switched to the hyperosmolar buffer during (c) tissue wet/dry wt ratios was restored to control levels.
Hyperosmolal coronary perfusion

Fig. 5. Effect of mannitol on myocardial water content (wet wt/dry wt ratios). 'Pre' = aerobic period before anoxia. 'Post' = reoxygenated recovery period. Hyperosmolal perfusion reduces the ratio during the aerobic period. The increase in ratio noted during and following anoxia was prevented by an increase in osmolality. III, KRB; □, KRB + mannitol.

Electron microscopic studies

(a) Control hearts

Electron microscopic examination of rat myocardium fixed immediately after removal revealed a submicroscopic appearance which did not differ significantly from that of other mammals. Both isosmolal and hyperosmolal perfused tissue showed identical electron microscopic anatomy (Fig. 6).

(b) Anoxic hearts

Hearts subjected to a 15-min anoxic period perfused with KRB alone showed the most severe submicroscopic pathology. Marked disorganization of most of the myocardial cell constituents was evident. The mitochondrial matrix was extracted (electron-lucent configuration) and these organelles were randomly dispersed in the sarcoplasm. Some of the mitochondria were markedly swollen and contained fewer cristae. Elements of the sarcoplasmic reticulum were dilated and disrupted, and vacuoles of various sizes were also prominent (Fig. 7). In some junctional areas, the intercalated discs were disrupted, and dense bodies of unknown composition were seen in the sarcoplasm. Myofibrillar alterations were also noted. Normal sarcolemmal and subsarcolemmal structure was altered owing to disruption and osmotic swelling of myocytes. Focal endothelial cell changes included swelling and a reduction in the number of pinocytic vesicles.

The presence of mannitol in the perfusate during the anoxic period significantly reduced the severity of these changes (Fig. 8). Approximately 50% of the mitochondria now remained in the condensed configuration and myofibrillar structure was generally maintained.

(c) Reoxygenated hearts

Reoxygenation of hearts perfused with KRB alone showed only minimal reversal of the submicroscopic alterations found in the anoxic hearts (Fig. 9). Most of the mitochondria examined continued to demonstrate an extracted, electron-lucent matrix, and various myofibrillar alterations such as fraying of myofilaments. The occurrence of contraction bands

Fig. 6. Rat heart fixed immediately after removal (no perfusion). Note the small capillary and intercalated disc. × 7400.
Fig. 7. Rat myocardium after 15 min anoxic perfusion with KRB. Note the disorganized, altered myofibrils and the electron-lucent mitochondria with disrupted cristae. $\times 15,000$.

Fig. 8. Rat myocardium after 15 min anoxic perfusion with hyperosmolar medium. Note that myocardial cell constituents show only slight alteration. $\times 7400$. 
Hyperosmolal coronary perfusion

FIG. 9. Reoxygenated rat myocardium perfused with isosmolal KRB buffer. Large areas of subcellular damage are still evident. × 12,000.

persisted despite reoxygenation. Focal areas of endothelial swelling and vacuolization were also apparent.

After reoxygenation with mannitol, a substantial reversal of the above mentioned anoxic changes was noted. Many of the myocardial and vascular endothelial cells were virtually identical in substructure to the controls (Fig. 10). Even in areas where contraction bands and myofibrillar fusion persisted, most of the mitochondria were in the condensed configuration. It should be pointed out, however, that scattered focal areas of severe, apparently irreversible, damage were still evident.

**Microcirculation studies**

The coronary microcirculation as studied by the silicone rubber injection technique visualized arteriolar and capillary vessels to a diameter of 7 μ. No significant areas of capillary obstruction were noted and no differential effect of mannitol on arteriolar or

FIG. 10. Rat myocardium after 15 min of reoxygenated perfusion (recovery) with hyperosmolal medium. Note that both myocardial cells and capillaries show essentially normal ultrastructure. × 6000.
capillary filling was observed after anoxia or during the reoxygenated recovery period.

Discussion

Leaf (1970) has recently emphasized the physiological significance of intracellular volume regulation. His studies of the heart and experiments with kidney and brain have demonstrated that intracellular oedema may accentuate or prolong the effects of oxygen deprivation after the stress of a reduction in flow has been relieved. Willerson’s studies (1972) in the post-coronary ligated dog heart demonstrated that elevated osmolality protected the ischaemic myocardium against the potentially toxic effects of increased cell volume. There was an improvement in left ventricular function curves, a lessening of ST segment elevation and an increase in total and collateral flow. Our studies have confirmed the positive inotropic effects of this agent when perfusate osmolality was raised by 60 mmol either before or after the anoxic stress.

It seems probable that induced obligatory extracellular hyperosmolality may: improve regional perfusion by reducing endothelial cell swelling and increasing patency of arterioles and capillaries or by directly reducing coronary arteriolar resistance (Willerson et al., 1972); protect mitochondria by maintaining an optimal ultrastructural configuration for oxidative phosphorylation and energy production (Caulfield et al., 1972); enhance inotropism by a relative increase in $Ca^{++}$ concentration or an increase in $Ca^{++}$ activity from small molecular weight complexes (Caulfield, 1973). An increase in extracellular osmotic potential would aid in the maintenance of a favourable intracellular environment by counteracting swelling induced by ischaemia (Jennings, Herdson and Sommers, 1969) and exploit the inverse relationship between ventricular wall compliance and an abnormal increase in myocardial cell volume.

We were unable to demonstrate an increase in total coronary flow during hyperosmolal perfusion at either high or low $pO_2$. In the absence of formed elements capillary entrapment was, of course, not possible. Furthermore, a reversal of capillary and arteriolar endothelial wall swelling sufficient to improve flow might well have been masked by the gross nature of coronary flow measurement utilized or by the marked arteriolar vasodilatation in response to anoxia. Silicone latex injection studies of the vascular tree performed and reported here also failed to demonstrate a difference in vessel calibre between these two types of buffer.

Ultrastructural alterations produced in mammalian myocardium following oxygen deprivation are well known and appear to be relatively independent of the means of inducing hypoxia. Mitochondria are extremely sensitive to anoxia and morphological changes in these structures were most striking in this study. We observed a loss of matrix constituents or a change from an electron-dense (condensed) configuration to an electron-lucent (extracted) configuration. Other mitochondrial alterations observed and also reported by other investigators included disappearance of intra-mitochondrial granules, swelling and fragmentation of cristae, increase in number and of size, and fusion. Severe mitochondrial changes induced by anoxia proved to be reversible during reoxygenated perfusion with hyperosmolal mannitol and were considerably reduced in myocardial samples taken immediately after anoxia. The degree of cellular oedema noted in isosmolar perfusates studied during anoxia was reduced by approximately 50% when buffer osmolality was elevated by addition of mannitol. Mannitol also appeared to decrease the amount of myofibrillar oedema, fusion, and contracture noted after anoxia. This protective effect correlated well with the enhanced haemodynamic recovery seen during the subsequent reoxygenation period.

The presence of mannitol in the perfusate did not significantly affect total coronary flow and no differences in arteriolar or capillary filling could be observed using the silicone rubber injection technique.

Although severe anoxic mitochondrial swelling was largely prevented during phases (b) and (c) of the hyperosmolal perfusion such histological improvement was not expressed biochemically by improvement in oxidative phosphorylation, enhanced utilization of glycogen or glucose, or maintenance or improvement in high energy phosphate stores.

Significant dissociation between preservation of mitochondrial and myofibrillar structure and their functional capability was evident in our model. The improvement in haemodynamic performance induced by hyperosmolality did not appear to be related to a similar improvement in myocardial energetics. There was a positive correlation between reduction in wet/dry wt ratios and enhanced mechanical performance during the improved recovery of phase (c) seen when mannitol was added. These findings suggest that increased compliance of the left ventricular wall due to a decrease in cell volume was of primary significance in explaining our data. Investigations continue in an effort to determine whether this simply represents a reversal of the detrimental effect of tissue fluid pools or a relative or absolute increase in intracellular $Ca^{++}$ concentration with enhanced inotropic response to this ion, already competing with increased $H^+$ ion for contractile binding sites during myocardial anoxia.
**References**


The effects of hyperosmolar coronary perfusion on the haemodynamic, metabolic and ultrastructural changes of myocardial anoxia

Norman Brachfeld, Robert Erlandson, James Christodoulou and Charles Smithen

doi: 10.1136/pgmj.51.595.299

Updated information and services can be found at:
http://pmj.bmj.com/content/51/595/299

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/