Evolution and resolution of oedema following severe temporary cerebral ischaemia in the gerbil

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SUMMARY Regional cerebral blood flow (rCBF) and oedema following profound temporary ischaemia were studied in the gerbil. Ninety-four per cent of animals died within 24 hours of reperfusion; 50% by 4 hours. Regional differences in oedema (specific gravity method), Evans blue (EB) staining and rCBF (hydrogen clearance technique) occurred. Oedema developed during arterial occlusion, being inversely proportional to residual flow and was markedly exacerbated during reperfusion. Reperfusion hyperaemia was maximal in the parietal and hippocampal regions (ischaemic rCBF 4 ml 100 g⁻¹ min⁻¹). Oedema was disappearing in all areas by 3 hours of reperfusion and autoregulation returned in the occipital region (mean ischaemia rCBF 8 ml 100 g⁻¹ min⁻¹). EB staining and haemorrhage appeared in the thalamus (rCBF 10 ml 100 g⁻¹ min⁻¹) as oedema was decreasing. It is suggested that the amount of oedema and hyperaemia during reperfusion are dependent on the severity of the ischaemia. Areas of moderate ischaemia (8–10 ml 100 g⁻¹ min⁻¹) show little hyperaemia and greater oedema resolution during reperfusion as compared to areas of severe ischaemia (circa 4 ml 100 g⁻¹ min⁻¹) where there is marked hyperaemia with less oedema resolution. Early in the reperfusion period, oedema is not associated with EB staining and indicates a cytotoxic mechanism. The vasogenic component, with macroscopic haemorrhages and leakage of EB occurs later. In this model it is concluded that the early cytotoxic oedema formation and hyperaemia are phenomena with little bearing on mortality, which correlates better with later vasogenic changes.

Oedema will develop in ischaemic brain tissue and clear thresholds have been demonstrated at which the process develops.1,2 The level of ischaemia at which oedema begins is similar to that at which electrical activity ceases.3,4 In the clinical situation, following a stroke, there may be a subsequent deterioration and the CT scan may demonstrate a brain shift and oedema surrounding the infarct. Following neonatal asphyxia the child’s condition may deteriorate after 18–36 hours and recent nuclear magnetic resonance studies suggest that water is accumulating within the brains. There is ample evidence in the literature to demonstrate the time course of oedema which develops following experimental ischaemia5 and this is in the same time period noted clinically. Clearly, the process is dynamic and not a static relationship implied in some work: little is known about resolution of oedema.

With reperfusion into the ischaemic area, another variable is introduced. The blood brain barrier may have been damaged. Intravascular fluid and “oedemagenic substances” may leak into the damaged cerebral tissue causing further deterioration. It is a situation with clear clinical implications such as that which would occur following a cardiac arrest, or cerebral embolus or surgical procedure. Iannotti and Hoff,6 studying reperfusion oedema in the gerbil, have shown that in areas of moderate ischaemia (rCBF, greater than 10 ml 100 g⁻¹ min⁻¹) oedema will rapidly resolve following restoration of flow. In severely ischaemic regions, less than 7 ml 100 g⁻¹ min⁻¹, oedema is aggravated during reperfusion. Whether the development and resolution of oedema has anything to do with mortality and morbidity has yet to be decided.

We have used a model of bilateral carotid occlusion in gerbils which reliably produces profound ischaemia in the anterior two-thirds of the cerebral
Reperfusion oedema following ischaemia

hemispheres, in order to study variations of flow and oedema, and correlated them with clinical survival following carotid occlusion. We have chosen a model of certain mortality to assess the effects of reperfusion through non viable cerebral tissue and to correlate this with our studies in brain tissue prosta
glandins. In this paper we describe the clinical effects and cerebral vascular changes consequent on 1 hour of bilateral carotid occlusion.

Methods and materials

Cerebral ischaemia was produced by bilateral common carotid occlusion, with or without reperfusion in 132 male mongolian gerbils (weight range 42–70 g). All animals received intraperitoneal pentobarbitone anaesthesia (60 mg kg⁻¹), with supplementation as necessary. Surgery was performed with an operating microscope, the common carotid arteries being carefully isolated and a sling passed under each vessel to ensure simultaneous bilateral occlusion with Scoville Lewis aneurysm clips. Temperature was maintained at 37°C by heating blanket and lamp. In ani
mals undergoing rCBF studies, a femoral artery catheter gave access for continual blood pressure (MABP) monitoring and intermittent blood gases.

The experimental protocol was divided into four groups:

- **Group A:** Mortality and morbidity studies (34 gerbils).
- **Group B:** Macroscopic morphological changes (27 gerbils plus 20 from Group A).
- **Group C:** Specific gravity studies (53 gerbils, including some from Group D).
- **Group D:** Regional cerebral blood flow (rCBF) (21 gerbils).

**Group A: Mortality and morbidity**

Thirty-four gerbils were observed following 60 minutes of temporary bilateral cerebral ischaemia. Note was made of paresis, seizures, conscious level and time of death. When breathing stopped, fixation-perfusion was carried out by infusion of saline, followed by 10% formalin, through the left ventricle. Brains were removed and immediately examined macroscopically after 12 coronal slices had been cut.

**Group B: Macroscopic and morphological changes**

Evans blue was given as a bolus intravenously (0.05 ml of 2% solution in saline pH 7-4) 30 minutes after application of the clip. Twenty-seven animals were sacrificed at set times to assess the evolution of Evans’s blue (EB) staining. These reperfusion time intervals (with numbers of animals) of 60 minutes (9), 120 minutes (10), and 180 minutes (8) were taken following the standard hour of ischaemia. Animals were kil-perfused with 10% formalin and the presence of EB discoloration and/or haemorrhages noted.

**Group C: Specific gravity (SG)**

Regional specific gravity was measured in 53 gerbils according to the techniques of Marmarou et al.** Slight adjustments were made to the column calibration span to allow for gerbil grey matter. Gerbils were sacrificed by decapitation, the brain being rapidly removed and placed in a kerosene filled petri dish. Six pieces 1 mm³ were cut from each of eight regions; frontal, parietal, occipital, cor
dus striatum, hippocampus, thalamus, cerebellum and pons. The linearly inhomogenous columns of kerosene and bromobenzene were calibrated with six known SG standards. Only columns with a linear correlation coefficient (r) better than 0.9950 were accepted for use. The samples were allowed to settle for 3 minutes, when the exact level was noted. The following groups were examined (number in brackets equals number of animals); control, with anaesthetic, but no surgery (4). Sham sampled 2 hours after operation, but without occlusion (4), 60 minutes ischaemia only (4), 120 minutes ischaemia only (4). The following reperfusion periods were chosen after 1 hour of ischaemia; 5 minutes (4), 30 minutes (8), 60 minutes (8), 120 minutes (8), and 180 minutes (9).

**Group D: Regional cerebral blood flow (rCBF)**

rCBF was measured by the hydrogen (H₂) clearance tech
nique of Aukland et al. (1964),¹⁰ with modifications¹¹ ² in 21 gerbils. Burr holes were made in the skull and six platinum-iridium teflon coated electrodes (175 µ max. diameter), with 1 mm of the tip exposed, were positioned with micro-manipulators and fixed with methylmethacrylate cement. Placement was confirmed after death: there was a mixture of cortical and deep electrode placements in each animal so that all sites were studied in the group as a whole. Flow was calculated by the initial slope technique. The slope of the log height—time curve, multiplied by 100 gave the rCBF in ml 100 g⁻¹ min⁻¹.¹² (See section on Validation of Technique.) Zero flow was defined as a lack of H₂ clearance, with no change upon further H₂ administration. Hyperaemia was defined as a rCBF measured during reperfusion that was greater than the control value. All rCBF studies had arterial blood gases measured prior to occlusion, and once or twice during reperfusion (total volume less than 410 µl).

**Validation of Technique**

Duplicate flows showed a ± 6% reproducibility. The semilogarithmic plots were monoe
xponential in the gerbil.¹² ² In 82% of flow estimations, the coefficient of determination (r²) for the exponential regres
sion was ≥ 0.9950, between 0.9900 and 0.9949 in 12% and ≥ 0.9850 and < 0.9899 in less than 5%. Fewer than 1% were non linear and in all these a brief variation in MABP was noted (for example seizure). The poorest values of r² were seen with slow flows in the immediate reperfusion phase.

**Analysis**

Chi squared testing with the Yates correction for con
tinuity was used in the analysis of the mortality data, inci
dence of haemorrhages and Evans blue staining. Flow and oedema studies were compared by the student’s t test (two tailed, unpaired). Results are expressed as means ± one standard deviation.

**Results**

Mean arterial blood pressure (MABP) was 61 ± 10
mm Hg (n = 21), and rose sharply with clamping (fig 1), but fell to a plateau above resting levels for the remainder of carotid occlusion. Blood pressure plunged to a nadir (40 mm Hg) within 2 minutes of clip release, recovering by 5 minutes and reaching control levels by 30 minutes. MABP stabilised at 80 mm Hg for the last 90 minutes of protocol. Brief transients of MABP were recorded with convulsions. PaCO₂ was 43.9 ± 5.0 mm Hg with a fall during the hyperventilation associated with carotid occlusion; statistically the levels did not alter subsequently.

GROUP A: MORTALITY AND MORBIDITY
In the 34 animals there was a 50% mortality by 4 hours of reperfusion, and only 6% survived to 24 hours (fig 2). During the second hour of reperfusion, signs of severe neurological damage were invariable, with circling or hemiparetic behaviour, abnormal posture and irritability to sound. Fits, both focal and generalised, began with arousal in the second hour of reperfusion in all animals recovering consciousness.

GROUP B: MACROSCOPIC MORPHOLOGICAL CHANGE
This group consisted of animals followed to death (Group A) and others killed at predetermined intervals. The earliest EB staining was seen at 60 minutes of reperfusion in one of nine gerbils. The prevalence of EB was 50% by 120 minutes and more pronounced in degree at 180 minutes. Leakage was usually bilateral and symmetrical, being most marked in the thalamus and lateral geniculate bodies. The parietal, hippocampal, lateral mid-brain and occipital areas were less often affected. Of the animals showing EB staining, half had haemorrhages in the thalamus. Haemorrhages were thalamic except for two cases (central pontine grey and bilateral hippocampal). Haemorrhage and EB occurred in similar proportions (X²; N.S.) (table).
Reperfusion oedema following ischaemia

Table Incidence of Evans blue staining and haemorrhage following reperfusion after 60 carotid occlusion

<table>
<thead>
<tr>
<th>Reperfusion time</th>
<th>1 hr</th>
<th>2 hrs</th>
<th>3 hrs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Evans Blue (EB)</td>
<td>11%</td>
<td>50%</td>
<td>62%</td>
</tr>
<tr>
<td>Haemorrhage (H)</td>
<td>0%</td>
<td>22%</td>
<td>28%</td>
</tr>
<tr>
<td>H + EB</td>
<td>0%</td>
<td>0%</td>
<td>25%</td>
</tr>
<tr>
<td>H - EB</td>
<td>0%</td>
<td>10%</td>
<td>13%</td>
</tr>
</tbody>
</table>

The incidence of Evans blue staining (noted on inspection) and haemorrhage into the deep grey structures in given. There is no gross EB staining before one hour or reperfusion and no haemorrhages until two hours. However, there is no obvious relationship between EB staining and the presence of haemorrhages.

H + EB Haemorrhage and Evans Blue in same animal.
H - EB Haemorrhage without Evans Blue in same animal.

although EB was more commonly bilateral, and tended to manifest earlier. Focal seizures did not correlate with asymmetries of EB or haemorrhage.

GROUP C: SPECIFIC GRAVITY
During the 60 minutes of ischaemia all areas in the carotid vascular territory showed a significant fall in SG (increase in water). There was further and more rapid accumulation of oedema within 5 minutes of clip removal (fig 3), which was greatest in areas with more severely impaired occlusion flows. By 5 minutes of reperfusion the degree of oedema was very similar throughout the brain and it continued to increase for a further 60 minutes. The thalamus showed the most pronounced oedema during the ischaemic phase and, uniquely, did not show an increase in water accumulation during reperfusion. All areas, apart from the hippocampus, showed a return towards normal SG in the third hour of reperfusion. Resolution of oedema was greatest in the areas of least severe occlusion ischaemia and, for example, in the occipital area where the SG returned to control levels (figs 1 and 3). Despite simultaneous bilateral occlusion and release of the arteries, differences in SG were found between the left and right hemispheres, being most pronounced in the parietal areas. There was no change in the SG in the pons or cerebellum following ischaemia and reperfusion.

GROUP D: REGIONAL CEREBRAL BLOOD FLOW
In spontaneously breathing intact animals under barbiturate anaesthesia, the mean CBF was 29.6 ml 100 g⁻¹ min⁻¹ (215 electrodes). Mean rCBF ranged from 23 (corpus striatum) to 38 (parietal) ml 100 g⁻¹ min⁻¹. Clipping produced a marked fall in rCBF to 3 ml 100 g⁻¹ min⁻¹ in the forebrain and 10 ml 100 g⁻¹ min⁻¹ in the thalamus and occiput. Following clip release, there was no significant mean flow change at 5 minutes. Regions of zero flow were seen occasionally in the forebrain during the early reperfusion phase, but these did not persist. Hyperaemia was then seen in all areas, with regional variations; flow returned to control values by 3 hours of reperfusion. Hyperaemia immediately after clip release was seen in the thalamus of only one animal. Hyperaemia was most marked in areas where the ischaemic flow was rCBF of 4.1 ± 1.5 ml 100 g⁻¹ min⁻¹, (parietal area, fig 1). There was no direct relationship between peak hyperaemia and peak oedema. MABF and rCBF in the first and third hour of reperfusion showed a passive pressure–flow relationship, suggesting a loss of autoregulation (fig 4).

Fig 3 Regional specific gravity measurements before, and after 60 min bilateral carotid occlusion are shown. The results are mean data obtained from groups of at each time point. The numbers are given in the text but was between 4 and 9 for each time point. X-axis displays sham animals, their dotted line shows changes during occlusion (Black Box) and solid line indicates regional specific gravity patterns during three hours of reperfusion. In the carotid territory there was significant increase in water, maximal at 1–2 h, which then subsided. In the non-ischaemic vertebrobasilar territory (Po and Cb) there is no change in specific gravity.

SG 1-0500 = normal. 1-0460 = oedema.
F = frontal; P = parietal; Oc = occipital; Cs = corpus striatum; H = hypothalamus; Th = thalamus; Po = pons; Cb = cerebellum.
Autoregulation in response to induced blood pressure changes is shown during the reperfusion phase in the occipital regions. Blood pressure changes and corresponding regional cerebral blood flow changes 1 hour and 3 hours after the removal of the occlusive carotid clips are shown. A linear regression analysis of the flow and blood pressure results allowed construction of the graphs. There is a passive pressure flow relationship in the first hour of occlusion, but by the third hour reperfusion correlation had altered. The mean rCBF in the occipital region was 10 ml gm⁻¹ min⁻¹ during occlusion. In the frontal and parietal areas where occlusive flows were lower, autoregulation did not return.

Discussion

The gerbil has been used as a model of cerebral ischaemia because the hemispheric blood supply is from the carotid arteries with incomplete anastomoses of the circle of Willis. Microvascular anastomoses exist but these cannot develop sufficiently to sustain life. Bilateral ligation results in 100% mortality within 4 hours, whereas a unilateral ligation results in a mortality of 20–65%. The reason for the wide range in mortality with a unilateral ligation is the side to side variations in the circle of Willis which have been elegantly demonstrated by Tamura et al. With temporary bilateral occlusion there is a mortality which varies with the period of occlusion. Fifteen minutes of bilateral occlusion rarely causes death; 30 minutes occlusion produces a 40% mortality at 24 hours. Restoration of circulation after 1 hour's carotid occlusion results in 94% mortality within 24 hours. These results are similar from different laboratories and under different anaesthetics, and provide a crude but reproducible index of the depth and duration of ischaemic tolerance. They provide a baseline for the assessment of possible therapeutic agents aimed at minimising the effect of the ischaemic insult.

Histology of a unilateral occlusion has been well described and has shown maximal damage within the thalamus and surrounding deep grey structures. In our bilateral occlusion animals, with reperfusion, there was maximal damage to blood vessels within the thalami with frank haemorrhage and Evans blue leakage. This is in spite of a blood flow during ischaemia comparable to the occipital region which appears to survive and in which autoregulation returns. Evans blue leakage appears to precede signs of frank haemorrhage in the area and both haemorrhage and Evans blue leakage are related to the length of the occlusive period, for we have not noted haemorrhages with occlusions lasting 30 minutes or less and Evans blue leakage is also less common with shorter occlusion times. These changes therefore would also appear to be related to duration of ischaemia.

To standardise the model, we have used adult male gerbils. Mortality is independent of sex in bilateral ischaemia, but uniformity in weight and maturity is important to exclude animals more resistant (young), or more sensitive (old) to ischaemia. We have chosen intraperitoneal barbiturate anaesthesia as it provides a reliable stable preparation even though there may be some reduction in mean CBF and an elevation in pCO₂. Its possible cerebral protective property is another source of artefact but as the technique is unchanged in all our studies the error introduced by this form of anaesthesia theoretically should be constant.

With occlusion, we have previously shown that there is an increase in cortical water as judged by a decrease in specific gravity following carotid occlusion. In this study we have extended our observations to the deep grey matter, cerebellum and pons. In the thalamus there is a progressive accumulation during ischaemia, but no change in specific gravities in the pons or cerebellum. During the restoration of
Reperfusion oedema following ischaemia

...flow there is a rapid accumulation of water which reaches a peak at 1–2 hours following the occlusion and then subsides. There is no change in water content during reperfusion in the pons and cerebellum where there was no ischaemia and none in the thalamus where there was no hyperaemia. Maximal resolution of oedema occurs in the occiput which has the highest cortical flow and in which autoregulation returned. The changes in specific gravity in general mirror the regional blood flow changes and the question arises as to whether the changes in specific gravity are real or artefacts due to an increase in local cerebral blood volume during the hyperaemia. This cannot be excluded in our experiments, nevertheless, our results are in broad agreement with those who have used differing methods for assessing water accumulation such as wet weight/dry weight and isotopes. The specific gravity technique assumes that there is no change in tissue solids during the measurements. If there is protein extravasation, a correction factor has been applied in the cat. We have been unable to apply the correction factor in gerbils, perhaps because of the distribution of grey and white matter within the gerbil brain. We do not consider that a change in tissue solids would significantly contribute to an error in our result prior to the leakage of Evans blue and a protein marker. The question arises as to whether the specific gravity changes are due to regional changes in blood volume. Shigeno et al have quantified this artefact and the magnitude of our changes are well in excess of those produced by blood volume. Our own studies on gerbils (unpublished) confirm this. Thus, the changes in specific gravity in the first hour of reperfusion would be generally reliable and in the areas where there was no Evans blue staining might be reliable for the total period of observation. The accumulation and resolution of oedema illustrates the dynamic situation that pertains following restoration flow. Preliminary studies showed that if the ischaemic flow was above 10 ml 100 min⁻¹, then the oedema accumulating during ischaemia would be resolved within an hour of reperfusion. In areas where the flow was less than 7 ml 100 min⁻¹ there was an increase in oedema after 1 hour’s restoration of flow. These time course studies demonstrate that within the severely ischaemic tissue there is resolution of the oedema after the initial increase associated with reperfusion. In the occipital regions with ischaemic flows in the region of 8 ml 100 min⁻¹ a borderline situation exists, and while the oedema resolves it does so more slowly in less ischaemic tissue.

The Evans blue extravasation demonstrating a loss of barrier to protein bound dye was later (1 hour after reperfusion), confined to the deep grey structures and over the short period of our reperfusion studies, progressive. It appeared in the same time course as the animal mortality and the question arises as to whether these observations are causally related or mere coincidence. Further work will be required to elucidate this.

In this study we have extended our observations on the regional cerebral blood flow, judged by hydrogen clearance technique in the gerbil. Flows in the parietal region (38 ml 100 min⁻¹) were well in accord with previous results. During bilateral occlusion, flow fell to practically zero in the frontal region with a gradation towards the occipital area. The thalamus had a surprisingly high flow but this is because of the contribution of the posterior circulation to that area. With restoration and flow there was a generalisation of hyperaemia which was maximal in areas with flow in the region of 4 ml 100 min⁻¹. In the frontal region, there was less hyperaemia than we anticipated and this may have been due to areas of microscopic impaired reperfusion. The hyperaemia corresponded roughly to the intensity and duration of the ischaemia, as reported by others. It was interesting that there was the maximal oedema in areas where the ischaemic flow was 4 ml 100 min⁻¹, in the same sort of values at which we noted maximal oedema during ischaemia and from which areas of intravascular potassium extravation has been reported.

Autoregulation is lost in all regions where flow is less than 13–15 ml 100 min⁻¹ in the primate, and in the gerbil Iannotti and Hoff have demonstrated a similar finding. In these studies we have shown that in the occipital region with a mean flow of 8 ml 100 min⁻¹, autoregulation returns 3 hours after the restoration of flow and this would suggest that in areas where the flow during ischaemia is above 8 ml 100 min⁻¹ autoregulation ultimately will return. We have also shown that in the same region, oedema will resolve and it might be concluded that even with this prolonged period of cerebral ischaemia ischaemic flow above 8 ml 100 min⁻¹ might be tolerated with ultimate recovery.

What conclusions might be drawn from our results? Firstly, that in ischaemic tissue there is a reproducible relationship between ischaemic blood flow and the development of oedema. During reperfusion, however, the situation is a dynamic one and while there is a rough correlation between the hyperaemia and the extent of cytotoxic oedema, the relationship is less close. The accumulation and resolution of oedema extends to tissues which will not survive and the time scale of the accumulation and resolution of cytotoxic oedema does not correlate with the time of the animal’s death. The latter is more closely related to signs of blood brain barrier...
and vascular endothelial damage. Lastly, we have shown that autoregulation will return in tissues where the flow is above 8 ml 100 g⁻¹ min⁻¹ for 1 hour and, with the exception of the thalamic area for which there may be local exceptional circumstances, this level of ischaemic flow might be compatible with ultimate survival and return to normal function.

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