Acute Ultrastructural Changes in the Middle Cerebral Artery Due to the Injury and Ischemia of Surgical Clamping

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SUMMARY: Ultrastructural changes in the zone of clamping of the middle cerebral artery of the squirrel monkey are described after application of a surgical clip. The experimental model utilized has been widely applied to the study of cerebral ischemia and possibly has relevance to clamps applied to the cerebral vessels during neurosurgical treatment of patients with cerebrovascular disorders.

INTRODUCTION
One of the most consistently reproducible animal models for studying experimental cerebral infarction is clamping of the middle cerebral artery (MCA) (Hudgins and Garcia, 1970; Sundt and Waltz, 1966). The clamps are usually those used in neurosurgery for treating cerebrovascular disorders such as ruptured aneurysm. After experimental clipping of the middle cerebral artery, the cerebrovascular changes mimic the hemodynamic, morphological and metabolic responses resulting from naturally occurring stroke in man. Numerous investigators have used this animal model for the study of ischemic effects on the brain.

Morphological changes of intimal (endothelial) involvement have been reported in studies of larger vessels (aorta and common carotid) following injury by suture (Webster, Bishop and Geer, 1974a, b.), ligation (Hoff, McDonald and Hays, 1968), and clipping (Kawamura et al., 1974; Nelson 1973; Nelson et al., 1975). However, no reports are available of the ultra-structural changes following injury to the entire wall of an artery due to clipping and the subsequent changes which occur in accompanying nerve fibers.

METHOD
Adult squirrel monkeys were used. After being lightly anesthetized (sodium pentobarbital), the animals were subjected to a transorbital surgical procedure (Hudgins and Garcia, 1970) to expose the right middle cerebral artery at the M-I segment (Krayenbuhl and Yasargil, 1968). A Heifetz clip was then placed on this area of sphenoidal segment for respective periods of...
either ½, 1, 2, 3, or 4 hours. Three animals were used for each time grouping with an additional three animals serving as controls. Controls were subjected to the surgical procedure but without arterial clamping.

The clip was removed at the end of the desired occlusive period and the animals were perfused through a cannula clamped into the left ventricle. The right atrium served as the site of drainage. Perfusate consisted of 3% glutaraldehyde in 0.1 M phosphate buffer. The perfusion pressure was maintained at systolic level in order to minimize the artifacts which occur in non-perfused arteries (Haudenschild, Baumgartner and Studer, 1972). Perfusate flow was maintained until 500 mls of solution had passed through the circulatory system of the animal. The middle cerebral arteries were carefully dissected from the clipped (right) side and the non-involved (left) side. The clipped artery was sampled at three locations — 6 mm proximal to the clipped area, 6 mm distal to the clipped area, and the immediate area of the clip.

The specimens were then placed in the glutaraldehyde/phosphate fixative for an additional 12 hours, washed in two buffer rinses, and post-fixed for 4 hours in a 1% osmium tetroxide /0.1 M phosphate solution. Following ethanol dehydration, embedment was carried out in Spurr embedding media (Spurr, 1969).

Semi-thin plastic sections were stained with Toluidine blue and scanned by light microscopy. Thin sections were evaluated with an RCA-EMU-4 electron microscope.

RESULTS

The arteries and associated nerve fibers from the left side of the occluded animals as well as from both sides in the sham operated animals contained no morphological variations from those of the normal M-I segments of the middle cerebral artery of the squirrel monkey (Dodson, Tagashira and Chu, 1975).

The tissue alterations for each clipped group will be reported as follows, based on the respective intervals of occlusion.

One-Half Hour Occluded Group

The predominant changes within the arterial wall were in the media. The two types of changes in the smooth muscle were increased density of the cytoplasmic ground substance, and/or large intramuscular vacuoles (Fig. 1). Changes in the medial layer were most prominent in the proximal and clipped areas and random degeneration of some myelinated fibers was present in neural compartments (Fig. 2). The periodicities of the myelin sheaths were often found to be replaced in part by a granular amorphic substance (Fig. 2). In the more advanced regions of degeneration, axonal shrinkage was also observed. Unmyelinated fibers were found to be stable, and neural changes were greater distal to the clip and were minimal in the proximal region.

One Hour Occluded Group

The proximal areas contained vacuoles within the medial layers similar to the ½ hour group (see above). The neuronal bundles contained Schwann cell changes (both intramyelin and intracellular vacuolization) as well as axonal shrinkage.

In the region of the clip, intimal changes consisted of endothelial breakdown and platelet adhesion. Extracellular pockets of edema were commonly found within the arterial wall. Intracellular vacuoles were seen in the media. Neuronal alterations varied, not only from bundle to bundle but also within the same bundle (Fig. 3). Neural degeneration was more advanced and widespread than in the ½ hour group.

The distal area of the artery contained comparatively fewer changes than the other two regions. A few vacuoles were observed in the medial layer. Alterations within the nerve bundles were limited to the...
myelin sheaths, consisting of bulges within the sheath as well as disruption of lamellar periodicity.

Two Hour Occluded Group
A few vacuoles were found in the medial layer of the proximal sample and only minimal myelin involvement was noted in the nerve bundles.

In the area of the clip, large intramuscular vacuoles were commonly observed on the adventitial side of the medial layer. Dense smooth muscle cells with or without vacuolar involvement were also seen. Platelet deposition was a common finding at the endothelial-intimal interface (Fig. 4). This relationship ranged from adherence in some regions to a continuous thin layer of platelets in other areas. Some subendothelial white blood cells were also evident.

Severe alterations were noted in some nerve bundles while varying morphological states were observed within the other bundles. Myelin sheath structures became rearranged into membrane whirls or separated into lamellar packets; intra-axonal reactions consisted of shrinkage and increased numbers of vacuolar constituents. Vacuoles were also common within the unmyelinated fibers and Schwann cell cytoplasm.

The arterial sections distal to the clip contained only a few abnormal features. Some vacuolar involvement was evident in the medial layers, and minimal alterations were seen within the myelin sheaths of the nerve bundles.

Three Hour Occluded Group
Changes in the arterial wall and in neural components were of the same nature and extent as reported in the proximal region of the 2 hour group.

The responses in the region of the clip were more evident than in the groups with shorter periods of occlusion. Platelet adhesion was widespread at the degenerating endothelial lining. Edematous pockets were present in the subendothelial regions as well as between some layers of the smooth muscle. The myofilaments and other structural elements of some muscle cells were displaced or replaced by large centrally located vacuoles which contained membranous swirls and granular material. Diapedesis was evident in the wall as reflected by the presence of numerous white blood cells.

The nerve bundles contained extensive alterations as did the arterial wall. Axonal compartments of myelinated and unmyelinated fibers were often represented by small clumps of dense material. Schwann cell involvement was expressed by myelin degeneration and by changes within their perikarya, such as cytoplasmic loss of organelles and peripheral clumping of chromatin within the karyoplasm.

The distal region was characterized by the same type of endothelial degeneration and platelet interaction as described in the clipped area (Fig. 5). Medial changes were not as widespread but were of the same nature (vacuolar and increased cytoplasmic density). Diapedesis was not as common in this area as it was in the clipped region.

Widespread degeneration was seen within the nerve bundles. As in the clipped area, both myelinated and unmyelinated fibers were replaced with amorphous material (Fig. 6). The epineurium, endoneurium, and Schwann cell components lost most of their recognizable integrity.

Four Hour Occluded Group
Proximal response was no more advanced in this group than in the 2 and 3 hour sections. Predominant features consisted of platelet interaction of the endothelial interface, vacuoles within the media, and myelin sheath alterations in the nerve bundles.

As in the 3 hour group, dramatic changes were evident in the region of the clip. Extensive platelet interaction with the remnants of a highly damaged endothelial region

**Figure 3**—Changes within a periaortic nerve bundle, which lies in the region of the clip, include axonal shrinkage (A), and/or myelin granulation (m) adjacent to morphologically intact fibers (X). One hour occluded animal (x2,500).

**Figure 4**—View of the wall from a two hour occluded animal shows endothelial damage (arrow), platelet deposition (P), as well as extensive vacuolization (X) of the media layer. Clipped region. (x7,200).
Figure 5—The region distal to the clip in this three hour occluded animal shows necrotic activity within all layers of the wall. A prominent layer of platelets (arrows) lines the luminal space (x5,000).

was commonly seen. Edematous pockets were found throughout the wall. Myofilaments were displaced by large intracellular vacuoles and loss of structural integrity was evident within the nerve bundles. Dense regions of crenated axons and scattered myelinated fragments were noted.

Greater alterations were present in the distal region. The luminal surface consisted of platelets and white blood cells which replaced the endothelial components in most areas (Fig. 7). Scattered, smooth muscle cells were identifiable in the medial layer, and the remainder of this layer was comprised of collagenous fibers, elastin components, and various phagocytic cell types. Neural components are recognizable only as dense regions with associated myelinated fragments (Fig. 8).

DISCUSSION

As a result of the technical approach used in this model, arterial regions were exposed to a combination of three insults — hemostasis and resulting hemoconcentration, trauma and ischemia (Meyer, 1958). It would appear from this study that some division of pathological events in each area can be related to the length of the occlusion. In those animals occluded for 1/2 hour or 1 hour, changes were more prominent in the proximal and clipped areas. These changes were consistent with the myonecrosis shown in experimental vasospasm (Alksne, 1974; Alksne and Greenhoot, 1974) while the dis-

Figure 7—Extensive alterations within the arterial wall are shown in this section. Abundant phagocytic cell types (P) and necrotic debris comprise most of the field. A few severely altered smooth muscle cells are identifiable (X). A dense border comprised of platelets and unidentifiable cell types line the lumen. Distal to the clip, four hour occluded animal (x4,200).

Figure 6—This field of an area distal to clip contains the remnants of an adventitial nerve bundle. Amorphous substance replaces the myelin lamella (X) while the axonal compartments (A) are also in a necrotic state. Three hour occluded animal (x15,000).

Figure 8—The neuronal elements in the region of Figure 7, are represented by the adventitial bundle in this field. Dense material characterizes the remains of the myelin sheaths (X) while amorphous granular regions are the presumed remnants of the axonal structures (A). Four hour occluded animal distal to the clip (X6,800).
tal regions did not contain intimal changes previously reported for acute response to ischemia (Hoff, McDonald and Hayes, 1968). It was concluded that the initial alterations were induced by chemical substances released from the damaged nerves and stagnant vascular elements and/or direct surgical trauma to the wall.

In the animals subjected to 2, 3, or 4 hour occlusions, major alterations were found in all layers of clipped and distal sections. The advanced ischemic changes of the intimal lining were in accord with intimal responses in distal areas of the surgically occluded carotid arteries (Nelson et al., 1975). The overall topography of the distally treated region of the arterial wall in the 2, 3, or 4 hour groups showed greater degenerative changes than in the proximal region. Findings in the distal areas were interpreted as the result of ischemic influence coupled with traumatically induced changes. This concept was reinforced since proximal and distal areas of the wall were sampled at equal distances from the clip, thus the traumatic changes should affect both equally.

Neural changes are most evident in the clipped and distal areas of the 2, 3, and 4 hour groups. Myelinated fibers show greatest susceptibility in the 3 and 4 hour groups where involvement is sufficiently extensive to include all elements (myelinated axons, unmyelinated axons, Schwann cells, and epineural sheath) of the nerve bundles.

In these acute studies, alterations in the arterial wall point out morphological changes which suggest certain undesirable features in the use of this model for acute and chronic experiments. Degeneration of the vascular innervation might be expected to significantly alter autoregulatory functions which are inherently important to such cerebral vessels.

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REFERENCES