neurogenic hyperventilation. However, in the presence of coexistent hypoxia in varying degree in many of these patients, it is difficult to differentiate "central neurogenic hyperventilation" from reflex hypopnea. All patients with respiratory pattern abnormalities showed varying degrees of respiratory alkalosis (most severe in patients with tachypnea, less severe in the presence of CSR and least obvious in the presence of CSV). The underlying mechanism for the development of respiratory alkalosis in acute stroke is not clear. However, it is probably related to a combination of both neurogenic factors and hypoxic drive.

Impedance pneumography appears to be a useful technique for detecting respiratory pattern abnormalities which would otherwise often not be detected in the usual clinical setting. The method permits constant monitoring of respiratory patterns but does not provide accurate information on volumetric pulmonary changes. However, in conjunction with intermittent measurements of pulmonary ventilation and arterial gas studies, this technique may prove to be useful in further elucidating complex neurogenic influences on respiration.

This study also suggests that the types of respiratory pattern changes seen in patients with acute brain stem infarction may have a prognostic significance. Quantitation of these abnormal respiratory patterns may thus provide useful prognostic information in the observation and management of patients with acute brain stem lesions.

References

Hemodynamic Derangement for the Induction of Cerebrovascular Fat Deposition in Normotensive Rats on a Hypercholesterolemic Diet

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SUMMARY Cerebrovascular ring-like fat deposition, which was noted only in hypertensive rats but never observed in normotensive rats even after they had been fed a high-fat cholesterol (HFC) diet for a long time, was successfully developed in the posterior communicating or other cerebrobasal arteries in normotensive rats fed an HFC diet for ten weeks after bilateral or unilateral carotid artery ligation or basilar artery ligation. These posterior communicating arteries with fat deposits were clearly dilated to a significant extent. These findings corroborated the fact that not only high blood pressure but also hemodynamic derangements induced by hypertension or other causes were important factors for the development of fat deposition in cerebral arteries.

Introduction
THE EXTREMELY RAPID development of ring-like arterial fat deposition in spontaneously hypertensive rats (SHR) fed a high-fat cholesterol (HFC) diet for two weeks was first reported by Yamori and its detailed mechanism was studied in SHR and other experimental hypertensive rats. Our previous studies further showed that fat depositions quickly developed even in the circle of Willis, especially in stroke-prone spontaneously hypertensive rats (SHRSP) and other experimental hypertensive rats on an HFC diet. Because normotensive Wistar-Kyoto (WK) rats and antihypertensive agent-treated experimental hypertensive rats on an HFC diet never had cerebrovascular fat deposition, hypertension was confirmed as the decisive factor for cerebrovascular fat deposition. We observed the dis-
tribution of cerebrovascular ring-like fat deposition in more than 300 hypertensive rats on an HFC diet. The initial ring-like deposits were located near such branchings as the carotid fork, basilar head and vertebrobasilar junction. Moreover, ring-like fat deposits in mesenteric arteries developed in relation to segmental increase in vascular permeability caused by vasoconstriction and dilation in hypertensive rats. These findings suggested that hemodynamic alteration induced by hypertension might be an important local factor for arterial fat deposition. Consequently, we experimentally tried to induce ring-like fat deposition in the cerebral arteries in "normotensive" rats by artificial hemodynamic derangements.

Methods

Twenty-four normotensive WK rats, 60 days old, and five WK rats, 40 days old, were operated on as follows under pentobarbital anesthesia (40 mg per kilogram, i.p.) to induce artificial hemodynamic derangements in the cerebrobasal arteries.

Carotid arteries were bilaterally or unilaterally ligated with great care in order to prevent vagal irritation or injury which might cause cerebral hypoxia due to respiratory disturbances. (All 40-day-old WK rats were used for bilateral carotid artery ligation.)

Basilar arteries were ligated after being exposed with a cervical transclival approach by a microsurgical technique. Sham operations were performed as follows: the pontine arteries were exposed through the cervical transclival approach and two pontine arteries were bilaterally electrocoagulated at their origin from the basilar artery. These operated rats were fed an HFC diet (20% suet, 5% cholesterol and 2% cholic acid) with 1% salt in the drinking water for ten weeks.

The serum cholesterol level (Zurkowski's method), tail blood pressure (a pulse-pick up method without anesthesia), and body weight were measured while these rats were on the HFC diet.

In order to observe the deposition in the cerebrobasal arteries, 75% barium sulfate with 10% gelatin was perfused through an aortic cannula after the preperfusion of 200 ml of heparinized physiological saline (at 38°C), with the pressure equal to the systolic pressure of individual rats (checked before perfusion). The brain was removed after being kept at -20°C in a freezer for one hour, fixed with 10% cold formalin and stained by Sudan III (barium contrasted Sudan staining method [BCSS method]). Fat deposits were observed not only macroscopically but also by light or electron microscopic techniques after the preparation of ultrathin sections from the specimens refixed in osmium tetroxide and embedded in Epon.

Diameters of the carotid and basilar portions in the posterior communicating arteries were measured with a micrometer by use of a microscope.

![FIGURE 1A. Ring-like fat deposits in the cerebrobasal arteries induced by artificial hemodynamic derangement in normotensive WK rats on an HFC diet for ten weeks from the age of 60 days. Encircled numbers (1 to 5) indicate internal carotid artery, the carotid portion of the posterior communicating artery, the basilar portion of the posterior communicating artery, the superior cerebellar artery and the basilar artery, respectively. Cerebrobasal arteries in non-operated control WK rats; no fat deposits are observed.](http://stroke.ahajournals.org/)

![FIGURE 1B. Cerebrobasal arteries in WK rats after bilateral carotid ligation. Multiple ring-like fat deposits are observed at the basilar portion of left posterior communicating artery. See legend of figure 1A for explanation of numbers.](http://stroke.ahajournals.org/)
Table 1: Acute Fat Deposition in the Circle of Willis Caused by Artificial Hemodynamic Derangements in WK Rats on an HFC Diet for Ten Weeks From the Age of 60 Days

<table>
<thead>
<tr>
<th>Fat deposition</th>
<th>Carotid artery ligation</th>
<th>Basilar artery ligation</th>
<th>Pontine A. electrocoagulation (sham operation)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Bilateral</td>
<td>Unilateral</td>
<td>Bilateral</td>
</tr>
<tr>
<td>Post. Comm. A.</td>
<td>+</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Bilateral</td>
<td>2</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Unilateral</td>
<td>0</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Diameters</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post. Comm. A.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carotid portion (mm)</td>
<td>4.0 ± 0.1*</td>
<td>2.7 ± 0.1*</td>
<td>1.8 ± 0.2†</td>
</tr>
<tr>
<td>Basilar portion (mm)</td>
<td>3.8 ± 0.1*</td>
<td>2.5 ± 0.1*</td>
<td>2.9 ± 0.1*</td>
</tr>
<tr>
<td>Serum cholesterol level (mg/dl)</td>
<td>533 ± 35</td>
<td>549 ± 21</td>
<td>540 ± 35</td>
</tr>
<tr>
<td>Blood pressure (mm Hg)</td>
<td>117 ± 6</td>
<td>116 ± 7</td>
<td>110 ± 4</td>
</tr>
<tr>
<td>Weight gain (gm)</td>
<td>56.7 ± 7.7</td>
<td>53.3 ± 7.3</td>
<td>53.3 ± 4.0</td>
</tr>
</tbody>
</table>

*Significant differences from the sham-operated group (P < 0.01).
†Significant differences from the sham-operated group (P < 0.05).

Results

Figure 1 shows acute fat deposition in the circle of Willis of WK rats on an HFC diet for ten weeks from the age of 60 days after operation. All numerical data are summarized in Table 1. All six bilateral ligations, four of six unilateral carotid ligations, and three of six basilar ligations resulted in the development of fat deposition in the posterior communicating artery.

These deposits were noted bilaterally (in four cases) or unilaterally (in two cases) after bilateral carotid ligation, unilaterally on the ligated side after unilateral carotid ligation, and unilaterally on either side after basilar artery ligation. However, electrocoagulation of pontine arteries as a sham operation never resulted in fat deposition.

The group of five young WK rats, in which carotid arteries were bilaterally ligated at the age of 40 days, showed more widespread distribution of fat deposition; three of them had ring-like fat deposits in the posterior cerebral and posterior communicating arteries bilaterally, one of them in the basilar artery, and the other one in the posterior cerebral artery bilaterally.

These fat depositions were ring-like and completely similar to those observed in hypertensive rats on an HFC diet.1,4 Histological study also confirmed that these arterial

Figure 1C. Exirpated cerebrobasal arteries of WK rats after bilateral carotid ligation. Some ring-like fat deposits are noted at the carotid and basilar portions of the right posterior communicating artery. See legend of figure 1A for explanation of numbers.

Figure 1D. Exirpated cerebrobasal arteries of WK rats after right carotid ligation. A clear ring of fat deposit is noted at the basilar artery. See legend of figure 1A for explanation of numbers.
lesions were similar to those in hypertensive rats\cite{3,4}; the intima was thickened and contained foam cells\cite{3} and the media consisted of markedly vacuolated smooth muscle cells (fig. 2).

It was clearly noted that the regions with ring-like fat deposition were dilated and frequently elongated into an S-shape. The diameters at the carotid and basilar portions of the posterior communicating artery in the groups of bilateral and unilateral carotid and basilar artery ligations were significantly larger than the diameters at the non-dilated communicating artery in the sham-operated group. Such marked dilatations were far beyond the normal variations, although the basilar portions of the posterior communicating arteries were variable in diameter. The fat deposits were located mainly at the extended wall of elongated and dilated arteries. Serum cholesterol level, blood pressure and gain in body weight showed no significant differences among four experimental groups, including the sham-operated group.

Discussion

An extremely rapid development of ring-like fat deposition "in the mesenteric arteries" of SHR on an HFC diet was reported first by Yamori,\cite{1} and subsequently Yamori et al.\cite{2,4} confirmed the same ring-like fat deposition "in the circle of Willis" by their new technique used to demonstrate fat deposition in the cerebral arteries as a whole (BCSS method). The significance of acute arterial fat deposition as a model for studies on the initiation mechanism of atherosclerosis or arteriosclerosis was further studied by our previous reports.\cite{8,9,10} The process of ring-like fat deposition was speculated upon from some experimental findings as follows: Repetition of vasoconstriction and vasodilatation causes a segmental increase in vascular permeability, which is followed by ring-like fat deposition, especially in the media of vasodilated regions. The fat deposition accelerates the degeneration of the media which further dilates the arterial wall and increases the permeability, and results in the rapid development of marked ring-like lesions. Moreover, our previous observations\cite{8,9,10} confirmed the following two findings concerning the pathogenetic factors for fat deposition in cerebrobasal arteries.

(1) The predilection sites of ring-like fat deposition in the cerebral arteries were arterial bifurcations in the cerebrobasal arteries.\cite{8,9} This finding suggested a relationship between hemodynamics of cerebral blood flow and fat deposition in the cerebral arteries.

\textbf{FIGURE 2.} Electron microscopic findings on the cerebrobasal artery with fat deposits. \textit{Left:} Thickening of intima with foam cell formation. \textit{Right:} Marked vacuolization of smooth muscle cells with fat deposition.
The decisive importance of hypertension in the arterial fat deposition was observed, especially in the cerebral arteries. Namely, it was noted that normotensive WK rats on an HFC diet never had fat deposition and that even SHR rarely had fat deposition in the cerebral artery under normotensive conditions (in the prehypertensive stage or under treatment by antihypertensive agents) in contrast to some grade of fat deposition noted in the mesenteric arteries in these SHR. In fact, many of our experimental cases revealed that blood pressure and the severity of fat deposition in the cerebral arteries showed a positive co-relation.

Based on these two experimental findings, we speculated about the possibility of inducing fat deposition in the cerebral arteries of "normotensive" rats, and succeeded in making ring-like fat depositions as observed in "hypertensive" rats and in the cerebral arteries of "normotensive" WK rats on an HFC diet after bilateral or unilateral carotid ligation or basilar ligation. The predilection sites of fat deposition in normotensive rats after these operations were the posterior communicating arteries, which frequently showed an elongation and were significantly dilated in the diameter because of the compensatory increase in blood flow. This experimental success clearly indicates the importance of hemodynamic derangement in arterial fat deposition. The hemodynamic concept of atherosclerosis was suggested from experiments by Texon et al., who observed atherosclerosis of the S-shaped carotid artery autotransplanted into the femoral artery in dogs fed cholesterol plus propylthiouracil. Our findings suggest that hemodynamic derangements induced by hypertension altered endothelial permeability which predisposed to lipid deposition and finally resulted in atherogenesis.

References
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