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A Standardized Method for Measuring Intracranial Arterial Stenosis

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BACKGROUND AND PURPOSE: Atherosclerosis of the major intracranial arteries is an important cause of ischemic stroke. We established measurement criteria to assess percent stenosis of a major intracranial artery (carotid, middle cerebral, vertebral, basilar) and determined the interobserver/intraobserver agreements and interclass/intraclass correlations of these measurements.

METHODS: We defined percent stenosis of an intracranial artery as follows: percent stenosis = \([1 - (D_{stenosis}/D_{normal})] \times 100\), where \(D_{stenosis}\) = the diameter of the artery at the site of the most severe stenosis and \(D_{normal}\) = the diameter of the proximal normal artery. If the proximal segment was diseased, contingency sites were chosen to measure \(D_{normal}\): distal artery (second choice), feeding artery (third choice). Using a hand-held digital caliper, three neuroradiologists independently measured \(D_{stenosis}\) and \(D_{normal}\) of 24 stenotic intracranial arteries. Each observer repeated the readings 4 weeks later. We determined how frequently two observers’ measurements of percent stenosis of each of the 24 diseased arteries differed by 10% or less.

RESULTS: Among the three pairs of observers, interobserver agreements were 88% (observer 1 versus observer 2), 79% (observer 1 versus observer 3), 75% (observer 2 versus observer 3) for the first reading and were 75% (observer 1 versus observer 2), 100% (observer 1 versus observer 3), and 71% (observer 2 versus observer 3) for the second reading. Intraobserver agreement for each of the observers was 88%, 83%, and 100%. Interclass correlation was 85% (first reading) and 87% (second reading). Intraclass correlation was 92% (first and second readings combined).

CONCLUSION: This method shows good interobserver and intraobserver agreements for the measurement of intracranial stenosis of a major artery. If validated in subsequent studies, this method may serve as a standard for the measurement of percent stenosis of an intracranial artery.

Atherosclerotic stenosis of the major intracranial arteries is an important cause of ischemic stroke. Currently, there are no standard methods for measuring the severity of intracranial arterial stenosis. The established methods for measuring extracranial carotid stenosis are not suitable for measuring percent stenosis of a major intracranial artery because the intracranial arteries have several branches, they become slightly narrower in their distal portions, and they are often tortuous (1–3). Additionally, extracranial carotid measurement techniques have to contend with the carotid bulb, which has no equivalent in any of the intracranial arteries.

In this study, we established measurement criteria for determining percent stenosis of the major intracranial arteries (carotid, middle cerebral, vertebral, basilar) and determined the interobserver/intraobserver agreements and interclass/intraclass correlations of these measurements.

Methods

Equation for Measuring Intracranial Arterial Stenosis

The equation used for determining percent stenosis of a major intracranial artery was as follows: percent stenosis = \([(1 - (D_{stenosis}/D_{normal})) \times 100\), where \(D_{stenosis}\) = the diameter of the artery at the site of the most severe degree of stenosis and \(D_{normal}\) = the diameter of the proximal normal artery. \(D_{normal}\) was determined by the following criteria: for the middle cerebral, intracranial vertebral, and basilar arteries, the diameter of the proximal part of the artery at its widest, non-tortuous, normal segment was chosen (first choice). If the
proximal artery was diseased (eg, middle cerebral artery origin stenosis), the diameter of the distal portion of the artery at its widest, parallel, non-tortuous normal segment was substituted (second choice). If the entire intracranial artery was diseased, the most distal, parallel, non-tortuous normal segment of the feeding artery was measured (third choice). For example, if the entire basilar artery was diseased, D_{normal} was measured at the most distal, parallel, non-tortuous normal segment of the dominant vertebral artery; if the entire middle cerebral artery was diseased, D_{normal} was measured at the most distal, parallel segment of the suprachiasmatic carotid artery; if the entire intracranial vertebral artery was diseased, D_{normal} was measured at the most distal, parallel, non-tortuous normal segment of the extracranial vertebral artery. Because of the variability of vasculature size, slight magnification differences, and subtle differences in anteroposterior projection, the contralateral circulation was not used as the “normal” reference measurements.

Measurement of the intracranial carotid artery required a slightly different approach because the caliber of this artery often gets slightly smaller after the origin of the ophthalmic artery, and measuring the normal cavernous portion of the intracranial carotid artery can be difficult because of the tortuosity of these segments. With this in mind, D_{normal} for the precavernous, cavernous, and postcavernous stenoses was measured at the widest, non-tortuous, normal portion of the petrous carotid artery that had parallel margins (first choice). If the entire petrous carotid artery was diseased, the most distal, parallel part of the extracranial internal carotid artery was substituted (second choice). Figure 1 illustrates how these rules were applied for measuring a stenosis of the carotid siphon and basilar artery.

If tandem intracranial lesions were present (eg, distal vertebral and mid-basilar), percent stenosis of both sites was measured at the most distal, parallel, non-tortuous normal segment of the feeding artery. If two vessels were diseased in tandem (eg, distal vertebral and mid-basilar), the percent stenosis of both sites was measured and the more severe stenosis was selected. When a “gap sign” was present (ie, the lumen of the vessel could not be visualized at the site of severe stenosis), D_{stenosis} could not be measured with calipers. In these cases, percent stenosis was defined as 99% luminal stenosis. Similarly, the interpretation of this measure is that the difference between two readings of the same film by the same reader would differ by no more than this amount, half the width of the confidence interval, 95%, 90%, or 85% of the time.

As a measure of reproducibility, we calculated confidence intervals (95%, 90%, 85%) for the difference between the two readings of the same film by the same reader. The interpretation of this measure is that the difference between two readings of the same film by the same reader would differ by no more than this amount, half the width of the confidence interval, 95%, 90%, or 85% of the time.

As a measure of reproducibility, we calculated confidence intervals (95%, 90%, 85%) for the difference between two readings of the same film by different readers. The interpretation of this measure is that the difference between two readings of the same film by different readers would differ by no more than this amount, half the width of the confidence interval, 95%, 90%, or 85% of the time.

The analysis of variance model also provided estimates of the interclass and intraclass correlation coefficients. As a de-
TABLE 1: Intraobserver agreement (% stenosis within 10%)

<table>
<thead>
<tr>
<th>Observer 1</th>
<th>Observer 2</th>
<th>Observer 3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1st vs 2nd</strong></td>
<td><strong>Reading</strong></td>
<td><strong>Reading</strong></td>
</tr>
<tr>
<td>First Reading</td>
<td>21/24(88%)*</td>
<td>20/24(83%)*</td>
</tr>
<tr>
<td>Second Reading</td>
<td>18/21(86%)*</td>
<td>17/21(81%)*</td>
</tr>
</tbody>
</table>

* % Agreement including all 24 arterial stenoses
† % Agreement excluding arterial stenoses with a gap sign

TABLE 2: Interobserver agreement (% stenosis within 10%)

<table>
<thead>
<tr>
<th>Obs 1 vs Obs 2</th>
<th>Obs 1 vs Obs 3</th>
<th>Obs 2 vs Obs 3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First Reading</strong></td>
<td><strong>Second Reading</strong></td>
<td></td>
</tr>
<tr>
<td>First Reading</td>
<td>21/24(88%)*</td>
<td>19/24(79%)*</td>
</tr>
<tr>
<td>Second Reading</td>
<td>18/21(86%)*</td>
<td>24/24(100%)*</td>
</tr>
</tbody>
</table>

* % Agreement including all 24 arterial stenoses
† % Agreement excluding arterial stenoses with a gap sign

Discussion

Angiographic measurement standards for determining percent stenosis of the extracranial internal carotid artery are well established (1–3, 6). These methods are routinely used in clinical practice to identify patients who may benefit from carotid endarterectomy. Currently, there is no equivalent method for measuring percent stenosis of the major intracranial arteries. This may be because until recently, there were limited data on the impact of the severity of intracranial stenosis on the risk of ischemic stroke. Recent studies suggest that the annual risk of stroke in patients with at least 50% stenosis of a major intracranial artery is 4% to 10% (7). Moreover, patients with severe intracranial stenosis (70% to 99%) have a higher risk of stroke than do patients with moderate intracranial stenosis (50% to 69%) (7). If the prognosis of intracranial arterial stenosis and the choice of therapy (eg, anticoagulation, angioplasty) for these patients are clearly shown to be based on the severity of intracranial stenosis, a reproducible and repeatable method for measuring percent stenosis of the major intracranial arteries will be required. In our study, intracranial arterial stenoses of more than 50% were chosen because of the significance of the stroke risk of high-grade intracranial stenosis as compared with lesions of lesser severity. Future studies may address low- and moderate-grade intracranial stenoses.

The method developed in this study seems to fulfill these criteria. Using a narrow agreement range (ie, within 10% of each reader’s measurement of percent stenosis), interobserver agreements for the three readers ranged from 71% to 100% and intraobserver agreements ranged from 83% to 100%. These rates of observer agreements are similar to those reported for the extracranial carotid artery (8) and exceed those reported for the coronary arteries (9). In a study assessing measurement of extracranial carotid stenosis, Rothwell et al (8) reported interobserver agreements of percent stenosis within 10% of two readers’ measurements as 70% for the European Carotid Surgery Trial (ECST) method, 74% for the North American Symptomatic Carotid Endarterectomy Trial (NASCET) method, and 75% for the Common Carotid method. Zir et al (9) reported a low interobserver agreement of four coronary angiographers in the interpretation of coronary artery stenosis. In this study, interobserver agreement for proximal or mid-left anterior descending stenosis (within 10%) ranged from 0% to 40%. The four observers in this study agreed regarding the presence of at least 50% stenosis in only 45% of left anterior descending...
lesions and 65% of right coronary artery lesions. The authors reported that a major reason for the poor results was disagreement regarding where to measure the normal vessel, $D_{normal}$. Thus, development of specific criteria for measuring $D_{normal}$ was suggested as one method of reducing interobserver variability. Hence, in the present study, we defined specific rules for determining the specific site at which to measure $D_{normal}$. Moreover, the three participating neuroradiologists in the present study underwent extensive training with reinforcement of the rules before obtaining their second measurements. We suspect that the success of this measuring technique depends in large portion on ability to follow these rules.

This study has shown that obtaining reproducible measurements of percent stenosis of an intracranial artery by using conventional cerebral angiography is possible using standardized measurement criteria. If validated in further studies, this method may serve as a standard for the measurement of percent stenosis of an intracranial artery obtained using conventional angiography.

References
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