Review Article

Asymptomatic and Symptomatic Carotid Stenosis: An Obsolete Classification?

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Since many years, clinical decisions about the management of patients with carotid stenosis have been based on the distinction between “asymptomatic” and “symptomatic” presentations. This was also reflected by the design of previous studies on the surgical versus conservative treatment and of current studies on interventional treatment versus surgery. Both terms, however, only address different phases of activity of the one and the same condition and blur the significant message that carotid stenosis is a most important marker of systemic atherosclerosis, which is accompanied by a much higher risk of cardiovascular events rather than stroke. As a consequence, early diagnosis and followup during best medical treatment, lifestyle management, regular cardiovascular assessment, and good control of all vascular risk factors should be recommended in all patients with carotid stenosis—whether identified in the long-lasting “silent” or short-lasting “vulnerable” period lasting only a few weeks after cerebral ischemia. Patients in this short time window benefit from additional carotid intervention, under the condition of an individually favorable benefit-risk ratio (“individual vulnerability”).

1. Introduction

Carotid stenosis is common, especially in patients with vascular risk factors or with coexistent pathology of coronary or peripheral arteries [1, 2]. Since many decades, the classification between “symptomatic” and “asymptomatic” carotid stenosis has dominated the management of affected patients. This distinction corresponds to the design and results of previous clinical trials on surgical versus medical treatment of carotid stenosis for stroke prevention as well as of current studies comparing surgical with interventional procedures (angioplasty with or without stent) [3–8].

However, careful review of these studies, as well as long-term natural history observations for many decades, shows that carotid stenosis is much more sensitive as marker of systemic atherosclerosis than a cause of stroke [9, 10]. Only within a small time window, when carotid stenosis shows progression and high plaque vulnerability, either in the presence of or without clinical signs or symptoms, there is a higher incidence of stroke.

First descriptions of carotid stenosis related to cerebrovascular events date back to T. Willis (1621–1675) and J. J. Wepfer (1620–1695). However, it was not before the 1950s when surgical interventions in the acute phase of stroke or for secondary stroke prevention were reported by DeBakey and Eastcott [11, 12].

In the 1980s, with the introduction of vascular ultrasound, the true dimensions of carotid disease became apparent. Quite unsurprisingly, a high number of patients were diagnosed with a so-called “asymptomatic” carotid stenosis [2], but the prognosis and management of these patients were totally uncertain. Studies performed since then showed already very early an eminent contrast between a very low incidence of stroke (1-2%/year) and a rather high cardiovascular morbidity and mortality (5–10%/year), in particular in patients with clinically silent presentation (so-called “asymptomatic carotid disease”) [9, 10]. The same is true for the “symptomatic” carotid stenosis too, but with one difference: there is a higher risk of stroke (10–20%) within the first 14–28 days following a cerebrovascular event (TIA
or stroke) [13]. After this vulnerable period, the stroke risk declines to that of the “asymptomatic” carotid stenosis.

2. Pathophysiology of Carotid Stenosis

In most cases, carotid stenosis is the result of atherosclerotic changes of the vessel wall. Other causes (e.g., dissection, often observed in patients under 55 years of age) only account for a small fraction of carotid lesions.

Atherosclerosis is a progressive disease with periods of stability and sometimes reparation, which begins with endothelial damage of the vessel wall already in infancy and adolescence. There are many factors promoting this damage: arterial hypertension, elevated blood lipid levels and problems of cholesterol or glucose metabolism, reduced release of nitrogen oxide, but also genetic components such as expression of vascular cell adhesion molecules (VCAM-1) [14–19]. Functional disturbance of laminar blood flow, particularly in arterial junctions, promotes the accumulation of LDL and consecutively lead to remodeling of the vessel wall. Monocyte migration and adhesion in the region of endothelial damage then follows, induced by proinflammatory cytokines [20, 21]. Identification of these mechanisms as biomarkers for the activity of atherosclerosis has confirmed the association of general risk factors with vascular degeneration but unfortunately did not contribute much to the sensitivity/specificity of a biomarker to predict the individual stroke risk in a “vulnerable patient.”

3. Diagnostic Methods

Significant improvements in diagnostic methods used for detection of carotid stenosis have influenced our knowledge about causes, spontaneous course, and risks of this disease. Initially, auscultation of carotid bruits was the gold standard in clinical practice. Systematic evaluation of this method, however, revealed a very low specificity and sensitivity [2]. Digital subtraction angiography (DSA) was for a long time the only method able to reliably identify a carotid stenosis, but it was only used in preselected patients because of its invasive nature and the risks associated with the procedure. It was the introduction of vascular ultrasound, with its noninvasive nature and good visualization, that made better carotid screening possible in a large number of patients, including many in clinically silent periods. Nowadays, highly developed ultrasound systems allow visualization of vessel wall changes in early stages, various grades of vessel stenosis and of intraluminal flow phenomena, as well as analysis of plaque texture and structure in real time and 3D or even 4D [22]. An experienced user can perform a highly sophisticated and reliable examination, which can be repeated during long-term monitoring without any risks or complications for the patient. Such technologies have recently been introduced in large clinical trials (e.g., PERFORM study) [23]. Ultrasound imaging can be used for screening and control of intima-media thickness (IMT), plaque morphology, grade and progression in clinical practice, as well as in clinical and epidemiological studies [24, 25]. Today, DSA is only rarely necessary—as opposed to the time when NASCET [3], ECST [4] and ACAS [5] were carried out. In addition to ultrasound, examinations such as CT and MR angiography allow collection of supplementary diagnostic insights, for example, perfusion studies, detection of vessel anomalies (aneurysms, angiomas), and plaque imaging, and can also be useful in cases where vessels are not well accessible via ultrasound or as an alternative investigation [26, 27].

Despite these technical approaches and changes in diagnostic practice, the current classification of the degree of stenosis still corresponds to the modalities of angiographic measurements used in the aforementioned clinical trials, because they formed the basis for evidence-based regimens. However, the information provided by ultrasound, CTA, and MRA today is by far better for an individual decision than the restricted use of only three stages of obstruction: high grade (>75%), moderate (60–75%), and low grade (<50% of local lumen reduction). If combined with other parameters such as individual clinical presentation (“the vulnerable patient”), modern brain imaging, and plaque structure analysis (“the vulnerable plaque”), a best estimation of the individual stroke risk associated with carotid disease can be made regarding the therapeutic consequences. These parameters allow a differentiation between “active” (vulnerable plaques) and “stable” disease. Echomorphologic features (plaque ulcerations, hemorrhage, lipid accumulation), which can be identified through high-definition ultrasound imaging and/or application of contrast agent, correlate with histopathologic characteristics and signalize a high risk for arterioarterial embolization [28]. HITS (high-intensity transient signals) registered over the middle cerebral artery indicate microembolization and can facilitate the decision of individual treatment strategies with suitable medication (CARESS [29]), as can detection of progressive carotid stenosis or silent infarcts in cerebral CT/MRT be interpreted as a sign of activity or acuity of a carotid stenosis [30–32].

4. Studies

Most of our knowledge on the topic of carotid disease, especially regarding treatment and course of the disease, was derived from studies done in the last 25 years. Some studied the natural history and reported clinical and vascular followup, as did Chambers and Norris (1986) [9] and Hennerici et al. (1987) [10]. As far as the so-called “symptomatic” carotid stenosis is concerned, two major, multicenter, randomized controlled studies are widely known: the North American Symptomatic Carotid Endarterectomy Trial (NASCET) with 2885 patients and the European Carotid Surgery Trial (ECST) with 3024 patients [3, 4].

The two randomized controlled studies mostly referenced on the subject of “asymptomatic” carotid stenosis are the Asymptomatic Carotid Atherosclerosis Study (ACAS) with 1662 and the Asymptomatic Carotid Surgery Trial (ACST) with 3120 patients [5, 6].

Table 1 summarizes the most important results of the aforementioned studies [3–6, 33–38].
Table 1: Outline of the most important characteristics and results of large carotid surgery trials. DSA: digital subtraction angiography.

|                        | NASCET | ECST   | ACAS   | ACST   |
|------------------------|--------|--------|--------|--------|
| $n$ (total)            | 2885   | 3024   | 1662   | 3120   |
| Observation period prior to inclusion (max.) | 120 days | 180 days | 120 days | 180 days |
| Ratio TIA/stroke as qualifying event prior to inclusion | 61%/39% | 50%/50% | —      | —      |
| Diagnostic method for determination of stenosis grade (SG) | DSA (distal SG) | DSA (local SG) | Doppler sonography/DSA | Duplex sonography |
| Significant correlation of stroke risk with SG | Yes | Yes | No | No |
| Controlled medical treatment | No | No | No | No |
| Indication for carotid surgery | Limited at 50–69% SG >70–99% SG | Limited at 50–69% SG >70–99% SG | No | Limited at >60–99% |
| Stroke risk (surgical treatment) (%) | 15.7 (5 yrs) 8.9 (3 yrs) | 15.0 (5 yrs) 10.5 (5 yrs) | 5.1 (5 yrs) | 6.4 (5 yrs) |
| Stroke risk (medical treatment) (%) | 22.2 (5 yrs) 28.3 (3 yrs) | 12.1 (5 yrs) 19.0 (5 yrs) | 11.0 (5 yrs) | 11.7 (5 yrs) |
| Absolute risk reduction (%) | 6.5 | 19.4 | — | 2.9 |
| Relative risk reduction (%) | 29 | 69 | — | 45 |
| Numbers needed to treat | 15 | 5 | — | 12 |

There is also a plethora of meta-analysis data available in the literature [34, 36–38].

5. “Symptomatic” Carotid Stenosis

In NASCET, the annual stroke rate under uncontrolled medical treatment (within 2 years of followup) was 13% for high-grade stenosis ($>70\%$ distal degree of stenosis) and about 7% for moderate stenosis. Carotid occlusions were found to be associated with a low risk of stroke (2.1%/year for ipsilateral and 5.5%/year for all strokes) [3]. Therefore, carotid endarterectomy was not recommended for low degree of stenosis, as well as for patients of subtotal carotid stenosis (“pseudocclusion”). The annual stroke rate in ECST was up to 7% for high-grade stenosis ($>90\%$ local degree of stenosis) in the—also uncontrolled—medical treatment branch within a 3-year followup and as low as 1.5% for 70–89% local degree of stenosis [36, 39]. Both studies recruited patients with a qualifying event (stroke, TIA or retinal TIA, the latter being the case in about 1/3 of NASCET patients) within a time window of 4 to 6 months prior to recruitment. Many of these cerebrovascular events were not evaluated with modern brain imaging (CT or MRI), and the association with the carotid stenosis was made only based on patients’ histories. Other coincident potential sources of stroke or TIA, such as lacunar infarcts due to small vessel disease or cardioembolic infarcts from atrial fibrillation, were not taken into consideration [40], which is a major drawback compared to modern clinical workup.

The results of both studies were interpreted as a significant advantage of carotid surgery over medical treatment, which mainly consisted of acetylsalicylic acid, but lacked systematic, prospective study design of other risk factor management and monitoring of compliance during the study.

Data in the literature indicates that relevant carotid stenosis ($>75\%$) is the underlying cause in only 5–12% of all cerebral ischemic events [3, 4]: about 20% of strokes in the territory of a “symptomatic” carotid stenosis cannot be without any doubt attributed to carotid stenosis, for example, in case of coexisting lacunar or cardioembolic causes [41].

The new ASCO stroke subtype classification [42] is suited to display such coexisting stroke causes. We prospectively studied 158 consecutive patients (89 men and 69 women, mean age 75 ± 11 years) with carotid stenosis $\geq 50\%$ as diagnosed on Doppler/Duplex vascular imaging admitted to our stroke unit in 2010 with the diagnosis of ischemic stroke (142; 89.9%) or TIA (16; 10.1%), based on clinical features and brain imaging (CT/MRI). Patients’ characteristics, risk factors, and the distribution of the grades of the 232 carotid stenosis detected are displayed in Table 2. The ASCO score (A: atherothrombosis; S: small vessel disease; C: cardioembolism; O: other causes with corresponding grades: (1) definitely a potential cause, (2) causality uncertain, (3) disease present, but unlikely a cause, (0) no disease present, and (9) no suitable tests performed) was used to classify stroke etiology after a thorough stroke workup was completed.

As determined by ASCO (Figure 1), carotid stenosis was the probable cause of the ischemic event only in less than half of the patients recruited (grade A1; 65; 41.1%). However, in 17 patients (10.8%) this was combined with other equally probable stroke causes (2; 3.1% S1 and 15; 9.5% C1). Interestingly, only if degrees of stenosis $\geq 70\%$ (78; 49.3%) were considered, the majority of patients were identified correctly. However, there were still 15 patients with A1 and
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![ASCO classification of stroke etiologies. The ASCO classification for "A" (atherothrombosis), "S" (small vessel disease), and "C" (cardioembolic). The "O" (other causes) group was omitted because only 4 patients received a grade other than "0." As all patients had Duplex/Doppler imaging and carotid stenosis, no patient had "A0" or "A9," so these groups were omitted. The figure visualizes competing/coexisting etiologies.](image)

![Table 2: Patient characteristics.](image)

| Parameter                        | Value | %  |
|----------------------------------|-------|----|
| Total of patients                | 158   | 100|
| Men                              | 89    | 56.3|
| Women                            | 69    | 43.7|
| Mean age (±SD)                   | 75 ±11| —  |
| Cerebral ischemia                | 142   | 89.9|
| TIA                              | 16    | 10.1|
| Unilateral stenosis              | 84    | 53.2|
| Bilateral stenosis               | 74    | 46.8|
| Carotid stenosis grades          | 232   |    |
| ≥50–59%                          | 74    | (31.9)|
| ≥60–69%                          | 68    | (29.3)|
| ≥70–79%                          | 24    | (10.3)|
| ≥80–89%                          | 28    | (12.1)|
| ≥90–99%                          | 29    | (12.5)|
| 100%                             | 9     | (3.9)|
| Concomitant vascular diseases    |       |    |
| Coronary artery disease          | 38    | 24.1*|
| Peripheral artery disease        | 18    | 11.4|
| Vascular risk factors            |       |    |
| Arterial hypertension            | 140   | 88.6|
| Hyperlipidemia                   | 77    | 48.7|
| Diabetes mellitus                | 54    | 34.2|

*Correlation with stenosis grade: $P < 0.05$.

6. “Asymptomatic” Carotid Stenosis

Both natural history studies, as well as several large, multicenter, randomized controlled studies, have shown that “asymptomatic” carotid stenosis is quite a benign disease, with an annual stroke rate of only 1–2% and a stroke-related annual mortality rate of merely <0.05% [5, 10].

Carotid stenosis, however, is an established risk factor for myocardial infarction. The annual mortality, mainly because of cardiovascular events, is about 6–8%. About 40% of patients with “asymptomatic” carotid stenosis have coincident coronary artery disease (CAD) [10, 43, 44]. In many patients, silent atherosclerotic alterations of coronary vessels lead to a significantly increased mortality. ACAS showed a mortality rate of 6% with one year of followup [5]. The highest prevalence (28%) of carotid stenosis was found in patients with peripheral artery disease (PAD); these patients also have the highest mortality [2].

In our patient group described above, about one-fourth of the patients (38; 24.1%) had CAD and 18 (11.4%) PAD. There was even a statistically significant correlation between grade of stenosis and presence of CAD ($P = 0.018$). Classic vascular risk factors were very common: arterial hypertension (140; 88.6%), hyperlipidemia (77; 48.7%), and diabetes mellitus (54; 34.2%).

ACAS and ACST could show a minor preventive effect of carotid surgery in carotid stenosis between 60% and 99%. A refined assessment of risk according to stenosis grade could not be made because of various methodological problems. In ACAS, absolute risk reduction of stroke and/or death was 5.9% and in ACST 5.4%, in a time frame of 5 years. However, it should be taken into consideration...
that the perioperative complication risk in both studies was extremely low (2.3% and 2.9%), because of a very strict selection of surgeons [5, 6]. Nevertheless, many patients were treated unnecessarily because of the large number needed to treat (17–19).

Control of risk factors was reported in ACST on a retrospective analysis suggesting that the already small benefit of surgery was minimized or lost if adequate monitoring of risk factor management had been achieved [6, 45].

Results from studies with very long followups from our department support these findings of low stroke mortality rate in these patients [10, 46].

### 7. New Assessment of Prognosis and Management

Based on the available data [47–51], a number of factors associated with a high risk of stroke on the basis of a carotid stenosis can be identified (Table 3).

- Acute hemispheric ischemia associated with carotid stenosis in the last 4 weeks
- Ipsilateral silent infarcts in CT/MRI
- Intracranial artery stenosis
- Contralateral carotid occlusion
- Insufficient medical treatment (risk factors)
- Insufficient collateralisation over the circulus Willisii
- Coexistent coronary/peripheral artery disease
- Leukoaraiosis
- HITS detection
- Rapid progression of carotid stenosis
- Plaque ulcerations
- Highly echolucent plaques in carotid duplexsonography
- High Lp-PLA2 Concentration

#### Table 3: Factors that increase stroke risk in patients with carotid stenosis.

| Factor                                      |
|--------------------------------------------|
| Acute hemispheric ischemia                  |
| Ipsilateral silent infarcts in CT/MRI       |
| Intracranial artery stenosis                |
| Contralateral carotid occlusion             |
| Insufficient medical treatment (risk factors)|
| Insufficient collateralisation over the circulus Willisii |
| Coexistent coronary/peripheral artery disease|
| Leukoaraiosis                               |
| HITS detection                              |
| Rapid progression of carotid stenosis       |
| Plaque ulcerations                          |
| Highly echolucent plaques in carotid duplexsonography |
| High Lp-PLA2 Concentration                  |

The problem rises at least partly from the different methods used to evaluate the stenosis grade (angiography versus ultrasound) and from the nonvalidated evaluation criteria. Therefore, ACAS and ACST could not show any connection between stenosis grade and stroke risk, whereas this connection was clearly shown not only in natural history studies but also in NASCET and ECST. Because patients nowadays do not usually undergo conventional angiography before carotid surgery, the direct justification for surgery by these studies’ results is actually missing.

Other problems and insecurities in the management of carotid stenosis result from the fact that a significantly positive effect of carotid surgery (in NASCET and ECST) was shown only in patients who underwent surgery within 2–4 weeks, but not in those operated upon later, up to 4–6 months. Furthermore, there are significant asymmetries in the recruitment of men/women in all 4 major studies. These asymmetries were not corrected in the analysis of the primary endpoints, thus undermining the results of ACAS and ACST in particular [62]. Possible overlapping of different stroke pathogenesis was not taken into consideration, which limits the direct transfer of study results to the individual patient in everyday clinical practice. In our patient collective, for example, carotid stenosis was the probable cause of stroke in less than half of patients and even in part of these patients there was an overlapping with other (mostly cardiac) causes. The ASCO score is especially suitable to demonstrate overlapping stroke etiologies and to quickly create a rudimental risk profile of the patient [42].

Uncritical usage of the results from the aforementioned studies often misguides and leads to false conclusions, generalizations, and uncertainty about best available modern treatment. A strict classification in “symptomatic” and “asymptomatic” carotid stenosis and decisions on the degree of carotid stenosis alone could therefore be misleading. Carotid disease should be rather regarded as an entity with active and stable phases. When deciding about treatment, one should differentiate between those two phases utilizing clinical findings, followups (e.g., rapid progression, repeated TIA), or other parameters (silent infarcts in brain imaging, HITS, plaque configuration). During an active phase, carotid surgery is advisable under the condition that it is performed early and that the perioperative complication risk is low (<5%) [47, 63]. Carotid angioplasty with or without stent is currently not a routine option but can be performed by highly experienced interventionalists in selected patients who cannot undergo a surgery. None of the major studies
of the medical treatment branch of ECST \[61\].

| Risk Factor                                      | Hazard Ratio | 95% CI    |
|-------------------------------------------------|--------------|-----------|
| Cerebral events versus ocular events            | 2.45         | 1.27–4.75 |
| Residual neurological signs after 7 days         | 1.30         | 1.08–1.57 |
| Diabetes                                        | 1.82         | 1.18–2.80 |
| Any ischemic event within the last 2 months     | 1.71         | 1.20–2.44 |
| Number of events within the last 3 months (per event) | 1.02     | 1.01–1.03 |
| Previous myocardial infarction                  | 1.31         | 1.04–1.65 |
| Degree of carotid stenosis                      | 1.34         | 1.30–1.38 |
| Plaque surface irregularity                     | 1.80         | 1.14–2.83 |
| Poststenotic collapse of the internal carotid artery | 0.40     | 0.17–0.94 |
| Age (per year)                                  | 1.01         | 0.98–1.03 |
| Male sex                                        | 1.23         | 0.83–1.82 |
| Systolic blood pressure (per 10 mmHg)           | 1.05         | 0.90–1.15 |
| Diastolic blood pressure (per 10 mmHg)          | 1.10         | 0.80–1.30 |
| Peripheral vascular disease                     | 1.03         | 0.65–1.63 |
| Angina without previous myocardial infarction   | 0.96         | 0.71–1.29 |
| ECG signs of left ventricular hypertrophy        | 1.07         | 0.40–2.10 |
| Cerebral infarct on symptomatic side on CT      | 1.32         | 0.88–1.96 |
| Occlusion of the contralateral internal carotid artery | 1.00   | 0.72–1.63 |

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