Ultrasound plaque characterisation, genetic markers and risks

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Abstract

The arterial wall changes detected by ultrasound are the end result of all risk factors (exogenous, endogenous and genetic) known and unknown and are better predictors of risk than any combination of conventional risk factors. However, ultrasound cannot be used in people younger than 45 because characteristic changes occur after this age. Nevertheless, it can be used in individuals over 45 to identify the genetic risk factors associated with atherosclerosis. The identification of genetic factors will subsequently provide a means of identifying individuals at risk at an early age, even childhood. In addition, knowledge of the genetic abnormalities associated with increased risk in a particular individual will provide a means of targeting prophylactic therapy.

Introduction

Extracranial atherosclerotic disease, known also as atherosclerotic disease of the carotid bifurcation has two main clinical manifestations (a) asymptomatic bruits and (b) cerebrovascular syndromes such as amaurosis fugax, transient ischaemic attacks (TIAs) or stroke which are often the result of plaque erosion or rupture with subsequent thrombosis producing occlusion or embolisation [1,2]. Internal carotid artery stenosis, the main consequence of atherosclerotic disease of the carotid bifurcation, remains the single preventable cause of ischaemic stroke.

Recent studies involving angiography, high-resolution ultrasound, thrombolytic therapy, plaque pathology, coagulation studies and more recently molecular biology have implicated atherosclerotic plaque rupture as a key mechanism responsible for the development of cerebrovascular events [3-5].

Atherosclerotic plaques consist of a lipid-rich core and a fibrous cap, which separates the core from the lumen. The lipid-rich core contains T cells and lipid-laden macrophages (foam cells), which are derived from blood monocytes [3]. T cells produce interferon-γ, which suppresses the production of collagen by the smooth muscle cells [4] and stimulates the macrophages to produce metalloproteinases (stromelysins, gelatinases, collagenases), which digest existing collagen and other extracellular matrix components. In addition foam cells produce tissue factor, which stimulates thrombus formation when in contact with blood after plaque rupture.

Rupture prone plaques tend to have a large lipid core and a fibrous cap, which separates the core from the lumen. The lipid-rich core contains T cells and lipid-laden macrophages (foam cells), which are derived from blood monocytes [3]. T cells produce interferon-γ, which suppresses the production of collagen by the smooth muscle cells [4] and stimulates the macrophages to produce metalloproteinases (stromelysins, gelatinases, collagenases), which digest existing collagen and other extracellular matrix components. In addition foam cells produce tissue factor, which stimulates thrombus formation when in contact with blood after plaque rupture.

Rupture prone plaques tend to have a large lipid core, a thin fibrous cap, few smooth muscle cells and an abundance of macrophages [5]. Ruptured plaques heal or enlarge by incorporating the thrombus formed on their surface. Some thrombi grow and occlude the lumen or produce emboli. Whether a
thrombus will occur or enlarge depends on the local blood flow and the hypercoagulable state.

Conventional arteriography has been used for several decades to investigate the presence and severity of internal carotid artery stenosis. Because it is invasive it cannot be repeated frequently and carries a risk of stroke of 1.2%; in addition angiography provides little information on plaque structure. The development and continuing technical improvement of non-invasive high-resolution vascular ultrasound has enabled us to study the presence, rate of progression or regression of plaques and most importantly their consistency. The ultrasonic characteristics of unstable (vulnerable) plaques have been determined [6,7] and populations or individuals at increased risk for cardiovascular events can now be identified [8]. In addition, high-resolution ultrasound has enabled us to identify the different ultrasonic characteristics of unstable carotid plaques associated with amaurosis fugax, TIA's, stroke and different patterns of CT-brain infarction [6,7]. This information has provided new insight into the pathophysiology of the different clinical manifestations of extracranial atherosclerotic cerebrovascular disease using noninvasive methods.

It has now been realised that there are many unknown risk factors for cardiovascular disease. The last few years have seen the emergence of some new ones such as oestrogen deficiency, hyper-homocysteinaemia, abnormal levels and function of fibrinogen, TPA, PAI-1, factor V, XII, VII and XIII, Lipoprotein(a) and C-reactive protein. In addition a number of cardiovascular genes have been identified with protective or harmful polymorphisms.

Some of the cardiovascular genes may influence lipid metabolism and the early formation of atherosclerotic plaques. Others may influence the rate of progression of plaques (rapid or slow) and yet others may determine whether plaques will be stable containing an abundance of collagen or unstable having a large lipid pool and a thin fibrous cap. Unstable plaques tend to ulcerate. Whether healing or thrombosis will subsequently occur is also influenced by the hypercoagulable state and thrombophilic genotypes.

The aim of this presentation is to highlight the advances of high-resolution ultrasound on carotid plaque characterisation, its clinical importance and the contribution of genetic epidemiology in understanding the basic pathophysiology involved.

The need for $\beta$-mode image normalisation - description of the method

High-resolution ultrasound provides information not only on the degree of carotid artery stenosis but also on the characteristics of the arterial wall including the size and consistency of the atherosclerotic plaque. Different classifications have been proposed in the literature, according to plaque consistency, resulting in considerable confusion. For example, plaques containing medium or high-level uniform echoes were classified as homogenous by Reilly [9] and correspond closely to Johnson's dense and calcified plaques, [10] to Gray-Weale's type 3 and 4 [11] and to Widder's type I and II plaques [12] (i.e. echogenic or hyperechoic). A recent consensus on carotid plaque characterisation has suggested that echodensity should reflect the overall brightness of the plaque with the term hyperechoic referring to echogenic and the term hypoechoic referring to echolucent plaques [13]. The reference structure, to which plaque echodensity should be compared with, is for hypoechoic plaques, blood; for the isoechoic, the sternomastoid muscle; and for the hyperechoic ones, the bone of the adjacent cervical vertebrae.

Measurements of texture should not be confused with measurements of echodensity. The term homogenous should refer to plaques of uniform consistency irrespective of whether they are predominantly hypoechoic or hyperechoic. The term heterogeneous should be used for plaques of non-uniform consistency, i.e. having both hypoechoic and hyperechoic areas. Although this may appear to be a relatively simplistic point of view, it is the basis for a standard classification of carotid plaques based on high-resolution ultrasound.

There is enough evidence published to support the clinical usefulness of ultrasonic plaque characterisation, patients with hypoechoic carotid plaques being at increased risk of stroke. Polak has recently investigated the association between stroke and internal carotid artery plaque echodensity [14]. In this study plaques were subjectively characterised as hypoechoic, isoechoic or hyperechoic in relation to the surrounding soft tissues. The stroke rate for hypoechoic plaques was 2.78 times higher than for isoechoic and hyperechoic plaques. The authors suggested that quantitative methods of grading carotid plaque echomorphology such as computer-assisted plaque characterisation, developed by our group, might be more precise in determining the association between hypoechoic (echolucent) plaques and the incidence of stroke.

Computer assisted plaque characterisation involves processing of digitised B-mode images of plaques taken from a duplex scanner with fixed instrument settings including gain and time control. The median of the frequency distribution of grey values of the pixels within the plaque (grey scale median-GSM, scale 0-255, 0 = black, 255 = white) is used as the measurement of echodensity. Early work has demonstrated that plaques with a GSM of less than 32, i.e. echoluent plaques have been found to have a five-fold increase in the prevalence of silent brain infarcts on CT-brain scans [15]. Soon it became apparent that ultrasonic image normalisation was necessary, so that images captured under different instrument settings, from different scanners, by different operators and through different peripherals such as video or magneto-optical disk could be comparable.

As a result a method has been developed to normalise...
images by means of digital image processing using blood and adventitia as the two reference points.[16] With the use of commercially available software (Adobe Photoshop version 3.0 or later, Adobe Systems Inc.) and the "histogram" facility, the gray scale median (GSM) of the 2 reference points (blood and adventitia) in the original B-mode image is determined. Algebraic (linear) scaling of the image is performed with the "curves" option of the software so that in the resultant image the GSM of blood equals 0 to 5 and that of the adventitia equals 185 to 195. Thus brightness of all pixels in the image including those of the plaque becomes adjusted according to the two reference points. This results in a significant improvement in the comparability of the ultrasonic tissue characteristics [16-18].

Accurate selection of the appropriate areas of blood and adventitia for image normalisation and the avoidance of areas of acoustic shadow in the selection of the plaque area are imperative. The duplex settings recommended are as follows: Maximum dynamic range, low persistence and high frame rate. A high frequency linear array transducer ideally 7-10 MHz should be used. A high dynamic range ensures a greater range of grey scale values. High frame rate ensures good temporal resolution. In addition to these pre-settings the time gain compensation curve should be positioned vertically through the lumen of the vessel, as there is little attenuation of the beam at this point. The overall gain should be adjusted to give optimum image quality (bright echoes with minimum noise in the blood). A linear post-processing curve should also be used and finally where possible the ultrasound beam should be at 90 degrees to the arterial wall.

The previously discussed guidelines should result in the following:
- An area of noiseless blood
- An echodense piece of adventitia in the vicinity of the plaque and
- Visualisation of the extent and borders of the plaque. It is here that colour images can provide further information about plaque outline.

Three major reproducibility studies have been performed in order to establish the validity of the method of image normalisation and the value of GSM measurements [16-18]. These three studies have demonstrated that GSM after image normalisation is a highly reproducible measurement that could be used in natural history studies of asymptomatic carotid atherosclerosis disease, aiming to identify patients at higher risk of stroke.

It should be pointed out that training is essential if the level of reproducibility reported above is to be achieved. Training is necessary not only in the use of the software but also in the appropriate scanning technique. For an experienced ultrasonographer training requires two days.

Clinical significance of carotid plaque echodensity and structure

The clinical importance of ultrasonic plaque characterisation could be focused on three main areas. First, cross-sectional studies aiming at better understanding of the pathophysiology of carotid disease. Second, natural history studies seeking to identify high and low risk groups for stroke; these studies should refine the indications on selection of symptomatic or asymptomatic patients not only for carotid endarterectomy but also for stenting. Third, ultrasonic carotid plaque characterisation can provide useful information for the endovascular specialist who is likely to use it to select patients for carotid angioplasty and stenting (CAS) [19].

Cross-sectional studies

The use of the above method of image normalisation and analysis has resulted in the identification of differences in carotid plaque structure – in terms of echodensity and degree of stenosis – not only between symtomatic and asymptomatic plaques but also between plaques associated with retinal and hemispheric symptoms [7]. In asymptomatic and symptomatic patients having amaurosis fugax, TIAs and stroke with good recovery having 50-99% stenosis on carotid duplex scan, plaques associated with symptoms were significantly more hypoechoic, with higher degrees of stenosis than those not associated with symptoms (mean GSM = 13.3 versus 30.5 and mean degree of stenosis = 80.5% versus 72.2%). Furthermore, plaques associated with amaurosis fugax were hypoechoic (mean GSM = 7.4) and severely stenotic (mean stenosis 85.6%). Plaques associated with TIAs and stroke had a similar echodensity and a similar degree of stenosis (mean GSM = 14.9 versus 15.8 and degree of stenosis = 79.3% versus 78.1%). These findings confirm previous reports, which have shown that hypoechoic plaques are more likely to be associated with symptoms [15, 16, 20]. In addition they support the hypothesis that amaurosis fugax has a different pathophysiological mechanism to that of TIAs and stroke.

Our group has found that GSM separates echomorphologically the carotid plaques associated with silent nonlacunar CT-demonstrated brain infarcts from plaques that are not so associated. The median GSM of plaques associated with ipsilateral nonlacunar silent CT-demonstrated brain infarcts was 14, and that of plaques that were not so associated was 30 (p=0.003) [18]. Additionally, emboli counted on transcranial Doppler (TCD) in the ipsilateral middle cerebral artery were more frequent in the presence of low-plaque echodensity (low GSM), but not in the presence of a high degree of stenosis. These data support the embolic nature of cerebrovascular symptomatology [21].

The role of biomechanical forces in the induction of plaque
fatigue and rupture has been emphasised [22-24]. In our group of patients, carotid plaques associated with amaurosis fugax were hypoechoic and were associated with very high-grade stenoses. It may well be that the plaques that are hypoechoic and homogenous, undergo low internal stresses and therefore do not rupture but progress to tighter stenosis with post-stenotic dilatation, turbulence, platelet adhesion in the post-stenotic area resulting in the eventual production of showers of small platelet emboli. Such small platelet emboli may be too small to produce hemispheric symptoms but are detected by the retina. In contrast plaques associated with TIAs and stroke were less hypoechoic and less stenotic than those associated with amaurosis fugax. These plaques are hypoechoic but more heterogeneous undergoing stronger internal stresses. Therefore, they may tend to rupture at an earlier stage (lower degrees of stenosis), producing larger particle debris (plaque constituents or thrombi) that deprives large areas of the brain of adequate perfusion.

Natural history studies

The first study, which has shown the value of ultrasonic characterisation of carotid bifurcation plaques in asymptomatic patients, was done by Johnson in the early 80’s. [10] In that study, hypoechoic carotid plaques in comparison with the hyperechoic or calcified ones increased the risk of stroke during a follow-up period of 3 years; this effect was prominent in patients with carotid stenosis more than 75% (as estimated by cross-sectional area calculations and spectral analysis), as stroke occurred in 19% of them. None of the patients with calcified plaques developed a stroke.

A second study performed in the 80’s by Sterpetti [25], has shown that the severity of stenosis (lumen diameter reduction greater than 50%, haemodynamic stenosis) and the presence of a heterogenous plaque were both independent risk factors for the development of new neurological deficits (TIA and stroke). Twenty seven percent of the patients with heterogenous plaques and haemodynamically significant stenosis developed new symptoms. Unfortunately, their study had mixed cases as 37% of the patients had a history of previous neurologic symptoms, mainly hemispheric ones. History of these neurologic symptoms was a risk factor for the development of new neurological symptoms during the follow-up period, although this was found only in the univariate analysis. Because no subgroup analysis was performed, no conclusion can be drawn regarding asymptomatic or symptomatic patients.

The last study published in the 80’s by Langsfeld [26] confirmed that patients with hypochoic plaques (type 1, predominantly echolucent raised lesion, with thin “egg shell” cap of echogenicity and type 2, echogenic lesions with substantial areas of echolucency) had a twofold risk of stroke of 15% in comparison with 7% of those having hyperechoic plaques (type 3, predominately echogenic with small area(s) of echolucency deeply localised and occupying less than a quarter of the plaque and type 4, uniformly dense echogenic lesions). Patients also with >75% stenosis, were at increased risk. However, the overall incidence of new symptoms was low, in contrast with the previous studies, perhaps because only asymptomatic patients were included in that study. Based on their results, the authors proposed an aggressive approach in those patients with >75% stenosis and heterogenous plaques. There is some confusion regarding the interchangeable use of the terms heterogenous and hypoechoic in that article. Additionally the authors raised the point that it is important for each laboratory to verify its ability to classify plaque types. The same group in another study published four years later reported a 5.7% annual vessel event rate (TIA and stroke) for echolucent carotid plaques versus 2.4% for the echogenic ones (p = 0.03) [27].

Given the fair interobserver reproducibility for type 1 plaques, the use of reference points was proposed: anechogenicity to be standardized against circulating blood, isoechogenicity against sternomastoid muscle and hyperechogenicity against bone (cervical vertebrae). A similar method has been used in the late 1990’s by Polak [14], who investigated the association between stroke and internal carotid artery plaque echodensity in 4886 asymptomatic individuals aged 65 years or older, who were followed up prospectively for 48 months. Some 68% of those had carotid artery stenosis, which exceeded 50% in 270 patients. In this study plaques were subjectively characterised as hypoechoic, isoechogenic or hyperechogenic in relation to the surrounding soft tissues. Hypoechoic plaques causing stenoses 50-100% were associated with a significantly higher incidence of ipsilateral, non-fatal stroke than iso- or hyperechogenic plaques of the same degree of stenosis (relative risk 2.78 and 3.08 respectively). The authors of this study suggested that quantitative methods of grading carotid plaque echomorphology such as computer-assisted plaque characterisation might be more precise in determining the association between hypoechoic (echolucent) plaques and the incidence of stroke.

The Tromsø study conducted in Norway in 223 subjects with carotid stenosis >35% has also found that subjects with echolucent atherosclerotic plaques have increased risk of ischaemic cerebrovascular events independent of degree of stenosis.[28] The authors give no details on patient’s neurological history. The adjusted relative risk for all cerebrovascular events in subjects with echolucent plaques was 4.6 (95% c.i. 1.1 to 18.9), and there was a significant linear trend (p=0.015) for higher risk with increasing plaque echolucency. Ipsilateral neurological events were also more frequent in patients with echolucent or predominantly echolucent plaques (17.4% and 14.7%, respectively). The authors concluded that evaluation of plaque morphology in addition to the grade of stenosis might improve clinical decision-making and differen-
tiate treatment for individual patients and that computer-quantified plaque morphology assessment, being a more objective method of ultrasonic plaque characterisation, may further improve this.

This method has been recently used by Grønholdt,[29] who found that echolucent plaques causing >50% diameter stenosis were associated with risk of future stroke in symptomatic (n=135) but not asymptomatic (n=111) individuals. Echogenicity of carotid plaques was evaluated with high-resolution B-mode ultrasound and computer-assisted image processing. The mean of the standardised median gray-scale values of the plaque was used to divide plaques into echolucent and echorich. Relative to symptomatic patients with echorich 50% to 79% stenotic plaques, those with echorich 80% to 99% stenotic plaques, echolucent 50% to 79% stenotic plaques, and echolucent 80% to 99% stenotic plaques had relative risks of ipsilateral ischaemic stroke of 3.1 (95% c.i., 0.7 to 14), 4.2 (95% c.i., 1.2 to 15), and 7.9 (95% c.i., 2.1 to 30), equivalent to absolute risk increase of 11%, 18%, and 28%, respectively. The authors suggested that measurement of echolucency, together with degree of stenosis, might improve selection of patients for carotid endarterectomy. The relatively small number of asymptomatic individuals was probably the reason explaining why plaque characterisation was not helpful in predicting risk in the asymptomatic group.

Genetic Epidemiology of Vascular Disease

One of the earliest identified and studied genes is the apolipoprotein E. ApoE which plays a central role in the regulation of lipid metabolism by mediating the binding of lipoprotein particles to the LDL receptor and ApoE(remnant) receptor. The ApoE gene has three alleles; E2, E3 and E4. These alleles interact differently with lipoprotein receptors and alter circulating lipid levels. Although 60% of variation in plasma cholesterol is genetically determined, ApoE polymorphism accounts for only 14% of this. This illustrates the important effect of other genes in lipid metabolism.

The frequency of the ApoE alleles in the European populations is as follows: E3 80%, E2 8% and E4 12%. The E2 allele is associated with low LDL, high triglycerides and low HDL. The E4 allele is associated with high LDL, high triglycerides and low HDL. Kinetic studies have shown that E4 allows the accumulation of LDL and E2 the accumulation of triglycerides. The presence of an E4 allele increases the risk of coronary heart disease and the genotype E3/E4 is associated with myocardial infarction at an early age [30]. In young men, under 40 years, undergoing coronary angioplasty the E4/E4 genotype was 16 times more common than in the general population [31]. The frequency of the E4 allele is increased in patients with ischaemic cerebrovascular disease [32] and in a recent case control study the E2 allele was associated with protection from stroke at age 70-79 but not in those over 80 years. Individuals not carrying an E2 allele had a relative risk of 2.6 for stroke [33].

Another recent case control study has investigated the frequency of the ApoE and beta-fibrinogen G/A-455 polymorphisms in ischaemic stroke [34]. After excluding intracranial haemorrhage, 277 patients with focal cerebral ischaemia were compared with 277 normal controls. There was no difference in the allele frequency between the groups but when stroke was classified according to type, the E4 allele and the A allele of the beta-fibrinogen genes were more frequent in those with stroke (n=70) due to large vessel disease. The key message is that clinical end-points can be the result of different pathologies and future studies should use end-points that are appropriate for the genes studied.

The realisation that clinical end-points can be the result of different pathologies has stimulated many research teams to use ultrasound in an attempt to produce better defined phenotypes. It has been demonstrated that carriers of E4 allele have carotid wall thickness greater than subjects with the E3 allele.[35] The Q451 allele of the CETP gene is associated with increased plasma CETP and lower carotid wall thickness.[36] This is an example of a protective gene polymorphism. The LL genotype of the PON1 gene is associated with presence and severity of carotid atherosclerosis on duplex (independent risk factor)[37] and so does the Hind III genotype of the lipoprotein lipase gene [38]. Individuals with carotid stenosis greater than 75% have a higher frequency (47%) of the V allele (A/V polymorphism) of the MTHFR gene than controls (27%) with carotid stenosis less than 25% [39]. The TT genotype of the p-fibronogen gene (C148T of polymorphism) is associated with the highest grades of carotid stenosis on duplex [40]. The D allele of the ACE gene (I/D polymorphism) and the D/D genotype are more frequent in patients with carotid stenosis or ischaemic cerebrovascular disease (independent risk factor) [41, 42].

Case control studies such as the above provide information on associations and do not allow inferences on causality. Only hypotheses can be made. What is needed now is prospective studies in which both clinical risk factors, ultrasound and genetic markers are included. With adequate follow-up of patients the causative link between molecular mechanisms and the effect of established risk factors on the formation, progression, type of and rupture of plaques and to clinical events can be obtained. This is our new challenge.
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