Recent Developments regarding Human Immunodeficiency Virus Infection and Stroke

Souvik Sena Alejandro A. Rabinsteinb Mitchell S.V. Elkindc William J. Powersd

a University of South Carolina, Columbia, S.C., b Mayo Clinic, Rochester, Minn., c Columbia University, New York, N.Y., and d University of North Carolina, Chapel Hill, N.C., USA

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Abstract
Human immunodeficiency virus (HIV) infection is strongly associated with ischemic stroke in the young. Data obtained from the Nationwide Inpatient Sample in the United States show an increase in the number of stroke hospitalizations in the HIV-infected population despite an overall decrease in the number of stroke hospitalizations. Few data exist, however, that address the mechanism of HIV-associated stroke. Recent studies have demonstrated that HIV may infect the endothelium and alter cerebrovascular functions. Whether the proposed mechanism alters the stroke risk is undetermined. Epidemiological studies suggest that HIV-related stroke is associated with a risk factor profile that differs from the HIV-negative young stroke population in that HIV-associated strokes are less likely to have hypertension, diabetes, hyperlipidemia and smoking as risk factors. A large population-based study, moreover, suggests an association between antiretroviral therapy and increased cardio- and cerebrovascular risks. Specific antiretroviral agents such as protease inhibitors and non-nucleoside reverse transcriptase inhibitors have been implicated in the metabolic syndrome, accelerated atherosclerosis and an increased risk for ischemic stroke. In addition to discussing these developments, this paper also discusses the implications of recent data for stroke prevention in HIV-infected patients.

Introduction

Stroke is a leading cause of adult disability in the United States [1]. Available data suggest that human immunodeficiency virus (HIV) infection is strongly associated with ischemic stroke in the young (15–44 years of age) [2]. Moreover, the number of stroke hospitalizations in the HIV population in the United States has increased by 61% from 1997 to 2006 despite an overall decrease in the number of stroke hospitalizations [3]. Few data exist, however, that address the mechanism of HIV-associated stroke. Recent studies have demonstrated that HIV may infect the endothelium and alter cerebrovascular functions, but whether this proposed mechanism increases stroke risk.
is uncertain. Epidemiological studies suggest that HIV-associated stroke is associated with a risk factor profile that differs from the HIV-negative young stroke population in that HIV-associated strokes are less likely to have hypertension, diabetes, hyperlipidemia and smoking as risk factors. Antiretroviral therapy may also be associated with an increased risk of cerebrovascular events. Specific antiretroviral agents such as protease inhibitors (PI) and non-nucleoside reverse transcriptase inhibitors (NNRTI) have been implicated in the metabolic syndrome, accelerated atherosclerosis and an increased risk for ischemic stroke. In addition to discussing these developments, this paper also discusses the implications for stroke prevention in HIV-infected patients.

**Epidemiology**

Epidemiological data on cerebrovascular disease in HIV-infected patients may differ depending on the type of the population (i.e. industrialized countries vs. Sub-Saharan Africa) and the date of the study period [i.e. before vs. after highly active antiretroviral therapy (HAART) implementation]. Consequently, the question of whether HIV infection is an independent risk factor for stroke is challenging. Necropsy studies of HIV-infected subjects have shown the prevalence of cerebral infarction ranging between 6 and 34% [4]. These pathological findings have often been considered symptomatic [4]. In fact, the rates of stroke-like presentations in different cohorts of patients with HIV infection have not exceeded 5%, supporting the subclinical nature of some of the lesions identified at necropsy [5, 6]. Many of these necropsy and clinical series date back to the pre-HAART era and include a majority of patients with advanced acquired immunodeficiency syndrome (AIDS). Yet, most contemporary clinical series consistently show that strokes continue to occur at young ages (<50 years) in HIV-infected patients [3, 7–9]. No population-based studies have been conducted to determine if the incidence of stroke is increased by HIV infection. Retrospective case-control studies, however, indicate that HIV infection and particularly the diagnosis of AIDS appear to be associated with an increased risk of stroke in industrialized countries [5, 10, 11]. More limited information is available from African cohorts [12, 13]. At this time, whether HIV infection is a risk factor for stroke in African patients remains unclear. A summary of data from industrialized countries and from African cohorts is displayed in table 1.

**Causes and Mechanisms**

Clinical series predating the broad use of HAART consistently reported that cerebral infarction was more frequent in patients with advanced immunosuppression, and cerebral infarction was often associated with opportunistic infections and tumors [14]. Marantic endocarditis, prothrombotic states and complications of intravenous drug abuse were also commonly implicated. A specific type of cerebral vasculopathy characterized by arterial ectasia and aneurysm formation was recognized first in HIV-infected children and then in young adults as a cause of cerebral infarction and brain hemorrhage [15]. Pathologically proven cases of vasculitis (i.e. inflammatory changes in the walls of small cerebral arteries) ascribed to HIV infection per se have been rarely described and their significance remains unclear. Evidence that HAART, particularly regimens containing PIs, produces metabolic changes that can accelerate atherosclerosis [16] and increase the risk of myocardial infarction [17] motivated the conduction of more recent studies evaluating the mechanisms of ischemia in HIV-infected patients.

In South Africa, during the years 2000–2006, only 12% of 67 HIV-infected patients treated for a first stroke had received antiretroviral therapy and at the time of their stroke, 46% of these patients had CD4 counts <200 cells/mm³ [7]. Ischemic strokes largely predominated (96%) and most patients (91%) were younger than 46 years. Recent or intercurrent opportunistic infections were noted in 25 cases (37%) and were deemed the cause of stroke in 18 patients (28%). The most common infection was tuberculosis (15%). Extracranial or intracranial (aneurysmal or non-aneurysmal) vasculopathy without documented risk factors for atherosclerosis (defined by the investigators as ‘HIV-associated vasculopathy’) was observed in 13 cases (20%). Anti-cardiolipin antibodies were present in 12 (19%) patients and cardioembolism in 9 (14%) patients. Traditional vascular risk factors were uncommon in these HIV-infected patients with stroke [7]. Among 82 patients with stroke [77 with ischemic infarction and 5 with intracerebral hemorrhage (ICH)] at a large inner-city hospital in the United States [8], 61% of patients (mean age 42 years) had been treated with HAART, but most patients were severely immunosuppressed at the time of the stroke (mean CD4 count was 113 cells/mm³ and 85% had counts <200 cells/mm³). The mechanism of ischemic stroke was classified as large artery atherosclerosis in 12%, cardiac embolism in 18%, small artery occlusion in 18%, other determined cause in
23% and cryptogenic in 29%. Confirmed or probable vasculitis was considered the cause of the stroke in 10 (13%) patients and stroke was attributed to hypercoagulability in 7 (9%) patients [8].

In summary, in contemporary clinical series, strokes tend to occur in young patients with uncontrolled HIV infection and more severe immunosuppression. These strokes may occur because of opportunistic infections associated with HIV and AIDS as there is evidence in non-HIV populations that chronic infections are non-specifically associated with stroke risk [18]. Atypical causes remain common and many events are classified as cryptogenic. The mechanisms implicated in HIV-related strokes are listed in Table 2.

### Cerebrovascular Function in HIV-Infected Patients

Studies of peripheral arterial function in HIV-infected patients have shown consistent reductions in flow-mediated arterial dilation indicating deficient generation of endothelium-dependent nitric oxide (NO). There appears to be a contribution from the infection as well as from the antiretroviral therapy [19]. The role played by NO in the regulation of cerebral blood flow (CBF) is under investigation. There is evidence NO is involved in vasodilatory CBF responses to hypercapnia, hypoxia and hypotension [20]. Inhibition of NO resulted in a greater reduction in CBF when systemic mean arterial pressure (MAP) was reduced from 100 to 70 mm Hg [20]. Impaired compensatory responses by the cerebral vasculature to reductions in systemic blood pressure could increase the

### Table 1. HIV infection and risk of stroke

<table>
<thead>
<tr>
<th>First author</th>
<th>Population</th>
<th>Methods/ study period</th>
<th>Rate</th>
<th>Comments</th>
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<tbody>
<tr>
<td><strong>Studies in HIV-infected patients</strong></td>
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<tr>
<td>Engstrom [6] (1989)</td>
<td>1,600 patients with AIDS</td>
<td>case series 1982–1987</td>
<td>12 (0.75%)</td>
<td>probable under-recognition</td>
</tr>
<tr>
<td>Connor [4] (2000)</td>
<td>183 necropsies of HIV cases</td>
<td>necropsy series</td>
<td>10 (5.5%)</td>
<td>excluded cases with CNS opportunistic infections</td>
</tr>
<tr>
<td>Evers [5] (2003)</td>
<td>772 patients with HIV</td>
<td>cohort study 1993–2001</td>
<td>15 IS/TIA (1.9%)</td>
<td>incidence of IS/TIA 216 per 100,000 patients/year</td>
</tr>
<tr>
<td>Corral [9] (2009)</td>
<td>2,012 patients with HIV treated with HAART</td>
<td>case series 1996–2008</td>
<td>27 IS/TIA in 25 patients (1.2%)</td>
<td>incidence of first-ever IS/TIA 189 per 100,000 patients/year</td>
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<tr>
<td><strong>Studies in stroke patients</strong></td>
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<tr>
<td>Hoffmann [12] (2000)</td>
<td>1,298 South African patients with stroke (151 blacks)</td>
<td>case-control</td>
<td>25 of 1,298 (1.9%); 24 of 151 blacks (15.9%)</td>
<td>no increase in the incidence rate of stroke compared with patients without HIV infection</td>
</tr>
<tr>
<td>Cole [2] (2004)</td>
<td>556 cases of stroke (385 IS, 171 ICH)</td>
<td>case-control 1988–1991</td>
<td>12 (2.2%)</td>
<td>incidence of IS and ICH in AIDS patients 0.2% per year for both ARR 9.1 for cerebral infarction and 12.7 for ICH</td>
</tr>
<tr>
<td>Tipping [7] (2007)</td>
<td>1,087 African patients with stroke</td>
<td>case series 2000–2006</td>
<td>67 (6.2%)</td>
<td>91% of HIV-infected patients with stroke were &lt;46 years</td>
</tr>
</tbody>
</table>

Each study has at least 100 patients included. Rate = Rate of stroke or HIV infection, respectively. CNS = Central nervous system; IS = ischemic stroke; OR = odds ratio; ARR = adjusted relative risk ratio.
risk of stroke, especially in the presence of underlying arterial stenosis.

Under normal conditions, changes in systemic blood pressure over a wide range have little effect on CBF due to autoregulation, which is mediated by changes in arteriolar diameter. When blood pressure increases, the small arterioles and arterioles constrict to maintain CBF at a normal level. Similarly, when cerebral perfusion pressure decreases, vasodilation of the small arteries or arterioles prevents CBF from falling. This mechanism is effective at maintaining CBF in normal human subjects until the capacity for blood vessels to dilate in response to reduced blood pressure is exhausted [21]. Within the limits of autoregulation, a 10% decrease in MAP produces only a slight (2–7%) decrease in regional CBF [22]. When blood pressure falls below the lower autoregulatory limit and the maximum compensatory vasodilatory capacity of the cerebral circulation has been exceeded, CBF will decline markedly with further reductions in cerebral perfusion pressure. Oxygen extraction, normally 30–40%, will progressively increase as CBF falls to maintain oxygen metabolism [23]. When the increase in oxygen extraction is maximal and is no longer adequate to supply the energy needs of the brain, further reductions in blood pressure disrupt normal cellular metabolism and may produce ischemic cell death [24].

Stenosis of the large cerebral arteries produces no hemodynamic effect until a critical reduction of 60–70% in vessel lumen occurs. At this point, the arterial blood pressure distal to the stenosis may be reduced [25]. Even with this or greater degrees of stenosis, however, distal intravascular pressure is variable and may even remain normal with stenosis exceeding 90% because the hemodynamic effect of carotid artery stenosis depends not only on the degree of stenosis but also on the adequacy of the collateral circulation [25]. Thus, even when systemic arterial pressure is normal or high, the smaller arteries and arterioles distal to a large artery stenosis may be dilated in response to the reduced local perfusion pressure. Any reduction in systemic blood pressure, even within the normal range, could further reduce the pressure beyond the stenosis enough to precipitate cerebral infarction. This phenomenon has been most clearly described in patients with internal carotid artery occlusion who have ipsilateral hemispheric increases in oxygen extraction, indicating severe reductions in distal intrarterial pressure. These patients are at high risk for stroke, but the stroke rarely occurs in the setting of any significant systemic hypotension [26].

Given the extensive literature about peripheral arterial function in HIV-infected patients, there are surprisingly few data on cerebrovascular function. Decreased common carotid artery distensibility or increased wall stiffness has been measured in HIV patients [27]. Decreased distensibility is present in early atherosclerosis; however, since the elastic properties of the common carotid artery are not important in the regulation of CBF,
the significance of this finding is uncertain. Among 31 HIV-infected subjects (aged 23–59 years) with no stenotic or occlusive large artery lesions and 10 similarly aged controls, middle cerebral artery blood flow velocity at baseline and after treatment with 1 g of acetazolamide (ACZ) was reduced in the HIV-infected subjects [28]. ACZ is a carbonic anhydrase inhibitor that dilates distal intracranial vessels by an unknown mechanism and is most commonly used in patients with large artery stenosis or occlusion to assess the hemodynamic effect on the distal circulation. A failure to normally increase CBF in response to ACZ is inferred to mean autoregulatory vasodilation has taken place in response to reduced downstream perfusion pressure, conferring an increased risk of stroke [29]. While abnormal ACZ reactivity in the absence of large artery disease presumably reflects smaller artery vasodilatory dysfunction, the clinical significance as far as conferring risk of stroke is unknown.

Direct (rather than inferred) determination of autoregulatory function requires measurement of the effects of changes in MAP on CBF. The autoregulatory index (AI) reflects the effectiveness of autoregulation: $AI = -\frac{\text{change in CBF}}{\text{change in MAP}}$.

The negative sign means that CBF decreases as MAP decreases. An AI of 0 would be perfect. In animal experiments, the normal AI varies by species, hypotensive agent and CBF method. AI is usually slightly negative, −0.06 to −0.29 [22]. There are no published normal data for humans.

In a pilot study of autoregulation in HIV-infected patients, CBF was measured by MRI with continuous arterial spin labeling in 5 subjects, 4 of whom were on antiretroviral treatment at baseline and after blood pressure reduction [unpubl. data, 2009]. Nicardipine infusion reduced MAP from 92 ± 11 to 83 ± 11 mm Hg. There was a statistically significant change in CBF from 51 ± 8 to 46 ± 8 ml/100 g/min ($p = 0.03$). AI was −1.2 ± 1.42. These pilot data suggest that autoregulatory vasodilatation in response to reduced MAP may be impaired in patients with HIV, but studies including more patients and normal controls are needed.

Although there is extensive literature about peripheral arterial function in HIV-infected patients, there are few data on cerebrovascular function, which has potential importance to the mechanism and prevention of stroke in these patients. The very small amount of data available suggests cerebrovascular function is impaired in HIV-infected patients. The relative contributions of infection and antiretroviral therapy as well as the relevance to HIV-related stroke are still to be determined.

**HAART and Stroke**

As discussed above, patients with HIV infection may suffer a stroke due to a wide spectrum of mechanisms and etiologies. HAART has been effective in reducing opportunistic infections in AIDS patients and has transformed HIV infection from a progressively fatal condition to a manageable chronic disease. Paradoxically, recent studies have found associations between HAART and vascular events, particularly myocardial infarction. In particular, there has been concern about a possible risk of accelerated atherosclerosis associated with HAART, particularly with PIs, which may also cause dyslipidemia and insulin resistance [30]. The specific association between HAART and stroke, however, is yet to be ascertained in a population-based study.

A retrospective review compared the incidence rate of various events, including stroke, among 506 patients treated with HAART to the incidence rate among patients treated before the HAART era [31]. During the study period, 6 of the 506 patients presented with cerebrovascular events. There were 1 subarachnoid hemorrhage, 1 ICH with cerebral toxoplasmosis and 4 ischemic events (3 instances of large vessel disease and 1 instance of lacunar infarction) [31]. The incidence rates of ischemic stroke and ICH were more than twofold higher than rates reported before the HAART era in different parts of the world. A more recent retrospective review of ischemic strokes and transient ischemic attacks (TIA) occurring in a cohort of HIV-infected patients treated with HAART from 1996 to 2008, compared with controls drawn from an unselected group in the same cohort, found an independent association of stroke with HAART, a prior diagnosis of AIDS and history of high alcohol consumption. The study concluded that stroke incidence is high in patients with HIV infection treated with HAART [9]. Refer to table 3 for a summary of studies examining HAART and the risk of stroke.

While these retrospective studies reveal higher stroke rates in HIV-infected subjects in the HAART era, they do not establish a specific association between HAART and stroke. In a large prospective cohort study, the incidence of cardio- and cerebrovascular events increased with longer exposure to HAART after excluding those patients associated with other concomitant central nervous system diseases [32]. In another study, however, the global incidence of cardio- or cerebrovascular disease was shown to be decreased in the short term in AIDS patients after treatment with HAART, excluding cases of cerebrovascular disease associated with drug dependence and

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**HIV and Stroke**

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HIV-related infections [33]. In neither study was the impact of HAART on the incidence of ischemic stroke analyzed as a specific endpoint. Moreover, mechanisms of stroke were not specified and while HAART may reduce stroke caused by AIDS-associated disorders, in the long term, HAART may increase athereothrombotic strokes. Nevertheless, an increase in the frequency of athereothrombotic strokes in patients receiving HAART could not be demonstrated in a recent clinical series [8]. Based on these large epidemiological studies, no conclusion can be drawn as to whether HAART is specifically associated with stroke. Whether such a differential association of HAART subtype exists with stroke also remains to be determined. If determined, the finding could have a major impact on stroke prevention in HIV-infected subjects.

One mechanism by which HAART may increase the risk of a vascular event, including stroke, is through accelerated atherosclerosis by inducing dyslipidemia and insulin resistance [16]. Other proposed mechanisms include HAART-induced inflammation, as evidenced by an increase in serum markers such as high-sensitivity C-reactive protein and interleukin-6, contributing to accelerated atherosclerosis [34]. There is also evidence HAART may reduce the effects of endothelial dysfunction induced by HIV. Despite the beneficial effect of HAART, the overall effect on atherosclerosis may be deleterious and translate into a clinical risk of stroke and cardiovascular disease. Data from a case-control study demonstrate a higher than expected prevalence of premature carotid lesions detected by ultrasound in PI-treated versus PI-naive patients [35]. These findings were confirmed in HIV-infected patients on combination antiretroviral therapy by comparing the risk of subclinical atherosclerosis between patients at low and high coronary risk. Combination antiretroviral therapy was found to be a strong, independent predictor for the development of subclinical atherosclerosis in HIV-infected patients, regardless of known major cardiovascular risk factors and atherogenic metabolic abnormalities induced by this therapy. The majority of these patients were treated with PI [16].

One plausible argument is HIV-associated strokes during the HAART era occur because HAART extends the life expectancy of HIV-infected individuals, therefore, increasing their risk of stroke. In a recent study, the median age for HIV-associated stroke was the 5th decade, which is much lower than in the non-HIV-infected population [3]. This finding suggests, during the HAART era, that HIV-associated stroke may not be due solely to longer exposure to vascular risk factors and/or HAART.

**Table 3.** HAART and the risk of stroke in HIV-infected patients

<table>
<thead>
<tr>
<th>First author</th>
<th>Population</th>
<th>Methods/study period</th>
<th>Effect size</th>
<th>Significance</th>
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<tbody>
<tr>
<td>D:A:D Study Group [32] (2004)</td>
<td>23,648 subjects in Europe, USA and Australia on HAART therapy for approximately 6 years</td>
<td>prospective cohorts 1999–2002</td>
<td>RR for cardiac and cerebrovascular events was 1.26/year of HAART exposure</td>
<td>p &lt; 0.0001 Poisson regression model</td>
</tr>
<tr>
<td>Subsai [31] (2006)</td>
<td>pre-HAART =155; HAART = 506, in Thailand followed over 2 years after initiation of HAART</td>
<td>retrospective case-cohort study 2002–2004</td>
<td>IS and HS incidence; pre-HAART: 0.2%/year; HAART: 0.4%/year</td>
<td>p = 0.001 Poisson regression analysis</td>
</tr>
<tr>
<td>Bozzette [58] (2008)</td>
<td>41,213 persons in the USA followed for 4 years</td>
<td>retrospective cohort 1993–2008</td>
<td>reduction in HR for death, cardiovascular events and stroke by 0.18 (PI or NNRTI) and over 72 months</td>
<td>95% CI 0.15–0.23</td>
</tr>
<tr>
<td>Corral [9] (2009)</td>
<td>2,012 HIV subjects in Spain initiated on HAART followed for a mean of 6.6 years; stroke/TIA cases = 25; control = 100</td>
<td>case-control study 1996–2008</td>
<td>fewer months under HAART was associated with cerebrovascular events (OR 0.97)</td>
<td>95% CI 0.96–0.99</td>
</tr>
</tbody>
</table>

D:A:D = Data collection on adverse events of anti-HIV drugs; RR = relative risk; IS = ischemic stroke; HR = hazard ratio; CI = confidence interval; OR = odds ratio.
Implications for Prevention of Stroke in HIV-Infected Patients

With the lack of clear specific mechanistic targets, prevention of stroke in HIV-infected patients is similar to prevention of stroke in patients without HIV, albeit with a few twists. As in patients without HIV, the first two steps of stroke prevention are the assessment of stroke risk and the treatment of conventional risk factors. The treatment of HIV can also be regarded as an important part of stroke prevention in these patients, as there is evidence that HIV may contribute to vasculopathy. Treatment using HAART, however, may produce or worsen vascular disease risk factors. Moreover, HAART requires particular choices of medications, monitoring of side effects and adjustments of medications, particularly in patients with a history of stroke or other vascular diseases.

Assessment of Risk

Current approaches to primary prevention of vascular disease and secondary stroke prevention rely on estimates of the absolute vascular risk [36]. The American Heart Association primary prevention guidelines state that those with ≥10% risk of myocardial infarction or coronary death over 10 years should be considered for aspirin therapy [37]. The NCEP (National Cholesterol Education Program) Adult Treatment Panel III guidelines state patients who have already experienced a cardiac event should be considered candidates for maximal medical therapy for secondary prevention, including statins with a goal LDL <100 mg/dl, or for those at ‘very high risk’ <70 mg/dl [38]. Included in the group of coronary risk equivalents are those with diabetes mellitus; patients with multiple risk factors whose composite risk score, based on the Framingham risk profile [39], puts them at a ≥20% 10-year risk, and patients with occlusive atherosclerotic disease, including those patients with symptomatic carotid artery disease. Although the NCEP guidelines do not explicitly address whether stroke patients should be included among risk equivalents, emerging data suggest that patients with stroke are at the same high level of risk as those with coronary artery disease, and that stroke patients could reasonably be included among coronary risk equivalents [40]. Given the clinical and public health importance of cerebrovascular disease, moreover, one can reasonably speak not only of ‘coronary risk equivalents’ but of ‘stroke and coronary risk equivalents’ [41]. Inclusion of stroke as the relevant and important outcomes is likely to be especially important among minority groups who are at an increased risk of stroke relative to coronary events [42]. Moreover, these minority groups are also likely to be overrepresented among those with HIV.

The first step in preventing a first stroke among HIV-infected patients, then, is to determine their absolute risk of stroke and other vascular events, using the Framingham risk score or a similar risk assessment tool, which can be easily found online [43]. While there is limited evidence that these risk scoring systems may underestimate the burden of cardiovascular disease in patients with HIV [44], at present, no other scoring systems have been generally accepted for this subgroup of patients [43, 45]. Those at high risk (>20% over 10 years) should then be considered for early treatment with antiretroviral therapy, as well as vascular preventive strategies [46].

Treatment of Conventional Risk Factors

Although there is limited evidence for secondary prevention of stroke in HIV patients, current guidelines regarding the use of antiplatelet agents, statins and blood pressure-lowering therapies should be used in patients with HIV as they are used in patients without HIV for the management of traditional risk factors in these patients [47]. For patients who have already experienced an ischemic stroke, identifying the specific proximate cause of the stroke is essential, when possible, to treat these patients accordingly. For instance, patients with atrial fibrillation will require anticoagulation and those with symptomatic carotid stenosis ≥70% most likely should undergo a carotid endarterectomy [48]. In the absence of these conditions, most patients with ischemic stroke will be candidates for antiplatelet therapy. Statin therapy and blood pressure-lowering therapy will also be indicated to control hypercholesterolemia and hypertension [46, 47].

Smoking is particularly prevalent among patients with HIV (47–71% of patients) and abstinence should be aggressively pursued [49]. Pharmacological therapy, including varenicline, bupropion, nortriptyline, clonidine and nicotine replacement, may all be tried. Currently, obesity is also common among patients with HIV, which represents a change from the early days of this disease, when many patients presented with wasting and cachexia. Obesity is present in >40% of women and >60% of men with HIV [50]. Therefore, encouraging weight loss and aerobic exercise 30–60 min at least 5 times weekly is important; in addition, patients should see a nutritionist when weight loss is not achieved [43].

There is also evidence that HIV, even before initiation of HAART, is associated with dyslipidemia [51].
Lipid abnormalities may occur in HIV-infected patients, including decreased total cholesterol, decreased LDL, decreased HDL, increased triglycerides and increased proatherogenic small LDL particles. Several of these abnormalities potentially have proatherogenic effects that may also increase stroke risk. These effects should be managed, as indicated, with statins, fibrates or other agents.

**Treatment of HIV**

There is some evidence that HIV, even before initiation of HAART, was associated with an increased risk of stroke. Some large trials, moreover, provide evidence that aggressive viral suppression with HAART reduces the risk of fatal or non-fatal cardiovascular events (annual event rate 1.3% among those randomized to drug conservation vs. 0.8% among those randomized to aggressive viral suppression, p = 0.05) [52]. For these reasons, treatment of HIV may be considered part of a strategy to reduce stroke risk, among other benefits. Current guidelines for the management of patients with HIV, therefore, include the presence of a high risk of vascular disease as a reason to initiate HAART, even among patients with CD4 counts >350/mm³ [45]. This guideline represents, in fact, one of the few official recommendations to use antiretroviral therapies to reduce the risk of vascular disease [53]. These vascular protective benefits of treatment, however, must be weighed against the risk of medication-induced metabolic abnormalities, including diabetes, which themselves can contribute to stroke risk.

**Adjusting HAART in the Setting of Vascular Disease, High Risk or Risk Factors**

The awareness of metabolic complications of specific antiretroviral agents has led to the search for agents with better metabolic profiles. The primary guide to choose the appropriate antiretroviral agent should, in general, be the efficacy of the agent in the individual patient against the virus. Treatment of metabolic complications can then be managed as needed. There are data to support the use of statins and fibrates for the management of lipid levels among patients on HAART, rather than to change to a different antiretroviral [54]. When choosing initial therapy, however, there are advantages to avoiding or choosing specific antiretroviral and lipid-lowering agents. For instance, the PI atazanavir appears less likely to be associated with hyperlipidemia and insulin resistance and should be considered, when possible, in patients who are known to be at high cardiovascular risk [55]. Abacavir has also been associated with increased cardiovascular risk and should be avoided among patients who are already at high risk [56]. PIs also downregulate the cytochrome P₄₅₀ enzymes (e.g. CYP3A4) involved in the metabolism of specific statins; thus they may increase the levels of those statins. Therefore, use of pravastatin, atorvastatin or rosuvastatin in preference to lovastatin or simvastatin is reasonable [51].

Insulin resistance and diabetes mellitus are also common side effects of HAART. Insulin resistance may occur in up to 90% of patients treated with PIs, and 1–11% may develop overt diabetes mellitus over the course of 5 years. Diabetes may be accompanied by dyslipidemia, hypertension and visceral fat accumulation (i.e. the metabolic syndrome). As previously mentioned, atazanavir is less likely to cause these side effects. Patients on HAART should be advised to follow a diet that contains 50–60% carbohydrates, 10–20% protein, <30% fat, <100 mg cholesterol and <10% of total calories from saturated fat. In addition, these patients should be advised to exercise. When needed, medications to control diabetes, including metformin, rosiglitazone, pioglitazone or insulin, should be prescribed to maintain a goal HbA₁C <6.5% and a fasting blood sugar of 73–110 mg/dl [43].

Lipodystrophy also occurs frequently among patients on HAART and may take the form of either fat accumulation, which occurs in 17–67% of patients on HAART, or lipoatrophy, which is found in 20–75% of patients on HAART. Fat accumulation is associated with the use of PIs, but may also be seen in HIV disease alone and with efavirenz, an NNRTI. Lipoatrophy is associated with NNRTI. The management of these conditions may include changing from a PI to an NNRTI, following a low-fat diet, exercise, use of testosterone and use of growth hormone. There is evidence the growth hormone-releasing hormone analog tesamorelin administered daily for 26 weeks decreases visceral fat by 18% and improves lipid profiles. These effects may be useful in HIV-infected patients who have treatment-associated central fat accumulation [57]. Further studies are needed to confirm these benefits and to demonstrate benefits in clinical outcomes among such patients.

**Conclusion**

Infection with HIV may contribute to an increased risk of stroke. The mechanism by which HIV infection leads to this increase in stroke risk needs to be determined. HAART, specifically PI and NNRTI, have been
associated with metabolic syndrome and accelerated ath erosclerosis, which may compound an increased stroke risk. These findings need to be confirmed in population-based studies. Once confirmed, these findings could have widespread implications for stroke prevention in HIV-infected subjects.

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Disclosure Statement

The authors have no conflicts of interest to declare.


