Chronic cerebrospinal venous insufficiency and the doubtful promise of an endovascular treatment for multiple sclerosis

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INTRODUCTION
Recently, a radically different concept regarding the pathogenesis of multiple sclerosis (MS) has been proposed. Termed chronic cerebrospinal venous insufficiency (CCSVI), it suggests that macro occlusive abnormalities of the extracranial venous drainage pathways of the brain and spinal cord can cause or contribute to MS. As a consequence of this theory, it has been suggested that angioplasty and possibly stenting of the internal jugular and/or azygos veins can improve the signs and symptoms of MS. These interventions have been performed sporadically across the globe in an open label fashion and never in the context of a well designed, controlled, randomized and blinded clinical trial. Despite this, the procedure has been labeled by some as ‘liberation procedure’ and caused a firestorm of interest in the medical and MS communities, both for and against its utilization. The arguments on all sides are passionate, ranging from the belief that venous intervention is a miracle cure that must not be withheld from the patients, to the feeling that the procedure is ineffective and unwarranted at best and dangerous at worst. The various camps commonly protest that those with differing views are not acting in the best interest of their patients.

THE CCSVI THEORY AND SUPPORTIVE DATA
In 2006, Zamboni, an Italian vascular surgeon, in an article titled ‘The big idea: iron-dependent inflammation in venous disease and proposed parallels in multiple sclerosis’ suggested that there were similarities between chronic venous disease of the extremities and MS.2 He raised the possibility that venous reflux or obstruction in cerebral and spinal veins might have a relationship to MS.5 Several years later, Zamboni et al reported on blinded transcranial and extracranial color Doppler sonographic findings in patients with MS and matched healthy controls and those with other neurological disorders.3 They focused on five findings: (1) reflux in the internal jugular vein (IJV) or vertebral veins >0.88 s; (2) reflux propagated in at least one out of the three deep cerebral veins >0.55 s; (3) high resolution B mode evidence of proximal IJV stenosis; (4) flow not Doppler detectable in the IJV or vertebral veins despite deep inspirations; and (5) negative difference of the cross sectional area of the IJV comparing the value obtained in the supine versus the sitting position. The authors concluded that detection of two or more of these findings constitutes the diagnosis of CCSVI. They found CCSVI in all MS patients and in none of the controls. The sensitivity, specificity, positive predictive value and negative predictive value of the test were all 100%. They concluded that there was CCSVI in MS patients.3

In a second paper, Zamboni et al published that catheter venography in patients who met CCSVI Doppler criteria showed stenosis in the azygos vein 86% of patients who met CCSVI Doppler criteria published that catheter venography in patients who met CCSVI Doppler criteria showed stenosis in the azygos vein.3

As neurointerventionalists interested in interventional treatment of neurological disorders, it is time to take a thorough and objective look at CCSVI. This commentary will examine the origin of the CCSVI theory and discuss the data supporting and refuting its existence. An attempt will be made to critically analyze the available data and provide constructive recommendations about whether or not endovascular therapy represents a reasonable option at this point in time for patients with MS.
IJV and proximal azygos vein stenosis; (C) both IJV and normal azygos system; and (D) multilevel azygos stenosis with or without IJV involvement.

Finally, in 2009, Zamboni et al reported their results on the endovascular treatment of 65 MS patients with CCSVI.7 No isolated venous lesion was found, and the distribution of venographic patterns was 30%, 38%, 14% and 18% of types A to D, respectively.5 They performed percutaneous transluminal angioplasty (PTA) on all but one azygos lesion that did not respond to PTA alone and required stent placement. Pretreatment pressures beyond the stenosis were not significantly different than normal venous pressure and there was no significant change in pressure after angioplasty. Mean follow-up using extracranial Doppler was 18 months, with an overall restenosis rate of 47%; more common in the jugular than azygos veins. Clinical outcome at 18 months was reported as showing relapse free of 50% versus 27% preoperatively. It is important to note that the interpretation of the clinical results of this uncontrolled study is confounded since patients were continued on ‘immunomodulating’ therapy after endovascular therapy. These medical therapies have been shown to significantly reduce relapse rates as well as the accumulation of MRI detectable enhancing lesions. Finally, there was no improvement in patients with primary progressive or secondary progressive MS.5

DATA AGAINST CCSVI ROLE IN MS

Although the Zamboni papers have been quite supportive of CCSVI, there are a growing number of papers that raise serious questions about its validity. In early 2010, Khan et al described a number of independently accepted characteristics of venous disease and MS that contradict the CCSVI theory.6

1. Similar to other autoimmune diseases, MS is more common in young women while chronic venous insufficiency syndromes are not.
2. There are well known strong epidemiological associations between MS and geography, ethnicity, sun exposure, low vitamin D levels, gender, genetics and immigration studies that are not mirrored by chronic venous insufficiency.
3. Central veno-occlusive disease can lead to syndromes of idiopathic intracranial hypertension, ischemic and hemorrhagic infarcts and edema, none of which is typically seen in MS patients.
4. Vascular abnormalities related to chronically diminished venous flow would be expected to increase over time, yet after the age of 50 years the incidence of MS is quite low.
5. There is no other model of decreased venous drainage and an organ specific immune response.
6. Transient global ischemia is known to occur with jugular insufficiency but this entity is not seen in MS.
7. Radical neck dissections remove all jugular veins but they have never been seen to cause MS.6

The above cited challenges to the Zamboni thesis are based on largely theoretical considerations. In an attempt to replicate the Doppler findings of Zamboni, Doeppe et al studied 56 MS patients and 20 controls using similar CCSVI criteria.7 The authors found no patients in either the MS or control groups who had the two or more criteria required for a diagnosis of CCSVI. They concluded based on these results as well as their extensive longitudinal experience with cranial venous Doppler ultrasound, that there is typically tremendous reserve capacity of the extrajugular pathways for cerebral venous drainage and that it is highly unlikely that IJV stenosis would cause central venous congestion. Furthermore, they went on to discourage interventional procedures for CCSVI outside of the context of appropriately designed clinical research studies.3

Additionally, Sundstrom et al looked at MRI of 21 patients with relapsing remitting MS and 20 healthy controls, and found no differences in internal jugular venous outflow between the two groups.9 Finally, preliminary data from Zivadinov et al, from the MS research group at the State University of New York in Buffalo, presented findings in the first 500 participants studied with venous Doppler looking at the prevalence of CCSVI in MS patients and controls. Using the requirement that ≥2 CCSVI Doppler criteria be met, CCSVI was found in 62.5% of MS patients, 25.9% of healthy controls and 45% of other neurological disorders.10 At least preliminarily, these results are different from the 100% sensitivity and specificity found by Zamboni and colleagues.3

COMMENTARY

There is little debate as to the potential ravages of MS and the sincere desire to improve outcomes in patients suffering from this terrible disease. As such, when seemingly miraculous cures are proffered, we believe that it is our responsibility as neurointerventionalists to rationally review its use.

There are few data to support the validity of CCSVI. The lack of data seems counterbalanced by the great hope for the miracle of an endovascular treatment for MS. The topic has caused widespread attention and debate in the media, medical literature and the internet.11–17 As of late August 2010, a Google search on ‘liberation procedure’ yielded about 2,650,000 results and approximately 181,000 for ‘CCSVI’. Sponsored links appear for treatment in Mexico, Poland, Costa Rica, India and other locations. At least one toll free telephone number akin to ‘1-800-I Treat MS’ has been created.18

The prospect of opening an open label, non-study related MS endovascular CCSVI practice can be very seductive. For physicians, the barriers to entry are small since most interventionalists are technically able to perform these procedures and the required devices are readily available. At the same time, there are many patients who are desperate for a procedure which might improve their condition despite the lack of evidence to support its benefits and almost regardless of its potential risks. Indeed, some might argue that because the procedure is safe, if there is any possibility of ameliorating some of the symptoms of MS patients the procedure should be offered to them. However, no invasive procedure is completely safe. In fact, there are increasing reports of complications related to PTA or stenting for CCSVI, including intracranial hemorrhage, stent migration into the heart and jugular vein thrombosis.19

The moniker, ‘liberation procedure’, is a marketer’s dream and by itself suggests unrealistic but compelling expectations. Many patients are willing to pay cash, sometimes tens of thousands of dollars, for a single procedure. Many patients rave about their procedures, yet outside of a well controlled trial, it is hard to disprove the placebo effect and prove the true clinical benefits.

In view of the foregoing, and in an attempt to help resolve the CCSVI conundrum, it would seem that the fundamental questions are:
1. Is there a cause and effect relationship between CCSVI and MS, and in which direction does this work?
2. If CCSVI does cause or worsen MS, should this be treated with endovascular therapies?
3. If endovascular treatment is contemplated, which therapy should be offered?
and under what technical and clinical circumstances should they be applied? There is paramount need for credible scientific evidence that will allow us to address these questions. Firstly, we should encourage trials using non-invasive studies to test if CCSVI–MS actually exists. At the current time, the corroboration evidence supporting Zamboni’s initial findings of an association between CCSVI and MS are limited. In fact, the majority of additional evidence—including the work of Doepp et al and Sundstrom et al, cited in this review—actually failed to replicate the findings of Zamboni and colleagues.7–9

Moreover, the early results of Zivadinov et al are also not very compelling.10 In addition, the initial claim by Zamboni et al that they had developed a perfect test for CCSVI–MS raises serious questions about the credibility of their evidence. As pointed out by Novella, few if any tests in medicine have 100% sensitivity and 100% specificity.20

Fortunately, the US and Canadian MS societies have undertaken seven studies to investigate the CCSVI–MS association.14,15 The necessity of requiring an invasive diagnostic study such as catheter venography to evaluate the CCSVI–MS association is more difficult to reconcile at this point, particularly since the seminal findings of Zamboni et al which initiated this entire controversy were based on non-invasive Doppler ultrasound.

If the association between CCSVI and MS cannot be confirmed, then further studies evaluating CCSVI treatment are unnecessary. While it could be argued that even if the prevalence of venous ‘abnormalities’ is similar in patients with MS and controls, venous intervention in MS should still be studied since MS patients might be more susceptible to the detrimental effects of CCSVI than normal patients, this position seems tenuous at best.

If an association between CCSVI and MS can be established, then the next logical step would be to design multicenter randomized clinical trials to assess the benefits of endovascular interventions.

CONCLUSION

More evidence is needed to establish the association between CCSVI and MS. If more solid clinical evidence can be provided that the CCSVI–MS relationship is real, randomized clinical trials will be required to assess the benefits of endovascular interventions. If these trials establish a benefit for endovascular therapy, then this point can be assessed in our opinion, there is no role for the endovascular treatment of CCSVI in the MS patient outside of approved clinical trials.

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Several author names were omitted from an article published in the December 2010 issue of the journal (Chronic cerebrospinal venous insufficiency and the doubtful promise of an endovascular treatment for multiple sclerosis. J NeuroIntervent Surg 2010;2:309–11. Published Online First: 23 October 2010. doi:10.1136/jnis.2010.003947). On page 310, second paragraph, fourth sentence should read: ‘Pretreatment pressures beyond the stenosis were not significantly different from normal venous pressure, however, a change in pressure was demonstrated after angioplasty’. J NeuroIntervent Surg 2011;3:97. doi:10.1136/jnis.2010.003947corr1