Manual Therapy and Cervical Arterial Dysfunction, Directions for the Future: A Clinical Perspective

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Abstract: This paper offers a contemporary, evidence-based perspective on the issue of adverse neurovascular events related to cervical spine manual therapy. The purpose of this perspective is to challenge traditional thought and practice and to recognize areas where practice and research should develop. By considering the themes presented in this paper, the clinician can broaden his or her approach to neurovascular assessment in line with contemporary evidence and thought. We present information based on clinically relevant questions. The nature of vertebrobasilar insufficiency and the utility of pre-treatment testing are examined in light of contemporary evidence. In addition, we report on internal carotid artery pathology, and the significance of appreciating atherosclerosis in clinical decision-making. These later two areas are not commonly recognized within manual therapy literature, and we suggest that their importance to differential diagnosis of head and neck pain, as well as estimating treatment related risk, is paramount. We propose that the term cervical arterial dysfunction is more appropriate than classically used nomenclature. This term refers more accurately and completely to the range of pathologies at different anatomical sites that manual therapists treating patients with head and neck pain are likely to encounter. Finally, we present a brief review of the medico-legal status pertaining to this area. Although this is English law-related, the themes derived from this section are of interest to all manual therapists.


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Manual therapy (MT) is a commonly used intervention for the management of cervico-cranial pain1. Despite evidence demonstrating potential efficacy of this intervention2,3, there are known risks that have been associated with some aspects of cervical MT4. Although numerous types of adverse events have been documented, e.g., injury to the intervertebral disc, ligaments, nerves etc., the cause for most concern is arguably cerebro-vascular events related to stresses on the arterial vessels around the neck; the most common of such events is arguably vertebrobasilar insufficiency (VBI). There is a lack of consensus on the best way to assess for the possibility of adverse events occurring as a result of treatment. Although a number of countries have produced guidelines to assist the clinician in their assessment process5, there is still debate on the utility of such protocols6,7. This paper provides a contemporary, evidence-based overview of relevant information to facilitate clinicians understanding of the effect of cervical MT on blood flow. It is hoped that this information will be used in conjunction with
existing guidelines and knowledge in developing as complete a clinical assessment as possible.

Based on the concepts presented here, we propose a revision of the nomenclature by which we refer to arterial problems of the cervical spine. We suggest the term cervical arterial dysfunction (CAD). We believe that this term embraces both the completeness of the arterial anatomy (i.e., the verteobasilar system, the internal carotid arteries, and the circle of Willis), and the range of pathologies that the manual therapist may encounter (e.g., local dissection, atherosclerotic events, vessel injury, non-ischemic events, ischemic events). The paper is presented in sections meaningful to clinical practice, and each sub-heading relates to questions typically posed by MT clinicians.

What is the Risk of Arterial Complications Following Manual Therapy?

The actual number of reported cases of arterial complications related to manual treatment of the cervical spine is relatively low. Many authors use the number of published case reports to calculate the ratio of incidents per estimated number of treatments, so they can pass judgment on the size of the risk. These calculations range from 1:9122 to 1:5 million (incidents: number of treatments)\(^8,9\). The process is, however, an inaccurate and misleading judgment on the size of the risk. Primarily, there is the argument of under-reporting (i.e., not all cases are reported or published). A recent review suggested a 100% under-reporting rate, based on findings of 32 cases within the UK in one year, none of which had been previously reported\(^10\).

Most structured attempts to calculate risk have been made using questionnaire surveys of clinicians. These have either tried to get the clinician to recall past experiences (retrospective surveys\(^6,8,11\)) or record future experiences over a set period of time (prospective surveys\(^12-16\)). These methodologies are susceptible to a number of flaws, e.g., recall bias, an unwillingness by the clinician to report catastrophic events, an inability (legal or practical) to record such events, data-capture periods that are too short, and a lack of appreciation or clarity as to what counts as an adverse event. These studies are, again, an inaccurate way to establish prevalence.

The paucity of methodologically appropriate studies makes the actual size of the risk impossible to calculate. Figures for prevalence (number of cases in existence) or incidence rate (number of new cases occurring in a year) cannot therefore be stated. All that can be stated is that there is a risk of arterial complications with cervical spine treatment. Large-scale, high-quality, prospective trials are needed to develop understanding of the size of the risk associated with MT treatment. At present, no such studies have been reported. With this in mind, it is necessary for clinical decisions to be made with the information that is available, and to embrace the uncertainty in which this area of practice is held\(^17\).

What is Vertebrobasilar Insufficiency?

Vertebrobasilar insufficiency (VBI) relates to the transient or permanent reduction or cessation of blood supply to the hindbrain through the left and right vertebral arteries (VA) and the basilar artery\(^18\). Manual therapists have been concerned about this concept for many decades because of the intimate relationship between the course of the VAs and the cervical vertebral column. Particular concern has been given to the effect that upper cervical spine rotation has on blood flow in the VAs. Figure 1 demonstrates the distortion that right upper cervical rotation can produce on the left VA. It is this transient stress during movement that is thought to be the mechanism by which VBI is caused—either through a single trauma event or through repeated movements. Further details of the effect of movement on blood flow are given below.

Traditionally, VBI was said to result in transient episodes of hindbrain ischemia manifesting in a number of cardinal signs conventionally referred to as Coman’s 5 Ds\(^19\). However, clinical presentations of transient hindbrain ischemia are more diverse than these classically quoted cardinal signs. Table 1 provides a more complete review of signs and symptoms known to be associated with both local verteobasilar vessel damage and hindbrain ischemia. Of great importance for the manual therapist is the fact that neck and head pain is a commonly reported symptom related to arterial dissection. It is with this in mind that the clinician must be vigilant to the fact that patients with an existing arterial dissection may seek care from a manual therapist.

Is Functional Pre-Treatment Screening Useful?

Functional positional tests are often used to screen for the presence of VBI. Recent Australian Physiotherapy Association guidelines\(^5,14\) reviewed existing guidance of the screening test and suggested that functional testing should consist of sustained cervical rotation, as a minimum requirement. The idea of this, and variations on this theme, is that if blood flow can be altered in a controlled way (by carefully and vigilantly rotating the neck), cardinal signs of insufficiency can be observed. If such signs manifest, then certain treatments are contra-indicated. Current literature provides little support for the validity or reliability of these tests.

Many blood flow studies have demonstrated a change (reduction) in blood flow in the contralateral VA during rotation\(^20,28\). Other studies, however, have found no change in blood flow\(^20,32\). Some authors have used the results of studies demonstrating a reduction in blood flow to support the validity of screening tests; i.e., one tests to assess blood flow
### TABLE 1. Reported signs and symptoms of vertebrobasilar insufficiency (hindbrain ischemia)

<table>
<thead>
<tr>
<th>Condition</th>
<th>Symptoms</th>
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</thead>
<tbody>
<tr>
<td>Local vertebrobasilar vessel damage and hindbrain ischemia</td>
<td>Anhidrosis (lack of facial sweating)</td>
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<tr>
<td></td>
<td>Ataxia</td>
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<td></td>
<td>Clumsiness and agitation</td>
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<td>Diplopia</td>
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<td>Dizziness</td>
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<td>Drop attacks</td>
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<td></td>
<td>Dysarthria</td>
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<td></td>
<td>Dysphagia</td>
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<tr>
<td></td>
<td>Facial numbness</td>
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<td></td>
<td>Hearing disturbances</td>
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<td></td>
<td>Hoarseness</td>
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<tr>
<td></td>
<td>Hypotonia/limb weakness (arm or leg)</td>
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<tr>
<td></td>
<td>Loss of short-term memory</td>
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<tr>
<td></td>
<td>Malaise</td>
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<td>Nausea</td>
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<td>Nystagmus</td>
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<td></td>
<td>Pallor/tremor</td>
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<td></td>
<td>Papillary changes</td>
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<tr>
<td></td>
<td>Perioral dysesthesia</td>
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<td></td>
<td>Photophobia</td>
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<td></td>
<td>Vagueness</td>
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<td></td>
<td>Vomiting</td>
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*Fig. 1: Vertebral (and internal carotid) arteries during upper cervical rotation (Reprinted with the permission of NCMIC Group, Inc. No further reproduction is allowed without the express permission of NCMIC.)*
changes, these studies demonstrate that rotation changes blood flow; therefore, the test is valid. The tests may be valid in that they may alter blood flow, but there is little consistent evidence relating these changes to changes in symptoms; e.g., a patient could have significant reduction in blood flow but no symptoms and vice versa. This makes the specificity and sensitivity of these tests poor and variable, and this has been mathematically demonstrated in probability calculations. On the basis of the inconsistency of the evidence, a recent paper has raised the issue of whether manual therapists should stop using functional pre-screening tests. The opinion that the use of functional pre-screening testing cannot be supported has been highlighted again in the evidence reviewed for this present project.

There is no evidence to support the construct validity of functional pre-screening testing in terms of its ability to identify patients who are more likely to have spontaneous dissection events. This test is, of course, specific to the vertebrobasilar system and unlikely to be sensitive to ICA blood flow changes.

Internal Carotid Artery (ICA) Dysfunction

Complications of MT treatment related to the ICAs have been reported. Consideration of the reported cases alone will not lead to a full understanding of the complete nature and details of ICA pathologies. However, there is a significant amount of information available that manual therapists can exploit to understand problems associated with the ICA. This is summarized below so that the clinician is able to make informed judgments on the differential diagnosis of presentations that may be associated with ICA dysfunction. Figure 2 shows the anatomical course of the ICA.

The ICA supplies the brain and the retina. The natural onset and progress of ICA dissection begins with local arterial trauma (the dissection event itself). This dissection event can manifest in a number of signs and symptoms that are non-ischemic (i.e., somatic pain related to local injury). These local signs and symptoms can precede cerebral ischemia (TIA or stroke) or retinal ischemia by anything from less than a week to beyond 30 days. There is, therefore, a

Fig. 2: Course of the vertebral and internal carotid arteries through the cervical spine. (Adapted with permission from, Drake et al, Gray’s Anatomy for Students, www.studentconsult.com, Elsevier Ltd.)
period of time when a patient with ICA dissection may present to the manual therapist with signs and symptoms that may mimic a neuromusculoskeletal presentation. Table 2 details classic ICA non-ischemic and ischemic manifestations of ICA dissection.

It is important to appreciate that similar to a VA dysfunction, headache and/or cervical pain can be the sole presentation of ICA dysfunction—particularly in the early stages of the pathology. Cranial nerve palsies and Horner's syndrome are often pathognomonic of ICA pathology, especially if the onset is acute. The hypoglossal nerve (CN XII) is the most commonly affected followed by the glossopharyngeal (CN IX), vagus (CN X), or accessory (CN XI). However, all cranial nerves (except the olfactory nerve) can be affected. Horner's syndrome—characterized by drooping eyelid (ptosis), sunken eye (enophthalmia), a small, constricted pupil (miosis), and facial dryness (anhidrosis)—has been found to be present in up to 82% of patients with known ICA dissection. Most commonly, this syndrome occurs with head, neck, or facial pain.

Can We Identify “At-Risk” Patients?

Cervical arterial dysfunction (CAD) refers to a wide variety of pathophysiological events affecting both the vertebrabasilar and the carotid vessels. At one extreme are actual cerebrovascular accidents (CVA) (strokes). In the middle of the continuum are the much more subtle dysfunctions relating to transient interruption of perfusion to particular sites in the head, while at the other end of the continuum is the consideration of the patient's likelihood to be at risk from a future cervico-cranial ischemic event and the need to calculate the chances of physiotherapy intervention contributing to such an event. It is essential that clinicians consider the full scope of this continuum. At-risk patients may, therefore, be either those presenting with frank arterial pathology or those with no clinical pathology but with a high number of vascular risk factors. A multi-factorial approach must therefore be taken in the assessment of these patients, and reliance on single, physical tests with known poor utility represents an incomplete and misleading examination.

History-taking may assist in establishing which patients are at risk of CAD and, therefore, at an increased risk from cervical MT. The latest Australian Physiotherapy Association guidelines emphasize this point. It has been suggested that, for VBI at least, there are no identifiable predictors of adverse outcome and the phenomenon is an unpredictable, idiosyncratic event whose pathology is possibly inherent. Despite this single study, there is a wealth of information outside the MT literature that may help us to begin to identify patients who are more at risk of CAD than others, e.g., those with the symptomology of pathologies as detailed above. More specifically, this information can help us differentiate those people presenting with pain (with or without other signs and symptoms) of a vascular origin.

With the exception of truly unidentifiable genetic vascular dysfunction (e.g. hypoplasia, fibrodysplasia), which may account for rare random spontaneous dissection events, the basis for almost all other cervical vascular conditions is pathology of the endothelium, specifically atherosclerosis.

Atherosclerosis is intrinsically related to endothelial dysfunction. It is a cascade of inflammatory events influenced by conventional inflammatory mediators and infectious micro-organisms. These events are triggered by endothelial changes induced by chemical, mechanical, or immunological insult. Evidence-based clinical athero-
sclerotic risk factors include hypertension; diabetes; family history of atherosclerotic-related pathologies (e.g. heart disease, stroke, transient ischemic attack, peripheral vascular disease); smoking, high serum low density lipoprotein (hypercholesterolemia/hyperlipidemia/high cholesterol-high fat diet); hyperhomocysteinemia; infection by escherichia coli, helicobacter pylori, chlamydia pneu-
moniae, streptococcus, staphylococcus, salmonella, clostrid-
ium, mycobacterium, fungi, yersenia, treponema; and me-
chamical trauma to vessel46-55.

The logic for considering this in cervical MT is simple and is based on the following premises:

1) Atherosclerosis is strongly linked with localized ICA56-59
and VA60-64 dysfunction, and ultimately stroke as a result of this localized dysfunction.
2) Cervical movement affects blood flow in these vessels (see above).
3) Cervical trauma is associated with the aetiology and pro-
gression of CAD65-69.
4) Neck and head pain is a common early manifestation of
dysfunctional cervical arteries (see above).
5) Manual therapists see patients with neck and head pain
and histories of cervical trauma.
6) Manual therapists have the ability to produce further
cervical trauma through various treatment interven-
tions.

This logic provides the basis for our suggestion that, de-
spite comments to the contrary44,20, consideration of ather-
sclerotic risk factors when profiling for high-risk patients, or
when differentially diagnosing pain syndromes, should re-
main a concern of the manual therapist. We propose that
until there is substantial evidence to refute this suggestion,
the overwhelming extent of pathophysiological and epide-
miological knowledge regarding cervical arterial atheroscle-
rosis would make it professionally irresponsible for clinicians
to ignore this dimension of CAD.

It is beyond the scope of this paper to detail the techni-
calities of assessment for atherosclerotic profiling. However,
consideration of the above risk factors in the history taking,
and the use of blood pressure testing, for example, are im-
portant components to be integrated into MT practice71,72.

What is the Relative Risk of Manipulating
the Upper Cervical Spine Versus the Lower
Cervical Spine?

There are no studies in the current literature that focus di-
rectly on this question. Traditionally, the upper cervical
spine is believed to carry a greater risk because of the tortu-
ous course of the VA between C2 and the occiput. Most blood
flow studies have concentrated on the C1/C2 and cervical
anatomical area25,29,30,32,73-75. Some flow studies have insonated
the vessel as it enters the lower part of the vertebral column
(the pre-transverse level)22,76-80. Few studies have measured
blood flow in the intracranial VA21,23,24,28 where it may be ar-
gued that VA blood flow related to the upper cervical spine
would be a more accurate measure of flow change24,30,32,73,74.

Is Physiotherapy Manual Therapy as Risky
as Manipulative Treatment used by other
Groups (e.g., Chiropractic)?

Most MT case reports and surveys involve chiropractic care
and refer to manipulative thrust techniques of various forms.
This, together with the fact that chiropractors arguably carry
out a far greater number of manipulations than osteopaths
or physiotherapists, leads many authors to conclude that
chiropractic treatment carries the greatest risk. Although
there is some logic in this conclusion, it does not support the
idea that less forceful, or non-chiropractic treatment, is nec-
essarily safer. Blood flow studies have demonstrated signifi-
cant flow changes during gentle passive positioning and not
fast, manipulative procedures of the cervical spine.

One cause of hindbrain ischemic events is believed to be
dissection of an atherosclerotic thrombus. Although a
quicker movement (manipulative thrust) is more likely to
dissect a thrombus, it is also feasible, based on the results of
flow studies, that gentler, repeated movements offer a poten-
tial dissection-inducing force. Due to the progressive nature
of atherosclerotic pathology, ischemic events (e.g., emboli-
ization following thrombus formation and thrombus dissec-
tion) may occur some time after treatment18,39,42. This latency
concept will affect the true incidence rate in all MT groups.

In summary, although there are more chiropractic-re-
lated cases reported, this evidence is arguably disproporti-
ionate. Non-manipulative, MT treatment, and other forms of
hands-off physiotherapy have not been subject to the same
degree of study and so no judgment can be made regarding
this risk.
What is the Medico-Legal Situation Regarding Treatment and the Role of Guidelines?

In English law, the mere fact that a procedure is unsuccessful or goes wrong does not necessarily mean that the duty of care a health care professional owes to the patient has been breached. The court informs itself regarding whether there has been a failure to provide the required standard of care by hearing expert testimony from the profession concerned. If it can be shown that what the defendant did was in accord with a practice currently accepted as proper within his or her profession at the time of the incident, it is unlikely the professional will be held to have acted negligently even if there is another currently accepted but contrary practice\(^{62}\). Expert opinions may, however, be scrutinized under cross-examination and the court may disregard a practice, despite it being currently accepted, if it considered it to be unreasonable, irresponsible, or logically indefensible. In particular it needs to be satisfied that in forming their views, witnesses “have directed their minds to the question of comparative risks and benefits”\(^{63}\). If the defendant cannot show that he or she has followed an accepted practice, it falls to him or her to justify what he or she did. The more serious the harm suffered, the more robust that defense will have to be.

Despite the fact that a primary aim of guidelines is to ensure that all the right things and none of the wrong things are done when a patient presents with a particular clinical problem\(^{84}\), a guideline has no particular status or automatic effect in English law, and if reliance is to be placed on it, its existence and content must be brought to the court’s notice as part of the expert testimony mentioned above\(^{85}\). Nor does the mere existence of a guideline guarantee that the quoted aim has been met since a lack of consistency in the quality of evidential bases and methodology can render its authority and clinical trustworthiness suspect\(^{86}\). The worth of a particular guideline remains a matter for individual clinical judgment as does the decision to follow or depart from it in practice.

Summary

Serious adverse neurovascular responses are a rare but as yet unquantified consequence of cervical MT. Although this number of reported cases of serious adverse effects is low, a number of factors prevent using these cases to estimate the prevalence or incidence of this phenomenon, e.g., latency of pathology, lack of clarity regarding presentations, ambiguity regarding what is an adverse neurovascular event, etc. Surveys of MT practitioners are an inaccurate method of attempting to establish size of risk. The true size of these adverse events cannot be estimated at this time.

Traditional clinical practice has focused on pre-manipulative screening for the presence or absence of vertebrobasilar insufficiency. Despite concerns about the utility of VBI screening tests, they are still relied upon in clinical decision-making. We have proposed that the clinical approach should be broadened to incorporate not only VBI but also pathologies affecting the ICAs. Furthermore, we suggest that this clinical approach embraces differential diagnosis of known arterial pathologies that commonly present with neck and/or head pain. The underlying pathologies affecting all parts of the cervical arterial system are varied, but most known cervical arterial complications are atherosclerotic in nature. The exception to this is rare spontaneous dissection. With this in mind, we suggest that consideration of atherosclerotic risk factors may enhance decision-making with regards to both pre-treatment screening and differential diagnosis of frank arterial pathology.

The evidence surrounding cervical arterial dysfunction and MT is varied and inconclusive. As clinicians responsible for the health of our patients, we must strive to use the best information available. This information must be incorporated into our own clinical reasoning processes. In the absence of strong evidence-based guidelines, we must embrace the uncertainty of this clinical area and use our reasoning to do the best for our patients. First, we must do no harm.

Key Messages

- Traditional cardinal signs and symptoms of VBI following MT are not supported by the literature.
- The real risk of arterial complications following MT is unknown and impossible to estimate, based on existing data.
- The results of blood flow studies are contradictory and inconclusive. Commonly used functional screening tests are not supported by the data available from these studies, nor from case reports.
- Consideration of haemodynamics related to the cervical region may enhance clinicians’ understanding of risks and mechanisms of vascular events.
- Adherence to guidelines might not necessarily be a defense, and expert opinion can be over-ruled in law.

Recommendations

- We propose the term **cervical arterial dysfunction (CAD)** to embrace the whole cervical arterial system and the range of pathologies that affect this system.
- Consideration of atherosclerotic risk factors may assist in identifying at-risk patients and patients presenting with
frank arterial pathology. We recommend the use of specific questioning and testing—specifically blood pressure measurement—to facilitate risk-factor profiling.

- Knowledge of known arterial presentations will strengthen the clinician’s differential diagnosis skills. We recommend the incorporation of this knowledge, and testing procedures such as cranial nerve examination, to facilitate differential diagnosis.

REFERENCES

Manual Therapy and Cervical Arterial Dysfunction, Directions for the Future: A Clinical Perspective

This paper is an excellent review article pertaining to the arterial system and manual therapy of the cervical spine (CMT). Clinicians are reminded to consider the cervical arterial system as a whole and, through a careful history and neurological assessment, be alert to signs and symptoms of central nervous system pathology. Also, the process of clinical decision making during CMT is not based on the results of any single test, but on the clinical picture created during the history and examination. The clinician should look for signs and symptoms that contraindicate manual therapy and those that do not fit the model of a musculoskeletal disorder. The authors also raise the familiar concern of manipulative therapists; the presence of risk of vascular injury with cervical manipulation or high velocity techniques, and the difficulty quantifying the risk.

However, this paper raises some clinical questions. Bear in mind that the vast majority of patients presenting for treatment, in the absence of contraindications found during evaluation, will have mechanical neck pain with or without headache, conditions that are perfectly amenable to manual therapy. What level of treatment caution must we exercise? Should we not mobilize or apply mechanical traction to any patient with risk factors for atherosclerosis but no contraindications found on examination? Should we apply the same level of caution about oscillatory mobilizations that we might to high velocity techniques? Clinicians may be prudent to refrain from high velocity techniques in patients with obvious risk factors for atherosclerosis. Nevertheless, the bigger dilemma remains: arterial intimal tears and death have occurred in relatively young people with minimal atherosclerotic change. To date, there is no clinical test to indicate the presence of an artery susceptible to tear.

I also question the clinical value of routine blood pressure screening in addition to the customary history and examination prior to CMT. If this is performed, how is the clinician to interpret the result and what action should be taken? Do we withhold manual treatment of any sort from a patient with elevated blood pressure? A patient may have an elevated reading for other reasons (e.g. pain). More guidance and evidence for inclusion in our manual therapy practice would have been beneficial.