The Role of MR Imaging in Assessment of Brain Damage from Carbon Monoxide Poisoning: A Review of the Literature

T. Beppu

ABSTRACT

SUMMARY: The aim of this article is to review how MR imaging and associated imaging modalities provide clinicopathologic information on brain damage after carbon monoxide poisoning. Initially, many authors documented typical findings of conventional MR imaging in the gray matter structures such as the globus pallidus and in various regions of cerebral white matter. The focus of investigation has since shifted to observation of cerebral white matter areas that are more frequently detected on MR imaging and are more responsible for chronic symptoms than the gray matter. DWI has dramatically contributed to the ability to quantitatively assess cerebral white matter damage. Subsequently, DTI has enabled more sensitive evaluation than DWI and can demonstrate progressive pathologic changes in the early stage, allowing prediction of chronic conditions. In addition, MR spectroscopy reveals changes in metabolite levels, offering quantitative clinicopathologic information on brain damage after carbon monoxide poisoning.

ABBREVIATIONS: CO = carbon monoxide; CWM = cerebral white matter; DNS = delayed neuropsychiatric sequelae; FA = fractional anisotropy; GP = globus pallidus; MBP = myelin basic protein

C O poisoning causes serious damage to the brain and cardiac muscles. Damage to the brain directly affects the prognosis for patients who survive CO poisoning. Some clinical obstacles exist to the assessment of brain damage among such patients. First, predicting clinical behaviors such as specific types of chronic neuropsychiatric symptoms is difficult. Surviving patients will display 1 of 3 clinical behavioral types in the chronic phase after CO poisoning: approximately 70% of survivors present with various transient symptoms only in the acute phase; 20% of patients present with symptoms persisting from the acute to the chronic phase; and the remaining 10% exhibit DNS, representing recurrent neuropsychiatric symptoms occurring after an interval of apparent normality (the so-called lucid interval; mean duration, 22 days) after the apparent resolution of acute symptoms.1-5 Acute conditions including level of consciousness and carboxyhemoglobin concentration have been considered and found to be unhelpful in the prediction of chronic clinical behaviors.6-7 In particular, predicting whether patients who exhibit resolved acute symptoms have escaped or will experience DNS represents a very important clinical issue.8 A second problem is the difficulty of assessing the severity of brain damage. Chronic symptoms include parkinsonism, dystonia or other motor impairments, cognitive or executive dysfunction, akinesia, mood disorder, and personality change. The severity of brain damage cannot be directly compared among patients with different symptoms. One of the strategies to resolve these clinical problems is to apply objective assessments of brain damage by using MR imaging, a readily available and minimally invasive procedure. Here, I review the literature on assessment of brain damage secondary to CO poisoning by use of conventional MR imaging in addition to other sequences, such as DWI, DTI, and MR spectroscopy, and summarize what MR imaging can currently accomplish in the assessment of brain damage after CO poisoning.

MECHANISMS CAUSING BRAIN DAMAGE AFTER CO EXPOSURE

Comprehension of the mechanisms causing brain damage after CO exposure is crucial to understanding the findings of MR imaging in the assessment of brain damage. Brain damage has been considered to result from complicated mechanisms, as follows.4 The uptake of CO by hemoglobin with an affinity 200 times that of oxygen leads to hypoxia in tissues, and a shift in the oxyhemoglobin dissociation curve to the left inhibits release of oxygen, further exacerbating tissue hypoxia.9 Whereas hypotension adds to hypoxia, the hypoxia-hypotension process leads to ischemic changes in the arterial border zones in the brain.10 Brain hypoxia increases the level of excitatory amino acids including glutamine, leading to...
A elevated nitrite levels and causing subsequent injuries to the cerebral cortex.\textsuperscript{11} Furthermore, hypoxia in the brain causes oxidative stress, cellular necrosis, apoptosis, and ongoing inflammation.\textsuperscript{12} Although it is important to emphasize the damage to the brain that results from hypoxia-related effects, brain damage can also be caused by mechanisms other than hypoxia. CO also combines with cytochrome A and A3 to cause direct poisoning effects, and decreases in levels of cytochrome C oxidase inhibit mitochondrial metabolism, leading to cellular respiratory dysfunction.\textsuperscript{13} Furthermore, CO-mediated brain damage associated with free radicals has been noticed. CO binds to platelet heme proteins, causing the release of nitric oxide. Excess nitric oxide produces peroxynitrite, impairing mitochondrial function.\textsuperscript{14} As another example, CO causes platelet-to-neutrophil aggregation, and neutrophil degranulation leads to release or production of myeloperoxidase, proteases, and reactive oxygen species, which cause oxidative stress, lipid peroxidation, and apoptosis.\textsuperscript{14,15} Lipid peroxidation alters the structure of MBP, triggering lymphocytic immunologic responses and increasing the activation of microglia. Immunologic responses to altered MBP cause progressive demyelination accompanied with ongoing inflammation in the CWM.\textsuperscript{4}

CONVENTIONAL MR IMAGING

As mentioned in the previous section, pathologic changes induced by complicated mechanisms progress in the brain from the time of CO inhalation to even some years later. The timing of MR imaging will thus have a major effect on the results of imaging.\textsuperscript{16} In this review, times between CO inhalation and MR imaging were strictly defined as the ultra-acute phase within 24 hours; the acute phase between 24 hours and 7 days; the subacute phase between 8 and 21 days; and the chronic phase from 22 days and thereafter.

Since the end of the 1980s, findings of damaged brain from CO poisoning by using conventional MR imaging instead of CT have been reported. In those days, most articles were case reports that described findings in the chronic phase and documented typical findings of bilateral hyperintensities in the GP and CWM on T2-weighted images.\textsuperscript{17,18} In these reports, the authors speculated that the main pathologic changes depicted on MR imaging were necrosis in the GP and demyelination in the CWM. These suppositions were based on a report of autopsy findings by Lapresle and Fardeau,\textsuperscript{19} which identified 1) necrosis in the GP, 2) demyelination in the CWM, 3) spondyloitic changes in the cerebral cortex, and 4) necrosis in the hippocampus. Damage in the GP is frequently seen in patients poisoned with CO.\textsuperscript{20} In contrast, damage in other basal ganglia such as the caudate nucleus, putamen, and thalamus has not been reported as frequently as that in the GP.\textsuperscript{21,22} Authors have generally adopted 1 of 2 reasons for selective damage to the GP: that this region is easily affected by the hypoxic-hypotension process because of a poor anastomotic blood supply,\textsuperscript{18,21,23} or that CO binds directly to heme iron in the GP, which happens to be a brain region with the highest iron content.\textsuperscript{24-26} Furthermore, some reports have described a “pallidoreticular pattern” of damage to the substantia nigra in addition to the GP, another region with high iron content.\textsuperscript{26,27} The GP often demonstrates different findings of hemorrhagic infarction, which match the phase in which MR imaging is performed.\textsuperscript{23,28-30} Because the main pathologic features are edema in the acute phase and necrosis in the chronic phase, the size of the GP lesion may shrink on consecutive evaluations.\textsuperscript{31} Parkinsonism is the most commonly seen symptom in patients with CO exposure.\textsuperscript{1,32} The possibility that damage to the GP relating to the extrapyramidal tract causes parkinsonism is readily apparent, but parkinsonism can also occur in patients without damage to the GP. Pavese et al\textsuperscript{32} hypothesized that lesions in the CWM containing tracts outputting and/or inputting to the basal ganglia might cause parkinsonism in patients without GP damage.

Regarding gray matter structures other than the GP, many previous reports have described damage to the hippocampus (Fig 1A, -B). Almost all of these reports documented findings of hippocampal atrophy in the chronic phase (2 months to 21 years) and patients experiencing severe persistent symptoms.\textsuperscript{22,33-35} On the contrary, a few reports have mentioned damage to the cerebral cortex.\textsuperscript{30,32,36} These studies documented cortical atrophy in patients with persistent symptoms in the chronic phase. Such damage to both the hippocampus and the cerebral cortex might create lesions responsible for persistent symptoms in the chronic phase. In a report regarding hippocampal lesions in the acute phase, 2 of 4 patients had died within a short period.\textsuperscript{28} Hippocampal lesions in the acute phase may be predictive of a very poor prognosis.

Gray matter could be predicted to be more vulnerable to hypoxia from poisoning than WM, as neurons in gray matter show high activity and a high blood demand. However, neurons may be able to tolerate pure hypoxia for longer than they can tolerate ischemia.\textsuperscript{37} Damage to gray matter should be caused by ischemic changes through the hypoxic-hypotension process. Some authors have pointed out that CWM damage is more frequently detected on MR imaging than damage in the gray matter. A prospective study in which 73 patients with CO poisoning were consecutively enrolled without any selection reported that 12% of these patients showed CWM damage, whereas damage to the basal ganglia was found in only 1 patient.\textsuperscript{3} CWM has been recognized to be more responsible for chronic symptoms than gray matter. The most common regions of CWM to be affected are the centrum semiovale and periventricular WM.\textsuperscript{31,38} On T2-weighted images, both lesions are depicted as hyperintense in the bilateral CWM either symmetrically\textsuperscript{21,39} or asymmetrically.\textsuperscript{36,40}
Damaged CWM may occasionally show hemosiderin deposit foci suggesting hemorrhagic infarctions or iron extravasation after CO poisoning. CWM damage can be observed in various regions other than the centrum semiovale and periventricular WM, such as the temporal lobe, occipital lobe, parietal lobe, and corpus callosum. However, few reports have provided detailed information regarding damage to the cerebellum and brain stem. This may result from a high tolerance of the posterior structures for hypoxia, and such damage would thus appear under severely hypotensive conditions in addition to hypoxia. Indeed, patients demonstrating cerebellar damage in the acute or subacute phase uniformly show a poor prognosis. Prognosis parallels the amount of damage to WM such as lesions involving the cerebellum. Consequently, many reports have indicated that the amount of WM damage closely relates to prognosis. Specific regions in the CWM corresponding to each symptom have also been investigated. Damage to the frontal lobe has been considered to be associated with apathy, loss of motivation, and mutism. Some prospective studies have examined the relationship between MR imaging findings and results of neurocognitive tests in the chronic phase after CO poisoning. Kesler et al performed consecutive MR imaging in 69 patients and reported that atrophy of the fornix was already apparent at 2 weeks after poisoning, and was associated with reduced verbal memory at 6 months. Parkinson et al performed serial MR imaging and cognitive tests on the day of onset, and 2 weeks and 6 months after poisoning in 72 patients, revealing that damage to the centrum semiovale was closely associated with cognitive dysfunction in all phases. Porter et al observed atrophy of the corpus callosum at 6 months after poisoning in 62 patients, but cognitive dysfunction in that study appeared independently of corpus callosum atrophy. As mentioned above, however, the brain is damaged more widely in patients with CO poisoning who have more severe and varied symptoms. Elucidating the specific regions responsible for each asymmetric hyperintensity has been speculated to be caused by differences in presumably original arterial blood supply among patients. Damaged CWM may occasionally show hemosiderin deposit foci suggesting hemorrhagic infarctions or iron extravasation after CO poisoning. CWM damage can be observed in various regions other than the centrum semiovale and periventricular WM, such as the temporal lobe, occipital lobe, parietal lobe, and corpus callosum. However, few reports have provided detailed information regarding damage to the cerebellum and brain stem. This may result from a high tolerance of the posterior structures for hypoxia, and such damage would thus appear under severely hypotensive conditions in addition to hypoxia. 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**FIG 2.** Hyperintensities in the deep WM involving the centrum semiovale in a 53-year-old man with DNS after CO poisoning. A, T2-weighted imaging within 24 hours after CO inhalation reveals slight hyperintensity in the bilateral deep WM and shows a necrotic focus in the left parietal lobe. B, At 2 days after the occurrence of DNS, hyperintense areas appear widespread in the bilateral deep WM. Brain atrophy is also apparent in the bilateral frontal lobes.
phenomena might indicate demyelination of myelinated CWM.49,54-56 Subacute or chronic phases after the appearance of DNS.49,54-56 Fusion because of demyelination, when DTI was performed in the subacute phase, did not demonstrate clear abnormalities in the basal ganglia within 24 hours after CO poisoning, but markedly high signals were depicted in the bilateral GP on DWI (C) at the same time. Clear hyperintense areas in the bilateral GP were found on T2-weighted imaging after 3 months (D).

Low FA values might continue until 3 months after poisoning,55 suggesting a reduction in the directionality of water molecule diffusion, and FA as its quantitative value have been recognized to occur when the extent of demyelination progresses beyond a certain threshold, use of DTI to predict the occurrence of DNS has been suggested. At our institute, FA at the centrum semiovale has been measured at 2 weeks after poisoning in consecutive patients. As a result, we observed that FA in the centrum semiovale had already decreased before the appearance of DNS, and reported the possibility that FA in the subacute phase could offer a predictor for the occurrence of DNS.8 The concentration of MBP in CSF, as a marker of demyelinating disease activity, has been proposed as a predictor of DNS.57 Another of our previous studies demonstrated that FA in the centrum semiovale roughly correlates with MBP concentration in the subacute phase (approximately 2 weeks) in patients with CO poisoning who present with chronic persistent symptoms or DNS.58 These reports suggest that FA in DTI must be able to sensitively and quantitatively indicate the extent of demyelination after CO poisoning and, in particular, could demonstrate progressive damage leading to DNS.

FIG 3. High-sensitivity DWI in the acute phase after CO poisoning in a 64-year-old woman. A, T2-weighted imaging; B, FLAIR; and C, DWI, all taken within 24 hours after CO inhalation. D, T2-weighted imaging at 3 months after CO poisoning. T2-weighted imaging (A) and FLAIR (B) did not demonstrate clear abnormalities in the basal ganglia within 24 hours after CO poisoning, but markedly high signals were depicted in the bilateral GP on DWI (C) at the same time. Clear hyperintense areas in the bilateral GP were found on T2-weighted imaging after 3 months (D).

Different areas where the region of interest is placed on the CWM might influence the FA values. Lo et al55 reported FA values measured by using regions of interest placed on the centrum semiovale and explained why they placed regions of interest on these areas as follows. All patients in their study showed abnormalities in the deep WM involving the centrum semiovale on T2-weighted imaging, and areas involving the centrum semiovale were relatively homogeneous WM zones related to worse cognitive performance. Selection of these areas was sensible because some reports have shown a relationship between damage in the centrum semiovale and cognitive disorder after CO poisoning.48,59 Our previous study in which we used DTI with voxel-based analysis also showed that deep WM areas including the centrum semiovale were the most widely damaged regions in those patients presenting with chronic symptoms (Fig 4).60 When the CO-damaged brain is assessed by DWI or DTI, it is important to emphasize that evaluations of deep WM, including the centrum semiovale, are indispensable.

MR SPECTROSCOPY

MR spectroscopy enables noninvasive monitoring of changes in metabolism from brain damage in a specified region. Although most previous studies regarding findings of MR spectroscopy in patients with CO poisoning were only case reports and differences existed in the regions used for placement of the voxel of interest in the brain, these reports have consistently shown increased levels...
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months after CO poisoning in patients with chronic symp-
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liest in metabolites and decreased in the late period of the chronic
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**CURRENT AND FUTURE**

So far, most investigations by use of MR imaging and its optional sequences have been rigorously focused on either the GP or CWM as the sites of typical lesions in the chronic phase after CO poisoning. In particular, findings of MR imaging and other sequences on CWM correlating with chronic symptoms allow us to monitor the condition of the patient in the chronic phase. However, brain damage after CO poisoning should be attributed not only to injury of the CWM, but also to injury of gray matter structures such as the hippocampus or cerebral cortex. In the chronic phase, investigations are needed regarding relationships between damage to both gray and WM and the severity of symptoms. On the contrary, few reports have described investigations of MR imaging in gray matter structures and CWM in the acute or subacute phase. DTI and MR spectroscopy in the subacute phase have already revealed slight pathologic changes from progressive de-
myelination starting immediately after CO inhalation. Interest will eventually migrate to whether findings of MR imaging in the acute phase offer a predictor of the occurrence, severity, and type of chronic symptoms, because DNS can occur as early as within days after CO inhalation. In our experience of using 3T MR imaging, however, progressively pathologic changes in the acute phase must be too small to observe on conventional MR imaging and even on DWI, DTI, or MR spectroscopy in most patients. In the future, new approaches to the use of MR imaging, such as new sequences or more sensitive imaging such as 7T MR imaging, may reveal and resolve these issues.

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**FIG 4.** Detection of damaged regions of the CWM by use of voxel-based analysis with DTI. When FA values in all voxels on the FA map were compared voxel by voxel between 2 patient groups (patients with and without chronic neuropsychiatric symptoms), voxels showing a significant difference in FA (P < .03) between groups were identified as reddish voxels on the FA template. The number of reddish voxels was greater in the region corresponding to the deep CWM, including the centrum semiovale, than in other regions.
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