Carbon monoxide poisoning: easy to treat but difficult to recognise

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Carbon monoxide poisoning (CO) is produced from incomplete combustion of fossil fuel, cars being the most important environmental source.\(^1\)\(^2\) Indoor air pollution with CO may result from faulty kerosene or gas heaters used in unventilated areas,\(^3\) gas water heaters, or fireplaces with blocked flues.\(^4\) More unusual causes include barbecue cubes burnt indoors,\(^5\) small petrol electricity generators without proper emission disposal, or the use of forklift trucks in warehouses.\(^6\)\(^7\)

Cigarette smoke contains small amounts of CO, and it has been shown that smokers have higher baseline carboxyhaemoglobin (CO-Hb) levels than nonsmokers.\(^8\)\(^9\)

CO has a much higher affinity for the haemoglobin molecule than oxygen, thus replacing it and making it unavailable for oxygen transport.\(^10\) The oxygen dissociation curve is shifted to the left, making unloading of oxygen at the tissue level problematic. Furthermore, myoglobin in muscle and cytochromes in cells are partially inactivated.\(^11\)\(^12\) These effects result in generalised tissue hypoxia, the more susceptible tissues of the brain\(^13\) and myocardium\(^14\)\(^15\) being most commonly affected. CO also appears to cause brain lipid peroxidation.\(^16\)

Clinical presentation

The clinical features of CO intoxication are very nonspecific and mimic common conditions causing, unconsciousness, epileptic fits,\(^16\)\(^17\) headache,\(^16\)\(^19\) flu-like illness,\(^20\) and unstable angina.\(^21\) It is important to recognise CO poisoning for two reasons. Firstly, patients presenting with coma or fits, or impairment of consciousness due to serious intoxication, if not treated promptly with hyperbaric oxygen or 100% oxygen therapy, may either not survive, or else survive to develop long-term neurologic and psychiatric problems.\(^22\) Secondly, recognition of the patient with less serious intoxication, presenting with minor complaints such as headache or flu-like illness, is important mainly to identify the source of the CO and thus prevent subsequent, potentially catastrophic, exposures in the household.\(^5\)\(^7\)\(^16\)\(^25\)\(^24\)

A number of studies screening emergency room populations have been published, suggesting a rational approach to the recognition of intoxication. Serious intoxication, on which hyperbaric oxygen is likely to have a clear therapeutic impact, presents with acute neurological features.\(^23\) Heckerling \textit{et al} screened 168 consecutive acute neurological admissions and found one serious intoxication which possibly needed hyperbaric oxygen, and four minor intoxications, two of which were identified out of 43 admissions for epileptic fits.\(^26\)

At our hospital (St Luke's, Malta) we screened 307 acute neurological admissions from December 1994 to April 1995. Out of 29 patients admitted with impaired consciousness and no lateralisering neurological signs, three had serious intoxication, all of whom had bilateral upgoing plantars. (Heckerling\(^26\) had observed two cases of intoxication out of 81 patients screened in this diagnostic category.) However, out of 141 patients with presumed stroke or transient ischaemic attack, none had evidence of intoxication. Furthermore, out of 87 patients admitted with episodes of loss of consciousness who had recovered completely in the emergency room, none had evidence of intoxication.

In another study, Heckerling \textit{et al} \(^27\) screened 753 acute surgical, medical, neurologic, and psychiatric admissions. Only two minor cases of intoxication were identified. None of the patients with stroke had CO intoxication. However, out of 20 patients admitted with epileptic fits, one had evidence of acute exposure, a similar proportion to that found in the study mentioned above.\(^26\) In our study, none of the 39 patients with epileptic fits had evidence of CO exposure.

In a third study, Heckerling showed that 3-5% of patients presenting to an emergency department with headache,\(^18\) had occult CO exposure. The author
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**Difficulties in the diagnosis of CO poisoning**

- gas is colourless, odourless and non-irritant
- patient may be unaware of the exposure
- ambulance staff, co-habitants unaware of exposure
- patient with impaired consciousness unable to give history
- no pathognomonic feature of the intoxication (textbook cherry red colour, a rare finding)
- mimics common conditions such as unstable angina, drug overdosage, status epilepticus
- patients with underlying cardiovascular and cerebrovascular disease can have an atypical presentation

**Tips for the clinical diagnosis of CO poisoning**

**Why is it important to diagnose?**

- to identify those patients likely to need 100% or hyperbaric oxygen treatment so as to prevent long-term neuropsychiatric sequelae
- to recognise a source of CO intoxication before fatal or life-threatening exposures can result

**When to suspect it?**

- impaired consciousness or coma with no focal or lateralising neurological signs
- epileptic fits
- headaches, or flu-like illness
- unstable angina

**What to ask about?**

- accident sites likely to result in exposure (garage, warehouse, bathroom, car, fire)
- multiple simultaneous complaints in co-habitants
- type of indoor heating used
- take CO-Hb levels when the patient is unconscious and no history is available

**Pitfalls**

- chronic metabolic conditions, eg, diabetes, uremia, opiate addiction, do not exclude CO intoxication
- patients being discharged from the emergency room for minor complaints, rather than being admitted to hospital, could also have CO intoxication

also validated a prediction model showing that a good history on indoor heating and the presence of symptomatic co-habitants correctly identified affected individuals. During the winter months, Dolan et al screened 55 inner-city inhabitants who presented with flu-like symptoms, ie, headache, nausea, and malaise. Thirteen had evidence of exposure to CO, although none needed hyperbaric oxygen. In our opinion, Heckerling's suggestion to screen by history is preferable to routine assessment of CO-Hb levels, as the expense of screening every individual with a common cold would be prohibitive. However, Dolan's data suggest that it is more likely that exposed patients who present to the emergency department will be discharged than admitted, so that a high index of suspicion must not be restricted to patients being admitted to hospital.

In a study screening 103 patients routinely admitted for coronary care with unstable angina, we found three cases of serious intoxication, and five of minor exposure. Although most emergency room physicians aim for quick, efficient, admissions, we suggest that information on the type of indoor heating used should be obtained in all cases so that treatment with 100% oxygen can be implemented when necessary.

When assessing patients presenting with impairment of consciousness, a common pitfall leading to a missed diagnosis of CO poisoning is the presence of underlying chronic cerebrovascular or metabolic problems. Not only may the clinical picture be atypical and occur at CO-Hb levels of 10-20%, but a readily available alternative explanation stops one from considering CO intoxication. For example, among our acute neurologic emergencies, was a frail 75-year-old woman with diabetic neuropathy and nephropathy who presented with unconsciousness. The preliminary diagnosis was hypo-, or hyperglycaemia and uraemia. However, screening showed she had CO intoxication which was later shown to be due to a faulty kerosene heater. In our opinion, information on indoor heating must be obtained from members of the household in all patients admitted to hospital with impaired consciousness and no focal neurological signs. If no information is available, CO-Hb levels should be obtained.

**Diagnosis and treatment**

Whenever CO intoxication is suspected CO-Hb levels should be determined using spectrophotometric analysis in a CO oximeter. This test is easy to perform and should be available to all emergency room physicians. The result must be delivered as soon as possible to allow rational planning of therapy. When a CO oximeter is not available a hand-held CO expired air detector can be used to confirm the diagnosis. Two important pitfalls must be kept in mind. Firstly, arterial pO2, which reflects oxygen physically dissolved in plasma, is normal unless lung complications arise. Secondly, pulse oximetry is unreliable because of falsely high readings due to the erroneous detection of CO-Hb as oxyhaemoglobin.

The two main treatment modalities are 100% oxygen at atmospheric or at hyperbaric pressure. The rationale of this therapy is that the hypoxic insult to the tissues is greatly diminished by increasing the amount of oxygen dissolved in the blood and by markedly shortening the half-life of CO-Hb.

Although hyperbaric oxygen offers theoretical advantages over 100% oxygen therapy, it is not so widely available, takes a considerable time to organise, and can have a number of adverse effects. Furthermore, a Japanese study in 1977 found that half of the patients treated for intoxication and coma with hyperbaric oxygen still had neurological sequelae. However, it is possible that by reducing mortality, hyperbaric oxygen actually increased overall morbidity. Clinical experience and a number of reported series have all documented a visible clear-cut improvement with hyperbaric oxygen in seriously intoxicated patients. Furthermore, the threat of litigation exists if hyperbaric oxygen is not delivered. A recent prospective randomised study comparing the outcome of patients treated at atmospheric pressure with those treated with hyperbaric oxygen has now provided convincing evidence for the utility of the latter in preventing neurological sequelae, even in moderate poisoning.

A number of guidelines have been developed for the use of hyperbaric oxygen. When the diagnosis of CO poisoning is suspected, 100% oxygen treatment should be started immediately, without waiting for the CO-Hb results. Patients with coma, fits, prolonged loss of consciousness or myocardial instability should receive hyperbaric oxygen. If the peak levels of CO-Hb are more than 25% hyperbaric oxygen is recommended, even for minor symptoms. In children and pregnant women, more liberal use of hyperbaric oxygen is recommended; a lower cut-off point of 20% and possibly less is suggested, as both children and the fetus have a greater susceptibility to hypoxia.
The recommended regime is 100% oxygen administered at 2-3 atmospheres for about 90-120 minutes, followed by four hours of 100% oxygen therapy. Further hyperbaric oxygen treatments at eight-hourly intervals are recommended if neurological symptoms persist. Indeed one study has shown that multiple sessions are superior to single sessions. A recent study using a two-hour hyperbaric regime used once showed a much lower frequency of residual neurological problems than after normobaric therapy. Levels of 10-20% CO-Hb rarely cause serious illness, so it is preferable to treat with 100% oxygen for two to four hours, as hyperbaric oxygen usually takes about an hour to organise by which time treatment with 100% oxygen would have rendered it unnecessary. When unstable angina is the result of low-grade intoxication superimposed on coronary artery disease, we feel that 100% oxygen at atmospheric pressure is sufficient. However, myocardial toxicity in young healthy patients indicates severe intoxication in which case hyperbaric oxygen is necessary.

Hyperbaric therapy has occasionally been shown to cause baro trauma to the tympanic membrane. It should be avoided after cardiopulmonary resuscitation with external chest compression, pulmonary embolism, and asthma, as a life-threatening pneumothorax may develop. Probably, frail elderly patients are more safely treated at atmospheric pressure.

Supportive therapy is very important in acute intoxication. Severely intoxicated patients may need intubation, mechanical ventilation, and intensive care. However the majority of patients do well with hyperbaric oxygen. We monitor patients who do not need intubation in a coronary care unit. The hypoxic injury to the brain commonly results in an anoxic encephalopathy with cerebral oedema. For this reason, intravascular dexamethasone with an H2 antagonist such as cimetidine is usually given soon after admission.

A number of patients with intoxication are found by a passer-by after a period of time. In a warm environment rapid dehydration can ensue, resulting in hypotension and peripheral vasoconstriction. In such circumstances it is possible that the resultant ischaemia and hypoxia in the skin, visceral, and muscular may actually be more severe and prolonged than in the brain and heart. For this reason a balance must be struck between rapid rehydration and the minimisation of cerebral oedema. A number of cases of abdominal visceral infarction have been documented. Skin lesions have also been reported, but the most commonly reported complication is muscle rhabdomyolysis due to skeletal muscle hypoxia. This can result in serious renal impairment.

Neurological sequelae: prognostic factors

Neurological deficits can be of two types, immediate, due to the acute anoxic encephalopathy, or delayed, caused by late demyelination secondary to the primary hypoxic neuronal injury. Min reviewed 2967 patients admitted to hospital with CO poisoning, 86 (2.7%) of whom had long-term neuropsychiatric symptoms. However, Choi reported a frequency of 11.8%, with a mean onset of symptoms 22 days post-exposure. A longitudinal study on 100 pure. using extensive neuropsychiatric testing showed a frequency of 63% in those treated with 100% oxygen and 13% in those treated with multiple sessions of hyperbaric oxygen. Clinically, a broad range of neuropsychiatric abnormalities have been observed, such as dementia, psychosis, personality change and concentration deficit. Gait disturbance, facal or urinary incontinence, mutism, memory loss, and visual disturbance appear to be the commonest neurological problems. Parkinsonism appears to be one of the rare sequelae.

Autopsy studies have shown that the classical lesion of acute CO intoxication is symmetrical necrosis of the globus pallidus. A magnetic resonance imaging (MRI) study of 15 patients with delayed encephalopathy suggests that it is probably due to a diffuse reversible demyelinating process of white matter. A significant correlation between cerebral white matter changes on computed tomography scan and the development of delayed neurological sequelae has been documented.

There are a number of clinical risk factors for delayed encephalopathy. The CO-Hb level does not seem to be related. However, old age, comatous lasting two or three days, and persistent dizziness and fatigue after regaining consciousness are thought to predict problems. It is advisable to give multiple cycles of hyperbaric oxygen in these cases unless contraindicated. It appears that proton magnetic resonance spectroscopy can determine neurological viability before actual demyelination is observed on MRT. However, the role of this technique and the value of hyperbaric oxygen at that stage still has to be determined in prospective studies.
The prognosis of encephalopathy due to CO poisoning appears to be unfavourable. In one study, 57 eight patients had a progressive deteriorating course after poisoning with a persistent akinetik mute state. Four died after one year. Out of 23 patients with delayed encephalopathy after initial improvement, 14 were bed-bound and with akinetik and minium could walk but had severe cognitive impairment. After one year three had died but 14 patients had improved. Anatomical lesions tend to be permanent, as shown on MRI on patients who had been intoxicated 25 years previously.

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