Demystifying Carboxyhemoglobinemia - ED

Detecting Carbon Monoxide Poisoning in the Emergency Department

Summary

Carbon Monoxide (CO) is a gas produced by the combustion of carbon-containing fuels or the inadequate ventilation of natural gas. Once in the bloodstream, CO prevents oxygen from reaching tissues. Undetectable by humans, exposure to CO is the leading cause of death by poisoning in industrialized countries. Still, the condition presents a constellation of symptoms that mimic other illnesses. As a result as many as half of all CO-poisoned patients may be misdiagnosed when presenting to emergency departments, delaying treatment and even returning vulnerable patients and their families to potentially lethal environments.

CO exposure may be detected by measuring the carboxyhemoglobin (COHb) levels in a person’s blood. In hospitals, the most common means of measuring COHb is through the analysis of an invasive blood sample using a laboratory CO-Oximeter. However, according to one recently published study, only about half of all hospitals have the devices onsite. For hospitals with a CO-Oximeter, results may be obtained in about 10 minutes but in hospitals that must send the samples elsewhere for testing, results require an average of 15 hours. Each additional reading requires another blood draw and analysis. Conventional pulse oximeters are unable to detect COHb but a new device, the Masimo SET with Rainbow Technology monitor allows clinicians to detect and continuously monitor CO levels in the bloodstream noninvasively. Using multiple wavelengths of light to distinguish the various forms of hemoglobin (oxy-, deoxy-, carboxy- and met-) the device is capable of measuring blood CO saturation (SpCO®) levels and methemoglobin saturation (SpMet™) levels, in addition to the conventional variables of pulse rate, perfusion index and arterial oxygen saturation.

The stakes for properly diagnosing and treating CO poisoning are high. Assessment of a patient’s COHb level first provides an accurate diagnosis of CO poisoning and then guides treatment especially in cases elevated to the range of 10 percent or greater. Some mistakenly believe that if a patient recovers from the initial CO poisoning, they have made a complete recovery. However, multiple studies show that patients with prolonged and untreated CO exposure have long-term side effects and increased risk. If untreated, CO exposure may damage the neurological, cardiac, metabolic, pulmonary and renal systems of the body. Organs with a high metabolic requirement for oxygen, such as the heart and brain, are most susceptible to injury from CO. Even at relatively low COHb levels, patients with underlying cardiovascular disease are especially at serious risk for cardiac complications including myocardial ischemia or infarction, and even cardiac arrest.

Fortunately, the effects of CO poisoning can be reversed if caught in time. The immediacy of results and ability to trend the results over time expedite efficacious treatment and may contribute to improved clinical outcomes. With lives and significant resources at stake, the speed at which suspicion evolves to diagnosis is critical. A quick noninvasive measurement of COHb using a new Rainbow Pulse CO-Oximeter™ device may contribute to better informed treatment decisions.
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The Guessing Game

CO poisoning is the leading cause of death by poisoning in industrialized countries and may be responsible for more than half of all fatal poisonings worldwide. It is estimated that approximately 43,000 emergency room visits are attributed to CO poisoning in the United States each year. At least 3,800 people die annually in the U.S. from the effects of CO poisoning, and 1,400 of these deaths are accidental. Unfortunately, for most patients poisoned by the colorless, odorless, tasteless gas, CO poisoning is not the immediate and obvious diagnosis. Variable symptoms, a wide range of patient sensitivity and unsophisticated detection systems often result in misdiagnosis and treatment delays.

Rapid Determination of Carbon Monoxide Poisoning (COHb)

- Emergency Departments
- Urgent Care Facilities
- Physicians Offices
- Inpatient / Outpatient Surgery Centers
- First – Responders/ Emergency Medical Services and Fire / Homeland Security
- Acute Care Hospitals
- Point of Care – Natural Disaster Zones
- Toll Booths / Parking Garages
- Airplanes
- Construction zone

Even in the case of Randal McCloy, the sole survivor of the Sago Mine tragedy, in which CO poisoning was the probable cause of illness (according to multiple published accounts of the incident), the first physician to attend to the miner reported that McCloy’s carbon monoxide levels were negative. “That means that as best as we can tell with somewhat primitive equipment that we have here for measuring those, his carboxyhemoglobin levels were negative, indicating no carbon monoxide in his system, as far as we could tell,” she told reporters.

When pressed for more information, the clinician told the media, “When you put the oxygen saturation monitor on their finger, it’s false [but] it doesn’t give you a true reading in somebody with carbon monoxide poisoning. So you really have to be able to run the blood and check for carboxyhemoglobin.” In fact, tests run in the subsequent days show McCloy to be suffering from brain hemorrhaging and edema, muscle injury, liver failure and faulty heart function due to severe CO poisoning.

Later, physicians at Allegheny General Hospital stated that after receiving three hyperbaric oxygen (HBO) treatments, McCloy was showing signs of improved brain stem and organ function. MRI scans illustrated the evidence of neurological damage but the clinical consequences remain to be seen.

Some people are more susceptible to long-term harm from CO exposure than others. It is possible that the same physiology that enabled McCloy to survive generally lethal CO levels for more than forty hours may also afford him a better clinical outcome than would be expected. While there are populations known to be highly susceptible to the negative effects of CO: children, pregnant women, adults with cardiac disease, individuals with increased oxygen demand and patients with chronic respiratory problems; it is not possible to assess a person’s CO resilience.

Elevated COHb: Patients at High Risk for Negative Outcome

- Children; elderly
- Adults with cardiac disease;
- Pregnant women
- Patients with increased oxygen demand or decreased oxygen-carrying capacity;
- Patients with chronic respiratory insufficiency.
CO poisoning is the single most common source of poisoning injury as treated in hospital emergency departments. While its presentation is not uncommon, the constellation of symptoms that manifest when a patient is poisoned with carbon monoxide do not prompt most clinicians to consider carboxyhemoglobinemia when attempting a diagnosis. The vague symptoms can be mistaken for those of many other illnesses including food poisoning, influenza, migraine headache, or substance abuse. In the attempt to find the causative agent for the symptoms, many unnecessary, potentially costly and sometime resource-intensive diagnostics may be ordered, to no avail. Because the symptoms of CO poisoning may mimic an intracranial bleed, time and cost for a negative result may precede proper diagnosis, unnecessarily increasing healthcare costs. During the delay associated with running unnecessary diagnostics, patients may find that their symptoms abate and their health improves as the hidden culprit, CO, is flushed from the blood during the normal ventilation patterns over time. Multiple reports have shown that the patients may be discharged and returned back to the environment where the poisoning occurred, only to once again be exposed to the silent killer, carbon monoxide.

There are two main types of CO poisoning: acute, which is caused by short exposure to a high level of carbon monoxide, and chronic or subacute, which results from long exposure to a low level of CO. Which symptoms appear depend on the level of CO in the environment and the length of exposure, as well as the patient’s state of health.

The general symptoms of CO poisoning, including headache, dizziness, nausea, fatigue, and weakness, are vague. See Table 1. Patients with acute CO poisoning are more likely to present with more serious symptoms, such as cardiopulmonary problems, confusion, syncope, coma, and seizure. Chronic poisoning is generally associated with the less severe symptoms. Low-level exposure can exacerbate angina and chronic obstructive pulmonary disease, and patients with coronary artery disease are at risk for ischemia and myocardial infarction even at low levels of CO.

Patients that present with low COHb levels correlate well with mild symptoms described in Table 2 as do cases that register levels of 50-70%, which are generally fatal. However, intermediate levels show little correlation with symptoms or with prognosis. It seems that the severity of clinical condition is not only related to CO concentration but also the duration of exposure and the prevailing clinical disposition of the patient. Some patients presenting with a carboxyhemoglobin level of 20% may be remarkably symptomatic, while others experiencing the same level of COHb% may exhibit only mild, equivocal symptoms. A patient exposed to high concentrations for a short time may be less symptomatic than a patient who reaches the same COHb level after a prolonged exposure.

<table>
<thead>
<tr>
<th>Severity</th>
<th>COHb Level</th>
<th>Signs &amp; Symptoms</th>
</tr>
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<tbody>
<tr>
<td>Mild</td>
<td>&lt;15 - 20%</td>
<td>Headache, Nausea, Vomiting, Dizziness, Blurred Vision.</td>
</tr>
<tr>
<td>Moderate</td>
<td>21 - 40%*</td>
<td>Confusion, Syncope, Chest Pain, Dyspnea, Weakness, Tachycardia, Tachypnea, Rhabdomyolysis</td>
</tr>
<tr>
<td>Severe</td>
<td>41 - 59%*</td>
<td>Palpitations, Dysrhythmias, Hypotension, Myocardial ischemia, Cardiac arrest, Respiratory arrest, Pulmonary edema, Seizures; Coma</td>
</tr>
<tr>
<td>Fatal</td>
<td>60+%</td>
<td>Death</td>
</tr>
</tbody>
</table>

* At moderate to severe levels of COHb poisoning the correlation between blood levels and symptomatology is poor.
Detection

Table 2: COHb Levels in Persons 3 - 74 Years of Age

<table>
<thead>
<tr>
<th>Smoking Status</th>
<th>Percent COHb (mean ± SD)</th>
<th>Percent COHb (98th percentile)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonsmokers</td>
<td>0.83 ± 0.67</td>
<td>&lt; 2.50</td>
</tr>
<tr>
<td>Current smokers</td>
<td>4.30 ± 2.55</td>
<td>≤ 10.00</td>
</tr>
<tr>
<td>All statuses combined</td>
<td>1.94 ± 2.24</td>
<td>≤ 9.00</td>
</tr>
</tbody>
</table>

One thing that is certain about COHb levels is that smokers present with higher levels than do non-smokers. As can be seen in Table 2, the COHb level in non-smokers is approximately one to two percent. In patients who smoke, a baseline level of nearly five percent is considered normal, although it can be as high as 13 percent. Although COHb concentrations between 11 percent and 30 percent can produce symptoms, it is important to consider the patient’s smoking status.

CO poisoning is known as the great imitator for its ability to present with equivocal signs and symptoms, many of which closely resemble other diseases. In particular, patients may be misdiagnosed with viral illness, acute myocardial infarction, and migraine. It is estimated that CO poisoning misdiagnosis may occur in up to 30-50 percent of CO-exposed patients presenting to emergency departments. As described below, failing to assess COHb levels early may return vulnerable patients and their families to potentially lethal environments.

Missing the Signs

A 67-year-old man sought medical help after three days of light-headedness, vertigo, stabbing chest pain, cough, chills and headache. His wife had experienced similar ailments over the past week. He was admitted, evaluated and discharged with a diagnosis of viral syndrome. Ten days later he returned to the ER with vertigo, palpitations and nausea but was sent home for outpatient follow-up. Four days later he again returned to the ER with diarrhea and severe chest pain, collapsing to the floor. He was admitted to the Coronary Care Unit with acute myocardial infarction. Among the results of a routine arterial blood gas analysis, it was found that his COHb levels were 15.6%. A COHb level then obtained on his wife was 18.1%. A rusted furnace was found to be the source.

A 69-year-old man came to the ER after days of confusion, nausea, vomiting, intermittent syncope, hallucinations and shortness of breath. An arterial blood gas measurement found an oxygen saturation of 89%. He was admitted to the coronary care unit with a diagnosis of acute myocardial infarction. The next day a COHb level was measured and normal. While the patient was hospitalized he invited his sister and daughter-in-law to stay in his home. They both arrived at the ER the next morning with headaches, vomiting, and vertigo. Their COHb levels on initial observation were 28% and 32%. The man’s gas water heater was faulty.

A 47-year-old male urologist and his wife attended a medical conference in Jackson, Wyoming. Both reported to a local emergency department with symptoms including headache, malaise, and metabolic acidosis. The husband and wife were sent back to the conference resort hotel with a diagnosis of gastroenteritis. The following day, they were both found unresponsive in their hotel. He died 3 hours later; and his wife has severe long-term neurocognitive sequelae. The cause of death and long-term morbidity was carbon monoxide poisoning due to a faulty boiler. A $17,000,000 verdict was awarded to the afflicted family against the resort hotel owners.
Regardless of the means of detection used in emergency department care, several factors make assessing the severity of the CO poisoning difficult. The length of time since CO exposure is one such factor. The half-life of CO is four to six hours when the patient is breathing room air, and 40–60 minutes when the patient is breathing 100 percent oxygen. If a patient is given oxygen during their transport to the emergency department, it will be difficult to know when the COHb level peaked.15

In addition, COHb levels may not fully correlate with the clinical condition of CO-poisoned patients because the COHb level in the blood is not an absolute index of compromised oxygen delivery at the tissue level. Furthermore, levels may not match up to specific signs and symptoms; patients with moderate levels will not necessarily appear sicker than patients with lower levels.31

In hospitals, the most common means of measuring CO exposure is through the use of a laboratory CO-Oximeter. A blood sample, under a physician order, is drawn from either venous or arterial vessel and injected into a lab CO-Oximeter. The laboratory device measures the invasive blood sample using a method called spectrophotometric blood gas analysis.24 Because the CO-Oximeter can only yield a single, discrete reading for each aliquot of blood sampled, the reported value is a noncontinuous snapshot of the patient’s condition at the particular moment that the sample was collected. To compound the difficulty of detecting CO exposure, when the laboratory calculates the patient’s oxygen saturation levels from the oxygen partial pressure (PO2), the arterial SaO2 may appear normal. The clinical usefulness of CO-Oximetry is inhibited further by the relative deficiency of devices currently installed in acute care hospitals. One recent study found that fewer than half of hospitals in the U.S. have the necessary equipment on site to diagnose CO poisoning.25 For those that did not have the testing equipment, the average time to receive results of a blood sample sent to another facility was over 15 hours. In hospitals that have CO-Oximetry equipment, results may be returned in an average of 10 minutes (see Table 4.)

Unfortunately, standard pulse oximeters are incapable of isolating the carbon monoxide contaminated hemoglobin from the oxyhemoglobin.26 Thus, pulse oximeters artificially overestimate arterial oxygen saturation in the presence of elevated blood carbon monoxide. Therefore, the readings will be falsely high when carbon monoxide is occupying binding sites on the heme molecule.

<table>
<thead>
<tr>
<th>Table 3: Comparison of Testing Methods Time to Results</th>
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<tbody>
<tr>
<td>Testing Method</td>
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<tr>
<td>Pulse CO-Oximeter</td>
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<tr>
<td>Onsite CO-Oximeter</td>
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<tr>
<td>Off-Site CO-Oximeter</td>
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</table>

The latest technology in CO poisoning detection employs a noninvasive and continuous platform. The Masimo SET® with Rainbow Technology Pulse CO-Oximeter Monitor [Masimo, Irvine, CA] is the first device that allows clinicians to detect and continuously monitor CO levels in the bloodstream noninvasively. Using 7+ wavelengths of light to distinguish the various forms of hemoglobin (oxy-, deoxy-, carboxy- and met-) the device is capable of measuring blood CO saturation (SpCO) levels, methemoglobin saturation (SpMet) levels, in addition to pulse rate, arterial oxygen saturation, and perfusion index. The device’s accuracy has been demonstrated accurate to 40 percent SpCO, with a range of ±3 percent around the measurement.27

Noninvasive monitoring reduces the opportunity for hospital acquired infection and overall patient discomfort. Needle-free testing means a safer environment for patients and caregivers alike. In addition, the immediacy of results available at the point-of-care results in less drain on resources while expediting efficacious treatment and better outcomes. The continuous nature of the noninvasive Rainbow Pulse CO-Oximeter device enables the ability to trend data over time while conventional CO-Oximetry requires a new blood sample each time the status of the dyshemoglobins is required.
Because clinicians traditionally order blood measurement of COHb only when the condition is suspected, there has been a tendency to diagnose only the most symptomatic patient whose exposure history is known. With noninvasive Pulse CO-Oximeter technology now available, one might expect that many instances of elevated COHb will be discovered among patients without a classic history of recognized exposure to CO.\textsuperscript{28}

**Treatment**

Due to the challenges of traditional COHb detection and the lack of correlation between levels and symptoms, most experts recommend using COHb level only as confirmation of the diagnosis in a patient with suspected CO exposure. Treatment is then based on the patient’s history and the severity of symptoms. Still, COHb levels are recommended to guide management especially in cases elevated to the range of 25 percent or greater.\textsuperscript{29} Because noninvasive and continuous Pulse CO-Oximetry allows real-time trend evaluation of the CO-poisoned patient, efficacious treatment protocols can be quickly identified and implemented. Carbon monoxide toxicity is traditionally treated with either 100% oxygen therapy by mask or high flow device, or by hyperbaric medicine (HBO). A significant relationship exists between the delay between efficacious treatment, the severity of the CO toxicity, and the potential for delayed neuropsychiatric and/or abnormal cardiac sequelae.

### Table 4: Carbon Monoxide: Half-life Elimination from Blood

<table>
<thead>
<tr>
<th>Condition</th>
<th>Half-life</th>
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</thead>
<tbody>
<tr>
<td>Room Air</td>
<td>240 - 360 minutes</td>
</tr>
<tr>
<td>Oxygen (100%)</td>
<td>80 minutes</td>
</tr>
<tr>
<td>Hyperbaric Oxygen (HBO)</td>
<td>22 minutes</td>
</tr>
</tbody>
</table>

With the half-life of COHb at four to six hours, a COHb level should be obtained soon after exposure is suspected. Noninvasive and continuous COHb measurements employing Rainbow technology provide an increasingly valued diagnostic methodology without delays and potentially costly missed diagnosis.

Noninvasive Pulse CO-Oximeters capable of immediately and accurately detecting COHb in patients presenting with a host of symptoms are likely to drastically reduce misdiagnosis and aid rapid treatment. However, emergency department clinicians will need a guidance protocol for patient management when they uncover patients suffering from CO poisoning. One such triage protocol was developed for first-responders\textsuperscript{28} but applies well for emergency department care. See Figure 1 (next page).
If the SpCO level is 3–12 percent, the elevation could be due either to smoking or another source. If the patient is experiencing such symptoms as headache, nausea or vomiting, they should receive 100 percent oxygen and undergo further evaluation and treatment as needed. If the SpCO level is 3–12 percent and the individual is asymptomatic, no further medical evaluation of the SpCO level is necessary (although the source of the exposure should be identified and fully understood such that it can be eliminated as a future etiologic agent of CO poisoning). Without this understanding a patient may be inadvertently sent from the ED back to the environment where the poisoning likely occurred.

Hyperbaric Oxygen Therapy (HBO) can decrease the half-life of CO to 22 minutes, induce cerebral vasoconstriction to reducing intracranial pressure and cerebral edema, and reduce the risk of long-term disability. In particular, HBO treatment is appropriate for patients who experience unconsciousness, neurological signs, cardiovascular dysfunction or severe metabolic acidosis, irrespective of their COHb levels.25

Figure 1: Hampson SpCO Triage Algorithm26
Clinical Effects

The stakes for diagnosing and treating CO poisoning are high. Fast, effective treatment can do much to improve clinical outcomes and contain damage to the neurologic, cardiac, metabolic, pulmonary and renal systems of the body as described in Table 5.

| Table 5: Impact of CO Poisoning on the Body Systems |
|-----------------|--------------------------------------------------------------------------------------------------|
| Neurologic      | CO poisoning causes central nervous system depression presenting in a host of impairments. In mild cases, patients report headaches, dizziness and confusion. In severe cases, patients may be comatose or develop seizures. Long-term neurocognitive and neuropsychiatric sequelae are reported even after moderate to severe single exposures. |
| Cardiac         | CO poisoning causes decreased myocardial function and vasodilatation and a decreased oxygen delivery to, and utilization of, oxygen by the myocardium. As a result, the patients may present hypotensive or with tachycardia, chest pain, arrhythmias or myocardial ischemia. Most deaths from CO poisoning ultimately result from ventricular dysrhythmias. Long-term cardiac sequelae are reported even after moderate to severe single exposures, increasing the odds ratio of premature cardiac death. |
| Metabolic       | Respiratory alkalosis (hyperventilation) is possible in mild cases. With severe exposure, metabolic acidosis may result in elevated levels of acid throughout the body. |
| Pulmonary       | Pulmonary edema occurs in 10 – 30 percent of acute CO exposures. This may be due to a direct effect on the alveolar membrane, left ventricular failure, aspiration or neurogenic pulmonary edema. |
| Multiple Organ Failure | At high levels, multiple organ failures are expected, with a lethal outcome likely without immediate treatment to remove the CO. |
The effects of CO are not confined to the period immediately after exposure. Persistent or delayed effects have been reported. In particular, a syndrome of delayed neurological effects, often referred to as DNS may manifest in a myriad of forms. DNS is experienced by 11 percent to 30 percent of patients who have had CO poisoning. The resultant sequelae—confusion, seizures, hallucinations, persistent vegetative state, parkinsonism, short-term memory loss, psychosis, and behavioral changes—may appear anywhere from three to 240 days after carbon monoxide exposure, even in patients in whom neurologic impairment isn’t initially recognized, and may be chronic. There is no way of predicting which patients will suffer such sequelae. In general, those with more severe initial symptoms are at highest risk. Most mild cases resolve within two months, although patients with severe exposure may never make a full recovery from delayed neuropsychiatric sequelae.

Patients with underlying cardiovascular disease are at risk for cardiac complications. Risk of sudden cardiac death increases with CO poisoning. Hypotension and inadequate oxygenation can cause myocardial ischemia or infarction, and even cardiac arrest. Even years after being treated for moderate to severe CO poisoning, patients who sustained myocardial injury as a result of exposure had an increased risk of death.

Metabolic disturbances such as respiratory alkalosis and metabolic acidosis, as well as pulmonary and renal maladies may also arise from CO exposure. Pulmonary edema which occurs in 10–30 percent of acute CO exposures. The build up of COHb in the blood stream may also cause rhabdomyolysis - the breakdown of muscle fibers resulting in the release of muscle fiber contents into the circulation. Some of these are toxic to the kidney and frequently result in kidney damage and renal failure.
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Causes of CO Poisoning

Carbon monoxide is a gas produced by the combustion of carbon-containing fuels (oil, kerosene, gasoline, coal, wood) or the inadequate ventilation of natural gas. It is undetectable by humans. Faulty furnaces, motor vehicles, motor boat docks with swimming platforms, portable generators, stoves, gas ranges, and gas heaters are the most common sources of carbon monoxide poisoning. At one time, it was estimated that 29 percent of unintentional CO-related deaths were due to motor vehicles exhaust. This rate has declined significantly since 1979, likely owing to improved emissions standards. Before catalytic converters, closed environment exposure to car exhaust could produce death within 30 minutes. CO poisoning occurs most often in the fall and winter months, when the use of gas furnaces and alternative heat sources increases. It also occurs when generators, which provide power to residences and businesses, are used in poorly ventilated areas. Portable generators are commonly used in situations where power interruptions are expected (hurricane zones, ice storms, flood regions). A lesser known source of carbon monoxide is the vapors from methylene chloride, a compound commonly found in paint strippers. When the fumes are inhaled, it is converted in vivo to carbon monoxide.

Pathophysiology

Once in the bloodstream, CO has a multi-prong deleterious effect on the body. But despite the many adverse mechanisms outlined in Table 6, each produces the same result: preventing oxygen from reaching tissues thus causing tissue hypoxia:

<table>
<thead>
<tr>
<th>Table 6: Oxygen-limiting mechanisms of CO</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Limits oxygen transport</strong></td>
</tr>
<tr>
<td>The affinity between carbon monoxide and hemoglobin is more than 200 times greater than that between oxygen and hemoglobin. As a result, CO more readily binds to hemoglobin, forming carboxyhemoglobin (COHb).</td>
</tr>
<tr>
<td><strong>Inhibits oxygen transfer</strong></td>
</tr>
<tr>
<td>CO further changes the structure of the hemoglobin molecule, which inhibits the already limited oxygen that has attached to be prematurely released into tissues.</td>
</tr>
<tr>
<td><strong>Tissue inflammation</strong></td>
</tr>
<tr>
<td>The tissue damage caused by poor perfusion and lack of oxygen attracts leukocytes to the damaged area. This initiates and sustains an inflammatory response, causing further tissue damage with increased capillary leakage and edema. This process results in a tissue reperfusion injury, similar to what is seen in patients who have suffered a myocardial infarction.</td>
</tr>
<tr>
<td><strong>Poor cardiac function</strong></td>
</tr>
<tr>
<td>Decreased oxygen delivery to, and utilization of, oxygen by the myocardium may cause tachycardia, arrhythmias or myocardial ischemia. Long-term damage to the heart has been demonstrated even after a single moderate to severe CO exposure.</td>
</tr>
<tr>
<td><strong>Increased activation of nitric oxide</strong></td>
</tr>
<tr>
<td>CO increases the production of nitric oxide, an important component in peripheral vasodilatation. In systemic inflammatory response syndrome (SIRS), nitric oxide can increase secondary to CO, or induce CO secondary to hemolysis.</td>
</tr>
<tr>
<td><strong>Vasodilatation</strong></td>
</tr>
<tr>
<td>CO increases the production of nitric oxide. Nitric oxide causes vasodilation. Vasodilatation is responsible for decreased cerebral blood flow and systemic hypotension. Nitric oxide is largely converted to methemoglobin in the body.</td>
</tr>
<tr>
<td><strong>Free radical formation</strong></td>
</tr>
<tr>
<td>Nitric oxide also accelerates free radical formation. Free radical formation causes endothelial damage and oxidative damage to the brain.</td>
</tr>
</tbody>
</table>
Conclusion

The effects of CO poisoning can be reversed if caught in time. Detection and diagnosis of CO poisoning is currently based upon clinical suspicion and confirmed by invasive blood sampling for COHb analyzed by CO-Oximetry. While many hospitals have blood gas machines with CO-Oximetry, many smaller hospitals do not, which makes timely confirmed diagnosis of CO poisoning in these situations impossible. Organs with a high metabolic requirement for oxygen, such as the heart and brain, are particularly susceptible to injury from CO. The resulting tissue ischemia can lead to organ failure, permanent changes in cognition, or death. Those that survive the initial poisoning may experience serious long-term neurological, cardiac, metabolic, pulmonary and renal impairment as a result of their CO exposure.

With lives and significant resources at stake, the speed at which suspicion evolves to diagnosis is critical. A quick noninvasive measurement of COHb using the new Masimo Pulse CO-Oximeter device may contribute to better informed treatment decisions ending the guessing game.

References
