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Toxic alcohols

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POISONINGS BY THE TOXIC ALCOHOLS (METHANOL, ETHYLENE GLYCOL, ISOPROPANOL, DIETHYLENE GLYCOL, AND PROPYLENE GLYCOL) can cause cellular dysfunction and death, but symptoms may be nonspecific. Delays in diagnosis increase the risk of irreversible organ damage and death. In this review, we discuss the mechanisms of toxicity, methods available for diagnosis, and current recommendations for therapy.

MECHANISMS OF TOXICITY

The toxic alcohols are inebriating but are not directly toxic, except for isopropanol. Their toxic effects result from their metabolites. A simplified schema depicting their primary metabolic pathways is shown in Figure 1A.

Alcohol dehydrogenase catalyzes the first oxidation of the toxic alcohols. The resulting aldehydes (except for acetone from isopropanol) undergo further oxidation by aldehyde dehydrogenase to form carboxylic acid metabolites: methanol is metabolized to formic acid, ethylene glycol to oxalic and glycolic acid, diethylene glycol to 2-hydroxyethoxyacetic acid and glycolic acid, and propylene glycol to \( \alpha \)-lactic and \( \beta \)-lactic acid. Alcohol dehydrogenase is the critical enzyme that modulates the production of the toxic metabolites. Coingested ethanol, a competitive substrate for alcohol dehydrogenase, delays production of the toxic metabolites. Increased production of lactic acid can result from exposure to the metabolites of methanol or ethylene glycol, but spurious increments in blood lactate may occur with exposure to ethylene glycol metabolites as a result of interference of glycolate with the lactate measurement by point-of-care instruments.

EPIDEMIOLOGIC FEATURES

The intoxications can occur through different means (Table 1). Methanol intoxication most commonly follows ingestion of automotive windshield-washer fluid, industrial products, or adulterated liquids, but exposure can also occur through pulmonary and cutaneous routes. Ethylene glycol is most commonly ingested by adults in antifreeze or in adulterated spirits in which ethylene glycol has been added in lieu of ethanol, in an attempt to commit suicide; in children, it is most commonly ingested unintentionally. Isopropanol intoxication usually results from ingestion of rubbing alcohol, hand sanitizer, and various industrial products, but intoxication can also be due to inhalation or absorption through dermal or rectal routes.

Diethylene glycol intoxication results from ingestion of automotive brake fluids or industrial products, but it usually occurs in outbreaks in which consumer products or oral medications for children contain diethylene glycol in lieu of propylene glycol...
Figure 1. Metabolic Pathways of Toxic Alcohols and Time Course of Changes in the Osmolal and Anion Gaps with and without Coingested Ethanol.

Panel A shows the metabolic pathways of toxic alcohols. Alcohol dehydrogenase and aldehyde dehydrogenase sequentially oxidize the toxic alcohols. Alcohol dehydrogenase catalyzes the first oxidation of the toxic alcohols and is an important target for antidotal therapy. The enclosed boxes highlight the putative toxic metabolites. Methanol is metabolized to formic acid, ethylene glycol to oxalic and glycolic acid, diethylene glycol to 2-hydroxyethoxyacetic acid and glycolic acid, and propylene glycol to D-lactic and L-lactic acid. Panel B shows the time course of changes in the osmolal and anion gaps with and without coingested ethanol. An increased osmolal gap is prominent early owing to the accumulation of the un-ionized alcohols. As metabolism proceeds, the osmolal gap declines with the formation of ionized metabolites. Conversely, the serum anion gap is lowest before the alcohol is metabolized and increases with the formation of ionized metabolites. The time course of these changes in both parameters varies among the alcohols. They typically evolve over several hours to over a day. Coingested ethanol impedes metabolism (dashed lines) and delays the onset of the high anion-gap acidosis.
### Table 1. Clinical and Laboratory Features of the Toxic Alcohols.

<table>
<thead>
<tr>
<th>Alcohol Type and Molecular Weight</th>
<th>Increase in Serum Osmolality per 10-mg/dl Increase in Serum Alcohol Concentration</th>
<th>Common Sources</th>
<th>Common Clinical Features</th>
<th>Major Laboratory Features</th>
<th>Onset of Clinical and Laboratory Features*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mOsm/kg of water</td>
<td><strong>Ethylene glycol, 62.07</strong></td>
<td>1.60</td>
<td>Automotive antifreeze, engine coolants, deicing fluids</td>
<td>Inebriation, acute kidney injury</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Methanol, 32.04</strong></td>
<td>3.09</td>
<td>Windshield-washer fluid, carburetor cleaner, octane boosters, racing fuels, camp stove fuel, adulterated ethanol (&quot;moonshine&quot;)</td>
<td>Inebriation, abdominal pain, decreased vision with blindness, Parkinson-like features (rare)</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Propylene glycol, 76.09</strong></td>
<td>1.31</td>
<td>Diluent in parenteral medications, automotive antifreeze (marketed as a safer alternative to ethylene glycol)</td>
<td>Hepatic and renal disease are uncommon but will predispose to more severe toxicity</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Diethylene glycol, 106.12</strong></td>
<td>0.9</td>
<td>Automotive brake fluids, hydraulic fluids, adulterated liquid medications (e.g., inappropriate substitution for propylene glycol or glycerin) (most common source)</td>
<td>Abdominal pain, nausea and vomiting, acute pancreatitis, acute kidney injury</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Isopropanol, 60.02</strong></td>
<td>1.66</td>
<td>Rubbing alcohol, hand sanitizers</td>
<td>Inebriation, depressed senso-rium, abdominal pain</td>
</tr>
</tbody>
</table>

* Without or with coingested ethanol indicates the impact of the simultaneous presence of ethanol in the patient. NA denotes not applicable.
as a diluent. In rare cases, dermal absorption across nonintact skin may produce toxic effects. Propylene glycol is present in numerous consumer products and in antifreeze, but intoxication is usually due to prolonged high-dose infusions of medications such as lorazepam that contain propylene glycol as a diluent.

**Clinical Findings**

The alcohols initially depress the sensorium and later produce organ dysfunction. Methanol is associated with decreased vision (in 29 to 72% of cases, occasionally producing blindness), pulmonary dysfunction, abdominal pain, coma, and, rarely, Parkinson-like symptoms. Clinical findings usually evolve over 6 to 24 hours but can be delayed as long as 72 to 96 hours if ethanol is coingested. Neurologic sequelae may ensue days or weeks after exposure.

Ethylene glycol poisoning leads to formation of oxalate crystals, which deposit in the lungs, heart, and kidney and produce organ dysfunction. Cranial nerve damage, sometimes delayed for days, can also occur. Neurologic dysfunction develops in the first 12 hours, followed by cardiac and pulmonary dysfunction 12 to 24 hours after exposure and acute kidney injury 48 to 72 hours after exposure. However, the organ dysfunction can occur concomitantly. Coingestion of ethanol can delay the appearance of clinical abnormalities.

Isopropanol intoxication depresses the sensorium and can produce respiratory dysfunction, cardiovascular collapse, acute pancreatitis, hypotension, and lactic acidosis. Serum isopropanol concentrations above 500 mg per deciliter (83 mmol per liter) are clinically significant, and those greater than 1500 mg per deciliter (250 mmol per liter) produce deep coma. Acetone can produce a spurious increase in serum creatinine concentration as a result of interference with laboratory measurement.

Diethylene glycol poisoning can cause abdominal pain, nausea, vomiting, diarrhea, acute pancreatitis, altered mental status, hepatic disease, central and peripheral neuropathy (sometimes leading to quadriplegia), and acute kidney injury. Acute kidney injury often appears 8 to 24 hours after exposure, can require dialysis, and is a major cause of death. Coiningestion of ethanol can delay toxicity by as long as 48 to 72 hours. Cranial nerve palsies and other neurologic complications can appear 5 days or longer after exposure.

Propylene glycol intoxication often produces only an increased osmolal gap, but it can produce lactic acidosis and acute kidney injury. Preexisting hepatic disease, renal disease, or both are predisposing factors. Patients who receive a continuous infusion of high-dose lorazepam (>10 mg per hour) for more than 48 hours are at high risk.

**Diagnosis**

Information from the medical history, physical examination, blood chemical profiles, and tests to identify the parent alcohol or its metabolites are helpful in diagnosis. History of exposure to one of the toxic alcohols is important, given the nonspecific clinical findings and the potential delay between exposure and their appearance.

**Blood Chemical Profiles**

Accumulation of the alcohol increases the serum osmolality and the osmolal gap (the difference between the serum osmolality measured by the freezing-point depression and the serum osmolality estimated from the equation given below). Later, accumulation of organic acid anions increases the serum anion gap. The serum osmolality can be estimated with various formulae, but the formula that is acceptable for clinical purposes is as follows: estimated serum osmolality = (2 × Na⁺ [in millimoles per liter]) + (blood urea nitrogen [in milligrams per deciliter] ÷ 2.8) + (glucose [in milligrams per deciliter] ÷ 18). The expected normal osmolal gap is 10 to 20 mOsm per kilogram of water. Higher levels reflect accumulation of osmotically active substances such as the toxic alcohols. The increase in the serum osmolal gap depends on the serum concentration and the molecular weight of the alcohols (Table 1).

The basal serum osmolal gap can be less than 10 mOsm per kilogram of water or even negative. A low basal serum osmolal gap might obscure any increase caused by accumulation of a toxic alcohol. A normal osmolal gap cannot be used to rule out toxic alcohol ingestion; in one study, some patients with toxic alcohol poisonings had osmolal gaps within the normal range.
The osmolal gap varies during the course of intoxication (Fig. 1B). Accumulation of the parent alcohol initially elevates the osmolal gap, but as metabolism progresses, the osmolal gap falls. Coingested ethanol (observed in 10 to 60% of cases of methanol and ethylene glycol intoxication) will contribute to the increase in the osmolal gap (increase of 2.17 mOsm per kilogram of water per 10-mg-per-deciliter increase in serum alcohol concentration) and will also slow metabolism of the alcohols, prolonging the duration of an increased osmolal gap.

The baseline value of the serum anion gap (i.e., before the accumulation of organic acid anions that results from metabolism of the alcohol) can vary by 10 mmol per liter from the lowest to the highest value. If the baseline anion gap is low, it might not rise above the upper limit of normal despite considerable accumulation of organic acid anions. Also, the anion gap rises as metabolism progresses (Fig. 1B). A poisoned patient can present with both a normal or high osmolal gap and a normal or elevated serum anion gap. An elevated osmolal or anion gap does not always indicate toxic alcohol poisoning. Lactic acidosis, ketoacidosis, chronic kidney disease, and the sick cell syndrome all may increase both gaps. In one study, a minority of the patients with increased osmolal and anion gaps had toxic alcohol poisoning. Some racing fuels containing methanol and nitromethane may falsely elevate creatinine concentration as a result of interference with laboratory testing by means of the Jaffe reaction.

Ethylene glycol is metabolized to oxalic acid, which leads to crystalluria and, at times, severe acute kidney injury. Dihydrate crystals appear early, and monohydrate crystals appear later. Coprecipitation of oxalate with calcium can occasionally produce hypocalcemia. Acetone resulting from isopropanol metabolism can produce a positive nitroprusside reaction for acetoacetate at high concentrations. Lactic acidosis due to accumulation of the l-isomer is most frequent in propylene poisoning, but d-lactic acidosis has occurred in some cases. Because d-lactic acidosis will not be detected by the usual method for measuring lactate that detects only the l-isomer, the presence of d-lactic acidosis could be missed. Gas or liquid chromatography most accurately detects and quantifies the toxic alcohols in body fluids but is laborious, expensive, and often unavailable. Table 2 shows other methods to identify toxic alcohols that are either in use or in development. Pitfalls in the interpretation of laboratory data include failure to recognize that concentrations may be expressed in milligrams per liter or milligrams per deciliter, with values higher than 200 mg per liter or 20 mg per deciliter indicating the need for treatment. Also, tests for volatile substances may detect methanol, ethanol, acetone, and isopropanol but not ethylene glycol or diethylene glycol.

**TREATMENT**

Delays in treating toxic alcohol poisonings lead to worse outcomes. Therefore, therapy should commence expeditiously when there is a strong suspicion of toxic alcohol poisoning or when metabolic acidosis of unknown cause is present. Although there will be variability in the approach to diagnosis and treatment of the toxic alcohol poisonings, an algorithm consistent with our experience and current literature is depicted in Figure 2.

**METHANOL AND ETHYLENE GLYCOL**

Gastrointestinal absorption of methanol or ethylene glycol is rapid, so gastric decontamination is usually not helpful. Treatment includes prevention of metabolism and removal of the alcohol and its metabolites from the body.

Intravenous administration of base solution corrects metabolic acidosis and increases the ionization of formate, which facilitates its urinary excretion and reduces its penetration into the optic nerve. Antidotal treatment should commence when the serum methanol or ethylene glycol concentration is higher than 20 mg per deciliter (methanol, 6 mmol per liter; and ethylene glycol, 3 mmol per liter) and the patient has a documented history of ingesting one of the alcohols or when there is strong suspicion that the patient ingested one of the alcohols and has an osmolal gap greater than 10 mOsm per kilogram of water or metabolic acidosis of unknown cause (in accordance with the guidelines from the American Academy of Clinical Toxicology [AACT]; Table S1 in Supplementary Appendix 1, available with the full text of this article at NEJM.org). Ethanol has a high affinity for alcohol dehydrogenase, and intravenous ethanol, although
not approved by the Food and Drug Administration (FDA), is often used for treatment. A serum ethanol concentration of 100 mg per deciliter (22 mmol per liter) provides competitive inhibition of the enzyme. Advantages include its ready availability and low cost. Disadvantages include the need for compounding by a pharmacist for intravenous use, the need for frequent monitoring of serum concentrations, its effect in blunting the sensorium, and the need for hospitalization in the intensive care unit.

Fomepizole (or 4-methylpyrazole) is a strong inhibitor of alcohol dehydrogenase (fomepizole has an affinity for alcohol dehydrogenase 8000 times that of ethanol) that received FDA approval for the treatment of ethylene glycol and methanol poisoning in 1997 and 2000, respectively, but it is not approved for the treatment of the other toxic alcohol poisonings. Fomepizole is effective at low concentrations, has minimal side effects, and does not require monitoring in an intensive care unit. For patients who are not undergoing dialysis, the loading dose is 15 mg per kilogram every 12 hours. The drug may be removed by dialysis, and therefore giving it immediately after dialysis is recommended. The package insert gives a more detailed schedule for patients who receive treatment with both fomepizole and hemodialysis (see Supplementary Appendix 1).

In the United States, fomepizole is frequently used to treat methanol and ethylene glycol poisonings. In 2012 and 2015, fomepizole was used in 90 to 94% of cases, and ethanol was used in 5 to 6% of cases. Outside the United States, fomepizole is less readily available and ethanol is used more frequently. Oral ethanol is effective when intravenous ethanol is not available.

<table>
<thead>
<tr>
<th>Test</th>
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<tr>
<td><strong>Wood’s lamp to detect urine fluorescence</strong></td>
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</table>
| Detects fluorescein in antifreeze (ethylene glycol); false positives and false negatives occur frequently, which makes the test unreliable
|
| **Alco-Screen (AlcoPro), a reagent strip containing alcohol oxidase** |
| Ethanol and methanol are detected at low serum concentrations (5 mg/dl), but ethylene glycol is detected only at high concentrations (>300 mg/dl)
|
| **Portable dry strip with enzyme formate dehydrogenase and formazan dye** |
| Useful in the diagnosis of methanol poisoning; serum methanol concentration is indirectly determined from assessment of serum formate concentration; positive results are detected by visual inspection and by means of a photometer
|
| **Modified rapid veterinary assay for the detection of ethylene glycol (Catachem)** |
| An ethylene glycol assay that has been modified for use with automated clinical analyzers; good correlation with ethylene glycol concentrations measured by means of gas chromatography; also detects propylene glycol, so it could be used in suspected cases of this poisoning
|
| **Liquid-based tests that use the enzymes alcohol oxidase or alcohol dehydrogenase or the oxidizing agents sodium periodate or potassium permanganate to detect toxic alcohols in saliva** |
| Tests performed with the use of saliva that has been spiked with various toxic alcohols; detects concentrations of alcohols as low as 5 mg per deciliter; requires confirmation of accuracy with the saliva from actual patients
|
| **Gas or liquid chromatography with flame-ionization detection** |
| Most precise method to detect toxic alcohols; laborious and expensive; not available in most clinical laboratories
|

* To convert the values for ethanol, methanol, and ethylene glycol to millimoles per liter, multiply by 0.2171 for ethanol, by 0.3121 for methanol, and by 0.1611 for ethylene glycol.
Figure 2. Algorithm for the Diagnosis and Treatment of Methanol, Ethylene Glycol, and Isopropanol Intoxications.

This algorithm provides an approach to the diagnosis and treatment of the three most common poisonings. A similar approach might be useful for diethylene glycol poisoning, although this poisoning is rare. One criterion for dialysis (serum ethylene glycol concentration, >300 mg per deciliter after antidote administration) reflects the practice of the second author. To convert the values for methanol, ethylene glycol, and isopropanol to millimoles per liter, multiply by 0.3121 for methanol, by 0.1611 for ethylene glycol, and by 0.1664 for isopropanol.
The toxic alcohols and their metabolites are small and water soluble and are removed during hemodialysis. Guidelines for the use of hemodialysis in the treatment of methanol or ethylene glycol intoxication from the AACT and for treatment of methanol poisoning from the Extracorporeal Treatments in Poisoning Workgroup (EXTRIP) are provided in Table S1 in Supplementary Appendix 1. In general, both guidelines recommend dialysis with severe metabolic acidosis, serum methanol and ethylene glycol concentrations higher than 50 mg per deciliter (methanol, 16 mmol per liter; ethylene glycol, 8 mmol per liter), deteriorating vital signs despite supportive care, and problems with vision (associated with methanol poisoning) or acute kidney injury. Intermittent hemodialysis (with a large-surface-area dialyzer and high-flux membrane) removes toxic alcohols more rapidly than continuous renal replacement therapy.

Treatment of methanol or ethylene glycol intoxication with fomepizole alone (without hemodialysis) with no adverse consequences has been reported. However, fomepizole prolongs the elimination half-lives of methanol and ethylene glycol to as high as 71 hours and 16 hours, respectively, as compared with 2.5 and 2.7 hours, respectively, with dialysis. A longer duration of exposure increases days of hospitalization and cost — reasons that are given by some experts to support the inclusion of hemodialysis. The comparative costs of the two treatments will depend on several factors, including exposure dose, relative costs of the drug, cost of dialysis, and room costs, and should be factored in the decision about therapy.

Treatment in children is similar to that in adults. Further examination of the value and limitations of hemodialysis with or without an alcohol dehydrogenase inhibitor in the treatment of these intoxications is warranted. An interactive program to predict the duration of dialysis required to lead to reductions in the parent alcohol and metabolites to safe levels is shown in Supplementary Appendix 2, available at NEJM.org. Body redistribution of the alcohol, metabolites, or both might require repeat dialysis.

In methanol poisoning, 1 mg per kilogram of folic acid every 4 to 6 hours promotes the conversion of formic acid to carbon dioxide and water. In ethylene glycol poisoning, pyridoxine and thiamine promote the metabolism of glycolic acid to less toxic compounds.

Diethylene Glycol
Several experts recommend the use of alcohol dehydrogenase inhibitors in the treatment of diethylene glycol poisoning. Fomepizole alone has been successful, but because acute kidney injury is common, it seems reasonable to treat patients with both fomepizole and hemodialysis.

Isopropanol
Supportive measures are often sufficient, but hemodialysis may be necessary if the serum isopropanol concentration is 500 mg per deciliter (83 mmol per liter) or more or if hypotension or lactic acidosis is present. Alcohol dehydrogenase inhibitors slow the removal of isopropanol and should not be used.

Propylene glycol
In most cases, the elevated serum osmolality resolves with discontinuation of the drug containing propylene glycol. There is no consensus regarding the use of fomepizole, but if lactic acidosis develops, hemodialysis has been recommended.

Monitoring of Patients
Patients with severe poisoning or hemodynamic instability or those who are receiving ethanol therapy warrant care in an intensive care unit, but patients with less severe poisoning or hemodynamic stability or those who are receiving fomepizole therapy can safely be cared for outside the intensive care unit. Measurements of acid–base variables, electrolytes, renal function, and serum osmolality are necessary to assess the response to therapy. Measurement of serum concentrations of the toxic alcohols would be ideal to monitor treatment; however, obtaining serum concentrations in a timely fashion is not often feasible. In their absence, the serum concentration of the toxic alcohol can be estimated from the osmolar gap. Therapy should continue until the serum concentration of ethylene glycol or methanol falls below 20 to 30 mg per deciliter.
Methanol, ethylene glycol, and diethylene glycol poisoning can cause severe cellular dysfunction and high mortality if not recognized and treated quickly. Isopropanol frequently causes medical complications but has a lower risk of death. A high anion-gap metabolic acidosis, an increased serum osmolar gap, or both can suggest that one of the toxic alcohols is present in the blood, but these abnormal laboratory results are not always present. One of the poisonings should be strongly suspected in persons with the clinical findings described previously, in all obtunded patients, or in those with an unexplained high osmolar gap, high anion-gap metabolic acidosis, or both. Definitive tests such as high-pressure liquid chromatography are not always available, even in developed countries but especially in undeveloped countries. Therefore, there is an unmet need for tests that are accurate and can be completed rapidly.

Treatment with alcohol dehydrogenase inhibitors and the use of dialysis are effective, but both methods are not always available. Also, there is no consensus on when one or both methods should be used. Despite much progress in our understanding of the pathogenesis of reactions to these toxic alcohols and despite the development of effective treatments, much remains to be done to eliminate the severe clinical disturbances that result from exposure to these substances. No potential conflict of interest relevant to this article was reported.

Disclosure forms provided by the authors are available with the full text of this article at NEJM.org.

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