Emerging and Underrecognized Complications of Illicit Drug Use

Alysse G. Wurcel, Elisabeth A. Merchant, Roger P. Clark, and David R. Stone

Illicit drug use can result in a wide range of medical complications. As the availability, synthesis, and popularity of illicit drugs evolve over time, new syndromes associated with their use may mimic infections. Some of these symptoms are anticipated drug effects, and others are complications of adulterants mixed with drugs or complications from the method of using drugs. Some illicit drugs are associated with rare infections, which can be difficult to diagnose with standard microbiological techniques. The goal of this review is to orient a wide range of clinicians—including general practitioners, emergency medicine providers, and infectious diseases specialists—to complications of illicit drug use that may be underrecognized. Improving awareness of infectious and noninfectious complications of illicit drug can expedite diagnosis and medical treatment of persons who use drugs and facilitate targeted harm reduction counseling to prevent future complications.

Keywords. illicit drug use; injection drug use; skin and soft-tissue infections; altered mental status; substance abuse.

In 2013, an estimated 24.6 million Americans aged ≥12 years (9.4% of the total population) were active illicit drug users [1]. Rates of morbidity and mortality associated with illicit drug use especially from use of opiates and heroin, is increasing in the United States [2, 3]. In contrast to demographics of drug use from 10 years ago, the most dramatic increases in illicit drug use have been recorded among young white persons living in rural areas [4, 5].

The methods of illicit drug use—including but not limited to smoking, sniffing, injecting, and “skin popping”—are risk factors for infectious diseases. There are several well-known infectious complications of injection drug use, including tissue and bloodstream infections, bacterial endocarditis, human immunodeficiency virus (HIV), and viral hepatitis [6]. There are also underrecognized (and some newly emerging) complications of commonly used illicit drugs, including cocaine and marijuana, which can cause patterns of multisystem organ dysfunction similar to infectious syndromes. Furthermore, several emerging illicit drugs, including synthetic cannabinoids, cathinone derivatives, and pipеразine derivatives, are not detected with simple urine or blood toxicology screens and can cause symptoms that mimic infections; synthetic cannabinoids, in particular, are increasing in popularity (38% of persons in substance abuse treatment programs report use), with major increases in hospitalizations and deaths reported in the United States since 2011 [7, 8].

Diagnosis and treatment of clinical syndromes related to illicit drug use are hampered by lack of validated toxicology screens, in addition to reluctance of persons who use drugs (PWUD) to report their drug use history and lack of knowledge among providers that these new illicit drugs and syndromes exist. The goal of this review is to give clinicians a foundation of knowledge to inform future clinical assessments and decision making in PWUD who present for medical care. We will discuss symptoms associated with emerging illicit drugs and underrecognized complications of commonly used drugs, and we will review these underrecognized complications in the context of the infectious disease syndromes they may mimic, including skin/soft-tissue infections,
pneumonia, meningitis, and gastroenteritis. Through out this paper we use the term illicit drug use because it is widely accepted and commonly used in the litera ture, but it does have negative connotations; providers should be aware of this and try to decrease stigmatization in patient interactions. As defined by the federal agency SAMHSA [Substance Abuse and Mental Health Services Administration], illicit drug use is the use of illegal drugs or the misuse or abuse of prescription drugs; drugs include marijuana/hashish, cocaine/crack, heroin, hallucinogens, inhalants, and prescription—type psychotherapeutics—pain relievers, tranquilizers, stimulants, and sedatives-used nonmedically.

METHODS

Literature Search
We developed a search strategy for Medical Subject Heading (MeSH) terms and free text key words relevant to illicit drug use, including the following terms: drug abuse, illicit drug use, endocarditis, polysubstance abuse, emerging drugs of abuse, rare complications, toxicities, novel psychoactive substances, bath salts, synthetic cathinones, synthetic cannabinoids, jelling-up, crush-resistant, krokodil, cocaine, heroin, marijuana, methamphetamine, adulterants, levamisole, black tar heroin, anthrax, and tetanus. We searched the PubMed database for all relevant articles, published until June 2015 containing individual key words and combinations of above key words.

Emerging Illicit Drugs and Adulterants
Before discussing the presenting symptoms that may be seen in PWUD, it is important to review some of the emerging illicit drugs and adulterants. Details about their history, methods of use, mechanisms of action, desired effects, and adverse effects are provided in Table 1.

Complications of Illicit Drug Use
Table 2 summarizes the complications of illicit drug use that can mimic infectious syndromes.

Skin and Soft-Tissue Infections and Manifestations
Injection drug use is a common cause of skin and soft-tissue infections with both gram-positive skin flora and gram-negative enteric flora, dependent on the materials used to process and use the drug. Gram-positive infections from Staphylococcus aureus and Streptococcus pyogenes are the most common and result from introduction of skin flora past the epidermis into deeper tissues (discussed below in Sepsis-Like Syndromes). The risk of infection increases with larger needles, which can sometimes be used if the drug is too thick and clots in the needle (“jelling up the dope”) [43]. Any clinical encounter with a PWUD should incorporate a review of harm reduction techniques, including cleaning the skin with alcohol before injecting and using bleach and water to disinfect syringes.

Two less common causes of severe skin-based infection are clostridial infections and anthrax. Clostridial infections can progress rapidly from cellulitis to necrotizing fasciitis or gas gangrene. Several outbreaks have been documented in the Western United States among black tar heroin users who inject using “skin popping,” the practice of injecting drugs subcutaneously, intradermally, or (rarely) intramuscularly [44]. Clostridial spores are introduced when black tar heroin is cut with brown material (eg, shoe polish, wood pulp, coffee grounds, or dirt) to increase bulk. Typically heroin is heated and dissolved in water before use, but clostridial spores can survive this process and even begin germination [6]. The most commonly involved clostridial species is Clostridium botulinum (also discussed below in the section below on neurological manifestations), but gangrene has also been reported from Clostridium perfringens, Clostridium sordellii, and Clostridium novyi [36, 37, 44–48]. Biopsy of the infected area can reveal black tar deposits in the tissue (Figure 1).

Injectional anthrax infection was first described in Norway in 2000 in a group of PWUD who injected heroin subcutaneously. There have been 2 subsequent outbreaks, one in 2009 in the United Kingdom and another in 2012 in Northern Europe, Germany, and the United Kingdom [49, 50]. Although injectional anthrax can progress to secondary bacteraemia, the initial presentation is a unique clinical syndrome lacking the classic black-crusted painless eschar of traditional cutaneous anthrax. Presenting symptoms are significant edema, necrosis, and blistering, potentially leading to compartment syndrome or necrotizing fasciitis. Imaging and soft-tissue exploration show edematous muscle and subcutaneous tissue with or without necrosis and without collections or abscesses. Other symptoms include nausea, vomiting, abdominal pain, and variable neurological syndromes, including meningitis and intracranial hemorrhage [49].

Two emerging illicit drugs should be considered when evaluating a PWUD with skin or soft-tissue symptoms. The first is the semisynthetic opioid derivative desomorphine, colloquially known as “krokodil.” The name derives from the green scaly appearance of the skin seen after injection, often accompanied by ulceration of skin, muscle, and cartilage. Tissue damage results from contamination with substances such as paint thinner, lighter fluid, gasoline, lead, zinc, and hydrochloric acid, which are used in the preparation of this drug. Damaged skin can become gangrenous, leading to sloughing of tissue down to the bone, often requiring extensive amputation [14]. Use of krokodil was first reported in Russia in 2003, and by 2013 cases were reported in Europe and the United States (Arizona, Utah, Oklahoma, and Illinois) [15].

The second drug that should be considered is levamisole, an increasingly common adulterant detected in the majority of cocaine seized in the United States and worldwide [18, 19].
<table>
<thead>
<tr>
<th>Name</th>
<th>History</th>
<th>Street Names</th>
<th>Method of Use</th>
<th>Mechanism of Actions</th>
<th>Desired Effects</th>
<th>Adverse Effects/Reactions</th>
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<tr>
<td>Cathinones: methcathinone, mephedrone, methylenedioxy-pyrovalerone, methylone, ethylone, butylone, methedrone [9–11]</td>
<td>Identified in 1975 as the principal psychoactive component of Khat (Catha edulis) leaves; first synthesized in 1928, explored for their stimulant, antidepressant and appetite suppressant properties</td>
<td>Khat, bath salts, ivory wave, white rush, white lightning, white dove, meow, meow, M-CAT, bubbles, cloud 9, explosion, impact, energy 1, bloom, blue silk</td>
<td>Oral; intranasal; can be inhaled, intravenous, intramuscular, or rectal</td>
<td>Similar to amphetamines and catecholamines; inhibition of vesicular monoamine transporters for serotonin, dopamine, and norepinephrine; release of intracellular neurotransmitter stores; inhibition of MAO</td>
<td>Stimulation, euphoria, friendliness, sexual arousal, and perceptual disturbances</td>
<td>Anxiety, paranoia, hallucinations, psychosis, aggression, impaired working memory, bruxism, seizures, hypertension, tachycardia, coronary vasospasm, dysrhythmia, rhabdomyolysis, acute kidney injury, hyponatremia, hyperkalemia, metabolic acidosis, hyperthermia serotonin syndrome, disseminated intravascular coagulation, multiorgan failure, death</td>
</tr>
<tr>
<td>Cannabinoids [9, 11–13]</td>
<td>Derivatives of THC, first synthesized in the 1940s and studied for treatment of pain, anxiety and nausea</td>
<td>K2, spice, happy tiger incense, smoke, aroma, Aztec fire, black mamba, blueberry posh, Bombay blue, blaze, bliss, eclipse, krypton, Mr Smiley, Yucatan fire, Zohar, sensation vanilla</td>
<td>Ingested orally or smoked</td>
<td>Alteration in sensory perception and processing of stimuli in the hippocampus, amygdala, and prefrontal cortex via reduction of GABA release and increase in dopamine and glutamate release</td>
<td>Euphoria, relaxation, disinhibition, altered perception and consciousness; similar to marijuana</td>
<td>Anxiety, confusion, agitation, mood dysregulation, paranoia, psychosis (including long term), perceptual disturbances, suicidal ideation, sedation, memory impairment, tremors, seizures, nausea, vomiting, diaphoresis, xerostomia, mydriasis, tachycardia, hypertension, chest pain, acute MI, acute kidney injury, respiratory depression, tachyphylaxis, death</td>
</tr>
<tr>
<td>Krokdil (desomorphine) [14, 15]</td>
<td>Introduced as an analgesic in the 1940s, withdrawn in 1952 over concerns about addictive potential</td>
<td>Krok, Russian magic</td>
<td>Injected subcutaneously or intravenously; also ingested orally</td>
<td>Potent mu-opioid agonist</td>
<td>Euphoria, relaxation, analgesia</td>
<td>Sedation, miosis, flushing, paresthesia, constipation, urinary retention, nausea/vomiting, allergic reactions, seizures, respiratory depression, pneumonia, septicemia, coronary artery burst, meningitis, ulcers and gangrene, rotting gums and tooth loss, bone infection</td>
</tr>
<tr>
<td>Piperazine derivatives [11, 16]</td>
<td>Initially developed as anthelminthics, also explored for their antidepressant properties</td>
<td>Legal ecstasy, party pills</td>
<td>Orally ingested</td>
<td>Central serotonergic effects; increased release and reuptake inhibition of dopamine, serotonin, and norepinephrine</td>
<td>Hallucinations, stimulation, euphoria</td>
<td>Hallucinations, confusion, anxiety, insomnia, dizziness, headaches, seizures, nausea/vomiting, shortness of breath, palpitations, sinus tachycardia, QT prolongation, seizures, hyponatremia, serotonin syndrome, hyperthermia, rhabdomyolysis, renal failure, disseminated intravascular coagulation</td>
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Levamisole is a popular adulterant because it is inexpensive, similar in appearance to cocaine, and enhances cocaine’s effects. Use of levamisole-adulterated cocaine, including snorting, injecting, or smoking, may lead to several complications related to an induced vasculopathy. Dermatological manifestations may include fixed drug eruptions, lichen planus, ulceration, nodules, erythema nodosum leprosum, nonspecific maculopapular rashes, and hemorrhagic bullae [18, 20]. The characteristic lesions are rapidly progressive cutaneous ecchymoses, raised purpura, and bullae resulting in a distinct stellate lesion with erythematous borders and a necrotic center, which have a predilection for the ears and cheeks (Figures 2 and 3) [18, 20, 21]. In addition to the dermatological manifestations, agranulocytosis, leukoencephalopathy, and acute kidney injury (suspected to be from renal tubular necrosis) can also be caused by levamisole [21]. Most patients with complications from levamisole have autoantibodies, including but not limited to pANCA, cANCA, antiphospholipid, antinuclear antibody, and anti–double-stranded DNA [20].

Vasculitis is another common skin and soft-tissue manifestation that can be related to illicit drug use. Cocaine-induced midline destructive lesions, such as nasal septal and palatal perforations, are well-recognized effects of snorting cocaine that can mimic granulomatosis with polyangiitis, characterized by ANCA positivity (specifically, anti–human neutrophil elastase) [27]. Cocaine has also been reported to cause urticarial vasculitis, Churg-Strauss vasculitis, necrotizing granulomatous vasculitis, palpable purpura, Buerger disease, and scleroderma, which are either induced or unmasked by unknown mechanisms [26]. A rare complication of marijuana use is local arteritis, most often affecting the lower limbs, and may present as Raynaud phenomenon or as claudication followed by development of ulcers or gangrene, resembling Buerger disease [51]. The proposed mechanisms for these manifestations are related to vasoconstriction or actions of a contaminant [26].

Clinical Mimics of Pneumonia
PWUD presenting with shortness of breath or pleuritic chest pain may have complications of drug use other than pneumonia. One common syndrome called “crack lung” is a characteristic acute pulmonary manifestation that occurs after smoking crack cocaine. It is characterized by fever, dyspnea, pleuritic chest pain, and hemoptysis that can progress to hypoxemia and respiratory failure. Symptoms usually occur within 48 hours of crack cocaine use [27, 28]. Imaging can show diffuse bilateral alveolar infiltrates, interlobular septal thickening, peribronchial nodules, ground glass opacities, or nonspecific consolidation. Bronchoalveolar lavage is similarly nonspecific [27, 28]. Inhalation of cocaine can also be associated with a chronic, often subclinical, diffuse alveolar hemorrhage, for which proposed mechanisms include vasoconstriction, pulmonary
<table>
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<tr>
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<th>Clinical Features</th>
<th>Laboratory and Imaging Findings</th>
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<tr>
<td><strong>Skin and soft-tissue syndromes</strong></td>
<td></td>
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<tr>
<td>Skin ulceration</td>
<td>Krokodil [14]</td>
<td>Rash with scaly, green appearance, ulcerations, destruction of skin, muscle, cartilage, gangrene and skin sloughing</td>
<td>Nonspecific</td>
</tr>
<tr>
<td><strong>Vasculitis</strong></td>
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<tr>
<td>Levasimole [18, 20, 21]</td>
<td></td>
<td>Characteristic lesions are stellate lesion with erythematous borders and a necrotic center frequent on ears and cheeks</td>
<td>Agranulocytosis; positive titers of pANCA, cANCA, anti-cardiolipin, ANA, and anti-dsDNA; evidence of acute kidney injury in acute tubular necrosis pattern</td>
</tr>
<tr>
<td>Cocaine [26]</td>
<td></td>
<td>Urticarial vasculitis, Churg-Strauss vasculitis, necrotizing granulomatous vasculitis, palpable purpura, Buerger disease</td>
<td>Nonspecific</td>
</tr>
<tr>
<td><strong>Pneumonia and other pulmonary syndromes</strong></td>
<td></td>
<td></td>
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<tr>
<td>Crack lung [27, 28]</td>
<td>Cocaine</td>
<td>Within 48 h of cocaine smoking: fever, dyspnea, pleuritic chest pain, hemoptysis, hypoxemia, respiratory failure</td>
<td>Imaging shows diffuse alveolar infiltrates, interlobular septal thickening, peribronchial nodules, ground glass opacities, nonspecific consolidation BAL with eosinophilia, hemosiderin-laden macrophages, IgE deposition, hyaline membrane formation peripheral eosinophilia</td>
</tr>
<tr>
<td>Subclinical alveolar hemorrhage [27, 28]</td>
<td>Cocaine (chronic use)</td>
<td>Largely asymptomatic; possible hemoptysis or nonspecific pulmonary symptoms</td>
<td>Diffuse alveolar hemorrhage with hemosiderin-laden macrophages in BAL fluid</td>
</tr>
<tr>
<td>Bronchospasm [27]</td>
<td>Cocaine</td>
<td>Bronchospasm with presentation resembling asthma; can be severe</td>
<td>Nonspecific</td>
</tr>
<tr>
<td><strong>Pulmonary edema</strong></td>
<td>Cocaine, amphetamine-like substances, heroin</td>
<td>Shortness of breath, cough</td>
<td>Radiographic evidence of bilateral or unilateral pulmonary edema; BAL fluid may reveal high protein concentrations</td>
</tr>
<tr>
<td>Pulmonary granulomatosis [27, 29]</td>
<td>Various drugs (intravenous drug injection or inhalation)</td>
<td>Pulmonary hypertension; interstitial fibrosis</td>
<td>Biopsy shows perivascular granulomatosis</td>
</tr>
<tr>
<td><strong>Sepsis-like syndromes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cotton fever [30–32]</td>
<td>Heroin</td>
<td>Fever, other typical sepsis features typically occurs 10–30 min after injection; usually self-limited</td>
<td>Leukocytosis, other sepsis features, negative blood cultures</td>
</tr>
<tr>
<td><strong>Sympathomimetic effects</strong> [9–12, 33, 34]</td>
<td>MDMA, piperazine derivatives, synthetic cannabinoids, cathinone derivatives</td>
<td>Tachycardia, hyperthermia, maybe be rhabdomyolysis with acute kidney injury</td>
<td>Synthetic cannabinoids, cathinone derivatives and piperazine derivatives not detected in standard urinary toxicology screens</td>
</tr>
<tr>
<td><strong>Neurological syndromes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stroke [11, 35]</td>
<td>Cocaine, methamphetamine, MDMA, synthetic cannabinoids, cathinone derivatives</td>
<td>Abrupt onset of focal neurological symptoms, possibly accompanied by change in mental status or loss of consciousness</td>
<td>Evidence of ischemic or hemorrhagic stroke on brain imaging</td>
</tr>
<tr>
<td>Cranial nerve palsies [36, 37]</td>
<td>Black tar heroin contaminated with botulism and tetanus</td>
<td>Cranial neuropathy (including diplopia, ophthalmoplegia, ptosis, and facial nerve palsy); may be accompanied by cellulitis, with possible necrotizing fasciitis and gangrene</td>
<td>Nonspecific</td>
</tr>
<tr>
<td>Seizures [9–11, 33]</td>
<td>Synthetic cannabinoids, cathinone derivatives, piperazine derivatives</td>
<td>Seizures; possibly status epilepticus</td>
<td>Synthetic cannabinoids and cathinone derivatives not detected in standard urinary toxicology screens</td>
</tr>
</tbody>
</table>
Bronchospasm is a separate and distinct entity from crack lunch which has been reported with use of freebase cocaine (cocaine heated to remove impurities) in both known asthmatics and nonasthmatics. There have also been several reported cases of new-onset severe acute asthma within a few months after the onset of heroin use [29].

In addition to crack lung, other complications of illicit drug use may present with respiratory symptoms. Intravenous stimulant use is associated with pulmonary foreign body granulomas, pulmonary artery muscular hypertrophy, and fibrous intimal proliferation and has been linked to idiopathic pulmonary hypertension [27, 52, 53]. Pulmonary vascular granulomatosis is caused by embolization or inhalation of the drug itself.

Table 2 continued.

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Agent</th>
<th>Clinical Features</th>
<th>Laboratory and Imaging Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leukoencephalopathy</td>
<td>Heroin [38]</td>
<td>Altered mental status, restlessness, apathy, cerebellar speech disturbance, ataxia, hyperactive reflexes, spasticity, tremor, choreoathetoid movements, hypotonia, areflexia, respiratory failure</td>
<td>Mild CSF pleocytosis; cerebellar signal abnormalities on MR images</td>
</tr>
<tr>
<td>Heroin, synthetic cannabinoids</td>
<td>Levamisole-adulterated cocaine [22]</td>
<td>Confusion, altered mental status, language impairment, visual changes, focal neurological deficits, white matter lesions on MR images, primarily affecting frontal lobes</td>
<td>Marijuana but not synthetic cannabinoids at standard toxicology screening</td>
</tr>
</tbody>
</table>

Abbreviations: ANA, antinuclear antibody; BAL, bronchoalveolar lavage; cANCA, cytoplasmic antineutrophil cytoplasmic antibodies; CSF, cerebrospinal fluid; IgE, immunoglobulin E; MDMA, 3,4-methylenedioxymethamphetamine; MR, magnetic resonance; pANCA, perinuclear anti-neutrophil cytoplasmic antibodies.
contaminants, or insoluble filler materials (such as silica, cellulose, and talc) into the lungs and other organs (Figure 4). Pulmonary vascular granulomatosis can progress to a pneumoconiosis-like interstitial fibrosis or pulmonary hypertension, both of which can be severe or fatal [27]. Pulmonary edema has been reported with use of stimulants (eg, cocaine and amphetamines), as well as heroin [27, 28]. Barotrauma-related injuries, including pneumomediastinum, pneumothorax, and pneumopericardium have been reported after the use of MDMA (3,4-methylenedioxy-methamphetamine), heroin, cocaine, and marijuana [28]. Direct puncture of the lung while performing "pocket shots" (ie, injection directly into the jugular or subclavian veins) can also lead to direct lung injury [27].

**Sepsis-Like Syndromes**

The first concern for PWUD presenting with a sepsis-like syndrome should be bacteremia. The most common routes of infection are direct injection of bacteria via dirty and/or shared equipment and local spread from a cellulitis or soft-tissue infection. The most common causes of bacteremia in PWUD are *S. aureus* and *S. pyogenes*, although other types of oral and gastrointestinal flora (including *Candida* species) from licking needles or using dirty water can also cause a septic presentation [54–57]. PWUD can also present with bloodstream infections caused by less common organisms, such as *Lactobacillus*, *Corynebacteria*, and *Bacillus* (very rarely, *Bacillus anthracis*), which should be considered as well as the more common bacterial causes and not discounted as a possible contaminant [58–62]. We have recently seen several cases of nontuberculous atypical mycobacterial bacteremias clustered among intravenous drug users hospitalized for drug-related bacterial infections and receiving antibiotics through peripherally inserted central catheters. These patients had fever and tachycardia, and Gram stains of their blood showed gram-positive rods that were originally mistaken for contaminant until sent to specialty laboratories for identification.

If a person who has a history of drug use presents with a sepsis-like syndrome with no growth in blood cultures, there are other illicit drug complications to consider. “Cotton fever” is a syndrome first characterized in 1975, which includes fevers and leukocytosis in the absence of positive blood cultures [30]. It characteristically occurs 10 to 30 minutes after injection of heroin. Its cause is related to the practice of using cotton to filter out particulate matter when drawing up liquid heroin and then heating the used cotton to extract residual drops of heroin that are injected, a practice colloquially referred to as “shooting the cotton” [31, 32]. Research has shown that a gram-negative bacterium, *Enterobacter agglomerans*, colonizes the cotton plant and produces an endotoxin, which may be responsible for the fevers [63]. The syndrome is usually self-limited, yet the symptoms may be worrisome and lead PWUD to seek emergency care [32, 64].

Tachycardia and hyperthermia are common adverse effects of MDMA (also known as Ecstasy or Molly). Several of the emerging drugs, including synthetic cannabinoids, cathinone derivatives, and piperazine derivatives, can also cause hyperthermia and tachycardia and have been further linked to cases of serotonin syndrome. Urine toxicology assays used at most hospitals will not detect these drugs, so if there is a concern about illicit drug use, it is important to specifically ask the patient about these substances [9, 10, 33]. One clue that a person with hyperthermia and tachycardia might be using these drugs is acute kidney injury secondary to rhabdomyolysis. In a similar manner to cocaine, MDMA, cathinone derivatives, synthetic cannabinoids, and piperazine derivatives may cause
rhabdomyolysis and acute kidney injury by increasing skeletal muscle use and metabolic demand while also causing vasoconstriction, resulting in hypoperfusion and poor heat dissipation [12, 34]. Tachycardia, agitation and chest pain can also be seen in persons exposed to clenbuterol ($\beta_2$-agonist similar in chemical structure ephedrine), which is used illicitly by bodybuilders to increase muscle mass and lose weight and also used as a bulking agent for heroin [23, 24].

Clinical Mimics of Meningitis, Encephalitis, and Other Neurological Syndromes

Altered mental status is common in PWUD presenting to the emergency room and often leads to concern about possible meningitis or encephalitis. In addition to central nervous infections, healthcare providers should consider other causes of altered mental status, including stroke, movement disorders, and seizures caused by illicit drug use or withdrawal syndromes.

Cocaine and methamphetamine use are well-known causes of cerebral vasospasm and stroke, but other illicit drugs, such as marijuana, synthetic cathinones, and cathinone derivatives, have also been linked with cerebrovascular complications. Ischemic and hemorrhagic strokes have also been observed with MDMA use, possibly related to similar sympathomimetic mechanisms or high fever triggering disseminated intravascular coagulation [35]. There are also a few distinct neurological syndromes that present with specific nerve involvement and could be mistaken for strokes. Although decreasing in incidence, clostridial infection from Clostridium tetani and C. botulinum should be considered in PWUD presenting with cranial nerve involvement or muscle weakness [44, 65, 66]. There have been several outbreaks of black tar heroin–related tetanus and botulism in California, China, Saudi Arabia, and Western Europe [66–69]. Withdrawal from heroin can cause acute esotropia, or strabismus, which typically resolves after the withdrawal period is over [70]. In persons who injected crush-resistant opiates, 2 cases have been reported of opiate-associated hearing loss, thought to be secondary to cochlear ischemia [71].

Common causes of seizures include benzodiazepine and alcohol withdrawal. Less recognized causes of seizures include sympathomimetic, drugs such as synthetic cannabinoids, cathinone derivatives, and piperazine derivatives [11]. Between 1 January and 22 April 2015, the American Association for Poison Control Centers released surveillance data showing dramatic increases (229%) in the number of calls related to exposure to synthetic cannabinoids compared with the same time frame in the previous year [72,73]. Several states health departments, including those in New York, Colorado, and Georgia, have issued alerts about symptoms that can result from ingestion, including severe agitation, hallucinations, seizures and tremors [74–76].

Both heroin and cocaine can be associated with leukoencephalopathy, although the imaging findings and clinical presentation may be slightly different. Leukoencephalopathy associated with heroin use is characterized by demyelination in the cerebellum or limbic system and white matter edema and is suspected to be due to oligodendrocyte mitochondrial dysfunction and apoptosis. The clinical presentation of heroin-associated leukoencephalopathy, also known as “chasing the dragon” is characterized by restlessness, apathy, cerebellar speech disturbances, ataxia, hyperactive reflexes, spasticity, tremors, and choreoathetoid movements, eventually leading to hypotonia, areflexia, and possible respiratory failure. This clinical picture is associated with a mild cerebrospinal fluid pleocytosis and cerebellar signal abnormalities at magnetic resonance imaging (Figure 5) [38]. Leukoencephalopathy from cocaine—thought to be mostly related to levamisole adulteration—has magnetic resonance imaging findings of spongiform leukoencephalopathy with white matter lesions similar to toxic-metabolic injury, primarily affecting the frontal lobe [22].

Clinical Mimics of Gastroenteritis

Illicit drug use can cause a variety of abdominal symptoms. Withdrawal from opiates may present like an infectious gastroenteritis with diaphoresis, nausea, vomiting, and diarrhea. Acute viral hepatitis B, C, or D can present with nausea, vomiting, and abdominal pain, and in the early stages patients may be seronegative despite high viral loads. “Body packing” or concealing illicit drugs...
such as narcotics or cocaine by swallowing them in bags can lead to intestinal obstruction in addition to acute overdose symptoms if the bags break. Both methamphetamine and cocaine are known to cause intestinal ischemia with subsequent infarct, necrosis, bleeding, or perforation as a result of splanchnic vasoconstriction and enhanced thrombosis [39, 40]. Chronic use of marijuana or synthetic cannabinoids is associated with cannabinoid hyperemesis syndrome, which presents with periods of nausea, vomiting, and abdominal pain, often associated with compulsive bathing or showering with hot water [41, 42].

CONCLUSIONS

It behooves all clinicians to stay up to date on complications of illicit drug use. Knowledge of emerging trends in illicit drug use can facilitate specific questions to assess both infectious and noninfectious risks and create a culturally sensitive and nonjudgmental atmosphere. In addition to addressing acute or subacute complications of drug use, encounters with PWUD represent a prime opportunity for counseling patients regarding the risks of continued drug use and making efforts to link them to treatment programs.

Notes

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References


