# Drug Testing in Pain Management and Substance Use Disorder Treatment

## Medical Policy

### Title:

**Drug Testing in Pain Management and Substance Use Disorder Treatment**

### Professional

Original Effective Date: May 21, 2015  
Revision Date(s): May 21, 2015; July 25, 2016; January 1, 2017; October 1, 2017; February 15, 2018; July 1, 2018; January 1, 2019  
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<table>
<thead>
<tr>
<th>Populations</th>
<th>Interventions</th>
<th>Comparators</th>
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</tr>
</thead>
</table>
| Individuals:  
- With chronic pain treated with opioids | Interventions of interest are:  
- Urine drug testing | Comparators of interest are:  
- No urine drug testing | Relevant outcomes include:  
- Test accuracy  
- Test validity  
- Health status measures  
- Resource utilization |
| Individuals:  
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| Individuals:  
- With chronic pain treated with opioids | Interventions of interest are:  
- Oral fluid or hair drug testing | Comparators of interest are:  
- No testing  
- Urine drug testing | Relevant outcomes include:  
- Test accuracy  
- Test validity  
- Health status measures  
- Resource utilization |
DESCRIPTION

Summary
Patients in pain management programs and substance use disorder treatment may misuse prescribed opioids and/or may use nonprescribed drugs. Thus, these patients are often assessed before treatment and monitored while receiving treatment. Urine drug testing (UDT) can be part of this monitoring strategy; it is most often used as part of a multifaceted intervention that includes other components, such as patient contracts.

Objective
The objective of this evidence review is to determine whether urine, oral fluid, and/or hair testing for drug use improve net health outcomes in individuals with chronic pain receiving opioid treatment or with a drug addiction and in substance use disorder treatment.

Background

Opioids
According to a 2012 evidence assessment by the American Society of Interventional Pain Physicians, approximately one-third of chronic pain patients do not use opioids as prescribed or may abuse them. In 2016, the International Narcotics Control Board reported that between 1999 and 2010, the number of deaths related to the use of prescription opioid painkillers increased 5-fold among United States women and increased by a factor of 3.6 among United States men. Additionally, studies have found that a substantial proportion of chronic pain patients inaccurately report nonadherence to prescribed medications and use of illicit drugs.

Monitoring Strategies
Various strategies are available to monitor pain management and substance use disorder treatment patients, and multicomponent interventions are often used. Many settings

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</tr>
<tr>
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<td>Interventions of interest are: • Hair drug testing</td>
<td>Comparators of interest are: • No testing • Urine drug testing</td>
<td>Relevant outcomes include: • Test accuracy • Test validity • Health status measures • Resource utilization</td>
</tr>
</tbody>
</table>
require patients to sign a contract before they are given a prescription for opioids. The contracts generally involve obtaining patients’ agreement on behaviors they will engage in during the treatment period (eg, taking medication as prescribed) and not engage in (eg, selling prescribed medication and/or obtaining additional prescriptions from other physicians).

Confirming whether patients follow these behavioral guidelines can be a challenge. Risk-assessment screening instruments, such as the Screener and Opioid Assessment for Patients with Pain, and the Opioid Risk Tool, can aid in the assessment of patients’ risk for inappropriate drug use. In addition, the presence of “aberrant behaviors” can be used as a marker for patients who are at high risk for deviating from treatment protocols. Aberrant behaviors include multiple lost prescriptions, obtaining prescriptions from other practitioners, and displaying evidence of acute intoxication during office visits.

**Testing Strategies**

Another strategy for monitoring patients is testing of biologic specimens for the presence or absence of drugs. Currently, urine is the most commonly used biologic substance. Advantages of urine drug testing (UDT) are that it is readily available and standardized techniques for detecting drugs in urine exist. Other biologic specimens (eg, blood, oral fluids, hair, sweat) can also be tested and may gain in popularity over time as techniques for collecting and analyzing these specimens become more standardized. In addition to urine testing, this review will address testing for oral fluids and hair.

**Urine Drug Testing**

There are 2 primary categories of UDT: immunotherapy and specific drug identification.

**Immunoassay Testing**

Immunoassay testing (also called presumptive testing or qualitative testing or screening) can be performed in a laboratory or at point-of-service. Immunoassay tests are based on the principle of competitive binding and use antibodies to detect a particular drug or drug metabolite in a urine sample. With competitive binding, a fixed amount of a labeled drug is added to the urine sample, and the drug or metabolite in the sample competes with the labeled drug for binding sites on the antibody. The amount of labeled antigen that binds with the antibody is inversely proportional to the amount of the drug or metabolite in the sample.

Immunoassay tests vary in the type of compounds they can detect. Some detect specific drugs and may fail to recognize similarly structured drugs within the same class. Other immunoassays identify only classes of drugs and thus results cannot be used to determine which drug a patient is taking. For example, a positive result to an opiate immunoassay can be due to morphine or hydromorphone. The degree of crossreactivity (ie, an antibody’s reactivity with a compound other than the target of the test) varies widely among immunoassays.
Immunoassay findings are generally reported qualitatively as either positive (drug level above a prespecified threshold) or negative (drug level below a prespecified threshold). Raising or lowering the threshold thus changes the proportion of positive tests. A negative test is interpreted as a level below the threshold and does not necessarily mean that the drug or metabolite is absent.

Immunoassays generally have a rapid turnaround time, to within minutes for on-site tests, and 1 to 4 hours for laboratory-based tests.4

**Specific Drug Identification**
Confirmatory tests are always performed in a laboratory. Gas chromatography/mass spectrometry (GC/MS) and liquid-chromatography/mass spectrometry (LC/MS) are considered to be the criterion standard for confirmatory testing. These techniques involve using GC or LC to separate the analytes in a specimen and for MS to identify the specific molecular structures of the drug and its metabolites. The tests are able to quantify the amount of drug or metabolite present in the urine sample. Definitive quantitative tests can be used to confirm the presence of a specific drug identified by a screening test and can identify drugs that cannot be isolated by currently available immunoassays. Results are reported as the specific levels of substances detected in the urine. GC/MS and LC/MS generally requires specification of the drug or drugs to be identified. Alternatively, “broad-spectrum screens” can be conducted. There is a several-day turnaround time for GC/MS and LC/MS testing.5

An issue with both types of UDT is the possibility of sample tampering to mask the presence of illegal drugs. A variety of products and techniques are available to patients and can be as simple as drinking a large amount of water to dilute the sample. There are also commercial dilution and cleaning products, additives, and urine substitutes. Some of these techniques can be detected by visual inspection of the sample (eg, color) or by on-site testing of sample characteristics including urine temperature, creatinine concentration, and specific gravity.

The correct interpretation of UDT results is very important. Knowledge of drug metabolites is essential for accurate interpretation. Accurate interpretation of test results also requires knowledge of the drug manufacturing process. For example, due to manufacturing impurities, a small amount of hydrocodone may be present in urine samples from patients prescribed oxycodone. Thus, it would be acceptable to detect a small amount of hydrocodone if high amounts of oxycodone were also present.

There are various approaches to incorporating UDT into pain management and substance use disorder treatment settings. Most commonly, patients undergo urine drug screening before beginning treatment to verify current drug use. Some clinicians believe that UDT should be routinely used to establish baseline information about substance use, but the optimal frequency and interval of testing remains uncertain. A universal approach to screening may uncover more inappropriate use and may reduce patients’ sense that
testing is being performed due to a lack of trust. However, routine universal screening may place an unnecessary burden on the health care system and on the doctor-patient relationship. An alternative approach is selective testing of patients who are judged to be at increased risk for drug misuse.

Existing protocols vary for the use of presumptive vs definitive tests. Some involve conducting routine confirmation of positive presumptive tests with definitive quantitative testing. Others use selective confirmation of positive presumptive tests, such as when an unexpected immunoassay result is not adequately explained by the patient. There is also a mixed approach, with routine confirmation of presumptive tests only for drugs with poor-performing immunoassays.

Full informed consent is a requirement before UDT. Patients should be informed of the specific drug testing protocol before treatment and should provide written agreement with the plan for monitoring. As stated in a joint U.S. Veterans Affairs/Department of Defense guideline, patients’ refusal to consent to urine testing should be considered a factor in the overall assessment of patients’ ability to adhere to treatment.6

Oral Fluid Drug Testing
Oral fluid (liquid samples obtained from the oral cavity) can potentially be used to test for drug use. Oral fluid contains secretions from several different sources, including secretions from the 3 pairs of major salivary glands (parotid, sublingual, and submandibular), secretions from the minor salivary glands, oro-naso-pharyngeal secretions, and cellular debris. The mixture of fluids obtained varies depending on the collection method used (eg, spitting, suctioning, draining, or collection on some type of absorbent material). Drug concentrations can be affected by the collection method and by the use of saliva stimulation methods. Several collection devices are commercially available in the United States, and they generally involve collection on an absorbent material, such as foam pads; pads are then placed in a container with a stabilizing buffer solution. Drug concentrations may also depend on how the oral fluid is recovered from the collection device (eg, by centrifugation or by applying pressure). Drug concentrations may not reflect blood levels because of residual amounts of drug (specifically those ingested or smoked) remaining in the oral cavity after recent use.

Analysis techniques must be able to detect drugs present in low concentration and in a small volume of fluid (often <1 mL). Immunoassay techniques are available to detect drugs in oral fluid; they require a small sample volume (≈25 μL). Immunoassays tend to be relatively sensitive techniques, but they have low specificity. Confirmation analysis is generally performed using MS-based methods. In recent years, advancements have been made in MS analysis techniques, including the development of multianalyte LC-MS methods.

A practical advantage of oral fluid collection compared with urine collection is that samples can be obtained under direct supervision and without loss of privacy. It has
been used in situations where urine sampling is impractical, such as testing drivers during traffic stops. Oral fluid sampling also has the potential to be useful in the pain management or substance use disorder treatment settings, particularly when substitution or tampering with urine drug samples is suspected.

**Hair Testing**

Hair is composed of protein that traps chemicals in the blood at the time the hair develops in the follicle. Hair on the human head grows at approximately 0.5 inch per month. Thus, a 1.5-inch hair sample could be used to detect drug use during the previous 90 days. Potential advantages of hair as a drug testing source include: noninvasive collection; ease of collection, storage, and shipping; availability of samples for testing and retesting; and difficulty in tampering. Potential disadvantages include: recent drug use (ie, within past 7 days) cannot be detected; difficulty in detecting very light drug use (eg, a single episode); and drug levels can be affected by environmental exposure. In addition, variation in hair texture as well as cosmetic hair treatments can affect drug incorporation into hair and the accuracy of drug tests on hair samples. As with other types of samples, hair can be initially tested using immunoassay techniques, with confirmation by MS-based methods. Hair testing has been used in a variety of situations where detection of drug use during the previous several months is desired (eg, pre-employment screening, post-drug-treatment verification of relapse).

**Regulatory Status**

Clinical laboratories may develop and validate tests in-house and market them as a laboratory service; laboratory-developed tests must meet the general regulatory standards of the Clinical Laboratory Improvement Amendments (CLIA). Testing with GC/MS and some immunoassays are performed in laboratory settings. Laboratories that offer laboratory-developed tests must be licensed by CLIA for high-complexity testing. To date, the U.S. Food and Drug Administration (FDA) has chosen not to require any regulatory review of this test.

A CLIA waiver is available for the use of certain point-of-care immunoassays. Tests eligible for a CLIA waiver are those considered to be simple, with low risk of error and low potential for harm. FDA is tasked with approving manufacturers’ applications for test system waivers. There are commercially available CLIA-waived urine tests for drugs such as cocaine, methadone, morphine/opiates, and oxycodone. Moreover, there are commercially available hair testing tests such as Quest Diagnostics ELISA tests for amphetamines, opiates, cocaine, marijuana metabolites, and phencyclidine. In addition, Omega Laboratories (Mogadore, OH) offers hair drug screening for cocaine and cocaine metabolites.
Several oral fluid drug test collection devices have been cleared for marketing by FDA through the 510(k) process. They include:

- Intercept™ Oral Fluid Drug Testing System (OraSure Technologies, Bethlehem, PA)
- Oral-Eze Saliva Collection System (Quest Diagnostics, Madison NJ)
- Quantisal® Oral Fluid Collection Device (Alere, Waltham, MA).

In addition to the oral fluid collection devices, the FDA has cleared a number of assays for analysis of oral samples. For example, there are FDA-cleared assays for 9 drugs collected with the Intercept™ device: amphetamines, methamphetamine, cocaine/metabolite, opiates, marijuana/THC, phencyclidine, barbiturates, benzodiazepines, and methadone.

**POLICY**

A. In outpatient pain management, presumptive (ie, immunoassay) urine drug testing may be considered _medically necessary_ for:

1. Baseline screening before initiating treatment or at the time treatment is initiated, when the following conditions are met:
   a. An adequate clinical assessment of patient history and risk of substance use disorder is performed
   b. Clinicians have knowledge of test interpretation
   c. There is a plan in place regarding how to use test findings clinically

2. Subsequent monitoring of treatment at a frequency appropriate for the risk level of the individual patient (see Policy Guidelines section)

B. In outpatient substance use disorder treatment, in-office or point-of-care presumptive (ie, immunoassay) urine drug testing may be considered _medically necessary_ under the following conditions:

1. Baseline screening before initiating treatment or at the time treatment is initiated (ie, induction phase), 1 time per program entry, when the following conditions are met:
   a. An adequate clinical assessment of patient history and risk of substance use disorder is performed
   b. Clinicians have knowledge of test interpretation
   c. There is a plan in place regarding how to use test findings clinically

2. Stabilization phase – targeted weekly presumptive screening for a maximum of 4 weeks (see Policy Guidelines section)

3. Maintenance phase – targeted presumptive screening once every 1 to 3 months (see Policy Guidelines section)
C. Definitive (ie, confirmatory) urine drug testing, in outpatient pain management or substance use disorder treatment, may be considered medically necessary under the following circumstances:
1. When immunoassays for the relevant drug(s) are not commercially available
2. In specific situations for which definitive drug levels are required for clinical decision making (see Policy Guidelines section)

D. In outpatient pain management and outpatient substance use disorder treatment, urine drug testing is considered not medically necessary when the above criteria are not met including but not limited to routine presumptive or definitive urine drug testing (eg, testing at every visit, without consideration for specific patient risk factors or without consideration for whether quantitative testing is required for clinical decision making)

E. In outpatient pain management and substance use disorder treatment, oral fluid drug testing and hair drug testing are considered experimental / investigational.

Policy Guidelines
Drugs or classes of drugs may be commonly assayed first by a presumptive screening method followed by a definitive drug identification method. Presumptive methods include, but are not limited to, immunoassays (IA, EIA, ELISA, RIA, EMIT, FPIA, etc), enzymatic methods (alcohol dehydrogenase, etc), chromatographic methods without mass spectrometry (TLC, HPLC, GC, etc), or mass spectrometry without adequate drug resolution by chromatography (MS-TOF, DART, DESI, LDTD, MALDI). LC-MS, LCMS/MS, or mass spectrometry without adequate drug resolution by chromatography may also be used for presumptive testing if the chromatographic phase is not adequate to identify individual drugs and distinguish between structural isomers or isobaric compounds. All drug class immunoassays are considered presumptive, whether qualitative, semi-quantitative, or quantitative. Methods that cannot distinguish between structural isomers (such as morphine and hydromorphone or methamphetamine and phentermine) are also considered presumptive.

Pain Management
The risk level for an individual patient should include a global assessment of risk factors and monitoring for the presence of aberrant behavior. Standardized risk-assessment tools are available, such as the 5-item Opioid Risk Tool (ORT). Another screening instrument is the Screener and Opioid Assessment for Patients in Pain, a 24-item tool
Aberrant behavior is defined by one or more of the following:

a. multiple lost prescriptions
b. multiple requests for early refill
c. obtained opioids from multiple providers
d. unauthorized dose-escalation, and
e. apparent intoxication during previous visits

Opinions vary on the optimal frequency of urine drug screening to monitor patients on opioid therapy for chronic pain. Screening frequency using a risk-based approach, as recommended by the Washington State interagency guideline (Washington State Agency Medical Directors’ Group, 2015) is as follows:

a. Low risk by ORT: Once a year
b. Moderate risk by ORT: Twice a year
c. High risk or opioid dose >120 mg MED/d: 3 to 4 times a year
d. Recent history of aberrant behavior: Each visit

Note that the ORT is a copyrighted instrument (http://www.opioidrisk.com/node/884). The Canadian Guideline for Safe and Effective Use of Opioids for Chronic Non-Cancer Pain does not include specific screening frequencies but states that an individual patient’s risk for opioid misuse and addiction should be considered when deciding when to order a urine drug screen (http://nationalpaincentre.mcmaster.ca/opioid).

**Substance Use Disorder**

**Stabilization phase**
Most patients are expected to be on a stable dose of opioid medication within 4 weeks of initiating treatment. In some complicated patients, the stabilization phase may last longer than 4 weeks.

**Maintenance phase**
For most patients, targeted presumptive screening once every 1 to 3 months is sufficient during the maintenance phase of treatment. More frequent testing may be appropriate for some complicated patients.

**Guidance On Definitive (Confirmatory) Testing**
Specific situations for definitive drug testing may include, but are not limited to the following:

a. Unexpected positive test inadequately explained by the patient
b. Unexpected negative test (suspected medication diversion)
c. Need for quantitative levels to compare with established benchmarks for clinical decision making

There may not be commercially available tests for certain synthetic or semisynthetic opioids.
The following information on immunoassay availability and diagnostic capacity is included in the Washington State interagency guideline (Washington State Agency Medical Directors’ Group, 2015):

a. Natural Opioids (eg, codeine, morphine)

"Immunoassays for 'opiates' are responsive for morphine and codeine but do not distinguish which is present. Confirmatory testing is required to reliably identify drug(s) present. Since codeine is metabolized to morphine and small quantities to hydrocodone, these drugs may be found in the urine. Also, morphine may metabolize to produce a small amount (<10%) of hydromorphone."

b. Semisynthetic Opioids (eg, hydrocodone, hydromorphone, oxycodone, oxymorphone)

"Opiates' immunoassays may also detect semisynthetic opioids depending on their crossreactivity pattern. However, a negative result does not exclude use of semisynthetic opioids. Confirmatory testing (GC/MS [gas chromatography/mass spectrometry] or LC/MS/MS [liquid chromatography tandem mass spectrometry]) is required to verify compliance with the prescribed semisynthetic opioid(s).

Since hydrocodone is metabolized in small amounts to hydromorphone, both may be found in the urine. Likewise, oxycodone is metabolized to oxymorphone, so these may both be present in the urine of oxycodone users. However, the reverse is not true. In other words, hydromorphone and oxymorphone use does not result in positive screens for hydrocodone and oxycodone, respectively."

c. Synthetic Opioids (eg, fentanyl, meperidine, methadone, propoxyphene)

"Current 'opiates' immunoassays do not detect synthetic opioids. Thus confirmatory testing (GC/MS or LC/MS/MS) is needed to identify these drugs. If the purpose is to document compliance with treatment, the laboratory can be instructed to remove the cutoff concentration so that the presence of lower concentrations can be identified."

Table PG1, on interpreting unexpected results of urine drug tests, was adapted from a table developed by the Canadian National Opioid Use Guideline Group that was cited by the American Society of Interventional Pain Physicians in its guideline on prescribing opioids for chronic noncancer pain.
Table PG1. Interpreting Unexpected Urine Drug Tests Results

<table>
<thead>
<tr>
<th>Unexpected Result</th>
<th>Possible Explanations</th>
<th>Possible Actions for the Physician</th>
</tr>
</thead>
</table>
| Test is negative for prescribed opioid | - False negative  
- Noncompliance  
- Diversion | - Conduct confirmatory testing, specifying the drug of interest (eg, oxycodone often missed by immunoassay)  
- Take a detailed history of patient’s medication use for the preceding 7 days (eg, could learn that patient ran out several days before test)  
- Ask patients if they’ve given the drug to others  
- Monitor compliance with pill counts |
| Test is positive for nonprescribed opioid or benzodiazepines | - False positive  
- Patient acquired opioids from other sources (double-doctoring, "street") | - Repeat urine drug testing regularly  
- Ask patients if they accessed opioids from other sources  
- Assess for opioid misuse/addiction  
- Review/revise treatment agreement |
| UDS positive for illicit drugs (eg, cocaine, cannabis) | - False positive  
- Patient is occasional user or addicted to the illicit drug  
- Cannabis is positive for patients taking certain medications (eg, dronabinol) | - Repeat urine drug test regularly  
- Assess for abuse/addiction and refer for addiction treatment as appropriate |

UDS: urine drug screen

Rationale
This evidence review has been updated with a literature review through October 16, 2017.

Evidence reviews assess whether a medical test is clinically useful. A useful test provides information to make a clinical management decision that improves the net health outcome. That is, the balance of benefits and harms is better when the test is used to manage the condition than when another test or no test is used to manage the condition.

The first step in assessing a medical test is to formulate the clinical context and purpose of the test. The test must be technically reliable, clinically valid, and clinically useful for that purpose. Evidence reviews assess the evidence on whether a test is clinically valid and clinically useful. Technical reliability is outside the scope of these reviews, and credible information on technical reliability is available from other sources.

This evidence review addresses urine drug testing (UDT) as a component of pain management and substance use disorder treatment. For each of these settings, the literature search focused on the accuracy of testing and on the clinical usefulness of testing (ie, the impact of test results on patient management and/or on health outcomes). When published studies were not identified, relevant national and regional clinical practice guidelines were sought. In 2016, testing oral fluids and hair was added to this evidence review. The following is a summary of the key literature to date.
Urine, Oral Fluid, and Hair Drug Testing

Clinical Context and Test Purpose
The purpose of drug testing in patients with chronic pain treated using opioids or patients with drug addiction in substance use disorder treatment is to determine drug use. This review evaluates drug testing on urine, oral fluids, and hair.

The question addressed in this evidence review is: Does UDT, oral fluid drug testing, or hair drug testing improve the net health outcome in individuals with chronic pain being treated with opioids or individuals with drug addiction being treated for a substance use disorder?

The following PICOTS were used to select literature to inform this review.

Patients
The populations of interest include the following: patients with chronic pain who are being treated with opioids and patients with a drug addiction who are being treated for a substance use disorder.

Interventions
The interventions of interest are UDT, oral fluid drug testing, and hair drug testing.

Comparators
For UDT, the comparator is no testing.
For oral fluid drug testing and hair drug testing, the comparators are UDT and no testing.

Outcomes
Outcomes of interest for all tests are test accuracy and validity, health status outcomes, and resource utilization.

A negative result for a prescribed opioid could indicate a false negative, noncompliance, or diversion. The physician may conduct confirmatory testing, obtain a detailed history of medication use to determine the cause of noncompliance, and/or discuss whether the prescribed medication was given to others.

A positive result for a nonprescribed opioid or benzodiazepine could indicate a false-positive or that the patient is obtaining medication from other sources. The physician may request regularly repeated drug testing, discuss how the patient is accessing medication, address medication addiction, and/or revise the treatment agreement.

A positive result for illicit drug use could indicate a false-positive, a patient who is an occasional user, or a patient who is addicted to the illicit drug. The physician may request regularly repeated drug testing and/or assess the patient for drug abuse or addiction.

Timing
In patients with chronic pain being treated with opioids, drug testing may occur at baseline prior to treatment initiation. Subsequent tests may be conducted at a frequency appropriate for the addiction risk level of the individual patient.
In patients with drug addiction being treated for substance use disorders, drug testing may occur at baseline screening. During the stabilization phase, drug testing should be performed weekly for 4 weeks. During the monitoring phase, drug testing may be performed every 1 to 3 months.

**Setting**
In patients with chronic pain being treated with opioids, drug testing can be conducted in outpatient settings.

In patients with drug addiction being treated for substance use disorders, drug testing may be conducted in outpatient substance use disorder centers.

**Urine Drug Testing**

**Technically Reliable**
Assessment of technical reliability focuses on specific tests and operators and requires review of unpublished and often proprietary information. Review of specific tests, operators, and unpublished data are outside the scope of this evidence review and alternative sources exist. This evidence review focuses on the clinical validity and clinical utility.

**Clinical Validity of Detecting Prescribed Opioids and/or Illicit Drugs**
A study evaluating the accuracy of UDT outside of the research setting was published in 2011 by Manchikanti et al. The investigators compared in-office immunoassay testing with gas chromatography/mass spectrometry (GC/MS) as the criterion standard comparison. The study recruited consecutive patients from a single pain management practice. Urine samples were tested for opioids and for illicit drugs. A total of 1000 patients had both the immunoassay and confirmatory tests performed on the same urine sample. Personnel analyzing the tests were blinded to the results of the other test and to patient demographics. The diagnostic accuracy of in-office immunoassays for detecting opioids compared with the reference standard are shown in Table 1. The most common illicit drugs identified were marijuana and amphetamines. Immunoassay sensitivity and specificity for detecting marijuana were 91% and 98%, and for amphetamines 47% and 99%, respectively. There were too few data to report diagnostic accuracy of other illicit drugs reliably.

To most effectively monitor medication compliance of patients being treated for chronic pain, an algorithm has been proposed. The algorithm would first test patients using immunoassays; next, samples with positive immunoassays plus pertinent negatives (samples with negative immunoassays for a medication that was prescribed) would undergo liquid-chromatography tandem mass spectrometry (LC-MS/MS, considered the criterion standard). A study by Snyder et al (2017) tested this algorithm on 530 urine samples of patients being treated for chronic pain. Urine samples were tested for amphetamines, buprenorphine, benzodiazepines, cocaine, opiates, and oxycodone. Overall sensitivity of the immunoassay tests for these 6 drugs/drug classes was 78.5%. When positive immunoassay samples and pertinent negative samples were further tested by LC/MS, the overall sensitivity was increased to 84.6%. Table 1 provides sensitivity results for immunoassays conducted on subsamples from patients prescribed specific drugs.
Table 1. Diagnostic Accuracy of Immunoassay Compared with Gas- or Liquid-Chromatography/ Mass Spectrometry

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Sensitivity (95% CI), %</th>
<th>Specificity (95% CI), %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manchikanti et al (2011)⁷</td>
<td>748</td>
<td>92.5 (90 to 94)</td>
<td>89.6 (82 to 95)</td>
</tr>
<tr>
<td>Patients prescribed opiatesᵃ</td>
<td>134</td>
<td>80.0 (71 to 87)</td>
<td>84.2 (60 to 96)</td>
</tr>
<tr>
<td>Patients prescribed methadone</td>
<td>46</td>
<td>97.8 (88 to 99)</td>
<td>100 (2 to 100)</td>
</tr>
<tr>
<td>Snyder et al (2017)⁸</td>
<td>284</td>
<td>79.4</td>
<td>NR</td>
</tr>
<tr>
<td>Patients prescribed oxycodone/oxymorphone</td>
<td>207</td>
<td>92.5</td>
<td>NR</td>
</tr>
<tr>
<td>Patients prescribed benzodiazepines</td>
<td>82</td>
<td>65.4</td>
<td>NR</td>
</tr>
</tbody>
</table>

CI: confidence interval; NR: not reported.
ᵃ Includes morphine, hydrocodone, codeine, or hydromorphone.

A 2016 retrospective analysis by Johnson-Davis et al evaluated the diagnostic accuracy of an in-house urine drug screen (UDS) panel at a national reference laboratory.⁹ Samples were from routine clinical testing in consecutive patients. The panel tested for 9 drug classes using immunoassay testing. Specimens that screened positive underwent confirmatory testing with GC/MS or LC-MS/MS. A shared confirmatory panel was used for samples testing positive to opiates or oxycodone. A total of 8825 samples were tested. Of them, 2642 (30%) tested positive for opiates and 1215 (14%) tested positive for oxycodone. Confirmatory testing identified 898 (34%) false-positive tests for opiates and 23 (2%) false-positives tests for oxycodone. Authors did not include information on what drugs, if any, were prescribed to patients.

A retrospective analysis by Bertholf et al (2016) evaluated the diagnostic accuracy of UDS in an outpatient setting by sending unexpected positive specimens to a reference laboratory for confirmatory testing.¹⁰ A total of 786 urine specimens with positive screening results were submitted for confirmation. Of the 387 amphetamine-positive specimens, 36 were confirmed by the reference laboratory as having amphetamine and/or methamphetamine, for a positive predictive value (PPV) of 9.3%. Of the 114 opiate-positive specimens, 99 were confirmed for a PPV of 86.8%. The PPV for methadone was 44.1% (45/102 positive specimens). The PPV for oxycodone and/or oxymorphone was 67.6% (38/74 positive specimens) and for both cannabinoid and cocaine was 100% (19/19 positive cannabinoid specimens and 27/27 positive cocaine specimens).

Section Summary: Clinical Validity of Detecting Prescribed Opioids and/or Illicit Drugs

Few studies have evaluated the accuracy of UDT outside of the research setting, either for pain management or substance use disorder treatment patients. Two studies conducted in pain management clinical settings reported that diagnostic accuracy varied by drug type. Among patients prescribed opiates, sensitivity of the UDT ranged from 79% to 93%, and among patients prescribed oxycodone, sensitivity ranged from 80% to 93%. A retrospective analysis of a urine drug panel found false-positive rates of 34% for opiates and 2% for oxycodone. A retrospective analysis focusing on unexpected positive samples reported a wide range of PPVs for UDS, from 100% for cocaine and cannabis to 87% for opioids.
Clinical Utility of Chronic Pain Patients Treated With Opioids

The preferred study design is a randomized controlled trial (RCT) comparing treatment decisions and/or health outcomes in patients managed with and without the use of UDT. When multifaceted interventions are used, it may be difficult to isolate the impact of drug testing from that of other components of the intervention. In that case, the preferred study design would include an arm with the full intervention and another arm with the same intervention but without UDT. In the absence of RCTs, the next most preferred study design is a nonrandomized controlled trial that adjusts findings for potential confounding factors.

Managing Patients With UDT vs Without UDT

Systematic Reviews

A systematic review of the available literature on UDT in the chronic pain management setting, alone or as part of a treatment agreement, was published in 2010 by Starrels et al. Studies were eligible for inclusion in the review if they enrolled patients with chronic non-cancer pain who were treated in an outpatient setting and measured opioid misuse outcomes after intervention implementation. Eleven studies met eligibility criteria; none was an RCT. Quality of the studies was assessed as poor to fair. Eight studies included UDT, with seven of the eight interventions also involving treatment agreements. The protocol and frequency of UDT varied in these studies. In three studies, UDTs were performed at baseline and annually, with additional testing performed according to the judgment of the treating clinician. One study performed testing at baseline and on a monthly basis. In the remaining four studies, the frequency was not specified explicitly, but was described as “regular” or “random.” Five studies described the type of testing used; four of them included confirmatory GC/MS testing.

Reviewers reported that four of the studies included a control or comparison group. However, two of those studies used historical comparison groups and one was a prospective single-arm study. Due to the heterogeneity of interventions, Starrels et al did not pool findings of the 4 studies, with individual studies reporting opioid misuse reductions from 7% to 23% after intervention compared with preintervention rates or historical controls. In the 7 uncontrolled studies in the systematic review, the proportion of patients with opioid misuse after treatment agreements, drug testing, or both ranged from 3% to 43%.

Only a single study included in the systematic review used a concurrent comparison group. Goldberg et al (2005) retrospectively reviewed data from a university-affiliated Veterans Affairs (VA) medical center database on 91 patients with a documented pain management contract. By signing the contract, the patient agreed to 8 provisions, one of which was “lab tests may be used to check opioid use.” Among the other 7 provisions was an agreement not to use illegal drugs and not to share or sell any medication, and an agreement that the patient would receive opioid medication only from a single primary care or pain clinic physician. The comparison group consisted of 224 patients who received opioid prescriptions without pain management contracts during the same time period, matched by age, sex, and race. Consumption of opioids was higher in the intervention group (average of 91 opioid units quarterly) than in the comparison group (average of 81 opioid units quarterly), though this difference was not statistically significant. An opioid unit was defined as 1 systematic administration of morphine sulfate 10 mg. Patients in the intervention group visited their primary care providers significantly more than the control group. Visits to the emergency department did not differ statistically between the groups.
In 2014, Dupouy et al published a systematic review of the literature on the impact of UDT on patient management. All study designs and clinical settings were eligible for inclusion. For a study to be included, the UDS had to be conducted using the enzyme immunoassay technique. For controlled studies, the comparison arm was patient management in the absence of urine testing. In addition, some type of medical management outcome had to be reported (eg, reassessment of treatment, referral for specialist visits, hospitalization). Eight studies met reviewers’ inclusion criteria. Five were rated as poor quality and three as fair quality. The studies consisted of 1 RCT, 2 quasi-randomized studies, 1 observational cohort study, and 4 cross-sectional studies. The RCT was a study of routine drug screening in a psychiatric emergency center, a setting not addressed in this evidence review. Other studies were either drug screening in an emergency department or parents requesting screening of their children. Only 2 studies evaluated relevant populations for this review: one was an uncontrolled evaluation of UDT of opioid-addicted patients, and the other was a quasi-randomized study conducted in U.S. pain centers. The latter study, by Manchikanti et al (2006), was included in the 2010 systematic review, previously described. Authors of the 2014 systematic review (Dupouy et al) did not pool study findings.

Observational Studies
In 2016, Krishnamurthy et al conducted a retrospective cohort study in a university-based pain clinic comparing no-show with dropout rates in chronic pain patients who did and did not receive UDT. Before each clinic visit, patients received a letter stating that their provider might monitor adherence to treatment, including UDT. Investigators used propensity score matching to adjust for potential selection bias and confounding. The sample included 723 patients with a total of 4448 clinic visits (all patients had at least 2 visits). Results showed that UDT at the first visit was significantly associated with a higher rate of no-shows at the second visit (odds ratio, 2.73; 95% confidence interval, 1.66 to 4.47; p<0.001). The no-show rate was 10.2% in patients without UDT and 23.8% in patients with UDT. Moreover, the no-show rate was higher in patients testing positive for illicit drugs (34.6%) than in those testing negative for illicit drugs (21.7%). In addition, the rate of dropout from treatment increased significantly with each additional UDT (95% CI of the hazard ratio, 1.54 to 2.61).

To address the increase in suicide and overdose events occurring with the increase in prescription opioid use, the VA and the Department of Defense formulated clinical practice guidelines for the safe use of opioid treatment for chronic pain. In the guideline, baseline and random UDS were recommended (see the Supplemental Information section). Brennan et al (2016) evaluated the effect of implementing these guidelines on suicide- and overdose-related events. Facility-level data and patient-level data were used in the analysis. From 141 VA healthcare facilities, the facility percentage of opioid-prescribed patients who obtained one or more UDS was calculated. The following data on 484,241 patients from the VA National Patient Care Data files was collected: age, sex, race (white or non-white), marital status, medical comorbidities, mental health comorbidities, UDS, and suicide or drug-related events. The statistical model estimated the effect of UDS practices from 2010 to 2013 on opioid-prescribed patients’ suicide/overdose risk in 2013. Patients’ average age was 60 years, and 7% were women, 19% non-white, and 50% unmarried. Twenty-nine percent had medical comorbidities, and 55% had mental health comorbidities. From 2010 to 2013, the percent of opioid-prescribed patients who received UDS increased from 29% to 42% in VA health care facilities. Patient-level analyses showed that younger, non-white, and unmarried patients had significantly higher risk of suicide or overdose events. Facility-level analyses showed that conducting more UDS in a facility...
was significantly related to a reduction in suicide and overdose events. Model estimates suggested that for every 1% increase in UDS, there was a 1% reduction in patient-level risk of suicide or overdose event.

Stammet et al (2016) evaluated treatment changes occurring after a pharmacist-run UDT e-consult service was implemented in a southeast VA health care system.\(^\text{19}\) During the 2-year pilot study period, 143 e-consults interpreting 190 UDT results were assessed. Based on VA prescription records, the UDS results were classified as: expected (18%), unexpected (28%), and not necessarily inappropriate (54%). In more than 50% of the unexpected results group, the e-consult service recommended immediate action to be taken and, in 35% of those situations, providers documented action within a 30 day period. Other recommendations by the pharmacist included: orders for immediate UDT or more frequent UDT, changes in drug prescriptions, and referrals to pain management services or substance use disorder treatment programs. Follow-up to the recommendations was not available.

**Managing Patients With Routine UDT vs Selective UDT**
No studies were identified.

**Managing Patients With Routine Confirmation of Positive Presumptive Tests vs Selective Confirmation of Positive Presumptive Tests**
No studies were identified.

**Section Summary: Clinical Utility of Chronic Pain Patients Treated With Opioids**
A single RCT in the systematic reviews was identified, though the setting was a psychiatric emergency facility and not relevant to this review. There are several nonrandomized studies with comparison groups and several observational studies. However, both the interventions and the outcomes differed among studies. The interventions often involved patient contract agreements in which UDTs were a component, and 1 intervention was an e-consult service that included UDT recommendations. UDTs were usually conducted randomly and at the discretion of the health care provider. Outcomes across the studies included: primary care visits, emergency room visits, suicide and overdose risk, and opioid misuse. Due to the heterogeneity across interventions and outcomes, pooling of results studies was not possible.

**Clinical Utility of Substance Use Disorder Treatment**

**Managing Patients With UDT vs Without UDT**
One RCT was identified that suggested UDT increases treatment compliance when receiving take-home methadone compared with no UDT. In 2001, Chutuape et al conducted a study of patients in a methadone treatment program who had submitted fewer than 80% positive opiate and/or cocaine-positive urine samples during a 5-week baseline period in which patients were tested every Monday, Wednesday, and Friday.\(^\text{20}\) These patients then participated in a methadone take-home program and were randomized into 1 of 3 groups: (1) continued permission to take-home methadone was contingent on 1 negative urine sample, randomly selected each week (n=16); (2) continued permission to take-home methadone was contingent on 1 negative urine sample, randomly selected each month (n=18); or (3) permission to take-home methadone was not based on results of urine testing, but on results of a random drawing in which half of the control group were given the take-home methadone (control group) (n=19). Ten subjects withdrew from the trial, six from the weekly-tested group, three from the monthly-tested group, and one from
the never-tested group. After participating in the intervention, the rate of sustained (≥8 weeks) opiate and cocaine abstinence was significantly higher in the groups receiving UDT. The percentages of patients with sustained (≥8 weeks) opiate and cocaine abstinence were 56.6%, 38.9%, and 10.5% in the weekly, monthly, and control groups, respectively (p<0.002).

In 2016, McDonell published an RCT evaluating a drug treatment intervention in primary care that included an analysis of whether UDT can detect underreporting of drug use.21 The trial included 829 patients with self-reported nonprescribed drug use or illegal drug use in the past 90 days. UDT was performed at baseline and at 3, 6, 9, and 12 months. Investigators found that 331 (40%) participants denied drug use but had a positive drug screen during at least 1 of the 5 assessments. Patients who denied opioid use but whose UDT was positive were more likely to be older, female, and have a higher Addiction Severity Index drug composite score. This trial was not designed to compare treatment success rates in patients managed with and without UDT.

Managing Patients With Routine UDT vs Selective UDT
No studies were identified.

Managing Patients With Routine Confirmation of Positive Presumptive Tests vs Selective Confirmation of Positive Presumptive Tests
No studies were identified.

Section Summary: Clinical Utility of Substance Use Disorder Treatment
One small RCT on UDT of patients in substance use disorder treatment focused on the specific situation of testing to determine eligibility for take-home methadone. The percentage of patients with 8 or more weeks of opiate or cocaine abstinence was significantly larger in the groups receiving UDT compared with the group not receiving UDT, though there was a large dropout rate in the groups receiving UDTs. Another RCT found UDT identified drug use in a substantial number of patients who denied illicit usage; the impact on treatment success was not addressed in this trial. A currently ongoing RCT sponsored by the French Ministry of Health is evaluating the effectiveness of performing on-site UDS on patients receiving drug addiction treatment in a general practitioner’s office.22 Patients were randomized to a control group (usual care) or to an intervention group in which the practitioners receive UDS supplies and a training session on the use and interpretation of UDS. The primary outcome is retention of opioid management treatment at 6 months. Secondary outcomes are patient adherence to buprenorphine, psychoactive substance use, patient acceptance of UDS, and practitioner acceptance of UDS. Data collection is expected to continue through July 2018.

Oral Fluid Drug Testing

Technically Reliable
Assessment of technical reliability focuses on specific tests and operators and requires review of unpublished and often proprietary information. Review of specific tests, operators, and unpublished data are outside the scope of this evidence review and alternative sources exist.

Clinically Valid
In 2011 Heltsley et al collected oral fluid samples from 231 pain clinics across 20 states, analyzed the samples for 40 drugs or metabolites, and ran confirmatory tests with LC-MS/MS.23 A total of 6441 samples were collected and 5401 tested positive for at least 1 drug category: 38% for 1 drug, 33% for 2 drugs, and 29% for 3 or more drugs. The highest drug prevalence rates
detected through immunoassay testing were: opiates (52%), oxycodone (41%), and benzodiazepines (16%). Confirmatory analyses showed the true positive rates for the oral fluid tests of these 3 highest detected drugs to be 76%, 88%, and 99%, respectively.

Several studies have compared oral fluid testing with UDT using paired samples collected concurrently. For example, in 2011, Vindenes et al in Norway published a study comparing drug detection in oral fluid with urine samples in the drug treatment setting. A total of 164 pairs of urine and oral fluid samples, obtained at the same time, were collected from 45 opioid-dependent patients participating in a drug treatment program. Oral fluid samples were collected using the Intercept device and analyzed using an LC-MS/MS method developed in Norway. Urine samples were screened using immunoassays and confirmed using LC-MS/MS. All patients were treated with buprenorphine or methadone, so it was expected that one of these drugs would be detected in each sample. Other than these 2 drugs, drugs most commonly detected were 7-aminoflunitrazepam (metabolite of flunitrazepam), amphetamine, and tetrahydrocannabinol. The sensitivity and specificity of the oral fluid samples compared with urine results were calculated. Key findings are shown in Table 2.

A 2012 study by Heltsley et al included 133 patients undergoing pain management treatment who consented to provide oral fluid and urine samples. Oral samples were collected with the Quantisal device and specimens were analyzed by LC-MS/MS. Urine specimens were screened by immunoassay procedures, and non-negative samples were confirmed by MS. Samples were tested for 34 drugs or drug metabolites, although in some instances different analyses were performed on urine and oral fluid specimens. A total of 1544 paired tests were performed; of these, 329 (21.3%) were positive, and 984 (63.7%) were negative in both matrices, for an overall agreement of 85%. Eighty-three (5.4%) findings were positive in oral fluid only, and 148 (9.6%) were positive in urine only. Authors conducted several analyses of the sensitivity and specificity of oral fluid samples using urinalysis as the reference standard (see Table 2).

Table 2. Sensitivity and Specificity of Oral Fluid Samples Using Urinalysis as the Reference Standard

<table>
<thead>
<tr>
<th>Drug</th>
<th>Sensitivity (95% CI), %</th>
<th>Specificity (95% CI), %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vindenes et al (2011)²⁴</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methadone</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Buprenorphine</td>
<td>75</td>
<td>Not applicable (analytic problems)</td>
</tr>
<tr>
<td>7-aminoflunitrazepam</td>
<td>76</td>
<td>97</td>
</tr>
<tr>
<td>Amphetamine</td>
<td>100</td>
<td>95</td>
</tr>
<tr>
<td>Tetrahydrocannabinol</td>
<td>82</td>
<td>98</td>
</tr>
<tr>
<td>6-MAM (heroin)</td>
<td>95</td>
<td>80</td>
</tr>
<tr>
<td>Heltsley et al (2012)²⁵</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All drugs</td>
<td>69.0 (64.6 to 73.1)</td>
<td>92.2 (90.4 to 93.7)</td>
</tr>
<tr>
<td>Four drug categories⁴</td>
<td>76.1 (60.9 to 86.9)</td>
<td>95.9 (92.0 to 98.0)</td>
</tr>
<tr>
<td>Six drug categories⁵</td>
<td>82.3 (75.0 to 87.9)</td>
<td>92.2 (88.7 to 94.7)</td>
</tr>
</tbody>
</table>

CI: confidence interval.

⁴ Categories include amphetamines, cannabis, cocaine, and opiates.

⁵ Includes the above categories plus hydrocodone and oxycodone.

In 2014, Conermann et al compared findings of oral fluid and urinalysis in 153 paired samples from patients attending a pain management clinic. This study focused on confirmation that a treatment drug was being taken and did not report the sensitivity and specificity of oral fluid...
samples compared with urine samples. Oral fluid samples were collected with the Quantisal device. All specimens were screened with immunoassays, and presumptive positive findings were confirmed using LC/MS. A total of 136 (89%) of the 153 paired samples tested positive for one or more treatment drugs (ie, opioids, benzodiazepines) in one or both matrices. After excluding 4 paired samples due to missing data, 101 (76.5%) of 132 positive specimen pairs had exact drug class matches. In another 21 paired samples, there was at least 1 (15.9%) drug class match. Thus, there was an overall agreement between samples of 92.4%. Two analyses were positive in oral fluid only, and eight were positive in urine only.

Kunkel et al (2015) conducted a retrospective analysis of 4560 unobserved urine collection samples and 2368 observed oral fluid collection samples of patients undergoing opioid addiction treatment.27 The samples were tested for 13 different illicit and prescription drugs. Oral fluid testing detected higher rates of most of the drugs compared with urine testing. For example, oral fluid tests detected 6.5% morphine, 5.2% oxycodone, 3.8% codeine, and 3% cocaine use, while urine tests detected 2.3% morphine, 0.6% oxycodone, 0.7% codeine, and 1.4% cocaine use. Interpretation of these results is limited because the urine and oral fluid samples were not paired.

**Clinically Useful**

No studies were identified that compared patient management decisions or health outcomes in patients managed using oral fluid drug testing vs UDT or no drug testing.

**Section Summary: Oral Fluid Testing**

The limited number of studies on the diagnostic accuracy of oral fluid testing compared with urine testing had variable findings. No studies were identified on the impact of oral fluid testing on health outcomes compared with UDT or no drug testing.

**Hair Testing**

**Technically Reliable**

Assessment of technical reliability focuses on specific tests and operators and requires review of unpublished and often proprietary information. Review of specific tests, operators, and unpublished data are outside the scope of this evidence review and alternative sources exist.

**Clinically Valid**

No studies were identified that compared the accuracy of hair and urine testing using paired samples collected concurrently in the pain management setting or drug use disorder treatment setting. One study using paired samples of urine and hair from patients in a psychiatric facility was identified (Musshoff et al, 2006).28 Patients underwent drug testing as part of the intake process for psychiatric treatment. Urine and hair samples (both head hair and pubic hair) from known drug users were analyzed. Fifty-one patients were included; all provided urine samples, 47 provided head hair samples (1-3 segments), and 36 provided pubic hair samples. Drug analysis was done using GC-MS methods. The hair test was considered positive if any segment had a positive finding. Urine samples were analyzed using standard immunoassays; positive findings were not confirmed. Prevalence rates of drugs identified in hair and urine samples, as well as self-report of drug use, are shown in Table 3.
Table 3. Prevalence Rates of Drug Use (N=47)

<table>
<thead>
<tr>
<th>Source</th>
<th>Opiates</th>
<th>Cocaine</th>
<th>Methadone</th>
<th>Cannabinoids</th>
<th>Amphetamines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Self-report, n (%)</td>
<td>42 (89)</td>
<td>18 (38)</td>
<td>15 (32)</td>
<td>26 (55)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>Urine test, n (%)</td>
<td>33 (70)</td>
<td>13 (28)</td>
<td>14 (30)</td>
<td>21 (45)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Hair test, n (%)</td>
<td>38 (81)</td>
<td>26 (55)</td>
<td>23 (49)</td>
<td>15 (32)</td>
<td>1 (2)</td>
</tr>
</tbody>
</table>

Hair tests revealed a higher prevalence of drug use than UDT for most drugs, with the exception of cannabinoids. The prevalence of amphetamines was too low to make meaningful comparisons. Cannabinoids are known to be excreted slowly in urine and to have a low incorporation rate into hair. It is important to note that the hair analysis was used to detect drug use anytime during the past several months and the urine analysis to detect drug use in the past several days.

**Clinically Useful**

No studies were identified that compared patient management decisions or health outcomes in patients managed using testing of hair vs UDT or no drug testing.

**Section Summary: Hair Testing**

Hair testing cannot detect recent drug use (ie, in the past few days). One study looked at this longer time frame in patients starting psychiatric treatment. It found a higher prevalence of drug use with hair testing vs UDT for most drugs; however, the implications of study findings for patients in pain management or substance use disorder treatment is unclear. No studies were identified on the diagnostic accuracy of hair testing vs UDT in patients with chronic pain or substance use disorder. In addition, no studies were identified on the clinical utility of hair testing in pain management or substance use disorder treatment.

**Summary of Evidence**

For individuals who have chronic pain treated with opioids who receive UDT, the evidence includes nonrandomized comparative studies and systematic reviews. Relevant outcomes are test accuracy and validity, health status measures, and resource utilization. The evidence on the diagnostic accuracy of urine immunoassay tests, as confirmed by gas- or liquid-chromatography/mass spectrometry, shows sensitivities ranging from about 80% to 93% for both opiates and oxycodone. No RCTs evaluating clinical utility were identified. Several nonrandomized comparative studies have been conducted, though interventions and outcomes have varied across the studies. Most interventions included patient contracts along with UDT, and therefore, the effect of UDT alone could not be determined. Most studies did not provide details on the frequency of UDTs and whether the testing was random or scheduled. As a result, these studies provided inconclusive evidence on whether UDT in the pain management setting improves patient outcomes. The evidence is insufficient to determine the effects of the technology on health outcomes.

For individuals who have a drug addiction who are in substance use disorder treatment who receive UDT, the evidence includes 2 RCTs. Relevant outcomes are test accuracy and validity, health status measures, and resource utilization. No studies were identified that evaluated the accuracy of UDT compared with a valid reference standard in individuals undergoing substance use disorder treatment. One small RCT focused specifically on UDT to determine eligibility for take-home methadone. The second RCT found that UDT identified drug use in a substantial number of patients who denied illicit usage; the impact on treatment success was not addressed. The evidence is insufficient to determine the effects of the technology on health outcomes.
For individuals who have chronic pain treated with opioids or with a drug addiction in substance use disorder treatment who receive oral fluid drug testing, the evidence includes diagnostic accuracy studies using UDT as the reference standard. Relevant outcomes are test accuracy and validity, health status measures, and resource utilization. The limited number of studies on the diagnostic accuracy of oral fluid testing compared with UDT have varied findings. No studies were identified assessing the impact of oral fluid testing on health outcomes compared with UDT or no drug testing. The evidence is insufficient to determine the effects of the technology on health outcomes.

For individuals who have chronic pain treated with opioids or with a drug addiction in substance use disorder treatment who receive hair drug testing, the evidence includes a diagnostic accuracy study in the psychiatric treatment setting. Relevant outcomes are test accuracy and validity, health status measures, and resource utilization. Hair testing cannot detect recent drug use (ie, in the past few days), and thus has limited applicability to pain management or substance use disorder treatment settings, except, perhaps, for initial intake. There are no studies comparing the diagnostic accuracy of hair testing with UDT in either setting. One relatively small study tested the hair and urine of known drug users recruited from a psychiatric clinic. The study looked for drug use over the past several months rather than the shorter timeframe generally needed in pain management or drug treatment settings. No studies were identified on the clinical utility of hair testing in pain management or substance use disorder treatment. The evidence is insufficient to determine the effects of the technology on health outcomes.

Clinical Input From Physician Specialty Societies and Academic Medical Centers
While the various physician specialty societies and academic medical centers may collaborate with and make recommendations during this process, through the provision of appropriate reviewers, input received does not represent an endorsement or position statement by the physician specialty societies or academic medical centers, unless otherwise noted.

In response to requests, input was received from 5 physician specialty societies and 8 academic medical centers while this policy was under review in 2014. There was near-consensus among reviewers that, in outpatient pain management, presumptive (ie, qualitative) urine drug testing may be considered medically necessary for patients who meet the stated criteria and that the frequency of repeat drug testing should depend on the risk level of the individual. There was also near-consensus among reviewers that, in substance abuse treatment, baseline presumptive drug testing may be considered medically necessary for patients who meet the stated criteria and that targeted weekly qualitative screening for a maximum of 4 weeks may be considered medically necessary during the stabilization phase. There was mixed input on the frequency of presumptive drug testing that may be considered medically necessary during the maintenance phase of substance abuse treatment. In addition, clinical input was mixed on confirmatory definitive (ie, quantitative) drug testing and particularly on whether definitive drug testing should only be performed on a drug-specific basis.

Practice Guidelines and Position Statements
Pain Management
In 2014, Nuckols et al published a systematic review of guidelines that addressed management of opioid use for chronic pain.29 Reviewers included guidelines from national organizations and specialty societies, as well as guidelines from state agencies and specific health systems. Moreover, reviewers identified 9 guidelines with recommendations on urine drug testing (UDT).
Recommendations varied widely; two recommended mandatory testing for all patients, another recommended testing only patients at increased risk of medication use disorder, and two stated that testing patients at low risk of abuse is not cost-effective. If UDT is used, the recommended frequency of follow-up testing was at least quarterly in 1 guideline, at least yearly in another, and randomly in two.

Centers for Disease Control and Prevention
In 2016, the Centers for Disease Control and Prevention published guidelines on opioids for chronic pain.30 The guidelines included the following recommendation on UDT: "When prescribing opioids for chronic pain, clinicians should use urine drug testing before starting opioid therapy and consider urine drug testing at least annually to assess for prescribed medications as well as other controlled prescription drugs and illicit drugs."

American Society of Interventional Pain Physicians
In 2017, the American Society of Interventional Pain Physicians issued guidelines for responsible, safe, and effective opioid prescribing for chronic non-cancer pain.31 The guidelines included the following recommendations on UDT (see Table 4).

Table 4. Recommendations on Urine Drug Testing for Chronic Non-Cancer Pain

<table>
<thead>
<tr>
<th>Recommendation</th>
<th>LOE</th>
<th>SOE</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Comprehensive assessment and documentation is recommended before initiating opioid therapy, with documentation of comprehensive history, general medical condition, psychosocial history, psychiatric status, and substance use history.”</td>
<td>I</td>
<td>Strong</td>
</tr>
<tr>
<td>“Screening for opioid abuse is recommended, as it will potentially identify opioid abusers and reduce opioid abuse.”</td>
<td>II-III</td>
<td>Moderate</td>
</tr>
<tr>
<td>“Presumptive UDT is implemented at initiation of opioid therapy, along with subsequent use as adherence monitoring, using in-office point of service testing, followed by confirmation with chromatography/mass spectrometry for accuracy in select cases, to identify patients who are not compliant or abusing prescription drugs or illicit drugs. UDT may decrease prescription drugs abuse of illicit drug use when patients are in chronic pain management therapy.”</td>
<td>III</td>
<td>Moderate</td>
</tr>
</tbody>
</table>

LOE: level of evidence; SOE: strength of evidence; UDT: urine drug testing.

American Pain Society and American Academy of Pain Medicine
In 2009, the American Pain Society and American Academy of Pain Medicine jointly published clinical guidelines on the use of opioid therapy in chronic non-cancer pain.32 The guidelines did not address UDT or other forms of monitoring adherence.

American College of Occupational and Environmental Medicine
The latest guidelines from the American College of Occupational and Environmental Medicine (ACOEM) on the use of opioids for the treatment of acute, subacute, chronic, and postoperative pain, were published in 2014.33 The following recommendations on UDT were made for subacute (1-3 months) and chronic pain (>3 months) (see Table 5).
Table 5. Recommendations on Opioid Use to Treat Acute, Subacute, Chronic, and Postoperative Pain

<table>
<thead>
<tr>
<th>Recommendation</th>
<th>SOR</th>
<th>CIR</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Baseline and random urine drug screening, qualitative and quantitative, for patients prescribed opioids for the treatment of subacute or chronic pain to evaluate presence or absence of the drug, its metabolites and other substance(s) use. In certain situations, other screenings (eg, hair particularly for information regarding remote use or blood) (for acute toxicity) may be appropriate.”</td>
<td>C</td>
<td>High</td>
</tr>
</tbody>
</table>

Recommendations rating schema: A: strongly recommended; B: moderately recommended; C: recommended. CIR: confidence in recommendation; SOR: strength of recommendation.

Urine drug screening was not recommended for acute pain (up to 4 weeks) or for postoperative pain (up to 4 weeks).

As a companion to the guidelines, ACOEM developed a combined Opioid Consent Form and Opioid Treatment Agreement.34 The form provides explanations of the potential benefits and harms to be expected from opioid treatment, and asks the patient to agree to numerous terms of opioid use, which include submitting to unscheduled urine, blood, saliva, or hair drug testing at the prescriber’s request and seeing an addiction specialist if requested.

Screening was recommended for all patients at baseline, and then randomly at least twice and up to 4 times a year, and at termination. Screening should also be performed if the provider suspects abuse of prescribed medication.

**National Opioid Use Guideline Group**

The National Opioid Use Guideline Group issued guidelines in Canada in 2010 on the safe and effective use of opioids for chronic non-cancer pain that included the following recommendation on urine drug screening5: “When using urine drug screening (UDS) to establish a baseline measure of risk or to monitor compliance, be aware of benefits and limitations, appropriate test ordering and interpretation, and have a plan to use results. (Grade C).”

The guidelines also stated that there was no “compelling evidence” to guide physicians on identifying patients who should have UDS, or on how often they should be tested. The document stated that the following factors should be considered when deciding whether to order a urine drug screen:

- patient’s risk for opioid misuse and addiction
- aberrant drug-related behaviors
- testing availability (note: this may be a Canadian-specific issue).

**Department of Veterans Affairs and Department of Defense**

In 2010, the Department of Veterans Affairs and Department of Defense issued clinical practice guidelines for managing opioid therapy for treatment of chronic pain.6 The recommendations on assessing adherence to prescribed opioids includes obtaining a urine drug test (with patient consent) before initiating opioid therapy, and then randomly at follow-up to confirm appropriate use. Other strategies recommended include clinical assessment and screening aids such as random pill counts, adherence checklists, and standardized instruments such as the Screener and Opioid Assessment for Patients with Pain.
The guidelines included the following specific recommendations on UDT:

“RECOMMENDATIONS

1. Inform patients that drug testing is a routine procedure for all patients starting or on opioid therapy [OT], and is an important tool for monitoring the safety of their treatment.

2. With patient consent, obtain a UDT in all patients prior to initiation of OT.

3. With patient consent, monitor all patients on OT with periodic random UDTs to confirm adherence to the treatment plan. Increase the frequency of UDTs based on risk level for aberrant drug-related behaviors and following each dose increase.

4. Take into consideration a patient’s refusal to take a UDT as part of the ongoing assessment of the patient’s ability to adhere to the treatment plan and the level of risk for adverse outcomes.

5. When interpreting UDT results take into account other clinical information (e.g., past SUD [substance use disorder], other risk factors, aberrant drug-related behaviors, and other conditions indicating risk.)

6. Understanding of lab methods for drug testing and reporting are necessary to interpret UDT results (i.e., screen versus confirmatory test, substances tested, cut-off levels for tests). Maintain a close working relationship with the clinical laboratory to answer any questions about the UDT or for confirming the results.”

Washington State Agency Medical Directors’ Group
In 2015, the Washington State Agency Medical Directors’ Group updated its interagency guidelines on opioid dosing for chronic non-cancer pain.35 The guidelines included recommendations on UDT. Recommendations on testing frequency differed depending on the patient risk of opioid addiction and opioid dosage, as listed below:

- Low risk: Once per year
- Moderate risk: Twice per year
- High risk or opioid dose over 120 mg MED/d: 3-4 times per year
- Aberrant behavior: Each visit.

No pain management guidelines were identified that had recommendations on oral fluid or hair testing.

Substance Use Disorder Treatment
American Society of Addiction Medicine
The American Society of Addiction Medicine (ASAM) has published several documents on drug testing: a public policy statement (2010),36 a white paper (2013), which provided background on the science and current practices of drug testing,37 and guidelines (2017) on the effective use of drug testing.38

ASAM’s public policy statement asserts that: “Urine drug testing is a key diagnostic and therapeutic tool that is useful for patient care and in monitoring of the ongoing status of a person who has been treated for addiction. As such, it is a part of medical care, and should not face undue restrictions.”36 ASAM recommended drug testing where medically appropriate in clinical diagnostic settings and clinical treatment settings. The term “drug testing” in this document was a broad term that included urine or other body fluids or tissues.

The ASAM White Paper concluded that “The most important challenge in drug testing today is not the identification of every drug that we are technologically capable of detecting, but to do
medically necessary and accurate testing for those drugs that are most likely to impact clinical outcomes. The paper acknowledged that more specific guidance on drug testing was needed, which led to the development of the 2017 guidelines, described below.

The 2017 ASAM guidance on appropriate drug testing in clinical addiction medicine advises health care providers that before choosing the type of drug test, they should first identify the questions they are seeking to answer and be aware of benefits and limitations of the various drug tests. Table 6 summarizes characteristics of urine, oral fluid, and hair drug tests that may inform the decision of what type of drug test to use.

**Table 6. Summary of Drug Testing Characteristics**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Urine</th>
<th>Oral Fluid</th>
<th>Hair</th>
</tr>
</thead>
<tbody>
<tr>
<td>General detection period</td>
<td>Hours to days</td>
<td>Minutes to hours</td>
<td>Weeks to months</td>
</tr>
<tr>
<td>Point-of-care testing</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Primarily detects</td>
<td>Drug metabolite</td>
<td>Parent drug compound</td>
<td>Parent drug compound</td>
</tr>
<tr>
<td>Best use in treatment setting</td>
<td>Intermediate-term detection in ongoing treatment</td>
<td>Short-term detection in ongoing treatment</td>
<td>Long-term monitoring, 3-month history</td>
</tr>
<tr>
<td>Ease of collection</td>
<td>Requires restroom</td>
<td>Easily collected</td>
<td>Easily collected</td>
</tr>
<tr>
<td>Resistance to tampering</td>
<td>Low</td>
<td>High, with some uncertainty</td>
<td>High when chemically untreated</td>
</tr>
<tr>
<td>Retesting same sample</td>
<td>Possible</td>
<td>Difficult</td>
<td>Easy</td>
</tr>
</tbody>
</table>

Adapted from Jarvis et al (2017).

**U.S. Preventive Services Task Force Recommendations**

Not applicable.

**Ongoing and Unpublished Clinical Trials**

A currently unpublished trial that might influence this review is listed in Table 7.

**Table 7. Summary of Key Trials**

<table>
<thead>
<tr>
<th>NCT No.</th>
<th>Trial Name</th>
<th>Planned Enrollment</th>
<th>Completion Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ongoing</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NCT02345655</td>
<td>Impact of On-site Evaluation of Substances Consumption on Opiate Maintenance in the Context of Family Practice</td>
<td>400</td>
<td>Jan 2018</td>
</tr>
</tbody>
</table>

NCT: national clinical trial.
CODING

The following codes for treatment and procedures applicable to this policy are included below for informational purposes. Inclusion or exclusion of a procedure, diagnosis or device code(s) does not constitute or imply member coverage or provider reimbursement. Please refer to the member's contract benefits in effect at the time of service to determine coverage or non-coverage of these services as it applies to an individual member.

CPT/HCPCS

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>80305</td>
<td>Drug test(s), presumptive, any number of drug classes, any number of devices or procedures; capable of being read by direct optical observation only (eg, utilizing immunoassay [eg, dipsticks, cups, cards, cartridges]), includes sample validation when performed, per date of service</td>
</tr>
<tr>
<td>80306</td>
<td>Drug test(s), presumptive, any number of drug classes, any number of devices or procedures; read by instrument assisted direct optical observation (eg, utilizing immunoassay [eg, dipsticks, cups, cards, cartridges]), includes sample validation when performed, per date of service</td>
</tr>
<tr>
<td>80307</td>
<td>Drug test(s), presumptive, any number of drug classes, any number of devices or procedures, by instrument chemistry analyzers (eg, utilizing immunoassay [eg, EIA, ELISA, EMIT, FPIA, IA, KIMS, RIA]), chromatography (eg, GC, HPLC), and mass spectrometry either with or without chromatography, (eg, DART, DESI, GC-MS, GC-MS/MS, LC-MS, LC-MS/MS, LDTD, MALDI, TOF) includes sample validation when performed, per date of service</td>
</tr>
<tr>
<td>80320</td>
<td>Alcohol(s)</td>
</tr>
<tr>
<td>80321</td>
<td>Alcohol biomarkers; 1 or 2</td>
</tr>
<tr>
<td>80322</td>
<td>Alcohol biomarkers; 3 or more</td>
</tr>
<tr>
<td>80323</td>
<td>Alkaloids, not otherwise specified</td>
</tr>
<tr>
<td>80324</td>
<td>Amphetamines; 1 or 2</td>
</tr>
<tr>
<td>80325</td>
<td>Amphetamines; 3 or 4</td>
</tr>
<tr>
<td>80326</td>
<td>Amphetamines; 5 or more</td>
</tr>
<tr>
<td>80327</td>
<td>Anabolic steroids; 1 or 2</td>
</tr>
<tr>
<td>80328</td>
<td>Anabolic steroids; 3 or more</td>
</tr>
<tr>
<td>80329</td>
<td>Analgesics, non-opioid; 1 or 2</td>
</tr>
<tr>
<td>80330</td>
<td>Analgesics, non-opioid; 3-5</td>
</tr>
<tr>
<td>80331</td>
<td>Analgesics, non-opioid; 6 or more</td>
</tr>
<tr>
<td>80332</td>
<td>Antidepressants, serotonergic class; 1 or 2</td>
</tr>
<tr>
<td>80333</td>
<td>Antidepressants, serotonergic class; 3-5</td>
</tr>
<tr>
<td>80334</td>
<td>Antidepressants, serotonergic class; 6 or more</td>
</tr>
<tr>
<td>80335</td>
<td>Antidepressants, tricyclic and other cyclicals; 1 or 2</td>
</tr>
<tr>
<td>80336</td>
<td>Antidepressants, tricyclic and other cyclicals; 3-5</td>
</tr>
<tr>
<td>80337</td>
<td>Antidepressants, tricyclic and other cyclicals; 6 or more</td>
</tr>
<tr>
<td>80338</td>
<td>Antidepressants, not otherwise specified</td>
</tr>
<tr>
<td>80339</td>
<td>Antiepileptics, not otherwise specified; 1-3</td>
</tr>
<tr>
<td>80340</td>
<td>Antiepileptics, not otherwise specified; 4-6</td>
</tr>
<tr>
<td>80341</td>
<td>Antiepileptics, not otherwise specified; 7 or more</td>
</tr>
<tr>
<td>80342</td>
<td>Antipsychotics, not otherwise specified; 1-3</td>
</tr>
<tr>
<td>80343</td>
<td>Antipsychotics, not otherwise specified; 4-6</td>
</tr>
<tr>
<td>80344</td>
<td>Antipsychotics, not otherwise specified; 7 or more</td>
</tr>
<tr>
<td>80345</td>
<td>Barbiturates</td>
</tr>
<tr>
<td>Code</td>
<td>Substance/Class</td>
</tr>
<tr>
<td>-------</td>
<td>---------------------------------------</td>
</tr>
<tr>
<td>80346</td>
<td>Benzodiazepines; 1-12</td>
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<tr>
<td>80347</td>
<td>Benzodiazepines; 13 or more</td>
</tr>
<tr>
<td>80348</td>
<td>Buprenorphine</td>
</tr>
<tr>
<td>80349</td>
<td>Cannabinoids, natural</td>
</tr>
<tr>
<td>80350</td>
<td>Cannabinoids, synthetic; 1-3</td>
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<tr>
<td>80351</td>
<td>Cannabinoids, synthetic; 4-6</td>
</tr>
<tr>
<td>80352</td>
<td>Cannabinoids, synthetic; 7 or more</td>
</tr>
<tr>
<td>80353</td>
<td>Cocaine</td>
</tr>
<tr>
<td>80354</td>
<td>Fentanyl</td>
</tr>
<tr>
<td>80355</td>
<td>Gabapentin, non-blood</td>
</tr>
<tr>
<td>80356</td>
<td>Heroin metabolite</td>
</tr>
<tr>
<td>80357</td>
<td>Ketamine and norketamine</td>
</tr>
<tr>
<td>80358</td>
<td>Methadone</td>
</tr>
<tr>
<td>80359</td>
<td>Methyleneedioxyamphetamines (MDA, MDEA, MDMA)</td>
</tr>
<tr>
<td>80360</td>
<td>Methylphenidate</td>
</tr>
<tr>
<td>80361</td>
<td>Opiates, 1 or more</td>
</tr>
<tr>
<td>80362</td>
<td>Opioids and opiate analogs; 1 or 2</td>
</tr>
<tr>
<td>80363</td>
<td>Opioids and opiate analogs; 3 or 4</td>
</tr>
<tr>
<td>80364</td>
<td>Opioids and opiate analogs; 5 or more</td>
</tr>
<tr>
<td>80365</td>
<td>Oxycodone</td>
</tr>
<tr>
<td>80366</td>
<td>Pregabalin</td>
</tr>
<tr>
<td>80367</td>
<td>Propoxyphene</td>
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<tr>
<td>80368</td>
<td>Sedative hypnotics (non-benzodiazepines)</td>
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<tr>
<td>80369</td>
<td>Skeletal muscle relaxants; 1 or 2</td>
</tr>
<tr>
<td>80370</td>
<td>Skeletal muscle relaxants; 3 or more</td>
</tr>
<tr>
<td>80371</td>
<td>Stimulants, synthetic</td>
</tr>
<tr>
<td>80372</td>
<td>Tapentadol</td>
</tr>
<tr>
<td>80373</td>
<td>Tramadol</td>
</tr>
<tr>
<td>80374</td>
<td>Stereoisomer (enantiomer) analysis, single drug class</td>
</tr>
<tr>
<td>80375</td>
<td>Drug(s) or substance(s), definitive, qualitative or quantitative, not otherwise specified; 1-3</td>
</tr>
<tr>
<td>80376</td>
<td>Drug(s) or substance(s), definitive, qualitative or quantitative, not otherwise specified; 4-6</td>
</tr>
<tr>
<td>80377</td>
<td>Drug(s) or substance(s), definitive, qualitative or quantitative, not otherwise specified; 7 or more</td>
</tr>
<tr>
<td>83992</td>
<td>Phencyclidine (PCP)</td>
</tr>
<tr>
<td>G0480</td>
<td>Drug test(s), definitive, utilizing (1) drug identification methods able to identify individual drugs and distinguish between structural isomers (but not necessarily stereoisomers), including, but not limited to, GC/MS (any type, single or tandem) and LC/MS (any type, single or tandem) and excluding immunoassays (e.g., IA, EIA, ELISA, EMIT, FPIA) and enzymatic methods (e.g., alcohol dehydrogenase)); (2) stable isotope or other universally recognized internal standards in all samples (e.g., to control for matrix effects, interferences and variations in signal strength), and (3) method or drug-specific calibration and matrix-matched quality control material (e.g., to control for instrument variations and mass spectral drift); qualitative or quantitative, all sources, includes specimen validity testing, per day, 1-7 drug class(xsch), including metabolite(s) if performed</td>
</tr>
</tbody>
</table>
G0481 Drug test(s), definitive, utilizing (1) drug identification methods able to identify individual drugs and distinguish between structural isomers (but not necessarily stereoisomers), including, but not limited to, GC/MS (any type, single or tandem) and LC/MS (any type, single or tandem) and excluding immunoassays (e.g., IA, EIA, ELISA, EMIT, FPIA) and enzymatic methods (e.g., alcohol dehydrogenase)); (2) stable isotope or other universally recognized internal standards in all samples (e.g., to control for matrix effects, interferences and variations in signal strength), and (3) method or drug-specific calibration and matrix-matched quality control material (e.g., to control for instrument variations and mass spectral drift); qualitative or quantitative, all sources, includes specimen validity testing, per day, 8-14 drug class(es), including metabolite(s) if performed

G0482 Drug test(s), definitive, utilizing (1) drug identification methods able to identify individual drugs and distinguish between structural isomers (but not necessarily stereoisomers), including, but not limited to, GC/MS (any type, single or tandem) and LC/MS (any type, single or tandem) and excluding immunoassays (e.g., IA, EIA, ELISA, EMIT, FPIA) and enzymatic methods (e.g., alcohol dehydrogenase)); (2) stable isotope or other universally recognized internal standards in all samples (e.g., to control for matrix effects, interferences and variations in signal strength), and (3) method or drug-specific calibration and matrix-matched quality control material (e.g., to control for instrument variations and mass spectral drift); qualitative or quantitative, all sources, includes specimen validity testing, per day, 15-21 drug class(es), including metabolite(s) if performed

G0483 Drug test(s), definitive, utilizing (1) drug identification methods able to identify individual drugs and distinguish between structural isomers (but not necessarily stereoisomers), including, but not limited to, GC/MS (any type, single or tandem) and LC/MS (any type, single or tandem) and excluding immunoassays (e.g., IA, EIA, ELISA, EMIT, FPIA) and enzymatic methods (e.g., alcohol dehydrogenase)); (2) stable isotope or other universally recognized internal standards in all samples (e.g., to control for matrix effects, interferences and variations in signal strength), and (3) method or drug-specific calibration and matrix-matched quality control material (e.g., to control for instrument variations and mass spectral drift); qualitative or quantitative, all sources, includes specimen validity testing, per day, 20 or more drug class(es), including metabolite(s) if performed

0006U Detection of interacting medications, substances, supplements and foods, 120 or more analytes, definitive chromatography, with mass spectrometry, urine description and severity of each interaction, identified, per date of service

0007U Drug test(s), presumptive, with definitive confirmation of positive results, any number of drug classes, urine, includes specimen verification including DNA authentication in comparison to buccal DNA, per date of service

0051U Prescription drug monitoring, evaluation of drugs present by LC-MS/MS, urine, 31 drug panel, reported as quantitative results, detected or not detected, per date of service

0082U Drug test(s), definitive, 90 or more drugs or substances, definitive chromatography with mass spectrometry, and presumptive, any number of drug classes, by instrument chemistry analyzer (utilizing immunoassay), urine, report of presence or absence of each drug, drug metabolite or substance with description and severity of significant interactions per date of service
ICD-10 Diagnoses (Effective October 1, 2015)

Any of a large number of diagnosis codes might apply to this policy, the following are just examples

- **F11.10** Opioid abuse, uncomplicated
- **F11.11** Opioid abuse, in remission
- **F11.120** Opioid abuse with intoxication, uncomplicated
- **F11.121** Opioid abuse with intoxication delirium
- **F11.122** Opioid abuse with intoxication with perceptual disturbance
- **F11.14** Opioid abuse with opioid-induced mood disorder
- **F11.150** Opioid abuse with opioid-induced psychotic disorder with delusions
- **F11.151** Opioid abuse with opioid-induced psychotic disorder with hallucinations
- **F11.181** Opioid abuse with opioid-induced sexual dysfunction
- **F11.182** Opioid abuse with opioid-induced sleep disorder
- **F11.188** Opioid abuse with other opioid-induced disorder
- **F11.19** Opioid abuse with unspecified opioid-induced disorder
- **F11.20** Opioid dependence, uncomplicated
- **F11.21** Opioid dependence, in remission
- **F11.220** Opioid dependence with intoxication, uncomplicated
- **F11.221** Opioid dependence with intoxication delirium
- **F11.222** Opioid dependence with intoxication with perceptual disturbance
- **F11.23** Opioid dependence with withdrawal
- **F11.24** Opioid dependence with opioid-induced mood disorder
- **F11.250** Opioid dependence with opioid-induced psychotic disorder with delusions
- **F11.251** Opioid dependence with opioid-induced psychotic disorder with hallucinations
- **F11.281** Opioid dependence with opioid-induced sexual dysfunction
- **F11.282** Opioid dependence with opioid-induced sleep disorder
- **F11.288** Opioid dependence with other opioid-induced disorder
- **F11.29** Opioid dependence with unspecified opioid-induced disorder
- **F11.90** Opioid use, unspecified, uncomplicated
- **F11.920** Opioid use, unspecified with intoxication, uncomplicated
- **F11.921** Opioid use, unspecified with intoxication delirium
- **F11.922** Opioid use, unspecified with intoxication with perceptual disturbance
- **F11.93** Opioid use, unspecified with withdrawal
- **F11.94** Opioid use, unspecified with opioid-induced mood disorder
- **F11.950** Opioid use, unspecified with opioid-induced psychotic disorder with delusions
- **F11.951** Opioid use, unspecified with opioid-induced psychotic disorder with hallucinations
- **F11.981** Opioid use, unspecified with opioid-induced sexual dysfunction
- **F11.982** Opioid use, unspecified with opioid-induced sleep disorder
- **F11.988** Opioid use, unspecified with other opioid-induced disorder
- **F11.99** Opioid use, unspecified with unspecified opioid-induced disorder
- **F14.10** Cocaine abuse, uncomplicated
- **F14.11** Cocaine abuse, in remission
- **F14.120** Cocaine abuse with intoxication, uncomplicated
- **F14.121** Cocaine abuse with intoxication with delirium
- **F14.122** Cocaine abuse with intoxication with perceptual disturbance
- **F14.14** Cocaine abuse with cocaine-induced mood disorder
- **F14.150** Cocaine abuse with cocaine-induced psychotic disorder with delusions
- **F14.151** Cocaine abuse with cocaine-induced psychotic disorder with hallucinations
- **F14.180** Cocaine abuse with cocaine-induced anxiety disorder
- **F14.181** Cocaine abuse with cocaine-induced sexual dysfunction
- **F14.182** Cocaine abuse with cocaine-induced sleep disorder
- **F14.188** Cocaine abuse with other cocaine-induced disorder
- **F14.19** Cocaine abuse with unspecified cocaine-induced disorder
F14.20  Cocaine dependence, uncomplicated
F14.21  Cocaine dependence, in remission
F14.220 Cocaine dependence with intoxication, uncomplicated
F14.221 Cocaine dependence with intoxication delirium
F14.222 Cocaine dependence with intoxication with perceptual disturbance
F14.23  Cocaine dependence with withdrawal
F14.24  Cocaine dependence with cocaine-induced mood disorder
F14.250 Cocaine dependence with cocaine-induced psychotic disorder with delusions
F14.251 Cocaine dependence with cocaine-induced psychotic disorder with hallucinations
F14.280 Cocaine dependence with cocaine-induced anxiety disorder
F14.281 Cocaine dependence with cocaine-induced sexual dysfunction
F14.282 Cocaine dependence with cocaine-induced sleep disorder
F14.288 Cocaine dependence with other cocaine-induced disorder
F14.29  Cocaine dependence with unspecified cocaine-induced disorder
F14.90  Cocaine use, unspecified, uncomplicated
F14.920 Cocaine use, unspecified with intoxication, uncomplicated
F14.921 Cocaine use, unspecified with intoxication delirium
F14.922 Cocaine use, unspecified with intoxication with perceptual disturbance
F14.929 Cocaine use, unspecified with intoxication, unspecified
F14.94  Cocaine use, unspecified with cocaine-induced mood disorder
F14.950 Cocaine use, unspecified with cocaine-induced psychotic disorder with delusions
F14.951 Cocaine use, unspecified with cocaine-induced psychotic disorder with hallucinations
F14.98  Cocaine use, unspecified with other specified cocaine-induced disorder
F14.980 Cocaine use, unspecified with cocaine-induced anxiety disorder
F14.981 Cocaine use, unspecified with cocaine-induced sexual dysfunction
F14.982 Cocaine use, unspecified with cocaine-induced sleep disorder
F14.988 Cocaine use, unspecified with other cocaine-induced disorder
F14.99  Cocaine use, unspecified with unspecified cocaine-induced disorder
F16.10  Hallucinogen abuse, uncomplicated
F16.11  Hallucinogen abuse, in remission
F16.120 Hallucinogen abuse with intoxication, uncomplicated
F16.121 Hallucinogen abuse with intoxication with delirium
F16.122 Hallucinogen abuse with intoxication with perceptual disturbance
F16.14  Hallucinogen abuse with hallucinogen-induced mood disorder
F16.150 Hallucinogen abuse with hallucinogen-induced psychotic disorder with delusions
F16.151 Hallucinogen abuse with hallucinogen-induced psychotic disorder with hallucinations
F16.180 Hallucinogen abuse with hallucinogen-induced anxiety disorder
F16.183 Hallucinogen abuse with hallucinogen persisting perception disorder (flashbacks)
F16.188 Hallucinogen abuse with other hallucinogen-induced disorder
F16.19  Hallucinogen abuse with unspecified hallucinogen-induced disorder
F16.20  Hallucinogen dependence, uncomplicated
F16.21  Hallucinogen dependence, in remission
F16.220 Hallucinogen dependence with intoxication, uncomplicated
F16.221 Hallucinogen dependence with intoxication with delirium
F16.24  Hallucinogen dependence with hallucinogen-induced mood disorder
F16.250 Hallucinogen dependence with hallucinogen-induced psychotic disorder with delusions
F16.251 Hallucinogen dependence with hallucinogen-induced psychotic disorder with hallucinations
F16.280 Hallucinogen dependence with hallucinogen-induced anxiety disorder
F16.283 Hallucinogen dependence with hallucinogen persisting perception disorder (flashbacks)
F16.288 Hallucinogen dependence with other hallucinogen-induced disorder
F16.29  Hallucinogen dependence with unspecified hallucinogen-induced disorder
F16.90  Hallucinogen use, unspecified, uncomplicated
F16.920 Hallucinogen use, unspecified with intoxication, uncomplicated
F16.921 Hallucinogen use, unspecified with intoxication with delirium
F16.94 Hallucinogen use, unspecified with hallucinogen-induced mood disorder
F16.950 Hallucinogen use, unspecified with hallucinogen-induced psychotic disorder with delusions
F16.951 Hallucinogen use, unspecified with hallucinogen-induced psychotic disorder with hallucinations
F16.980 Hallucinogen use, unspecified with hallucinogen-induced anxiety disorder
F16.983 Hallucinogen use, unspecified with hallucinogen persisting perception disorder (flashbacks)
F16.988 Hallucinogen use, unspecified with other hallucinogen-induced disorder
F16.99 Hallucinogen use, unspecified with unspecified hallucinogen-induced disorder
F45.42 Pain disorder with related psychological factors
G89.21 Chronic pain due to trauma
G89.22 Chronic post-thoracotomy pain
G89.28 Other chronic postprocedural pain
G89.29 Other chronic pain
G89.3 Neoplasm related pain (acute) (chronic)
G89.4 Chronic pain syndrome

REVISIONS
05-21-2015 Policy added to the bcbsks.com web site.
07-25-2016 Published 06-24-2016. Effective 07-25-2016.
Description section updated
In Policy Section:
• Added Item E "In outpatient pain management and substance abuse treatment, hair
drug testing and oral fluid drug testing are considered experimental / investigational."
• Updated Policy Guidelines
Rationale section updated
In Coding section:
• Added CPT Codes: 80370, 80371, 80372, 80373, 80374
• Added HCPCS Codes: G0477, G0478, G0479, G0480, G0481, G0482, G0483 (Effective
January 1, 2016)
• Removed HCPCS Codes: G0431, G0434, G6030, G6031, G6032, G6034, G6035, G6036,
G6037, G6038, G6039, G6040, G6041, G6042, G6043, G6044, G6045, G6046, G6047,
G6048, G6049, G6050, G6051, G6052, G6053, G6054, G6055, G6056, G6057, G6058
(Effective January 1, 2016)
References updated
01-01-2017 In Coding section:
• Added CPT Codes: 80305, 80306, 80307 (Effective January 1, 2017)
• Removed CPT Codes: 80300, 80301, 80302, 80303, 80304 (Effective December 31,
2016)
10-01-2017 In Coding section:
• Added a Coding notation that 0020U (Drug test(s), presumptive, with definitive
confirmation of positive results, any number of drug classes, urine, with specimen
verification including DNA authentication in comparison to buccal DNA, per date of service
– Effective 10-01-2017) is a non-covered service.
• Added ICD-10 Codes: F11.11, F14.11, F16.11
02-15-2018 Revised Title replacing "Abuse" with "Use Disorder" to read "Drug Testing in Pain
Management and Substance Abuse Use Disorder Treatment"
Description section updated
In Policy section:
• In Items A 1 a, B, B 1 a, C, D, E and Policy Guidelines, replaced "abuse" with "use
disorder".
REVISIONS

- In Items A, B, B 2, B 3 and Policy Guidelines, replaced "qualitative" with "presumptive".
- In Items C C 2 and Policy Guidelines, replaced "quantitative" with "definitive".

Rationale section updated

In Coding section:
- Updated Nomenclature for CPT and HCPCS codes: 80305, 80306, G0480, G0481, G0782, G0783
- Corrected Nomenclature for CPT code: 80374
- Added CPT codes: 80375, 80376, 83377
- Removed HCPCS codes: G0477, G0478, G0479
- Added 0006U, 0007U to Coding notations.

References updated

07-01-2018
In Coding section:
Added PLA Code: 0051U

01-01-2019
In Coding section:
- Added PLA Code: 0082U
- Deleted PLA Code: 0020U
- Revised PLA Code: 0006U

REFERENCES