

Payment Policy: Urine Specimen Validity Testing

Reference Number: LA.PP.056

Effective Date: 08/2020 Coding Implications
Last Review Date: 07/2024 Revision Log

See Important Reminder at the end of this policy for important regulatory and legal information.

Policy Overview

Urine specimen testing is necessary to treat patients for specific medical problems. Providers use the results to detect and monitor drug levels for medical treatment purposes.

The purpose of this policy is to define payment criteria for urine specimen validity testing to be used in making payment decisions and administering benefits.

Application

Physician Office Laboratory, Independent Laboratories, Qualified Hospital Laboratory, Referring Laboratory, Reference Laboratory

Policy Description

Adulteration testing is the tampering or manipulation of a urine specimen with the intention of altering the test results. This tampering can cause false negative results by destroying drugs present in the urine sample and/or interfering with drug screening results.

CMS guidelines for Drug Testing documented in the National Correct Coding Initiative Policy Manual, Chapter X Pathology and Laboratory Services states, "Providers performing validity testing on urine specimens utilized for drug testing should not separately bill the validity testing. For example, if a laboratory performs a urinary pH, specific gravity, creatinine, nitrates, oxidants, or other tests to confirm that a urine specimen is not adulterated, this testing is not separately billed."

Providers that perform validity/adulteration testing on urine specimens should not separately bill for this testing. Confirmation testing is considered incidental to the more complex procedure (urine definitive drug test) and is clinically integral to the successful outcome of the primary procedure. Laboratory procedure codes in the 80305-80377 and G0480-G0483 ranges, along with 83992 and G0569 include sample validation when performed.

Reimbursement

Louisiana Healthcare Connections will disallow separate reimbursement for testing to confirm that a urine drug specimen is unadulterated. Validity testing is an internal control process that is not separately reportable.

Utilization

The health plan's code editing software will deny laboratory procedure codes 82570 (Creatinine other source) when billed with 80305-80307, 80320-80377, 83992, G0480-G0483, G0659 and will also deny 83986 (Ph Body Fluid, Not Elsewhere Specified) when billed with 80305-80307, 80320-80377, 83992, G0480-G0483 and G0659.



Documentation Requirements

Not applicable

Coding and Modifier Information

This payment policy references Current Procedural Terminology (CPT®). CPT® is a registered trademark of the American Medical Association. All CPT® codes and descriptions are copyrighted 2024, American Medical Association. All rights reserved. CPT codes and CPT descriptions are from current manuals and those included herein are not intended to be all-inclusive and are included for informational purposes only. Codes referenced in this payment policy are for informational purposes only. Inclusion or exclusion of any codes does not guarantee coverage. Providers should reference the most up-to-date sources of professional coding guidance prior to the submission of claims for reimbursement of covered services.

Specimen Validity Codes Which Are Not Covered

CPT/HCPCS Code	Descriptor
82570	Creatinine; other source
83986	pH; body fluid, not otherwise specified

Definitive Urine Drug Testing Procedure Codes

CPT/HCPCS Code	Descriptor	
80305	Drug test(s), presumptive, any number of drug classes, any number of devices or procedures (e.g., immunoassay); capable of being read by direct optical observation only (e.g., dipsticks, cups, cards, cartridges) includes sample validation when performed, per date of service	
80306	Drug test(s), presumptive, any number of drug classes, any number devices or procedures (e.g., immunoassay); read by instrument assisted direct optical observation (e.g., dipsticks, cups, cards, cartridges), includes sample validation when performed, per date of service	
80307	Drug test(s), presumptive, any number of drug classes, any number of devices or procedures, by instrument chemistry analyzers (e.g., utilizing immunoassay [e.g., EIA, ELISA, EMIT, FPIA, IA, KIMS, RIA]), chromatography (e.g., GC, HPLC), and mass spectrometry either with or without chromatography, (e.g., DART, DESI, GC-MS, GC-MS/MS, LC-MS, LC-MS/MS, LDTD, MALDI, TOF) includes sample validation when performed, per date of service	
80320	Alcohols	
80321	Alcohol biomarkers; 1 or 2	
80322	Alcohol biomarkers; 3 or more	
80323	Alkaloids, not otherwise specified	
80324	Amphetamines; 1 or 2	
80325	Amphetamines; 3 or 4	
80326	Amphetamines; 5 or more	
80327	Anabolic steroids; 1 or 2	



CPT/HCPCS Code	Descriptor	
80328	Anabolic steroids; 3 or more	
80329	Analgesics, non-opioid; 1 or 2	
80330	Analgesics, non-opioid; 3-5	
80331	Analgesics, non-opioid; 6 or more	
80332	Antidepressants, serotonergic class; 1 or 2	
80333	Antidepressants, serotonergic class; 3-5	
80334	Antidepressants, serotonergic class; 6 or more	
80335	Antidepressants, tricyclic and other cyclicals; 1 or 2	
80336	Antidepressants, tricyclic and other cyclicals; 3-5	
80337	Antidepressants, tricyclic and other cyclicals; 6 or more	
80338	Antidepressants, not otherwise specified	
80339	Antiepileptics, not otherwise specified; 1-3	
80340	Antiepileptics, not otherwise specified; 4-6	
80341	Antiepileptics, not otherwise specified; 7 or more	
80342	Antipsychotics, not otherwise specified; 1-3	
80343	Antipsychotics, not otherwise specified; 4-6	
80344	Antipsychotics, not otherwise specified; 7 or more	
80345	Barbiturates	
80346	Benzodiazepines; 1-12	
80347	Benzodiazepines; 13 or more	
80348	Buprenorphine	
80349	Cannabinoids, natural	
80350	Cannabinoids, synthetic; 1-3	
80351	Cannabinoids, synthetic; 4-6	
80352	Cannabinoids, synthetic; 7 or more	
80353	Cocaine	
80354	Fentanyl	
80355	Gabapentin, non-blood	
80356	Heroin metabolite	
80357	Ketamine and norketamine	
80358	Methadone	
80359	Methylenedioxyamphetamines (MDA, MDEA, MDMA)	
80360	Methylphenidate	
80361	Opiates, 1 or more	
80362	Opioids and opiate analogs; 1 or 2	
80363	Opioids and Opiate analogs; 3 or 4	
80364	Opioids and Opiate analogs; 5 or more	
80365	Oxycodone	
80366	Pregabalin	
80367	Propoxyphene	
80368	Sedative hypnotics (non-benzodiazepines)	
80369	Skeletal muscle relaxants; 1 or 2	
80370	Skeletal muscle relaxants; 3 or more	



CPT/HCPCS Code	Descriptor		
80371	Stimulants, synthetic		
80372	Tapentadol		
80373	Tramadol		
80374	Stereoisomer (enantiomer) analysis, single drug class		
80375	Drug(s) or substance(s), definitive, qualitative or quantitative, not otherwise specified; 1-3		
80376	Drug(s) or substance(s), definitive, qualitative or quantitative, not otherwise specified; 4-6		
80377	Drug(s) or substance(s), definitive, qualitative or quantitative, not otherwise specified; 7 or more		
83992	Phencyclidine (PCP)		
G0480	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(es), including metabolite(s) if performed		
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		



CPT/HCPCS Code	Descriptor
	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; 22 or more drug class(es), including metabolite(s) if performed
G0659	Drug test(s), definitive, utilizing 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), excluding immunoassays (e.g., IA, EIA, ELISA, EMIT, FPIA) and enzymatic methods (e.g., alcohol dehydrogenase), performed without method or drug-specific calibration, without matrix-matched quality control material, or without use of stable isotope or other universally recognized internal standard(s) for each drug, drug metabolite or drug class per specimen; qualitative or quantitative, all sources, includes specimen validity testing, per day, any number of drug classes

Modifier	Descriptor
NA	Not Applicable

ICD-10 Codes	Descriptor
NA	Not Applicable

Definitions

Physician's Office Laboratory

A physician office laboratory is a laboratory maintained by a physician or group of physicians for performing diagnostic tests in connection with the physician practice.



Qualified Hospital Laboratory

A qualified hospital laboratory is one that provides some clinical laboratory tests 24 hours a day, 7 days a week, to serve a hospital's emergency room that is also available to provide services 24 hours a day, 7 days a week. For the qualified hospital laboratory to meet this requirement, the hospital must have physicians physically present or available within 30 minutes through a medical staff call roster to handle emergencies 24 hours a day, 7 days a week; and hospital laboratory technologists must be on duty or on call at all times to provide testing for the emergency room.

Independent Laboratory

An independent laboratory is one that is independent both of an attending or consulting physician's office and of a hospital that meets at least the requirements to qualify as an emergency hospital as defined in §1861(e) of the Social Security Act (the Act.)

Referring Laboratory

A Medicare-approved laboratory that receives a specimen to be tested and that refers the specimen to another laboratory for performance of the laboratory test.

Reference Laboratory

A Medicare-enrolled laboratory that receives a specimen from another, referring laboratory for testing and that actually performs the test.

Definitive Drug Testing

Drug testing identification methods that can identify individual drugs and detect possible use or non-use of a drug. These tests may be either qualitative or quantitative in nature.

Additional Information

Not Applicable

Related Documents or Resources

Policy Number	Policy Name
LA.CP.MP.50	Outpatient Testing for Drugs of Abuse

References

- 1. Current Procedural Terminology (CPT)®, 2024
- 2. HCPCS Level II, 2024
- 3. Centers for Medicare and Medicare Services (CMS) National Correct Coding Initiative (NCCI) Policy Manual, Chapter X, Pathology and Laboratory Services

Revision History	Review Date	Approval Date
Converted corporate to local policy.	8/15/2020	
Annual Review;	8/30/2022	
Updated dates in the reference section from 2018 to 2020		



Removed clinical and added payment policy in "Important		
Reminder" section		
Annual review; updated copyright and reference dates.	8/01/2023	9/12/23
Annual review, verified codes, dates updated.	7/26/24	7/30/24
Did not send to LDH due to non-material revisions.		

Important Reminder

This payment policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this payment policy; and other available clinical information. LHCC makes no representations and accepts no liability with respect to the content of any external information used or relied upon in developing this payment policy. This payment policy is consistent with standards of medical practice current at the time that this payment policy was approved.

The purpose of this payment policy is to provide a guide to medical necessity, which is a component of the guidelines used to assist in making coverage decisions and administering benefits. It does not constitute a contract or guarantee regarding payment or results. Coverage decisions and the administration of benefits are subject to all terms, conditions, exclusions and limitations of the coverage documents (e.g., evidence of coverage, certificate of coverage, policy, contract of insurance, etc.), as well as to state and federal requirements and applicable LHCC administrative policies and procedures.

This payment policy is effective as of the date determined by LHCC. The date of posting may not be the effective date of this payment policy. This payment policy may be subject to applicable legal and regulatory requirements relating to provider notification. If there is a discrepancy between the effective date of this payment policy and any applicable legal or regulatory requirement, the requirements of law and regulation shall govern. LHCC retains the right to change, amend or withdraw this payment policy, and additional clinical policies may be developed and adopted as needed, at any time.

This payment policy does not constitute medical advice, medical treatment or medical care. It is not intended to dictate to providers how to practice medicine. Providers are expected to exercise professional medical judgment in providing the most appropriate care, and are solely responsible for the medical advice and treatment of members. This payment policy is not intended to recommend treatment for members. Members should consult with their treating physician in connection with diagnosis and treatment decisions.

Providers referred to in this payment policy are independent contractors who exercise independent judgment and over whom LHCC has no control or right of control. Providers are not agents or employees of LHCC.

This payment policy is the property of LHCC. Unauthorized copying, use, and distribution of this payment policy or any information contained herein are strictly prohibited. Providers,



members and their representatives are bound to the terms and conditions expressed herein through the terms of their contracts. Where no such contract exists, providers, members and their representatives agree to be bound by such terms and conditions by providing services to members and/or submitting claims for payment for such services.

©2024 Louisiana Healthcare Connections. All rights reserved. All materials are exclusively owned by Louisiana Healthcare Connections and are protected by United States copyright law and international copyright law. No part of this publication may be reproduced, copied, modified, distributed, displayed, stored in a retrieval system, transmitted in any form or by any means, or otherwise published without the prior written permission of Louisiana Healthcare Connections. You may not alter or remove any trademark, copyright or other notice contained herein. Louisiana Healthcare Connections is a registered trademark exclusively owned by Louisiana Healthcare Connections.