

State of New Mexico Medical Assistance Program Manual

Supplement



SPECIAL COVID-19 SUPPLEMENT #18

DATE: JUNE 2, 2022

TO: MEDICAID PROVIDERS

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BUREAU

SUBJECT: COVID-19 TESTING AND TREATMENT SERVICES AND CODES FOR NEW MEXICO

MEDICAID PROVIDERS

This Supplement supersedes the guidance that was provided in SPECIAL COVID-19 SUPPLEMENT #14, dated DECEMBER 30, 2020. The purpose of this Supplement is to provide guidance and directives for modification of services and program standards related to the national public health emergency associated with the 2019 Novel Coronavirus (COVID-19) outbreak. The purpose of these changes is to assure the continuation of essential services to Medicaid patients without disruption or delay while following Centers for Disease Control and Prevention (CDC) direction to maximize social distancing for the duration of the public health emergency. This supplement adds additional and updated information to Special COVID-19 Supplement #14.

COVID-19 Testing and Treatment Services:

 New Billing Codes for Testing – HSD has added new laboratory billing codes as directed by the Centers for Medicare and Medicaid Services (CMS) for COVID-19 lab testing. These codes are identified in Table 1 below and do not require a NM Medicaid Provider Identification Number and/or National Provider Identification (NPI) number for a referring, rendering/administering, or ordering provider(s).

Providers should follow CMS guidelines and timeframes related to diagnosis coding and code claims https://www.cms.gov/files/document/covid-dear-clinician-letter.pdf.

2. Claims Processing- New Mexico Medicaid has updated their Fee for Service (FFS) claims processing systems to bypass rendering/administering, referring, and ordering exceptions and adjust claims according to the rates listed in Table 1 below for dates of service on March 18, 2020

and subsequent to that date, unless noted otherwise.

Pharmacy Claims Processing:

Effective March 31, 2022, the Department of Health (DOH) DOH issued a standing order to bill Medicaid claims for over-the-counter (OTC) At-home COVID-19 Antigen Tests. Submission of a pharmacist's NPI shall be permitted as the ordering provider under their pharmacy's Medicaid enrollment.

For OTC At-Home COVID-19 Antigen Test billing, the pharmacy must populate the following National Council for Prescription Drug Programs (NCPDP) fields:

Field #	NCPDP Field Name	Value
405-D5	Days' Supply	A valid day supply
407-D7	Product Service ID	A valid NDC#
409-D9	Ingredient Cost Submitted	Lesser of methodology
411-DB	Prescriber ID	National Provider Identifier (NPI)
414-DE	Date Prescription Written	MMDDYYYY
419-DJ	Prescription Origin Code	1=Written, 2=Telephone, 3=Electronic,
		4=Facsimile, 5=Transfer
442-E7	Quantity Dispensed	Metric Decimal Quantity

- **3.** Confirmed COVID-19 Diagnosis- Effective for both diagnostic and screenings services on and after April 1, 2020, a confirmed diagnosis of COVID-19 (2019 novel coronavirus disease) should be reported with a diagnosis code **U07.1**, COVID-19. Assignment of this code is applicable to positive COVID-19 test results and presumptive positive COVID-19 test results.
- 4. Drive-through Testing/Screening HSD requests that contracted providers continue to work with their contracted MCOs, in coordination with DOH, to operate "drive-up" or "drive-through" COVID-19 testing and screening services, including the use of this strategy in rural/frontier areas, to the greatest extent possible. This strategy will help to alleviate the impact of crowding in medical clinics and facilities and mitigate the spread of COVID-19. Drive-through testing will be billed in accordance with current rules dependent on provider type and the associated facility where the testing is performed.
- 5. Diagnostic and Screening Testing Coverage: Without cost sharing, all types of Food and Drug Administration (FDA)-authorized COVID-19 diagnostic and screening tests consistent with the CDC recommendations will be covered, including "point of care" or "at-home" COVID-19 antigen testing provided to a Medicaid or CHIP beneficiary. Coverage of screening for testing to return to school or work or to meet travel requirements will be included.

6. Antibody Testing for COVID-19

HSD will only pay for FDA-authorized serologic testing that has been shown to be reliable based on independent testing. Serological tests for COVID-19 are used to detect antibodies against the SARS-CoV-2 virus and are intended for use in the diagnosis of the disease or condition of having current or past infection with SARS-CoV-2, the virus which causes COVID-19. The FDA currently believes such tests should not be used as the sole basis for diagnosis. The FDA has advised the

Departments that serological tests for COVID-19 meet the definition of an in vitro diagnostic product for the detection of SARS-CoV-2 or the diagnosis of COVID-19. Therefore, plans and issuers must provide coverage for a serological test for COVID-19 that otherwise meets the requirements of section 6001(a)(1) of the Families First Coronavirus Response Act (FFCRA), as amended by section 3201 of the CARES Act https://www.cms.gov/files/document/FFCRA-Part-42-FAQs.pdf.

Please note that serological antibody tests should not be used as the sole basis for obtaining a COVID- 19 diagnosis.

7. Lateral Flow Testing

Please note that HSD is not covering lateral flow testing devices at this time, until further evidence is available regarding their effectiveness.

8. Modification of payment for clinical diagnostic laboratory tests (CDLTs) for the detection of SARS—CoV—2 or the diagnosis of the virus that causes COVID-19 making use of high throughput technologies for HCPCS code U0003, U0004, and U0005 effective January 2, 2021

CMS has established a payment amount of \$75 per test for CDLTs making use of high throughput technologies for the detection of SARS-CoV-2 or the diagnosis of the virus that causes COVID-19, as identified by HCPCS codes U0003 and U0004.

CMS has established a new add-on payment of \$25, as identified by HCPCS code U0005. As required by the HCPCS code U0005 descriptor, this add-on payment may be billed with either HCPCS code U0003 or HCPCS code U0004 when the applicable test is completed within 2 calendar days of the specimen being collected.

Laboratories that do not complete the CDLT making use of high throughput technologies for the detection of SARS-CoV-2 or the diagnosis of the virus that causes COVID-19 within 2 calendar days may not bill HCPCS code U0005 and will not receive the \$25 add-on payment. Payment for these CDLTs will be \$75.

Under the Medicare guidance, the responsibility is with providers to determine their eligibility to bill for this additional payment and is subject to audit or medical review (see page 7 of the CMS policy) https://www.cms.gov/files/document/cms-ruling-2020-1-r2.pdf.

In the event of an audit or medical review, laboratories will need to produce documentation to support the add-on payment established in this ruling, even if such documentation would not otherwise be required under Medicare regulations.

For additional guidance please visit https://www.cms.gov/files/document/cms-ruling-2020-1-r2.pdf.

Table 1. Authorized COVID-19 Laboratory and Other Related Codes

Code	Description	Medicaid FFS Rate		
Laboratory Codes				
0223U (Effective 06/25/2020)	Infectious disease (bacterial or viral respiratory tract infection), pathogen-specific nucleic acid (DNA or RNA), 22 targets including severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), qualitative RT-PCR, nasopharyngeal swab, each pathogen reported as detected or not detected	\$416.78		
0224U (Effective 6/25/2020)	Antibody, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), includes titer(s), when performed	\$42.13		
0225U (Effective 08/10/2020)	Infectious disease (bacterial or viral respiratory tract infection) pathogen-specific DNA and RNA, 21 targets, including severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), amplified probe technique, including multiplex reverse transcription for RNA targets, each analyte reported as detected or not detected	\$416.78		
0226U (Effective 08/10/2020)	Surrogate viral neutralization test (sVNT), severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), ELISA, plasma, serum	\$42.28		
0240U (Effective 10/06/2020)	Infectious disease (viral respiratory tract infection), pathogen- specific RNA, 3 targets (severe acute respiratory syndrome coronavirus 2 [SARS-CoV-2], influenza A, influenza B), upper respiratory specimen, each pathogen reported as detected or not detected	\$142.63		
0241U (Effective 10/06/2020)	Infectious disease (viral respiratory tract infection), pathogen- specific RNA, 4 targets (severe acute respiratory syndrome coronavirus 2 [SARS-CoV-2], influenza A, influenza B, respiratory syncytial virus [RSV]), upper respiratory specimen, each pathogen reported as detected or not detected	\$142.63		
86318 (Effective 07/01/2020)	Immunoassay for infectious agent antibody, qualitative or semiquantitative single step method (e.g., reagent strip)	\$18.09		
86328 (Effective 04/10/2020)	Immunoassay for infectious agent antibody(ies), qualitative or semiquantitative, single step method (eg, reagent strip); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19])	\$45.28		
86769 (Effective 04/10/2020)	Antibody; severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19])	\$42.13		

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87426 (Effective 6/25/2020)	Infectious agent antigen detection by immunoassay technique, (eg, enzyme immunoassay [EIA], enzyme-linked immunosorbent assay [ELISA], fluorescence immunoassay [FIA], immunochemiluminometric assay [IMCA]) qualitative or semiquantitative; severe acute respiratory syndrome coronavirus (eg, SARS-CoV, SARS-CoV-2 [COVID-19])	\$45.23
87428 (Effective 11/10/2020)	Infectious agent antigen detection by immunoassay technique, (eg, enzyme immunoassay [EIA], enzyme-linked immunosorbent assay [ELISA], fluorescence immunoassay [FIA], immunochemiluminometric assay [IMCA]) qualitative or semiquantitative; severe acute respiratory syndrome coronavirus (eg, SARS-CoV, SARS-CoV-2 [COVID-19]) and influenza virus types A and B	\$73.49
87635	Infectious agent detection by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), amplified probe technique	\$51.31
87636	Infectious agent detection by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [Covid-19]) and influenza virus types A and B, multiplex amplified probe technique."	\$142.63
87811 (Effective 10/6/2020)	Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19])	\$41.38
C9803	Hospital Outpatient Clinic Visit Specimen Collection for Severe Acute Respiratory Syndrome Coronavirus2 (SARS-CoV-2) (Coronavirus Disease [COVID-19]), Any Specimen Source	• \$25.46 (rate prior to 1/1/2021) • \$24.67 (effective 1/1/2021)

G2023	Specimen collection for severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), any specimen source	• \$25.46 (rate prior to 1/1/2021) • \$23.46 (effective 1/1/2021)
G2024	Specimen collection for severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), from an individual in a skilled nursing facility or by a laboratory on behalf of a home health agency, any specimen source	\$25.46
U0001	CDC 2019 novel coronavirus (2019-nCoV) real-time RT-PCR diagnostic panel	\$35.92
U0002	2019-nCoV coronavirus, SARS-CoV-2/2019-nCoV (COVID-19), any technique, multiple types of subtypes (includes all targets), non-CDC	\$51.31
U0003	Infectious agent detection by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), amplified probe technique, making use of high throughput technologies as described by CMS-2020-01-R	 \$100.00 (current rate) \$75.00 (effective 1/1/2021)
U0004	2019-nCoV Coronavirus, SARS-CoV-2/2019-nCoV (COVID-19), any technique, multiple types or subtypes (includes all targets), non-CDC, making use of high throughput technologies as described by CMS-2020-01-R	 \$100.00 (current rate) \$75.00 (effective 1/1/2021)
U0005 (Effective 1/1/2021)	Infectious agent detection by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19]), amplified probe technique, CDC or non-CDC, making use of high throughput technologies, completed within 2 calendar days from date and time of specimen collection.	\$25.00

MANAGED CARE ORGANIZATONS:

The above guidance relates to fee-for-service claims submission. For managed care claims, please follow guidance provided by each MCO.

Thank you for your service to New Mexicans during this emergency pandemic. This COVID-19 Supplement will sunset when the Human Services Department determines that the national public health emergency associated with the 2019 Novel Coronavirus (COVID-19) outbreak has been contained. Please contact the Medical Assistance Division at (505) 827-6252 or MADInfo.HSD@state.nm.us if you have any questions regarding this guidance.