Illinois Medicaid COVID-19 Fee Schedule

PLEASE NOTE: New COVID-19 related codes will be added to the HFS system as they are released by the Centers for Medicare and Medicaid Services (CMS) in accordance with the <u>December 8, 2020 provider notice</u>. Claims containing new codes which do not have a Medicare or National Government Services (NGS) rate will temporarily suspend until a code rate is assigned. Once a rate is assigned, the HFS system will be updated with that rate and any held claims released into processing.

COVID-19 Vaccines and Vaccine Administration

COVID-19 vaccine product procedure codes are included as a reference but should not be billed when obtained at zero cost to the provider. COVID-19 vaccine administration procedure codes are billable by Physicians, Advance Practice Nurses (APNs), Physician Assistants (PAs), Local Health Departments, Federally Qualified Health Centers (FQHCs), Rural Health Clinics (RHCs), Encounter Rate Clinics (ERCs), and School-Based Health Centers (SBHCs).

<u>*Please Note*</u>: FQHCs, RHCs and ERCs must submit COVID-19 vaccine administration codes fee-forservice separately from an encounter claim, even if the vaccine was administered during a face-to face encounter with a practitioner.

Procedure	Effective	Description	State Max Amount
Code	Date		
91300	12/11/2020	Pfizer-BioNTech Covid-19 Vaccine (Aged 12 years and older) (Purple Cap)	N/A (currently government supplied at no cost to the provider)
0001A	12/11/2020 - 3/14/2021	Pfizer-BioNTech Covid-19 Vaccine (Purple Cap) Administration – First Dose	16.94
0001A	3/15/2021	Pfizer-BioNTech Covid-19 Vaccine (Purple Cap) Administration – First Dose	42.14
0002A	12/11/2020 - 3/14/2021	Pfizer-BioNTech Covid-19 Vaccine (Purple Cap) Administration – Second Dose	28.39
0002A	3/15/2021	Pfizer-BioNTech Covid-19 Vaccine (Purple Cap) Administration – Second Dose	42.14
0003A	8/12/2021	Pfizer-BioNTech Covid-19 Vaccine (Purple Cap) Administration – Third Dose	42.14
0004A	9/22/2021	Pfizer-BioNTech Covid-19 Vaccine (Purple Cap) Administration – Booster	42.14
91301	12/18/2020	Moderna Covid-19 Vaccine (Aged 12 years and older) (Red Cap)	N/A (currently government supplied at no cost to the provider)

0011A	12/18/2020 -	Moderna Covid-19 Vaccine (Red	16.94
	3/14/2021	Cap) Administration – First Dose	
0011A	3/15/2021	Moderna Covid-19 Vaccine (Red Cap) Administration – First Dose	42.14
0012A	12/18/2020 - 3/14/2021	Moderna Covid-19 Vaccine (Red Cap) Administration – Second Dose	28.39
0012A	3/15/2021	Moderna Covid-19 Vaccine (Red Cap) Administration – Second Dose	42.14
0013A	8/12/2021	Moderna Covid-19 Vaccine (Red Cap) Administration – Third Dose	42.14
91303	2/27/2021	Janssen Covid-19 Vaccine (Aged 18 years and older)	N/A (currently government supplied at no cost to the provider)
0031A	2/27/2021 – 3/14/2021	Janssen (Johnson & Johnson) COVID- 19 Vaccine Administration – Single Dose	28.39
0031A	3/15/2021	Janssen (Johnson & Johnson) COVID- 19 Vaccine Administration – Single Dose	42.14
0034A	10/20/2021	Janssen (Johnson & Johnson) COVID- 19 Vaccine Administration - Booster	42.14
91304	7/13/2022 – 8/21/2022	Novavax Covid-19 Vaccine, Adjuvanted (Aged 18 years and older)	N/A (currently government supplied at no cost to the provider)
91304	8/22/2022	Novavax Covid-19 Vaccine, Adjuvanted (Aged 12 years and older)	N/A (currently government supplied at no cost to the provider)
0041A	7/13/2022	Novavax Covid-19 Vaccine, Adjuvanted Administration – First Dose	41.80
0042A	7/13/2022	Novavax Covid-19 Vaccine, Adjuvanted Administration – Second Dose	41.80
91305	1/3/2022	Pfizer-BioNTech Covid-19 Vaccine Pre-Diluted (Aged 12 years and older) (Gray Cap)	N/A (currently government supplied at no cost to the provider)

0051A	1/3/2022	Pfizer-BioNTech Covid-19 Vaccine Pre-Diluted (Gray Cap) Administration - First dose	41.80
0052A	1/3/2022	Pfizer-BioNTech Covid-19 Vaccine Pre-Diluted (Gray Cap) Administration - Second dose	41.80
0053A	1/3/2022	Pfizer-BioNTech Covid-19 Vaccine Pre-Diluted (Gray Cap) Administration - Third dose	41.80
0054A	1/3/2022	Pfizer-BioNTech Covid-19 Vaccine Pre-Diluted (Gray Cap) Administration - Booster	41.80
91306	10/20/2021	Moderna Covid-19 Vaccine (Aged 18 years and older) (Red Cap) (Low Dose)	N/A (currently government supplied at no cost to the provider)
0064A	10/20/2021	Moderna Covid-19 Vaccine (Red Cap) (Low Dose) Administration - Booster	42.14
91307	10/29/2021	Pfizer-BioNTech Covid-19 Pediatric Vaccine (Aged 5 years through 11 years) (Orange Cap)	N/A (currently government supplied at no cost to the provider)
0071A	10/29/2021	Pfizer-BioNTech Covid-19 Pediatric Vaccine (Orange Cap) - Administration - First dose	42.14
0072A	10/29/2021	Pfizer-BioNTech Covid-19 Pediatric Vaccine (Orange Cap) - Administration - Second dose	42.14
0073A	01/03/2022	Pfizer-BioNTech Covid-19 Pediatric Vaccine (Orange Cap) - Administration - Third dose	41.80
0074A	05/17/2022	Pfizer-BioNTech Covid-19 Pediatric Vaccine (Orange Cap) - Administration - Booster	41.80
91308	06/17/2022	Pfizer-BioNTech Covid-19 Pediatric Vaccine (Aged 6 months through 4 years) (Maroon Cap)	N/A (currently government supplied at no cost to the provider)
0081A	06/17/2022	Pfizer-BioNTech COVID-19 Pediatric Vaccine (Aged 6 months through 4 years) (Maroon Cap) - Administration - First dose	41.80
0082A	06/17/2022	Pfizer-BioNTech COVID-19 Pediatric Vaccine (Aged 6 months through 4 years) (Maroon Cap) - Administration - Second dose	41.80

41.80	Pfizer-BioNTech COVID-19 Pediatric	06/17/2022	0083A
	Vaccine (Aged 6 months through 4		
	years) (Maroon Cap) -		
	Administration - Third dose		
N/A	Moderna Covid-19 Vaccine (Aged 6	3/29/2022	91309
(currently government supplied	years through 11 years or aged 18+)		
at no cost to the provider)	(Blue Cap with purple border)		
	50MCG/0.5ML		
41.80	Moderna Covid-19 Pediatric Vaccine	6/17/2022	0091A
	(Aged 6 years through 11 years)		
	(Blue Cap with purple border)		
	Administration - First dose		
41.80	Moderna Covid-19 Pediatric Vaccine	6/17/2022	0092A
	(Aged 6 years through 11 years)		
	(Blue Cap with purple border) -		
	Administration - Second dose		
41.80	Moderna Covid-19 Pediatric Vaccine	6/17/2022	0093A
	(Aged 6 years through 11 years)	0, _,	
	(Blue Cap with purple border) -		
	Administration - Third dose		
41.80	Moderna Covid-19 Vaccine (Aged 18	3/29/2022	0094A
+1.00	years and older) (Blue Cap with	5/25/2022	0054/1
	purple border) 50MCG/0.5ML		
	Administration - Booster		
N/A	Moderna Covid-19 Pediatric Vaccine	6/17/2022	91311
-	(Aged 6 months through 5 years)	0/1//2022	91311
currently government supplied) at no cost to the provider)	(Blue Cap with magenta border)		
at no cost to the provider)	250MCG/0.25ML		
41.80	Moderna Covid-19 Pediatric Vaccine	6/17/2022	0111A
41.00	(Aged 6 months through 5 years)	0/1//2022	
	(Blue Cap with magenta border) -		
	Administration - First dose		
41.80	Moderna Covid-19 Pediatric Vaccine	6/17/2022	0112A
41.00		0/1//2022	UTIZA
	(Aged 6 months through 5 years) (Blue Cap with magenta border) -		
44.00	Administration - Second dose	6/17/2022	01124
41.80	Moderna Covid-19 Pediatric Vaccine	6/1//2022	0113A
	(Aged 6 months through 5 years)		
	(Blue Cap with magenta border) -		
	Administration - Third dose	0.10.1.10.000	
N/A	Pfizer-BioNTech COVID-19 Vaccine,	8/31/2022 -	91312
(currently government supplied	Bivalent Product (Aged 12 years and	10/11/2022	
at no cost to the provider)	older) (Gray Cap)		
N/A	Pfizer-BioNTech COVID-19 Vaccine,	10/12/2022	91312
(currently government supplied	Bivalent Product (Aged 5 years and		
(currently government supplied	Divalent i Touuci (Ageu 5 years and		

0124A	8/31/2022	Pfizer-BioNTech COVID-19 Vaccine, Bivalent (Gray Cap) Administration – Booster Dose	41.80
91313	8/31/2022 - 10/11/2022	Moderna COVID-19 Vaccine, Bivalent Product (Aged 18 years and older) (Dark Blue Cap with gray border)	N/A (currently government supplied at no cost to the provider)
91313	10/12/2022	Moderna COVID-19 Vaccine, Bivalent Product (Aged 6 years and older) (Dark Blue Cap with gray border)	N/A (currently government supplied at no cost to the provider)
0134A	8/31/2022 – 10/11/2022	Moderna COVID-19 Vaccine, Bivalent (Aged 18 years and older) (Dark Blue Cap with gray border) Administration – Booster Dose	41.80
91314	10/12/2022	Moderna COVID-19 Vaccine, Bivalent Product (Aged 6 years through 11 years) (Dark Blue Cap with gray border)	N/A (currently government supplied at no cost to the provider)
0144A	10/12/2022	Moderna COVID-19 Vaccine, Bivalent (Aged 6 years through 11 years) (Dark Blue Cap with gray border) Administration – Booster Dose	41.80
91315	10/12/2022	Pfizer-BioNTech COVID-19 Vaccine, Bivalent Product (Aged 5 years through 11 years) (Orange Cap)	N/A (currently government supplied at no cost to the provider)
0154A	10/12/2022	Pfizer-BioNTech COVID-19 Vaccine, Bivalent Product (Aged 5 years through 11 years) (Orange Cap) Administration – Booster Dose	41.80
M0201	6/8/2021	COVID-19 Vaccine Administration Inside a Patient's Home Note : please see the HFS 7/2/21 provider notice for information ; however, per <u>CMS</u> billing of this code is no longer limited to once per DOS, per home effective 8/24/21	37.05

Vaccine Counseling

Billable by Local Health Departments, Physicians, Advance Practice Nurses (APNs), Physician Assistants (PAs), and School-Based Health Centers (SBHCs). Intended to provide reimbursement for the additional time needed for parental/caregiver counseling and informed consent for the COVID-19 vaccination of children ages 6 months through 20. **Note*: this code is *not billable as a telehealth service.*

Procedure Code	Effective Date	Ages	Description	State Max Amount
99402	10/29/2021 for ages 5y-20y 6/17/2022 for ages 6mos-5y	6 months – 20 years	Preventive medicine counseling and/or risk factor reduction intervention(s) provided to an individual (separate procedure); approximately 30 min.	30.00

Virtual Healthcare/Telehealth Expansion

Billable by Local Health Departments, Physicians, Advance Practice Nurses (APNs), and Physician Assistants (PAs) – including physicians, APNs, and PAs rendering the service in a Federally Qualified Health Center (FQHC), Rural Health Clinic (RHC), Encounter Rate Clinic (ERC), or School Based Health Center (SBHC):

**Note*: all virtual healthcare/telehealth codes must be billed with place of service 02 (or place of service 10 if applicable and date of service is on/after 7/1/2022), and modifier GT (or modifier 93 if applicable and date of service is on/after 7/1/2022).

Procedure Code	Effective Date	Description	State Max Amount
G0406	3/9/2020	Follow-up inpatient consultation, limited, physicians typically spend 15 minutes communicating with the patient via telehealth	39.17
G0407	3/9/2020	Follow-up inpatient consultation, intermediate, physicians typically spend 25 minutes communicating with the patient via telehealth	72.13
G0408	3/9/2020	Follow-up inpatient consultation, complex, physicians typically spend 35 minutes communicating with the patient via telehealth	103.70
G0425	3/9/2020	Telehealth consultation, emergency department or initial inpatient, typically 30 minutes communicating with the patient via telehealth	100.35
G0426	3/9/2020	Telehealth consultation, emergency department or initial inpatient, typically 50 minutes communicating with the patient via telehealth	136.14
G0427	3/9/2020	Telehealth consultation, emergency department or initial inpatient, typically 70 minutes or more communicating with the patient via telehealth	201.99
G2010	3/9/2020	Remote evaluation of recorded video and/or images submitted by an established patient (e.g., store and forward), including interpretation with follow-up with the patient within 24 business hours, not originating from a related e/m service provided within the previous 7 days nor leading to an e/m service or procedure within the next 24 hours or soonest available appointment	9.24

G2012	3/9/2020	Brief communication technology-based service, e.g. virtual check-in, by a physician or other qualified health care professional who can report evaluation and management services, provided to an established patient, not originating from a related e/m service provided within the previous 7 days nor leading to an e/m service or procedure within the next 24 hours or soonest available appointment; 5-10 minutes of medical discussion	13.19
G2061	3/9/2020 – 12/31/2020	Qualified nonphysician healthcare professional online assessment, for an established patient, for up to seven days, cumulative time during the 7 days; 5-10 minutes	12.10
G2062	3/9/2020 – 12/31/2020	Qualified nonphysician healthcare professional online assessment service, for an established patient, for up to seven days, cumulative time during the 7 days; 11-20 minutes	21.37
G2063	3/9/2020 – 12/31/2020	Qualified nonphysician qualified healthcare professional assessment service, for an established patient, for up to seven days, cumulative time during the 7 days; 21 or more minutes	33.14
G2250	1/1/2021	Remote assessment of recorded video and/or images submitted by an established patient (e.g., store and forward), including interpretation with follow-up with the patient within 24 business hours, not originating from a related service provided within the previous 7 days nor leading to a service or procedure within the next 24 hours or soonest available appointment	9.24
G2251	1/1/2021	Brief communication technology-based service, e.g. virtual check-in, by a qualified health care professional who cannot report evaluation and management services, provided to an established patient, not originating from a related service provided within the previous 7 days nor leading to a service or procedure within the next 24 hours or soonest available appointment; 5-10 minutes of clinical discussion	13.05
G2252	1/1/2021	Brief communication technology-based service, e.g. virtual check-in, by a physician or other qualified health care professional who can report evaluation and management services, provided to an established patient, not originating from a related e/m service provided within the previous 7 days nor leading to an e/m service or procedure within the next 24 hours or soonest available appointment; 11-20 minutes of medical discussion	25.14
98970	1/1/2021	Qualified nonphysician health care professional online digital assessment and management, for an established patient, for up to 7 days, cumulative time during the 7 days; 5-10 min.	11.36
98971	1/1/2021	Qualified nonphysician health care professional online digital assessment and management, for an established patient, for up to 7 days, cumulative time during the 7 days; 11-20 min.	20.31

98972	1/1/2021	Qualified nonphysician health care professional online digital assessment and management, for an established patient, for up to 7 days, cumulative time during the 7 days; 21+ min.	32.41
99421	3/9/2020	Online Digital Evaluation and Management Service, for an established patient, for up to 7 days, cumulative time during the 7 days; 5-10 minutes	13.19
99422	3/9/2020	Online Digital Evaluation and Management Service, for an established patient, for up to 7 days, cumulative time during the 7 days; 11-20 minutes	27.14
99423	3/9/2020	Online Digital Evaluation and Management Service, for an established patient, for up to 7 days, cumulative time during the 7 days; 21 or more minutes	43.23

Please Note: Evaluation and management services rendered by Physicians, Advance Practice Nurses, and Physician Assistants to new or existing patients using audio only telephonic equipment may be billed as a distant site telehealth service so long as the E/M service is of an amount and nature that would be sufficient to meet the key components of a face-to-face encounter. In this scenario, the claim must be submitted with place of service 02 (or 10 if applicable and the date of service is on/after 7/1/2022) and modifier GT (or 93 if applicable and the date of service is on/after 7/1/2022) appended to the applicable procedure code.

If an audio only telephonic interaction cannot meet key components of a face -to-face encounter, the provider may instead seek reimbursement for virtual check-in services using CPT code G2012. FQHCs/RHCs/ERCs will be reimbursed at the above rates (not their medical encounter rate) for virtual check-in and E-visit codes. Virtual check-in and e-visit/online portal services must be submitted fee-for-service without the T1015 encounter code.

Virtual Healthcare/Teledentistry Expansion

Billable by Dentists:

Procedure Code	Effective Date	Description	State Max Amount
D9995	3/9/2020	Teledentistry, synchronous; real-time encounter	13.19
D9996	3/9/2020	Teledentistry asynchronous; information stored and forwarded to dentist for subsequent review	9.24

Note: The above codes must be billed in addition to D0140, with place of service 02 (or 10, if applicable and date of service is on/after 7/1/2022). Do not include the GT or 93 modifier.

COVID-19 Treatment

COVID-19 antibody product procedure codes are included as a reference but should not be billed when obtained at zero cost to the provider. Only Q0247 was previously practitioner-purchased until it became unauthorized effective April 5, 2022.

Antibody treatment administration codes are billable by Physicians, Advance Practice Nurses (APNs), and Physician Assistants (PAs). Federally Qualified Health Centers (FQHCs), Rural Health Clinics (RHCs), and Encounter Rate Clinics (ERCs) may bill antibody treatment administration codes as detail codes with an encounter claim.

Hospitals may bill the antibody treatment administration codes marked with a double asterisk () using revenue code 0771. Reimbursement is based on DRG (inpatient setting) or EAPG (outpatient setting) methodology.

Procedure Code	Effective Date	Description & Labeler Name	State Max Amount
J0248	12/23/2021	Injection, REMDESIVIR, 1 mg	**Billable only by hospitals on
		Please reference the <u>10/21/22</u>	the 8371. Reimbursed using
		provider notice for details	EAPG methodology.
Q0220	12/8/2021	Tixagev and Cilgav, 300mg	N/A
			(currently government supplied
			at no cost to the provider)
M0220	12/8/2021	Injection, Tixagevimab and	150.50
		Cilgavimab, for the pre-exposure	
		prophylaxis only, for certain adults	
		and pediatric individuals (12 years	
		of age and older weighing at least	
		40kg) with no known sars-cov-2	
		exposure, who either have	
		moderate to severely compromised	
		immune systems or for whom	
		vaccination with any available	
		covid-19 vaccine is not	
		recommended due to a history of	
		severe adverse reaction to a covid-	
		19 vaccine(s) and/or covid-19	
		vaccine component(s), includes	
		injection and post administration	
		monitoring	
Q0221	2/24/2022	Injection, Tixagevimab and	N/A
		Cilgavimab, for the pre-exposure	(currently government supplied
		prophylaxis only, for certain adults	at no cost to the provider)
		and pediatric individuals (12 years	
		of age and older weighing at least	
		40kg) with no known sars-cov-2	
		exposure, who either have	
		moderate to severely compromised	
		immune systems or for whom	
		vaccination with any available	
		covid-19 vaccine is not	
		recommended due to a history of	

		severe adverse reaction to a covid- 19 vaccine(s) and/or covid-19 vaccine component(s), 600 mg	
M0221**	12/8/2021	Injection, Tixagevimab and Cilgavimab, for the pre-exposure prophylaxis only, for certain adults and pediatric individuals (12 years of age and older weighing at least 40kg) with no known sars-cov-2 exposure, who either have moderate to severely compromised immune systems or for whom vaccination with any available covid-19 vaccine is not recommended due to a history of severe adverse reaction to a covid- 19 vaccine(s) and/or covid-19 vaccine component(s), includes injection and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	**Billable only by hospitals on the 837I. Reimbursed using EAPG methodology.
Q0222	2/11/2022	Injection, Bebtelovimab, 175 mg	N/A (currently government supplied at no cost to the provider)
M0222	2/11/2022	Intravenous injection, Bebtelovimab, includes injection and post administration monitoring	350.50
M0223**	2/11/2022	Intravenous injection, Bebtelovimab, includes injection and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	**Billable only by hospitals on the 837I. Reimbursed using EAPG methodology.
Q0243	11/21/2020 – 1/23/2022	Casirivimab and Imdevimab, 2400 mg (Regeneron)	N/A (currently government supplied at no cost to the provider)
M0243**	5/6/2021 – 1/23/2022	Intravenous infusion, Casirivimab and Imdevimab, includes infusion	450.00

		and post administration monitoring (Regeneron)	**Hospitals billing on the 837I will be reimbursed using EAPG methodology.
M0244**	5/6/2021 – 1/23/2022	Intravenous infusion or subcutaneous injection, Casirivimab and Imdevimab includes infusion or injection, and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	**Billable only by hospitals on the 837I. Reimbursed using EAPG methodology.
Q0245	2/9/2021 – 1/23/2022	Injection, Bamlanivimab and Etesevimab, 2100 mg (Eli Lilly)	N/A (currently government supplied at no cost to the provider)
M0245	5/6/2021 – 1/23/2022	Intravenous infusion, Bamlanivimab and Etesevimab, includes infusion and post administration monitoring (Eli Lilly)	450.00
M0246**	5/6/2021 – 1/23/2022	Intravenous infusion, Bamlanivimab and Etesevimab, includes infusion and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	**Billable only by hospitals on the 837I. Reimbursed using EAPG methodology.
Q0247	10/1/2021 - 4/4/2022	Injection, Sotrovimab, 500 mg	2394.00
M0247	7/1/2021 - 4/4/2022	Intravenous infusion, Sotrovimab, includes infusion and post administration monitoring	450.00
M0248**	5/26/2021 – 4/4/2022	Intravenous infusion, Sotrovimab, includes infusion and post administration monitoring in the home or residence; this includes a beneficiary's home that has been made provider-based to the hospital during the covid-19 public health emergency	**Billable only by hospitals on the 837I. Reimbursed using EAPG methodology.
Q0249	6/24/2021	Injection, Tocilizumab, for hospitalized adults and pediatric patients (2 years of age and older) with covid-19 who are receiving systemic corticosteroids and require	N/A (currently government supplied at no cost to the provider)

	1		1
		supplemental oxygen, non-invasive or invasive mechanical ventilation,	
		-	
		or extracorporeal membrane	
		oxygenation (ECMO) only, 1 mg	
M0249**	6/24/2021	Intravenous infusion, Tocilizumab,	**Billable only by hospitals on
		for hospitalized adults and pediatric	the 837I. Reimbursed using DRG
		patients (2 years of age and older)	methodology.
		with covid-19 who are receiving	
		systemic corticosteroids and require	
		supplemental oxygen, non-invasive	
		or invasive mechanical ventilation,	
		or extracorporeal membrane	
		oxygenation (ECMO) only, includes	
		infusion and post administration	
		monitoring, first dose	
M0250**	6/24/2021	Intravenous infusion, Tocilizumab,	**Billable only by hospitals on
	0, = ., =0==	for hospitalized adults and pediatric	the 8371. Reimbursed using DRG
		patients (2 years of age and older)	methodology.
		with covid-19 who are receiving	methodology.
		systemic corticosteroids and require	
		supplemental oxygen, non-invasive	
		or invasive mechanical ventilation,	
		or extracorporeal membrane	
		oxygenation (ECMO) only, includes	
		infusion and post administration	
		monitoring, second dose	

Laboratory Services

Billable by Local Health Departments, Physicians, Advance Practice Nurses (APNs), Physician Assistants (PAs), and Independent Laboratories. Federally Qualified Health Centers (FQHCs), Rural Health Clinics (RHCs), and Encounter Rate Clinics (ERCs) may bill the following laboratory services as detail codes with an encounter claim when the laboratory service is performed on-site. Please note that providers normally subject to a SMART Act rate reduction (e.g. Independent Labs) must include modifier CR to exempt the COVID-related service from the rate reduction when billing outside HFS' COVID portal. It is not necessary to include the modifier when billing through the portal.

These rates also apply to hospitals who choose to have the Department generate their claim(s) within the HFS COVID Portal per the <u>February 8, 2021 provider notice</u> for uninsured individuals. For Medicaid-eligible participants, hospitals must bill on an institutional invoice and will be reimbursed via the EAPG methodology.

Procedure Code	Effective Date	Description	State Max Amount
0202U	5/20/2020	Infectious disease (bacterial or viral respiratory tract infection), pathogen specific nucleic acid (DNA or RNA), 22	250.07

		targets including severe acute respiratory syndrome	
		coronavirus 2 (SARS-CoV-2),	
		qualitative RT-PCR, nasopharyngeal swab, each pathogen	
		reported as detected or not detected	
0223U	6/25/2020	Infectious disease (bacterial or viral respiratory tract	416.78
		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	
0224U	6/25/2020	Antibody, severe acute respiratory syndrome coronavirus 2	42.13
		(SARS-CoV-2) (Coronavirus disease [COVID-19]), includes	
		titer(s), when performed	
0225U	8/10/2020	Infectious disease (bacterial or viral respiratory tract	416.78
		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	
0226U	8/10/2020	Surrogate viral neutralization test (sVNT), severe acute	42.28
		respiratory syndrome coronavirus 2 (SARS-CoV-2)	
		(Coronavirus disease [COVID-19]), ELISA, plasma, serum	
0240U	10/6/2020	Infectious disease (viral respiratory tract infection),	142.63
		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	
0241U	10/6/2020	Infectious disease (viral respiratory tract infection),	142.63
		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	
86318	4/10/2020	Immunoassay for infectious agent antibody (ies), qualitative	16.90
		or semiqualitative, single step method (e.g. reagent strip)	
86328	4/10/2020	Immunoassay for infectious agent antibody (ies), qualitative	45.23
		or semiquantitative, single-step method (e.g., reagent strip);	
		severe acute respiratory syndrome coronavirus 2 (SARS-CoV-	
		2) (Coronavirus disease [COVID-19])	
86408	8/10/2020	Neutralizing antibody, severe acute respiratory syndrome	42.13
		coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-	
		19]); screen	
86409	8/10/2020	Neutralizing antibody, severe acute respiratory syndrome	105.33
		coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-	
		19]); titer	

86413	9/8/2020	Severe acute respiratory syndrome coronavirus 2 (SARS-CoV- 2) (Coronavirus disease [COVID-19]) antibody, quantitative	51.43
86769	4/10/2020	Antibody; severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (Coronavirus disease [COVID-19])	42.13
87426	6/25/2020	Infectious agent antigen detection by immunoassay technique, (eg, enzyme immunoassay [EIA], enzyme-linked immunosorbent assay [ELISA], immunochemiluminometric assay [IMCA]) qualitative or semiquantitative, multiple-step method; severe acute respiratory syndrome coronavirus (eg, SARS-CoV, SARS-CoV-2 [COVID-19])	35.33
87428	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 SD: SARSCOV & INF VIR A&B AG IA	63.59
87635	3/13/2020	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	10/6/2020	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
87637	10/6/2020	Infectious agent detection by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV- 2) (Coronavirus disease [COVID-19]), influenza virus types A and B, and respiratory syncytial virus, multiplex amplified probe technique	142.63
87811	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
U0001	2/4/2020	CDC 2019-Novel Coronavirus real-time RT-PCR diagnostic panel	35.91
U0002	2/4/2020	Coronavirus (COVID-19) SARS-COV-2/2019-NCOV, Non-CDC Lab Test	51.31
U0003	4/14/2020 through 2/28/2021	Infectious Agent Detection by Nucleic Acid (DNA or RNA); SARS-COV-2, COVID-19, Amplified Probe Technique, High Throughput Technologies	100.00
U0003	3/1/2021	Infectious Agent Detection by Nucleic Acid (DNA or RNA); SARS-COV-2, COVID-19, Amplified Probe Technique, High Throughput Technologies	75.00

U0004	4/14/2020	2019-NCOV Coronavirus, SARS-CoV-2/2019-nCoV (COVID-	100.00
	through	19), Any Technique, Multiple Subtypes, Non-CDC, High	
	2/28/2021	Throughput Technologies	
U0004	3/1/2021	2019-NCOV Coronavirus, SARS-CoV-2/2019-nCoV (COVID-	75.00
		19), specimen collection. (⁺ add-on to U0003 or U0004 Any	
		Technique, Multiple Subtypes, Non-CDC, High Throughput	
		Technologies	
+U0005	3/1/2021	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; List separately in addition to either HCPCS code U0003 or U0004) <u>NOTE:</u> certain conditions must be met to bill this code; refer to the <u>02/26/2021</u> provider notice for billing guidelines	25.00

COVID-19 Diagnostic Testing Specimen Collection

*Billable by Local Health Departments, Physicians, Advance Practice Nurses (APNs), Physician Assistants (PAs), and Federally Qualified Health Centers (FQHCs) with drive-thru testing sites. FQHCs may bill fee-for-service when there is not a billable medical encounter. Please note that providers normally subject to a SMART Act rate reduction (e.g. Independent Labs) must include modifier CR to exempt the COVID-related service from the rate reduction when billing outside HFS' COVID portal. It is not necessary to include the modifier when billing through the portal.

**Billable by Local Health Departments, Physicians, Advance Practice Nurses (APNs), Physician Assistants (PAs), and Independent Labs.

Procedure Code	Effective Date	Description	State Max Amount
99000*	3/18/2020	Handling of Specimen for Transfer from Office to a Lab	23.46
G2023**	3/1/2020	Specimen Collection, SARS-CoV-2, COVID-19, any specimen source	23.46
G2024***	3/1/2020	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

***Billable by Independent Labs only.

COVID-19 Testing and Testing-Related Services for the Uninsured Population

The following procedure codes are covered for the uninsured population for the purposes of COVID-19 testing and testing-related services for dates of service beginning March 18, 2020. Testing-related services include those directly related to the administration of an in vitro diagnostic product described in section 1905(a)(3)(B) of the Social Security Act, or to the evaluation of a patient for purposes of determining the need for such product. *PLEASE NOTE: All claims for the uninsured population must contain a diagnosis code indicating the patient encounter was for the purposes of COVID testing or COVID vaccine administration. Paid claims with no COVID or COVID-related diagnosis code are subject to post-payment review and recoupment.*

HCPCS codes: G2010, G2012, G2023, G2024, G2061, G2062, G2063, G2250, G2251, G2252, T1015

CPT codes: All COVID-19 laboratory testing and vaccine administration codes*, 71045, 71046, 71047, 71048, 99000, 99201, 99202, 99203, 99204, 99205, 99211, 99212, 99213, 99214, 99215, 99421, 99422, 99423

**Note:* All COVID-19 laboratory testing and vaccine administration codes are billable for the uninsured population, effective with the specific code effective date.

Rates for these codes may be found on the COVID-19 Fee Schedule above, on the <u>Practitioner Fee</u> <u>Schedule</u> or, in the case of T1015, will be at the FQHC/RHC/ERC provider-specific medical encounter rate. Please note, effective dates for codes found on the COVID-19 fee schedule above are applicable, and CPT 99201 is an obsolete code effective with dates of service beginning 1/1/2021. Providers normally subject to a SMART Act rate reduction must include modifier CR to exempt the COVIDrelated service from the rate reduction when billing outside HFS' COVID portal. It is not necessary to include the modifier when billing through the portal.

As a reminder, Federally Qualified Health Centers (FQHCs) may bill specimen collection procedure code 99000 – Handling of Specimen for Transfer from Office to Lab, as fee-for-service utilizing the full functionality of the COVID portal. However, if an FQHC needs to bill a T1015 encounter for a testingrelated service, it can only use the portal to obtain a RIN. The FQHC will have to submit the encounter claim through its normal billing process outside the portal once a RIN is obtained.