

Concert Infectious Disease: Respiratory Testing

Reference Number: LA.CG.CP.MP.2701
implications

[Coding](#)

Date of Last Revision 02/25

[Revision Log](#)

See [Important Reminder](#) at the end of this policy for important regulatory and legal information.

OVERVIEW

~~Respiratory illnesses cause significant morbidity and mortality within the United States and around the world. Seasonal influenza, respiratory syncytial virus (RSV), and SARS-CoV-2 infect many individuals each year, and while most will recover with no complications, a significant number will be hospitalized or die. Diagnostic testing for upper respiratory tract infections can be very useful for clinicians, as clinical signs and symptoms of these infections can have significant overlap between pathogens. Accurate and rapid testing techniques may aid clinicians, via identification of a specific pathogen, in selecting the best course of treatment for patients. Optimally, treatment is started within 48-72 hours of diagnosis. Testing methods range from culture and microscopy to immunoassays and advanced molecular diagnostic techniques; technology in this space is evolving rapidly and clinical guidelines can lag as a result.~~

~~This policy is intended for use in the outpatient setting.~~

This policy addresses the use of tests for upper respiratory tract infections including multi-pathogen panels. These criteria are intended for use in the outpatient setting.

For additional information see the Rationale and References section.

POLICY REFERENCE TABLE

Criteria <u>SectionsCriteria</u> <u>Sections</u>	Example Tests (Labs)	References <u>Support</u>
<u>Respiratory Pathogen Panel Tests</u>		
<u>Syndromic/Multiplex</u> <u>Respiratory Panels</u> <u>with 6 or More</u>	Respiratory Pathogen Panel, Quest Diagnostics	<u>3Rationale/</u> <u>References</u>
	<u>Biofire Filmarray Pneumonia (PN) Panel (bioMérieux)</u>	

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<u>Targets Syndromic/Multiplex Respiratory Panels with 6 or More Targets</u>	ePlex Respiratory Pathogen Panel (GenMark Diagnostics, Inc)	
	Biofire FilmArray Respiratory Panel 2.1 (Biofire Diagnostics)	
	QIAstat-Dx Respiratory SARS-CoV-2 Panel (QIAGEN Sciences)	
	ePlex <u>Biofire Spotfire</u> Respiratory Pathogen/Sore Throat (R/ST) Panel 2 (GenMark Diagnostics, Inc, Respiratory Menu (bioMérieux))	
	Respiratory Pathogen with ABR (RPX) (Lab Genomics LLC, ThermoFisher Scientific)	
	Biofire Spotfire Respiratory Virus PCR /Sore Throat (R/ST) Panel IV (Quest Diagnostics- Sore Throat Menu (bioMérieux))	
	ePlex Respiratory Viral Pathogen Panel, PCR (Quest 2 (GenMark Diagnostics, Inc))	
	HealthTrackRx Bronchitis - 0556U (HealthTrackRx)	
<u>SARS-CoV-2, RSV, or Influenza A/B, OR Multiplex Respiratory Viral Panels with 5 or Fewer Targets</u> <u>SARS-CoV-2, RSV, or Influenza A/B, OR Multiplex Respiratory Viral Panels with 5 or Fewer Targets</u>	Xpert Xpress SARS-CoV-2/Flu/RSV for SARS-CoV-2 and Flu targets only (Cepheid)	<u>3, 6, 7 Rationale/References</u>
	Xpert Xpress SARS-CoV-2/Flu/RSV for all targets (Cepheid)	
	Infectious Agent Antigen Detection by Immunoassay	

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	Infectious Agent Antigen Detection by Immunoassay, Qualitative or Semiquantitative	
	Infectious Agent Antigen Detection by Immunoassay, Qualitative or Semiquantitative, SARS-CoV-2 and Flu A/B	
	Influenza A and B and RSV RNA, Qualitative, Real-Time RT-PCR (Quest Diagnostics)	
	SARS-CoV-2 RNA (COVID-19), Qualitative NAAT (Quest Diagnostics)	
	SARS-CoV-2 RNA (COVID-19) and Influenza A and B, Qualitative NAAT (Quest Diagnostics)	
	Infectious Agent Antigen Detection by Nucleic Acid (DNA or RNA) SARS-CoV-2/Flu/RSV Multiplex Amplified Probe Technique	
	Infectious Agent Antigen Detection by Immunoassay with Direct Optical Observation	
	<u>Influenza, Single Type, Nucleic Acid Detection</u>	
	<u>Influenza A and B Virus with Subtyping, Real-Time PCR (Quest Diagnostics)</u>	
<u>Bacterial Respiratory Infection/Pneumonia Panels</u> <u>Bacterial Respiratory Infection/Pneumonia Target Panels</u>	Infectious Agent: Chlamydia pneumoniae Detection by Nucleic Acid (DNA or RNA), Direct Probe Technique	<u>3Rationale/References</u>
	Chlamydomphila pneumoniae, DNA, Qualitative, Real-Time PCR (Quest Diagnostics)	
	Infectious Agent: Chlamydia pneumoniae Detection by Nucleic Acid (DNA or RNA), Quantification	
	Legionella DNA, Qualitative, Real-	

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	Time PCR (Quest Diagnostics)	
	Infectious Agent: Mycoplasma pneumoniae Detection by Nucleic Acid (DNA or RNA), Direct Probe Technique	
	Mycoplasma pneumoniae, DNA PCR (Labcorp)	
	Infectious Agent: Mycoplasma pneumoniae Detection by Nucleic Acid (DNA or RNA), Quantification	
<u>Influenza A and B</u>		
Influenza A and B Antibody Tests <u>Influenza A and B Antibody Tests</u>	Influenza Type A and Type B Antibody, Serum (Quest Diagnostics)	1 <u>Rationale/References</u>
<u>Streptococcus</u>		
Group A Streptococcus Pharyngitis Tests <u>Group A Streptococcus Pharyngitis Tests</u>	Streptococcus Group A Antigen Detection by Immunoassay	2 <u>Rationale/References</u>
	Streptococcus Group A Antigen Detection by Nucleic Acid Direct Probe Technique	
	Group A Streptococcus Detection, NAA (Labcorp)	
	Streptococcus Group A Antigen, Adult (Quest Diagnostics)	
Group A Streptococcus Pharyngitis Cultures <u>Group A Streptococcus Pharyngitis Cultures</u>	Streptococcus Group A Culture (Quest Diagnostics)	2, <u>4</u> <u>Rationale/References</u>
Group A Streptococcus Antibody Tests <u>Group A Streptococcus Antibody Tests</u>	Antistreptolysin O (ASO) Antibodies (Labcorp)	2 <u>Rationale/References</u>

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CRITERIA

It is the policy of Louisiana Healthcare Connections that the specific tests noted below are **medically necessary** when meeting the related criteria:

RESPIRATORY PATHOGEN PANEL TESTS

Syndromic/Multiplex Respiratory Panels with 6 or More Targets

I. ~~Syndromic Multiplex Respiratory Panels~~ multiplex respiratory panels with ~~six~~ 6 or more targets, ~~when performed in the outpatient setting,~~ are considered **medically necessary** ~~for~~ when:

A. ~~The member/enrollees meeting any~~ enrollee presents in the outpatient setting with signs or symptoms of an acute respiratory infection, AND

1. The member/enrollee meets at least one of the following criteria:

a) Immunocompromised, OR

b) Has severe pneumonia, OR

c) Has exacerbations of airway disease, AND

B. Results of the testing will influence the member's/enrollee's clinical management.

~~1. With serious or critical illness, OR~~

~~2. At imminent risk of becoming seriously or critically ill, OR~~

~~3. With immunodeficiency, AND/OR~~

~~4. With a severe underlying condition.~~

II. Current evidence does not support the use of ~~Syndromic Multiplex Respiratory Panel~~ syndromic multiplex respiratory panels with 6 or more targets for all other indications.

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SARS-CoV-2, RSV, or Influenza A/B, OR Multiplex Respiratory Viral Panels with 5 or Fewer Targets

- I. SARS-CoV-2, RSV, or Influenza A/B, **OR** ~~Multiplex Respiratory Viral Panels~~multiplex respiratory viral panels with 5 or fewer targets, ~~other than CPT 87631 which is out of the scope of this policy,~~ are considered **medically necessary** when:
 - A. The member/enrollee presents in the outpatient setting with ~~signs or symptoms of an acute respiratory infection~~signs or symptoms of an acute respiratory infection,
AND
 - B. Results of the testing will influence the member's/enrollee's clinical management.
 - ~~A. Results of the testing will influence the member's/enrollee's clinical management.~~
- II. Current evidence does not support the use of SARS-CoV-2, RSV, or Influenza A/B, **OR** ~~Multiplex Respiratory Viral Panels~~multiplex respiratory viral panels with 5 or fewer targets, for all other indications.

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Bacterial Respiratory Infection/Pneumonia Panels

- I. Bacterial ~~Respiratory Infection/Pneumonia Panels~~respiratory infection/pneumonia panels are considered **medically necessary** when:
 - A. The member/enrollee presents in the outpatient setting with ~~signs or symptoms of an acute respiratory infection~~signs or symptoms of an acute respiratory infection,
AND
 - B. The member/enrollee meets any of the following criteria:
 1. New or worsening lung infiltrates, **OR**
 2. ~~Moderate to severe upper respiratory illness~~Moderate to severe upper respiratory illness, **OR**
 3. Has received empiric antibiotics before obtaining cultures, **OR**
 4. Has possible multidrug-resistant bacteria or polymicrobial infection, **AND**
 - C. Results of the testing will influence the member's/enrollee's clinical management.
- II. Current evidence does not support the use of ~~Bacterial Respiratory Infection/Pneumonia Panels~~bacterial respiratory infection/pneumonia panels for all other indications.

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Influenza A and B Antibody Tests

- I. Current evidence does not support the use of ~~Influenza~~influenza A and B ~~Antibody Tests~~antibody tests for the purpose of diagnosing influenza.

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Group A Streptococcus Pharyngitis Tests

- I. Group A ~~Streptococcus Pharyngitis Tests~~streptococcus pharyngitis tests are considered **medically necessary** when:
 - A. The member/enrollee presents in the outpatient setting with at least one of the following:
 1. Acute pharyngitis, **OR**
 2. Fever, **OR**
 3. Tonsillopharyngeal inflammation, **OR**
 4. Patchy tonsillopharyngeal exudates, **OR**
 5. Palatal petechiae, **OR**
 6. Anterior cervical lymphadenitis, **OR**
 7. Scarletiform rash, **AND**
 - B. The member/enrollee does **NOT** have clinical and epidemiological features that strongly suggest a viral etiology (e.g., cough, rhinorrhea, hoarseness, and oral ulcers), **AND**
 - C. Results of the testing will influence the member's/enrollee's clinical management.
- II. Current evidence does not support the use of ~~Group~~group A ~~Streptococcus Pharyngitis Tests~~streptococcus pharyngitis tests for all other indications.

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Group A Streptococcus Pharyngitis Cultures

- I. Group A ~~Streptococcus Pharyngitis Culture~~ streptococcus pharyngitis culture is considered **medically necessary** when:
 - A. The member/enrollee is between the ages of 3 years and 18 years, **AND**
 - B. The member/enrollee had a negative group A ~~Streptococcus~~ streptococcus rapid antigen detection test (RADT), **AND**
 - C. The member/enrollee presents in the outpatient setting with at least one of the following:
 1. Acute pharyngitis, **OR**
 2. Fever, **OR**
 3. Tonsillopharyngeal inflammation, **OR**
 4. Patchy tonsillopharyngeal exudates, **OR**
 5. Palatal petechiae, **OR**
 6. Anterior cervical lymphadenitis, **OR**
 7. Scarletiform rash, **AND**
 - D. The member/enrollee does **NOT** have clinical and epidemiological features that strongly suggest a viral etiology (e.g., cough, rhinorrhea, hoarseness, and oral ulcers), **AND**
 - E. Results of the testing will influence the member's/enrollee's clinical management.
- II. Current evidence does not support the use of ~~Group A Streptococcus Pharyngitis Culture~~ group A streptococcus pharyngitis culture for all other indications.

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Group A Streptococcus Antibody Tests

- I. Current evidence does not support the use of ~~Group~~ group A Streptococcus Antibody Tests streptococcus antibody tests for the purpose of evaluating a member/enrollee with acute pharyngitis for a possible group A streptococcus infection.

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NOTES AND DEFINITIONS

1. **Moderate to severe ~~upper~~ upper respiratory illness** includes one or more clinical findings of lower respiratory illness (e.g., pneumonia, severe cough/bronchitis, shortness of breath, difficulty breathing).
2. **Severe pneumonia** is defined by the Infectious Diseases Society of America/American Thoracic Society Criteria as: the presence of one major criterion or at least three minor criteria.
 - a. Minor criteria: respiratory rate $\geq \geq 30$ breaths/min, PaO₂/FiO₂ ratio $\leq \leq 250$, multilobar infiltrates, confusion/disorientation, uremia (blood urea nitrogen level $\geq \geq 20$ mg/dl), leukopenia (white blood cell count $\leq \leq 4,000$ cells/ μ l), thrombocytopenia (platelet count $< 100,000/\mu$ l), hypothermia (core temperature $< 36^{\circ}\text{C}$), and hypotension requiring aggressive fluid resuscitation.
 - b. Major criteria: septic shock with need for vasopressors and respiratory failure requiring mechanical ventilation.
- ~~1. **Airway disease** is a nonspecific clinical term for a heterogeneous group of conditions including chronic obstructive pulmonary disease (COPD), emphysema, cystic fibrosis, asthma, and bronchiectasis.~~
3. **Signs ~~and/or~~ symptoms of an acute respiratory infection** include upper or lower respiratory tract symptoms (cough, runny nose, sore throat, bronchitis, pneumonia, bronchiolitis), with or without fever, influenza-like illness (ILI) (fever and either cough or sore throat), and respiratory distress (difficulty in breathing; often characterized by increased respiratory rate and use of accessory muscles of breathing).
4. **Immunocompromised** refers to patients with primary immune deficiency, recent or active cancer, organ or stem cell transplantation, HIV, chronic kidney disease, cystic fibrosis, sickle cell disease, or using immunosuppressive therapy (e.g., corticosteroids, biologics, or chemotherapy). These conditions increase infection risk and impact treatment decisions.

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BACKGROUND AND RATIONALE

Syndromic/Multiplex Respiratory Panels with 6 or More Targets

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Infectious Diseases Society of America

The IDSA published clinical and diagnostic recommendations in 2020 regarding molecular testing for acute respiratory tract infections (RTIs). These recommendations state that it is appropriate to use multiplex viral nucleic acid amplification tests in the following circumstances:

- For immunocompromised and critically ill patients with pneumonia
- In patients experiencing exacerbations of airway disease, defined in the supporting literature as asthma or chronic obstructive pulmonary disease (COPD) complicated by a respiratory infection (p. 2748).

Hanson KE, Azar MM, Banerjee R, et al. Molecular testing for acute respiratory tract infections: clinical and diagnostic recommendations from the IDSA's Diagnostics Committee. Clinical Infectious Diseases. 2020;71(10):2744-2751.

American Society of Transplantation (AST)

According to AST, broad-range diagnostic methods should be considered for identifying respiratory viral infections, as their clinical presentations are indistinguishable. This is especially important early after transplantation, during periods of increased immunosuppression, and throughout respiratory virus season, particularly for lung transplant recipients (p. 2).

Manuel O, Estabrook M. RNA respiratory viral infections in solid organ transplant recipients: Guidelines from the American Society of Transplantation Infectious Diseases Community of Practice. Clinical Transplantation. 2019;33(9). doi:doi:/10.1111/ctr.13511

American Academy of Pediatrics (AAP)

The AAP guidelines on bronchiolitis advise against routinely ordering radiographic or laboratory tests when diagnosis is based on clinical history and physical examination. However, an exception is made for infants and children who experience unexpected worsening of their condition, where additional testing may be warranted (p. 1474).

Ralston SL, Lieberthal AS, Meissner HC, et al. Clinical Practice Guideline: The Diagnosis, Management, and Prevention of Bronchiolitis. PEDIATRICS. 2014;134(5):e1474-e1502. doi:/10.1542/peds.2014-2742

SARS-CoV-2, RSV, or Influenza A/B, OR Multiplex Respiratory Viral Panels with 5 or Fewer Targets

Infectious Diseases Society of America (IDSA)

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The IDSA published clinical and diagnostic recommendations in 2020 regarding molecular testing for acute respiratory tract infections (RTIs). These recommendations state the following:

- ~~● “Multiplex viral NAAT [nucleic acid amplification tests] (potentially combined with bacterial NAAT) also make clinical sense for immunocompromised and critically ill patients with pneumonia as well as for those with exacerbations of airway disease.” (p. 2748).~~

~~SARS-CoV-2, RSV, or Influenza A/B, OR Multiplex Respiratory Viral Panels with 5 or Fewer Targets~~

Infectious Diseases Society of America

~~The IDSA published clinical and diagnostic recommendations in 2020 regarding molecular testing for acute respiratory tract infections (RTIs). These recommendations state the following:~~

~~“Molecular testing for multiple respiratory viruses simultaneously may also be more cost-effective than traditional antigen- or culture-based methods from a laboratory perspective, especially given certain thresholds of disease prevalence.” (p. 2744).~~

Centers for Disease Control and Prevention

~~The CDC states the following on their website discussing RSV: “Healthcare providers should consider RSV in patients with respiratory illness, particularly during the RSV season.”~~

~~The CDC states the following on their website discussing COVID-19: “Key times to get tested: if you have symptoms, test immediately.”~~

~~Hanson KE, Azar MM, Banerjee R, et al. Molecular testing for acute respiratory tract infections: clinical and diagnostic recommendations from the IDSA's Diagnostics Committee. Clinical Infectious Diseases. 2020;71(10):2744-2751.~~

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Bacterial Respiratory Infection/Pneumonia Panels

Infectious Diseases Society of America

~~The (IDSA)~~

~~The IDSA published clinical and diagnostic recommendations in 2020 regarding molecular testing for acute respiratory tract infections (RTIs). published clinical and diagnostic recommendations in 2020 regarding molecular testing for acute respiratory tract infections (RTIs). These recommendations state the following:~~

~~“...that bacterial NAAT may prove most nucleic acid amplification tests (NAATs) are useful in specific clinical situations where, including patients havewith new or worsening lung infiltrates, are moderately to severely ill, have received moderate to severe illness, recipients of empiric antibiotics beforeprior to obtaining cultures, and/or there is concern for if multidrug- resistant bacteria or a polymicrobial infection.”infections are a concern (p. 2747).~~

~~Hanson KE, Azar MM, Banerjee R, et al. Molecular testing for acute respiratory tract infections: clinical and diagnostic recommendations from the IDSA's Diagnostics Committee. Clinical Infectious Diseases. 2020;71(10):2744-2751.~~

~~*The European Respiratory Society (ERS), European Society of Intensive Care Medicine (ESICM), European Society of Clinical Microbiology and Infectious Diseases (ESCMID), and Latin American Thoracic Association (ALAT)*~~

~~In their 2023 guidelines on managing severe community-acquired pneumonia (sCAP), ERS/ESICM/ESCMID/ALAT recommend performing multiplex PCR testing on lower respiratory tract samples, such as sputum or endotracheal aspirates, when non-standard antibiotics for sCAP are being considered or prescribed (p. 4).~~

~~Martin-Loeches I, Torres A, Nagavci B, et al. ERS/ESICM/ESCMID/ALAT guidelines for the management of severe community-acquired pneumonia. Eur Respir J. 2023;61(4):2200735. doi:10.1183/13993003.00735-2022~~

~~*Centers for Disease Control and Prevention (CDC)*~~

~~The CDC recommends nucleic acid amplification testing (NAAT), such as real-time PCR or a respiratory pathogen panel, as the preferred method for diagnosing acute Chlamydia pneumoniae infection.~~

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Centers for Disease Control and Prevention. [Laboratory testing for Chlamydia pneumoniae. Chlamydia pneumoniae Infection. Published January 30, 2024.](https://www.cdc.gov/cpneumoniae/php/laboratories/index.html)
<https://www.cdc.gov/cpneumoniae/php/laboratories/index.html>

Influenza A and B Antibody Tests

Infectious Diseases Society of America ([IDSA](#))

The IDSA published clinical practice guidelines in 2018 which addressed testing criteria for seasonal influenza A and B viruses. These guidelines state that serologic testing for the diagnosis of influenza should not be used by clinicians, because the results from a single serum specimen cannot be reliably interpreted. (p. 898).

Uyeki TM, Bernstein HH, Bradley JS, et al. [Clinical practice guidelines](#) by the Infectious Diseases Society of America: 2018 Update on Diagnosis, Treatment, Chemoprophylaxis, and Institutional Outbreak Management of Seasonal Influenza. Clin Infect Dis. 2019;68(6):895-902.

Group A Streptococcus Pharyngitis Tests

Infectious Diseases Society of America

~~The [IDSA](#) published clinical practice guidelines in 2012 which addressed testing criteria for group A Streptococcal pharyngitis.~~

The IDSA published clinical practice guidelines in 2012 which addressed testing criteria for group A Streptococcal pharyngitis.

~~“Swabbing the throat and testing [Testing](#) for GAS [group A Streptococcus] pharyngitis [via throat swab and testing](#) by rapid antigen detection test (RADT) and/or culture should be performed [because the clinical](#). [Clinical](#) features alone do not reliably discriminate between GAS and viral pharyngitis [except when, unless](#) overt viral features [like are present, such as](#) rhinorrhea, cough, oral ulcers, and/or hoarseness [are present.](#)” (p. e87).~~

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~~“Patients with GAS pharyngitis commonly present with sore throat (generally of sudden onset), pain on swallowing, and fever. Headache, nausea, vomiting, and abdominal pain may also be present, especially in children. On examination, patients have tonsillopharyngeal erythema, with or without exudates, often with tender, enlarged anterior cervical lymph nodes (lymphadenitis). Other findings may include a beefy, red, swollen uvula; petechiae on the palate; excoriated nares (especially in infants); and a scarlatiniform rash.”~~
If a patient has GAS pharyngitis, they will commonly present with sore throat (generally of sudden onset), painful swallowing, and fever. Other symptoms may include headache, nausea, vomiting, and abdominal pain especially in children. Patients have tonsillopharyngeal erythema, with or without exudates on exam. Patients will also usually have tender, enlarged anterior cervical lymph nodes (lymphadenitis). Other findings may include a beefy, red, swollen uvula; petechiae on the palate; raw, irritated nares (especially in infants); and a scarlatiniform rash (a red, bumpy, sandpaper-like rash) (p. e91).

Shulman ST, Bisno AL, Clegg HW, et al. Clinical practice guideline for the diagnosis and management of group A streptococcal pharyngitis: 2012 update by the Infectious Diseases Society of America. *Clin Infect Dis.* 2012;55(10):e86-102.

2012;55(10):e86-102.

Group A Streptococcus Pharyngitis Culture

Infectious Diseases Society of America (IDSA)

The IDSA published clinical practice guidelines in 2012 which addressed testing criteria for group A Streptococcal pharyngitis.

~~The IDSA published clinical practice guidelines in 2012 which addressed testing criteria for group A Streptococcal pharyngitis.~~

“In children and adolescents, negative RADT [rapid antigen detection test] tests should be backed up by a throat culture...Routine use of back-up throat cultures for those with a negative RADT is not necessary for adults in usual circumstances, because of the low incidence of GAS [group A Streptococcus] pharyngitis in adults and because the risk of subsequent acute rheumatic fever is generally exceptionally low in adults with acute pharyngitis.” (p. e87).

“Swabbing the throat and testing for GAS [group A Streptococcus] pharyngitis by rapid antigen detection test (RADT) and/or culture should be performed because the clinical features alone do

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not reliably discriminate between GAS and viral pharyngitis except when overt viral features like rhinorrhea, cough, oral ulcers, and/or hoarseness are present.” (p. e87).

“Patients with GAS pharyngitis commonly present with sore throat (generally of sudden onset), pain on swallowing, and fever. Headache, nausea, vomiting, and abdominal pain may also be present, especially in children. On examination, patients have tonsillopharyngeal erythema, with or without exudates, often with tender, enlarged anterior cervical lymph nodes (lymphadenitis). Other findings may include a beefy, red, swollen uvula; petechiae on the palate; excoriated nares (especially in infants); and a scarlatiniform rash.” (p. e91).

Shulman ST, Bisno AL, Clegg HW, et al. Clinical practice guideline for the diagnosis and management of group A streptococcal pharyngitis: 2012 update by the Infectious Diseases Society of America. Clin Infect Dis. 2012;55(10):e86-

~~“Patients with GAS pharyngitis commonly present with sore throat (generally of sudden onset), pain on swallowing, and fever. Headache, nausea, vomiting, and abdominal pain may also be present, especially in children. On examination, patients have tonsillopharyngeal erythema, with or without exudates, often with tender, enlarged anterior cervical lymph nodes (lymphadenitis). Other findings may include a beefy, red, swollen uvula; petechiae on the palate; excoriated nares (especially in infants); and a scarlatiniform rash.” (p. e91)~~

102.

American Academy of Family Physicians (AAFP)

The ~~2024~~ American Academy of Family Physicians published an expert-authored evidence review in 2024 concerning the diagnosis and management of streptococcus pharyngitis ~~states,~~ stating the following:

“Rapid antigen testing may be omitted for patients at low clinical risk, including children younger than 3 years.” (p. 345).

“The Centers for Disease Control and Prevention and the American Academy of Pediatrics recommend obtaining a throat culture for all children and adolescents after a negative result on rapid antigen testing because of the higher risk of complications.” (p. 345).

Hamilton JL, Leon McCrea II. Streptococcal Pharyngitis: Rapid Evidence Review. Am Fam Physician. 2024;109(4):343-349.

Group A Streptococcus Antibody Tests

Infectious Diseases Society of America (IDSA)

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~~The IDSA published clinical practice guidelines in 2012 which addressed testing criteria for group A Streptococcal pharyngitis.~~

The IDSA published clinical practice guidelines in 2012 which addressed testing criteria for group A Streptococcal pharyngitis.

Per these guidelines, it is not recommended that individuals undergo anti-streptococcal antibody titers for the purpose of routine diagnosis of acute pharyngitis, as these results indicate a past infection and therefore do not aid in the diagnosis of the present illness. (p. e87).

“Measurement of anti-streptococcal antibody titers is often useful for diagnosis of the nonsuppurative sequelae of GAS pharyngitis, such as acute rheumatic fever and acute glomerulonephritis. However, such testing is not useful in the diagnosis of acute pharyngitis because antibody titers of the 2 most commonly used tests, antistreptolysin O (ASO) and antiDNase B, may not reach maximum levels until 3–8 weeks after acute GAS pharyngeal infection and may remain elevated for months even without active GAS infection.” (p. e93-94).

Shulman ST, Bisno AL, Clegg HW, et al. Clinical practice guideline for the diagnosis and management of group A streptococcal pharyngitis: 2012 update by the Infectious Diseases Society of America. Clin Infect Dis. 2012;55(10):e86-102.

Coding Implications

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NOTE: Coverage is subject to each requested code’s inclusion on the corresponding LDH fee schedule. Non-covered codes are denoted (*) and are reviewed for Medical Necessity for members under 21 years of age on a per case basis.

CPT [®] Code	Description
<u>0109U*</u>	<u>Infectious disease (Aspergillus species), real-time PCR for detection of DNA from 4 species (A. fumigatus, A. terreus, A. niger, and A. flavus), blood, lavage fluid, or tissue, qualitative reporting of presence or absence of each species</u>
<u>0115U*</u>	<u>Respiratory infectious agent detection by nucleic acid (DNA and RNA), 18 viral types and subtypes and 2 bacterial targets, amplified probe technique, including multiplex reverse transcription for RNA targets, each analyte reported as detected or not detected</u>
<u>0202U*</u>	<u>Infectious disease (bacterial or viral respiratory tract infection), pathogenspecific 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 (For additional PLA code with identical clinical descriptor, see 0223U. See Appendix O or the most current listing on the AMA CPT website to determine appropriate code assignment)</u>
<u>0223U*</u>	<u>Infectious disease (bacterial or viral respiratory tract infection), pathogenspecific 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 (For additional PLA code with identical clinical descriptor, see 0202U. See Appendix O or the most current listing on the AMA CPT website to determine appropriate code assignment)</u>
<u>0224U*</u>	<u>Antibody, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]), includes titer(s), when performed</u>
<u>0225U*</u>	<u>Infectious disease (bacterial or viral respiratory tract infection) pathogen-specific DNA and RNA, 21 targets, including severe acute respiratory syndrome coronavirus 2 (SARSCoV-2), amplified probe technique, including multiplex reverse transcription for RNA targets, each analyte reported as detected or not detected</u>
<u>0226U*</u>	<u>Surrogate viral neutralization test (sVNT), severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]), ELISA, plasma, serum</u>
<u>0240U*</u>	<u>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</u>
<u>0241U*</u>	<u>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</u>
<u>0442U*</u>	<u>Infectious disease (respiratory infection), Myxovirus resistance protein A (MxA) and C-reactive protein (CRP), fingerstick whole blood specimen, each biomarker reported as present or absent</u>
<u>0556U*</u>	<u>Infectious disease (bacterial or viral respiratory tract infection), pathogen-specific DNA and RNA by real-time PCR, 12 targets, nasopharyngeal or oropharyngeal swab, including multiplex reverse transcription for RNA targets, each analyte reported as detected or not detected</u>
<u>0563U*</u>	<u>Infectious disease (bacterial and/or viral respiratory tract infection), pathogen-specific nucleic acid (DNA or RNA), 11 viral targets and 4 bacterial targets, qualitative RT-PCR, upper respiratory</u>

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CPT® Code	Description
	<u>specimen, each pathogen reported as positive or negative</u>
<u>0564U*</u>	<u>Infectious disease (bacterial and/or viral respiratory tract infection), pathogen-specific nucleic acid (DNA or RNA), 10 viral targets and 4 bacterial targets, qualitative RT-PCR, upper respiratory specimen, each pathogen reported as positive or negative</u>
<u>0574U*</u>	<u>Mycobacterium tuberculosis, culture filtrate protein-10-kDa (CFP-10), serum or plasma, liquid chromatography mass spectrometry (LC-MS)</u>
86060	Antistreptolysin O; titer
86328	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])
86408	Neutralizing antibody, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]); screen
86409	Neutralizing antibody, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]); titer
86413	Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]) antibody, quantitative
<u>86486</u>	<u>Skin test; unlisted antigen, each</u>
<u>86510</u>	<u>Skin test; histoplasmosis</u>
<u>86590</u>	<u>Streptokinase, antibody</u>
<u>86602</u>	<u>Antibody; actinomyces</u>
<u>86603</u>	<u>Antibody; adenovirus</u>
<u>86606</u>	<u>Antibody; Aspergillus</u>
<u>86609</u>	<u>Antibody; bacterium, not elsewhere specified</u>
<u>86615</u>	<u>Antibody; Bordetella</u>
<u>86628</u>	<u>Antibody; Candida</u>
<u>86635</u>	<u>Antibody; Coccidioides</u>
<u>86638</u>	<u>Antibody; Coxiella burnetii (Q fever)</u>
<u>86641</u>	<u>Antibody; Cryptococcus</u>
<u>86648</u>	<u>Antibody; Diphtheria</u>
<u>86658</u>	<u>Antibody; enterovirus (eg, coxsackie, echo, polio)</u>

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CPT [®] Code	Description
<u>86671</u>	<u>Antibody; fungus, not elsewhere specified</u>
<u>86689</u>	<u>Antibody; HTLV or HIV antibody, confirmatory test (eg, Western Blot)</u>
<u>86698</u>	<u>Antibody; histoplasma</u>
86710	Antibody; influenza virus
<u>86711</u>	<u>Antibody; JC (John Cunningham) virus</u>
<u>86713</u>	<u>Antibody; Legionella</u>
<u>86720</u>	<u>Antibody; Leptospira</u>
<u>86723</u>	<u>Antibody; Listeria monocytogenes</u>
<u>86727</u>	<u>Antibody; lymphocytic choriomeningitis</u>
<u>86732</u>	<u>Antibody; mucormycosis</u>
<u>86735</u>	<u>Antibody; mumps</u>
<u>86738</u>	<u>Antibody; mycoplasma</u>
<u>86756</u>	<u>Antibody; respiratory syncytial virus</u>
86769	Antibody; severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19])
<u>86784</u>	<u>Antibody; Trichinella</u>
<u>86790</u>	<u>Antibody; virus, not elsewhere specified</u>
<u>86793</u>	<u>Antibody; Yersinia</u>
<u>87015</u>	<u>Concentration (any type), for infectious agents</u>
87040	Culture, bacterial; blood, aerobic, with isolation and presumptive identification of isolates (includes anaerobic culture, if appropriate)
87070	Culture, bacterial; any other source except urine, blood or stool, aerobic, with isolation and presumptive identification of isolates
87071	Culture, bacterial; quantitative, aerobic with isolation and presumptive identification of isolates, any source except urine, blood or stool
87073	Culture, bacterial; quantitative, anaerobic with isolation and presumptive identification of isolates, any source except urine, blood or stool
87075	Culture, bacterial; any source, except blood, anaerobic with isolation and presumptive identification of isolates
87076	Culture, bacterial; anaerobic isolate, additional methods required for definitive identification, each

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CPT® Code	Description
	isolate
87077	Culture, bacterial; aerobic isolate, additional methods required for definitive identification, each isolate
87081	Culture, presumptive, pathogenic organisms, screening only;
87084	Culture, presumptive, pathogenic organisms, screening only; with colony estimation from density chart
87101	Culture, fungi (mold or yeast) isolation, with presumptive identification of isolates; skin, hair, or nail
87102	Culture, fungi (mold or yeast) isolation, with presumptive identification of isolates; other source (except blood)
87103	Culture, fungi (mold or yeast) isolation, with presumptive identification of isolates; blood
87106	Culture, fungi, definitive identification, each organism; yeast
87107	Culture, fungi, definitive identification, each organism; mold
87109	Culture, mycoplasma, any source
87116	Culture, tubercle or other acid-fast bacilli (eg, TB, AFB, mycobacteria) any source, with isolation and presumptive identification of isolates
87118	Culture, mycobacterial, definitive identification, each isolate
87140	Culture, typing; immunofluorescent method, each antiserum
87143	Culture, typing; gas liquid chromatography (GLC) or high pressure liquid chromatography (HPLC) method
87147	Culture, typing; immunologic method, other than immunofluorescence (eg, agglutination grouping), per antiserum
<u>87149</u>	<u>Culture, typing; identification by nucleic acid (DNA or RNA) probe, direct probe technique, per culture or isolate, each organism probed</u>
<u>87150</u>	<u>Culture, typing; identification by nucleic acid (DNA or RNA) probe, amplified probe technique, per culture or isolate, each organism probed</u>
<u>87153</u>	<u>Culture, typing; identification by nucleic acid sequencing method, each isolate (eg, sequencing of the 16S rRNA gene)</u>
<u>87154*</u>	<u>Culture, typing; identification of blood pathogen and resistance typing, when performed, by nucleic acid (DNA or RNA) probe, multiplexed amplified probe technique including multiplex reverse transcription, when performed, per culture or isolate, 6 or more targets</u>
87158	Culture, typing; other methods

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CPT® Code	Description
<u>87168</u>	<u>Macroscopic examination; arthropod</u>
<u>87169</u>	<u>Macroscopic examination; parasite</u>
<u>87176</u>	<u>Homogenization, tissue, for culture</u>
<u>87181</u>	<u>Susceptibility studies, antimicrobial agent; agar dilution method, per agent (eg, antibiotic gradient strip)</u>
<u>87184</u>	<u>Susceptibility studies, antimicrobial agent; disk method, per plate (12 or fewer agents)</u>
<u>87185</u>	<u>Susceptibility studies, antimicrobial agent; enzyme detection (eg, beta lactamase), per enzyme</u>
<u>87186</u>	<u>Susceptibility studies, antimicrobial agent; microdilution or agar dilution (minimum inhibitory concentration [MIC] or breakpoint), each multi-antimicrobial, per plate</u>
<u>87187</u>	<u>Susceptibility studies, antimicrobial agent; microdilution or agar dilution, minimum lethal concentration (MLC), each plate (List separately in addition to code for primary procedure)</u>
<u>87188</u>	<u>Susceptibility studies, antimicrobial agent; macrobroth dilution method, each agent</u>
<u>87190</u>	<u>Susceptibility studies, antimicrobial agent; mycobacteria, proportion method, each agent</u>
<u>87205</u>	<u>Smear, primary source with interpretation; Gram or Giemsa stain for bacteria, fungi, or cell types</u>
<u>87206</u>	<u>Smear, primary source with interpretation; fluorescent and/or acid fast stain for bacteria, fungi, parasites, viruses or cell types</u>
<u>87207</u>	<u>Smear, primary source with interpretation; special stain for inclusion bodies or parasites (eg, malaria, coccidia, microsporidia, trypanosomes, herpes viruses)</u>
<u>87210</u>	<u>Smear, primary source with interpretation; wet mount for infectious agents (eg, saline, India ink, KOH preps)</u>
<u>87220</u>	<u>Tissue examination by KOH slide of samples from skin, hair, or nails for fungi or ectoparasite ova or mites (eg, scabies)</u>
<u>87230</u>	<u>Toxin or antitoxin assay, tissue culture (eg, Clostridium difficile toxin)</u>
<u>87250</u>	<u>Virus isolation; inoculation of embryonated eggs, or small animal, includes observation and dissection</u>
<u>87252</u>	<u>Virus isolation; tissue culture inoculation, observation, and presumptive identification by cytopathic effect</u>
<u>87253</u>	<u>Virus isolation; tissue culture, additional studies or definitive identification (eg, hemabsorption, neutralization, immunofluorescence stain), each isolate</u>
<u>87254</u>	<u>Virus isolation; centrifuge enhanced (shell vial) technique, includes identification with immunofluorescence stain, each virus</u>
<u>87255</u>	<u>Virus isolation; including identification by non-immunologic method, other than by cytopathic</u>

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CPT® Code	Description
	<u>effect (eg, virus specific enzymatic activity)</u>
<u>87260</u>	<u>Infectious agent antigen detection by immunofluorescent technique; adenovirus</u>
<u>87265</u>	<u>Infectious agent antigen detection by immunofluorescent technique; Bordetella pertussis/parapertussis</u>
87275	Infectious agent antigen detection by immunofluorescent technique; influenza B virus
87276	Infectious agent antigen detection by immunofluorescent technique; influenza A virus
<u>87278</u>	<u>Infectious agent antigen detection by immunofluorescent technique; Legionella pneumophila</u>
<u>87279</u>	<u>Infectious agent antigen detection by immunofluorescent technique; Parainfluenza virus, each type</u>
<u>87280</u>	<u>Infectious agent antigen detection by immunofluorescent technique; respiratory syncytial virus</u>
<u>87281</u>	<u>Infectious agent antigen detection by immunofluorescent technique; Pneumocystis carinii</u>
<u>87299</u>	<u>Infectious agent antigen detection by immunofluorescent technique; not otherwise specified, each organism</u>
<u>87300</u>	<u>Infectious agent antigen detection by immunofluorescent technique, polyvalent for multiple organisms, each polyvalent antiserum</u>
<u>87301</u>	<u>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; adenovirus enteric types 40/41</u>
<u>87305</u>	<u>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; Aspergillus</u>
<u>87385</u>	<u>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; Histoplasma capsulatum</u>
87400	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; Influenza, A or B, each
87420	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; respiratory syncytial virus
87426	Infectious agent antigen detection by immunoassay technique (eg, enzyme immunoassay [EIA],

CPT [®] Code	Description
	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])
87428	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
87430	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; Streptococcus, group A
<u>87449*</u>	<u>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; not otherwise specified, each organism</u>
<u>87451</u>	<u>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; polyvalent for multiple organisms, each polyvalent antiserum</u>
87480	Infectious agent detection by nucleic acid (DNA or RNA); Candida species, direct probe technique
87481	Infectious agent detection by nucleic acid (DNA or RNA); Candida species, amplified probe technique
87482	Infectious agent detection by nucleic acid (DNA or RNA); Candida species, quantification
87485	Infectious agent detection by nucleic acid (DNA or RNA); Chlamydia pneumoniae, direct probe technique
87486	Infectious agent detection by nucleic acid (DNA or RNA); Chlamydia pneumoniae, amplified probe technique
87487	Infectious agent detection by nucleic acid (DNA or RNA); Chlamydia pneumoniae, quantification
87498	Infectious agent detection by nucleic acid (DNA or RNA); enterovirus, amplified probe technique, includes reverse transcription when performed
87500	Infectious agent detection by nucleic acid (DNA or RNA); vancomycin resistance (eg, enterococcus species van A, van B), amplified probe technique
87501	Infectious agent detection by nucleic acid (DNA or RNA); influenza virus, includes reverse transcription, when performed, and amplified probe technique, each type or subtype

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CPT® Code	Description
87502	Infectious agent detection by nucleic acid (DNA or RNA); influenza virus, for multiple types or sub-types, includes multiplex reverse transcription, when performed, and multiplex amplified probe technique, first 2 types or sub-types
87503	Infectious agent detection by nucleic acid (DNA or RNA); influenza virus, for multiple types or sub-types, includes multiplex reverse transcription, when performed, and multiplex amplified probe technique, each additional influenza virus type or sub-type beyond 2 (List separately in addition to code for primary procedure)
87540	Infectious agent detection by nucleic acid (DNA or RNA); Legionella pneumophila, direct probe technique
87541	Infectious agent detection by nucleic acid (DNA or RNA); Legionella pneumophila, amplified probe technique
87542	Infectious agent detection by nucleic acid (DNA or RNA); Legionella pneumophila, quantification
87550	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria species, direct probe technique
87551	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria species, amplified probe technique
87552	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria species, quantification
87555	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria tuberculosis, direct probe technique
87556	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria tuberculosis, amplified probe technique
87560	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria avium-intracellulare, direct probe technique
87561	Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria avium-intracellulare, amplified probe technique
<u>87562</u>	<u>Infectious agent detection by nucleic acid (DNA or RNA); Mycobacteria avium-intracellulare, quantification</u>
87580	Infectious agent detection by nucleic acid (DNA or RNA); Mycoplasma pneumoniae, direct probe technique
87581	Infectious agent detection by nucleic acid (DNA or RNA); Mycoplasma pneumoniae, amplified probe technique
87582	Infectious agent detection by nucleic acid (DNA or RNA); Mycoplasma pneumoniae, quantification
<u>87631</u>	<u>Infectious agent detection by nucleic acid (DNA or RNA); respiratory virus (eg, adenovirus,</u>

CPT [®] Code	Description
	<u>influenza virus, coronavirus, metapneumovirus, parainfluenza virus, respiratory syncytial virus, rhinovirus), includes multiplex reverse transcription, when performed, and multiplex amplified probe technique, multiple types or subtypes, 3-5 targets</u>
87632	Infectious agent detection by nucleic acid (DNA or RNA); respiratory virus (eg, adenovirus, influenza virus, coronavirus, metapneumovirus, parainfluenza virus, respiratory syncytial virus, rhinovirus), includes multiplex reverse transcription, when performed, and multiplex amplified probe technique, multiple types or subtypes, 6-11 targets
87633	Infectious agent detection by nucleic acid (DNA or RNA); respiratory virus (eg, adenovirus, influenza virus, coronavirus, metapneumovirus, parainfluenza virus, respiratory syncytial virus, rhinovirus), includes multiplex reverse transcription, when performed, and multiplex amplified probe technique, multiple types or subtypes, 12-25 targets
87634	Infectious agent detection by nucleic acid (DNA or RNA); respiratory syncytial virus, amplified probe technique
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
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
87637	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
87640	Infectious agent detection by nucleic acid (DNA or RNA); Staphylococcus aureus, amplified probe technique
87641	Infectious agent detection by nucleic acid (DNA or RNA); Staphylococcus aureus, methicillin resistant, amplified probe technique
87650	Infectious agent detection by nucleic acid (DNA or RNA); Streptococcus, group A, direct probe technique
87651	Infectious agent detection by nucleic acid (DNA or RNA); Streptococcus, group A, amplified probe technique
87652	Infectious agent detection by nucleic acid (DNA or RNA); Streptococcus, group A, quantification
87653	Infectious agent detection by nucleic acid (DNA or RNA); Streptococcus, group B, amplified probe technique
87797	Infectious agent detection by nucleic acid (DNA or RNA), not otherwise specified; direct probe technique, each organism
87798	Infectious agent detection by nucleic acid (DNA or RNA), not otherwise specified; amplified probe

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CPT® Code	Description
	technique, each organism
87799	Infectious agent detection by nucleic acid (DNA or RNA), not otherwise specified; quantification, each organism
87800	Infectious agent detection by nucleic acid (DNA or RNA), multiple organisms; direct probe(s) technique
87801	Infectious agent detection by nucleic acid (DNA or RNA), multiple organisms; amplified probe(s) technique
<u>87802</u>	<u>Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; Streptococcus, group B</u>
87804	Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; Influenza
87807	Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; respiratory syncytial virus
<u>87809</u>	<u>Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; adenovirus</u>
87811	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])
87880	Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; Streptococcus, group A
<u>87899</u>	<u>Infectious agent antigen detection by immunoassay with direct optical (ie, visual) observation; adenovirus</u>
<u>87900</u>	<u>Infectious agent drug susceptibility phenotype prediction using regularly updated genotypic bioinformatics</u>
<u>87905</u>	<u>Infectious agent enzymatic activity other than virus (eg, sialidase activity in vaginal fluid)</u>
87913	Infectious agent genotype analysis by nucleic acid (DNA or RNA); severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) (coronavirus disease [COVID-19]), mutation identification in targeted region(s)
0202U*8799 <u>9</u>	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 (For additional PLA code with identical clinical descriptor, see 0223U. See Appendix O or the most current listing on the AMA CPT website to determine appropriate code assignment)
<u>U0001*</u>	<u>CDC Test</u>

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CPT® Code	Description
U0002*	Non-CDC Viral identification test, amplified probe
0223U*U0003	Infectious disease (bacterial or viral respiratory tract infection), pathogen specific agent detection by 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) (coronavirus disease [COVID-19]), amplified probe technique, making use of high throughput technologies as detected or not detected (For additional PLA code with identical clinical descriptor, see 0202U. See Appendix O or the most current listing on the AMA CPT website to determine appropriate code assignment) described by CMS-2020-01-R
0225U*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. Infectious disease (bacterial or viral respiratory tract infection) pathogen-specific DNA and RNA, 21 targets, including severe acute respiratory syndrome coronavirus 2 (SARSCoV-2), amplified probe technique, including multiplex reverse transcription for RNA targets, each analyte reported as detected or not detected
0240U*	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
0241U*	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
0528U	Lower respiratory tract infectious agent detection, 18 bacteria, 8 viruses, and 7 antimicrobial resistance genes, amplified probe technique, including reverse transcription for RNA targets, each analyte reported as detected or not detected with semiquantitative results for 15 bacteria
U0001*	CDC Test
U0002*	Non-CDC Viral identification test, amplified probe
U0003*	High throughput Viral identification test, amplified probe
U0004*	High throughput Viral identification test, other than amplified probe
U0005*	Infectious agent detection by nucleic acid (dna <u>DNA</u> or rna <u>RNA</u>); severe acute respiratory

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CPT® Code	Description
	syndrome coronavirus 2 (sars-cov SARS-CoV-2) (coronavirus disease [eovid COVID-19]), amplified probe technique, ede CDC or non- ede CDC, making use of high throughput technologies, completed within 2 calendar days from date of specimen collection (list separately in addition to either hepes HCPCS code u0003 U0003 or u0004 U0004) as described by ems CMS-2020-01- r2R2

Reviews, Revisions, and Approvals	Revision Date	Approval Date	Effective Date
Converted corporate to local policy.	03/24	5/1/24	
Added (*) to codes 0202U, 0223U, 0225U, 0240U and 0241U per LDH’s IB 24-16. Did not send to LDH for review as revisions were per IB 24-16.	07/24	7/10/24	
Removed (*) from codes 87631, 87632, and 87633. Added footnote to page 16. Added section “ <u>Coverage Specific Guidelines with information regarding coverage of the codes and meeting medical necessity</u> ” from IB 24-31.	10/24	1/3/25	
Annual review. References and background reviewed and updated. Changed verbiage in policy statements from “may be considered medically necessary” to “are considered medically necessary. Noted that CPT 87631 is out of scope for this policy in criteria for SARS-CoV-2, RSV, or Influenza A/B, or Multiplex Respiratory Viral Panels with 5 or Fewer Targets and removed from the CPT table. In criteria for Syndromic/Multiplex Respiratory Panels with 6 or More Targets, clarified that criteria applies when performed in the outpatient setting and replaced prior criteria with the following options: “With serious or critical illness, OR At imminent risk of becoming seriously or critically ill, OR With immunodeficiency, AND/OR With a severe underlying condition”, per Louisiana Department of Health Informational Bulletin 24-31. Added 0528U as an in-scope code in the CPT table.	2/25	4/28/25	5/29/25

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REFERENCES

<p><u>Annual review. Updated policy number from LA.CP.CG.12 to align with Corporate. Added hyperlinked definition of “immunocompromised” to criteria for syndromic/multiplex respiratory panels with six or more targets. Added rationale/references to policy reference table. Updated revision and copyright dates. Added procedure codes 86486, 86510, 86590, 86602, 86603, 86606, 86609, 86615, 86628, 86635, 86638, 86641, 86648, 86658, 86671, 86689, 86698, 86711, 86713, 86720, 86723, 86727, 86732, 86735, 86738, 86756, 86759, 86762, 86765, 86784, 86790, 86793, 87015, 87149, 87150, 87153, 87154, 87168, 87169, 87176, 87181, 87184, 87185, 87186, 87187, 87188, 87190, 87205, 87206, 87207, 87210, 87220, 87230, 87250, 87252, 87253, 87254, 87255, 87260, 87265, 87278, 87279, 87280, 87281, 87299, 87300, 87301, 87305, 87385, 87449, 87451, 87562, 87631, 87632, 87633, 87802, 87809, 87899, 87900, 87905, 87999, 0109U, 0115U, 0224U, 0226U, 0373U, 0442U, 0556U, 0563U, 0564U, 0574U, U0003, U0004, U0005 and deleted 0528U, 0373U from Coding Implications table. Updated Notes and Definitions section. Updated all Rationale sections. Updated example tests on Policy Reference Table.</u></p>	<p><u>04/26</u></p>		
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- ~~3. Hanson KE, Azar MM, Banerjee R, et al. Molecular Testing for Acute Respiratory Tract Infections: Clinical and Diagnostic Recommendations From the IDSA's Diagnostics Committee. Clin Infect Dis. 2020;71(10):2744–2751. doi:10.1093/cid/ciaa508~~
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CLINICAL POLICY

Concert Infectious Disease: Respiratory Testing

Important Reminder

This clinical 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 clinical 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 clinical policy. This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved.

The purpose of this clinical 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 clinical policy is effective as of the date determined by LHCC. The date of posting may not be the effective date of this clinical policy. This clinical 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 clinical 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 clinical policy, and additional clinical policies may be developed and adopted as needed, at any time.

This clinical 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/enrollees. This clinical policy is not intended to recommend treatment for members/enrollees. Members/enrollees should consult with their treating physician in connection with diagnosis and treatment decisions.

Providers referred to in this clinical 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 clinical policy is the property of LHCC. Unauthorized copying, use, and distribution of this clinical policy or any information contained herein are strictly prohibited. Providers, members/enrollees and their representatives are bound to the terms and conditions expressed herein through the terms of their contracts. Where no such contract exists, providers,

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members/enrollees and their representatives agree to be bound by such terms and conditions by providing services to members/enrollees and/or submitting claims for payment for such services.

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