CONCERT GENETIC TESTING: HEARING LOSS

OVERVIEW

Hereditary hearing loss can be classified as syndromic or nonsyndromic. Syndromic hearing loss refers to hearing loss associated with other medical or physical findings, including visible abnormalities of the external ear. Because syndromic hearing loss occurs as part of a syndrome of multiple clinical manifestations, it is often recognized more readily as hereditary. Nonsyndromic hearing loss is defined as hearing loss not associated with other physical signs or symptoms. Nonsyndromic hearing loss accounts for 70% to 80% of genetically determined deafness, and it is more difficult to determine whether the etiology is hereditary or acquired.

This policy primarily focuses on the use of genetic testing to identify a cause of suspected hereditary hearing loss. The diagnosis of syndromic hearing loss can be made on the basis of associated clinical findings. However, at the time of hearing loss presentation, associated clinical findings may not be apparent; furthermore, variants in certain genetic loci may cause both syndromic and nonsyndromic hearing loss. Given this overlap, the policy focuses on genetic testing for hereditary hearing loss more generally.

POLICY REFERENCE TABLE

Below is a list of higher volume tests and the associated laboratories for each coverage criteria section. This list is not all inclusive.

Coding Implications

This clinical policy references Current Procedural Terminology (CPT®). CPT® is a registered trademark of the American Medical Association. All CPT codes and descriptions
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.

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**OTHER RELATED POLICIES**

This policy document provides coverage criteria for genetic testing for hereditary hearing loss. Please refer to:
• Genetic Testing: Prenatal and Preconception Carrier Screening for coverage criteria related to carrier screening for hereditary hearing loss.

• Genetic Testing: Multisystem Inherited Disorders, Intellectual Disability, and Developmental Delay for coverage criteria related to genetic disorders that affect multiple organ systems

• Genetic Testing: General Approach to Genetic Testing for coverage criteria related to genetic testing for hearing loss that is not specifically discussed in this or another non-general policy.

CRITERIA

It is the policy of Louisiana Healthcare Connections that the specific genetic testing noted below is medically necessary when meeting the related criteria:

HEREDITARY HEARING LOSS

Known Familial Variant Analysis for Hereditary Hearing Loss

I. Targeted variant analysis for known familial variant(s) to establish a diagnosis of hereditary hearing loss (81253*, 81403*) is considered medically necessary when:

   A. The member/enrollee has a close relative with pathogenic or likely pathogenic variant(s) in GJB2, GJB6, or another gene known to cause hereditary hearing loss.

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GJB2 and GJB6 Sequencing and/or Deletion/Duplication Analysis or Multigene Panel Analysis

I. GJB2 sequencing and/or deletion/duplication (81252*, 81479) and/or GJB6 sequencing and/or deletion/duplication analysis (81254*, 81479) or multigene panel analysis (81430*, 81431*) to establish a diagnosis of hereditary hearing loss is considered medically necessary when:

   A. The member/enrollee has hearing loss, AND

   B. There is no known acquired cause of the hearing loss (i.e., TORCH infections [Toxoplasma gondii, other agents, rubella, cytomegalovirus, and herpes simplex virus], bacterial infection, age-related or noise-related hearing loss).
II. *GJB2* sequencing and/or deletion/duplication (81252*, 81479) and/or *GJB6* sequencing and/or deletion/duplication analysis (81254*, 81479) or multigene panel analysis (81430*, 81431*) to establish a diagnosis of hereditary hearing loss is considered investigational for all other indications.

NOTES AND DEFINITIONS

1. Close relatives include first, second, and third degree blood relatives:
   a. First-degree relatives are parents, siblings, and children
   b. Second-degree relatives are grandparents, aunts, uncles, nieces, nephews, grandchildren, and half siblings
   c. Third-degree relatives are great grandparents, great aunts, great uncles, great grandchildren, and first cousins

CLINICAL CONSIDERATIONS

If there is not a high suspicion for a specific hearing loss etiology, ideally the evaluation should occur in a stepwise fashion. About 50% of individuals with autosomal recessive hereditary hearing loss have pathogenic variants in the *GJB2* gene, in the other 50% of patients with apparent autosomal recessive hereditary hearing loss, numerous other genes are implicated. There is no single identifiable gene responsible for most cases of autosomal dominant hereditary hearing loss.

If there is suspicion for autosomal recessive congenital hearing loss, it would be reasonable to begin with testing of *GJB2* and *GJB6* and if testing is negative, screening for the other genes associated with hearing loss with a multigene panel would be efficient. An alternative strategy for suspected autosomal recessive or autosomal dominant hearing loss would be to obtain a multigene panel that includes GJB2 and GJB6 as a first step.

Given the extreme heterogeneity in genetic causes of hearing loss, these 2 strategies may be considered reasonably equivalent.
BACKGROUND AND RATIONALE

Known Familial Variant Analysis for Hereditary Hearing Loss

Genetic Support Foundation

The Genetic Support Foundation’s Genetics 101 information on inheritance patterns says the following about testing for familial pathogenic variants:

Genetic testing for someone who may be at risk for an inherited disease is always easier if we know the specific genetic cause. Oftentimes, the best way to find the genetic cause is to start by testing someone in the family who is known or strongly suspected to have the disease. If their testing is positive, then we can say that we have found the familial pathogenic (harmful) variant. We can use this as a marker to test other members of the family to see who is also at risk.

GJB2 and GJB6 Sequencing and/or Deletion Duplication Analysis or Multigene Panel Analysis

American College of Medical Genetics and Genomics (ACMG)

The American College of Medical Genetics and Genomics (ACMG) published guidelines in 2022 to guide genetics evaluation for individuals with hearing loss: “For individuals lacking physical findings suggestive of a known syndrome, a tiered diagnostic approach should be implemented. Unless clinical and/or family history suggests a specific genetic etiology, comprehensive HL gene panel testing should be initiated.” (page 9) The guidelines state the following: “Although nonsyndromic HL [hearing loss] demonstrates high genetic heterogeneity, the DFNB1 locus, which includes the GJB2 gene encoding the gap junction protein connexin 26 and the GJB6 gene encoding the gap junction protein connexin 30, accounts for an estimated 50% of all autosomal recessive nonsyndromic HL and 15% to 40% of all deaf individuals in a variety of populations.” (page 3)

GeneReviews: Hereditary Hearing Loss and Deafness Overview

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. In the section that discusses possible differential diagnosis for hereditary hearing loss and deafness, it states:

In developed countries approximately 80% of congenital hearing loss is due to genetic causes and the remainder to environmental (acquired) causes. Acquired causes should be differentiated from genetic causes to inform the evaluation and required ancillary testing. Acquired hearing loss in children commonly results from prenatal infections from ‘TORCH’
organisms…or postnatal infections (such as CMV). Acquired hearing loss in adults is most often attributed to environmental factors.

### Reviews, Revisions, and Approvals

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<td>Converted corporate to local policy.</td>
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### REFERENCES


### 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.

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