Your children have a 25% chance of inheriting GJB2 Related hearing loss

Candace and Christopher are both carriers for GJB2 Related hearing loss

Carrier

GJB2 Related hearing loss

Hearing loss that varies from mild to profound, is typically present at birth, and does not worsen over time.

Carrier

Usher Syndrome Type 1 (USH1)

Complete hearing loss from birth, problems with balance and progressive loss of sight starting before puberty.

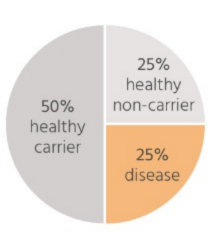
Your children are unlikely (0.0004%) to have this disease.

Both of you are unlikely to be a carrier for any of the others in the 5 disease panel.

Each child has a 25% chance of having the disease and 75% chance not having the disease.

Each child also has a 50% chance of being a carrier of any of the same diseases as you and your partner. When your children grow up it will be helpful to inform them of this possibility.

Learn why



A counselor is always available

Still have questions? Review your results with a licensed genetic counselor. We are available 9am-5pm Monday through Friday.

Contact Care Team

Send your report to your family or doctor

If you have siblings, they have 50% chance of being a carrier of the same diseases as you. Invitees will receive an email allowing them to view your report. You can revoke this access at any time.

Some siblings appreciate knowing this information, but some do not. Consider these reasons before telling family.

Grant report access to_

Limitations

This test is not able to guarantee your children will or will not inherit a disease but rather provide awareness to the likelihood of a child having a disease tested for.

Read the limitations of this test to understand what is and is not covered.

Test Details

[Patient, sample, and laboratory information go here.]