



Patient Name: \_\_\_\_\_ DOB: \_\_\_\_\_ MRN: \_\_\_\_\_

**MORL – Hearing Loss Testing Requisition Form**

**Part E) Pertinent Clinical History and Findings (Required for ALL testing).**

**Pertinent Clinical History and Findings:** (check)

**Audiologic History:**  
 (Audiograms or ABRs from all affected individuals)

Does this person have hearing loss:  Yes  No

Congenital hearing loss:  Yes  No  
 If **NOT** congenital, date/age of onset of hearing loss: \_\_\_\_\_

Early childhood  
 Late

**Inheritance Pattern**

Recessive  
 Dominant  
 X-linked  
 Other \_\_\_\_\_

**Degree of Hearing Loss**

Mild  
 Moderate  
 Severe  
 Profound  
 Symmetric  
 Asymmetric

**Type of Hearing Loss**

Conductive  
 Sensorineural  
 Syndromic  
 Nonsyndromic

**Family History:**

Other persons with hearing loss (if yes you **MUST** attach a pedigree):  
 Yes  No  
 If yes, relationship to person named on this sheet \_\_\_\_\_

**Family History Comments:** \_\_\_\_\_

**Pertinent Clinical History and Findings:** (check)

**Physical Examination:**  
 (provide clinic notes pertaining to hearing loss)

Excluding hearing loss, physical examination entirely normal:  
 Yes  No

If no, were features of syndromic hearing loss present? Describe features or name syndrome: \_\_\_\_\_

**Vision:**

Does your patient wear corrective lenses?  Yes  No

If yes, please provide diopter of vision correction:  
 Left: \_\_\_\_\_ / Right: \_\_\_\_\_

Eye findings (eg. cataracts)?  Yes  No  
 If yes, please describe: \_\_\_\_\_

**Motor Milestones:**

Delayed sitting:  Yes  No  
 If yes, approximate age patient sat: \_\_\_\_\_

Delayed walking:  Yes  No  
 If yes, approximate age patient walked: \_\_\_\_\_

**Radiologic Studies:**

CT scan / MRI of temporal bones (If yes, a disk of the temporal bone images or copy of report **MUST** be provided):  
 Yes  No

Report read as Dilated Vestibular Aqueducts?  
 Yes  No

**OR** Mondini malformation?  
 Yes  No

**Comments:**

\_\_\_\_\_

## Molecular Otolaryngology & Renal Research Laboratories Sample Requirements

Questions? Contact Amy Weaver at 319-335-6623 or [amy-weaver@uiowa.edu](mailto:amy-weaver@uiowa.edu)

### MORL – Hearing Loss Testing Requisition Form

**DNA Testing Sample requirement** Lavender (EDTA) top tubes (*for whole blood*)

**Minimum sample volume** 8-10 cc. whole blood is sufficient to run all tests requested  
(We are not responsible for broken tubes.)  
OR  
10 µg DNA, minimum concentration 50ng/ul - (A260/A280 1.8-2) resuspended in  
0.1mM EDTA (10mM Tris HCl, 0.1mM EDTA, pH 8, Teknova Cat# T0220)

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***Please note if you send less than the minimum volume/concentration of blood and/or DNA your request for testing will be rejected and a new sample will be required to perform the testing indicated on page 1 of this testing requisition form.***

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#### **Blood or DNA Shipping requirements:**

Overnight delivery, Room temperature (**DO NOT FREEZE**)

Delivery: Monday-Friday.

Samples may be refrigerated if delivery is delayed.

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**Information to Include:** Testing will not be performed unless ALL of the following information is provided and the minimum sample requirement (found on page 3) is met.

**Specimen information:** Patient identifiers (**full name, date of birth, sex and medical record number**)  
Patient address, necessary for receipt and/or reporting results  
Pertinent history and clinical findings (found on page 2)  
Date of collection & Sample type  
Ordering physician

**Billing information:** We will NOT bill insurance, Medicare or patient directly.  
**Institutional billing accepted. Visa and Master Card accepted.**  
**Personal checks NOT accepted.**  
*Please include contact information including phone & fax number for billing questions.*

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#### **DISCLAIMER:**

*This request to order molecular diagnostic tests from the MORL certifies to the MORL that the ordering physician has obtained informed consent from the patient as required by applicable state or federal laws for each test ordered, that the ordering physician has authorization from the patient permitting the MORL to report results for each test ordered to the ordering physician, and that the ordering physician assumes responsibility for providing the patient with all associated guidance and counseling regarding the test results.*

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For more information about the tests we offer at the MORL please visit our website at:

<https://morl.lab.uiowa.edu>

**PLEASE INCLUDE A PEDIGREE, AUDIOGRAMS, TEMPORAL BONE IMAGING STUDIES (if performed) & CLINIC NOTES REGARDING HEARING LOSS FOR ALL TESTING REQUESTS**

### **MORL – Hearing Loss Testing Requisition Form**

#### **IMPORTANT INFORMATION FOR PHYSICIAN OR GENETIC COUNSELOR:**

DNA tests may detect an abnormality. Detection methods are greater than 99% accurate. Many of these tests are new. The analysis and interpretation represents our best knowledge and understanding of the genetics of deafness.

There is a small possibility that a test may not work properly or an error may occur. You may be asked for an additional sample if it is felt that confirmatory testing is needed.

An error in diagnosis may occur if incorrect information is provided with the sample.

Hearing loss is a complex disorder and penetrance of a phenotype (the degree of hearing loss, for example) may be variable. Research to determine genotype-phenotype correlations is ongoing.

Unexpected test results may be detected. These results may occur with screening tests that evaluate many different genetic regions. From these tests, information may be learned about you, your child/children or your family that is not directly related to the clinical reason for ordering the test (secondary results). This information may provide data about the risk for a different genetic disease with symptoms that may or may not be currently evident.

A risk associated with this testing is that genetic testing has the potential to unintentionally reveal information regarding a child's paternity. Although the goal of this testing does not relate to paternity, during testing we may find genetic changes in both parents that may indicate possible non-paternity. In some cases non-paternity may be confused with genetic changes that arise spontaneously, called de novo variants. We will not attempt to resolve or disclose issues of non-paternity during the course of this testing.

Because of the complexity of DNA testing, results should be discussed with a genetic counselor or physician.

We follow guidelines proposed by the American College of Medical Genetics and do not offer carrier testing for autosomal recessive deafness in minors.

**Note: Hearing loss is a very complex disorder. This complexity means that variants in many different genes can lead to deafness. It is possible that no variants will be detected in the variant screens (the genes) you have requested.**

If your patient is interested in joining research studies ongoing in the MORL laboratories to investigate other genetic causes of deafness please ask your patient to email Dr. Richard JH Smith ( [richard-smith@uiowa.edu](mailto:richard-smith@uiowa.edu)), Carla Nishimura ( [carla-nishimura@uiowa.edu](mailto:carla-nishimura@uiowa.edu)), Amanda Schaefer ( [amanda-schaefer-1@uiowa.edu](mailto:amanda-schaefer-1@uiowa.edu)) or Amy Weaver ( [amy-weaver@uiowa.edu](mailto:amy-weaver@uiowa.edu)).