

# REPROXPANDED TEST REQUISITION FORM

| PATIENT INFORMATION   |  |
|---|--|
| First name  | Last name  |
| Sex <input type="radio"/> Male <input type="radio"/> Female<br>Gender identification (optional): _____  | Date of birth (mm/dd/yy)   |
| Ancestry <input type="radio"/> White/Caucasian <input type="radio"/> Hispanic <input type="radio"/> Black/African American<br><input type="radio"/> Native American <input type="radio"/> East Asian <input type="radio"/> South Asian<br><input type="radio"/> Middle Eastern <input type="radio"/> Ashkenazi Jewish<br><input type="radio"/> Other: _____ |  |
| Email   |  |
| Address   |  |
| City  | State Zip code   |
| Primary phone   | Is this patient deceased? <input type="radio"/> Yes <input type="radio"/> No<br>Deceased date: _____ |

| SAMPLE INFORMATION   |                                 |
|--|---------------------------------|
| Date sample obtained (mm/dd/yy)  | Medical record #                |
| <input type="radio"/> Blood <input type="radio"/> Buccal swab<br><input type="radio"/> Other: _____<br><input type="radio"/> DNA: tissue source _____  |                                 |
| Patient has had a blood transfusion <input type="radio"/> Yes <input type="radio"/> No<br>(2-4 weeks of wait time is required for some testing)  | Date of last transfusion: _____ |
| Patient has had an allogenic bone marrow transplant <input type="radio"/> Yes <input type="radio"/> No<br>Fibroblasts are recommended for patients who had an allogenic bone marrow transplant.<br>See <a href="http://www.genedx.com/specimen-requirements">www.genedx.com/specimen-requirements</a> for details. |                                 |
| <input type="radio"/> Treatment-Related RUSH   | Date: _____                     |

| PATIENT CONSENT  |      |
|--|------|
| By signing this form I acknowledge as the patient that I have read the attached informed consent document and that I authorize GeneDx to perform genetic testing as described. I have been informed that GeneDx may contact me or my healthcare provider about research opportunities in the future. More information, including the GeneDx Notice of Privacy Policies, is available on GeneDx's website: <a href="http://www.genedx.com">www.genedx.com</a> |      |
| <input type="radio"/> By checking this box, I confirm that I am a New York state resident, and I give permission for GeneDx to retain any remaining sample longer than 60 days after the completion of testing.  |      |
| <input type="radio"/> Check this box if you wish to opt out of being contacted for research studies.   |      |
| Signature of patient (required)  | Date |

| ACCOUNT INFORMATION   |                                   |
|---|-----------------------------------|
| Account number  | Account name                      |
| Phone   | Fax                               |
| Address City  |                                   |
| State   | Zip code Country                  |
| Ordering provider   |                                   |
| Name  | Role/Title                        |
| Phone   | NPI                               |
| Email address (for report access)   |                                   |
| Reporting Preference: <input type="radio"/> Portal <input type="radio"/> Fax <input type="radio"/> Email <input type="radio"/> Care Evolve <small>If unmarked, we will use the account's default preferences or fax to new clients.</small> |                                   |
| Additional Reporting Providers <input type="radio"/> Same as ordering provider  |                                   |
| Name  | Role/Title                        |
| Phone   | NPI                               |
| Email address (for report access)   |                                   |
| Additional clinical or laboratory contact (optional)  |                                   |
| Name  | Email address (for report access) |
| SEND ADDITIONAL REPORT COPIES TO  |                                   |
| Healthcare provider/Acct #  | Fax #/Email                       |

| STATEMENT OF MEDICAL NECESSITY  |      |
|---|------|
| By submission of this test requisition and accompanying sample(s), I: (i) authorize and direct you to perform the testing indicated; (ii) certify that I am authorized by state law to order the test(s) requested herein; (iii) certify that any custom panel and/or ordered test(s) requested on this test requisition form are reasonable and medically necessary for the diagnosis and/or treatment of a disease, illness, impairment, symptom, syndrome or disorder; (iv) the test results will determine my patient's medical management and treatment decisions of this patient's condition on this date of service; (v) have obtained this patient's written informed consent to undergo any genetic testing requested; and (vi) that the full and appropriate diagnosis code(s) are indicated to the highest level of specificity. |      |
| Signature of provider (required)  | Date |

|                     |                              |
|---------------------|------------------------------|
| ICD-10 codes:       |                              |
| Clinical diagnosis: | Age at initial presentation: |

|  |
|--|
| PATIENT STATUS – ONE MUST BE CHECKED: <input type="radio"/> Hospital outpatient <input type="radio"/> Hospital inpatient Date of discharge: _____ <input type="radio"/> Not a hospital patient |
|--|

| PAYMENT OPTIONS   |  |
|---|--|
| <input type="radio"/> Patient Bill<br>Amount: _____<br>If Patient Bill is selected, I am electing to be treated as a self-pay patient for this testing. I agree that neither GeneDx nor I will submit a claim to my insurance for this testing, if I have insurance. GeneDx will send an invoice to the patient listed above. | <input type="radio"/> Institutional Bill<br>GeneDx account # Hospital/Lab name |
| <input type="radio"/> GeneDx Affiliate Code:  | Place sticker/stamp here   |
|   |  |

| TEST MENU   |  |
|---|--|
| TEST CODE   | TEST NAME  |
| <input type="radio"/> J776  | ReproXpanded Individual  |
| <input type="radio"/> J842  | ReproXpanded Couple<br>Partner's information: <input type="radio"/> To be sent within three weeks* |
| First Name  | Last Name DOB  |
| *PARTNER'S SAMPLE MUST BE RECEIVED WITHIN 3 WEEKS. If not received, testing will convert to J776 ReproXpanded Individual test.<br>Please submit a separate requisition for partner. |  |

# CLINICAL INFORMATION

|            |              |               |
|------------|--------------|---------------|
| Account #  | Account Name |               |
| First Name | Last Name    | Date of Birth |

**CLINICAL INFORMATION CAN AID IN THE ACCURATE INTERPRETATION OF RESULTS. PLEASE ATTACH RELEVANT MEDICAL RECORDS.  
PLEASE CHECK ALL THAT APPLY.**

## CLINICAL INFORMATION

Is the patient/partner currently pregnant?  No  Yes (If Yes, EDD: \_\_\_\_\_)

Prior genetic testing performed on patient: \_\_\_\_\_

Personal and/or family history of known genetic condition?  No  Yes

If Yes, please indicate relationship to patient being testing and diagnosis, including gene mutation if known: \_\_\_\_\_

## PREGNANCY HISTORY (CHECK ALL THAT APPLY)

Infertility:  No  Yes

Recurrent pregnancy loss:  No  Yes

Prior intrauterine fetal demise (IUD):  No  Yes

Current and/or prior pregnancy with anomalies:  No  Yes

If Yes, please describe and/or provide diagnosis: \_\_\_\_\_

## Additional relevant clinical information:

\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_



Signature of provider (required)

Date

|            |              |               |  |
|------------|--------------|---------------|--|
| Account #  | Account Name |               |  |
| First Name | Last Name    | Date of Birth |  |

## General Information About Genetic Testing

### What is genetic testing?

DNA provides instructions for our body's growth and development. Genes are distinct sequences of DNA, and are arranged on chromosomes. The DNA in a gene contains instructions for making proteins, which determine things like growth and metabolism as well as traits like eye color and blood type. Genetic disorders are caused by certain changes in DNA affecting the structure or number of chromosomes. Genetic testing is a laboratory test that tries to identify these changes in chromosomes or the DNA. Genetic testing can be a diagnostic test, which is used to identify or rule out a specific genetic condition. Genetic screening tests are used to assess the chance for a person to develop or have a child with a genetic condition. Genetic screening tests are not typically diagnostic and results may require additional testing.

The purpose of this test is to see if I, or my child, may have a genetic variant or chromosome rearrangement causing a genetic disorder or to determine the chance that I, or my child, will develop or pass on a genetic disorder in the future. 'My child' can also mean my unborn child, for the purposes of this consent.

If I/my child already know the specific gene variant(s) or chromosome rearrangement that causes the genetic disorder in my family, I will inform the laboratory of this information.

### What could I learn from this genetic test?

The following describes the possible results from the test:

**1) Positive:** A positive result indicates that a genetic variant has been identified that explains the cause of my/my child's genetic disorder or indicates that I/my child am at increased risk to develop the disorder in the future. It is possible to test positive for more than one genetic variant.

**2) Negative:** A negative result indicates that no disease-causing genetic variant was identified by the test performed. It does not guarantee that I/my child will be healthy or free from genetic disorders or medical conditions. If I/my child test negative for a variant known to cause the genetic disorder in other members of my/my child's family, this result rules out a diagnosis of the same genetic disorder in me/my child due to this specific change.

**3) Inconclusive/Variant of Uncertain Significance (VUS):** A finding of a variant of uncertain significance indicates that a genetic change was detected, but it is currently unknown whether that change is associated with a genetic disorder either now or in the future. A variant of uncertain significance is not the same as a positive result and does not clarify whether I/my child is at increased risk to develop a genetic disorder. The change could be a normal genetic variant or it could be disease-causing. Further analysis may be recommended, including testing parents and other family members. Detailed medical records or information from other family members also may be needed to help clarify results.

**4) Unexpected results:** In rare instances, this test may reveal an important genetic change that is not directly related to the reason for ordering this test. For example, this test may tell me about the risk for another genetic condition I/my child is not aware of or it may indicate differences in the number or rearrangement of sex chromosomes. This information may be disclosed to the ordering health care provider if it likely impacts medical care.

Result interpretation is based on currently available information in the medical literature, research and scientific databases. Because the literature, medical and scientific knowledge are constantly changing, new information that becomes available in the future may replace or add to the information GeneDx used to interpret my/my child's results. Providers can contact GeneDx at any time to discuss the classification of an identified variant. In addition, I or my/my child's health care providers may monitor publicly available resources used by the medical community, such as ClinVar ([www.clinvar.com](http://www.clinvar.com)), to find current information about the clinical interpretation of my/my child's variant(s).

For tests that evaluate data from multiple family members, my spouse, or partner concurrently, results may be included in a single comprehensive report.

### What are the risks and limitations of this genetic test?

- Genetic testing is an important part of the diagnostic process. However, genetic tests may not always give a definitive answer. In some cases, testing may not identify a genetic variant even though one exists. This may be due to limitations in current medical knowledge or testing technology.
- Accurate interpretation of test results may require knowing the true biological relationships in a family. Failing to accurately state the biological relationships in my/my child's family may result in incorrect interpretation of results, incorrect diagnoses, and/or inconclusive test results. In some cases, genetic testing can reveal that the true biological relationships in a family are not as they were reported. This includes non-paternity (the stated father of an individual is not the biological father) and consanguinity (the parents of an individual are related by blood). It may be necessary to report these findings to the health care provider who ordered the test.
- Genetic testing is highly accurate. Rarely, inaccurate results may occur for various reasons. These reasons include, but are not limited to: mislabeled samples, inaccurate reporting of clinical/medical information, rare technical errors, or unusual circumstances such as bone marrow transplantation, or the presence of change(s) in such a small percentage of cells that the change(s) may not be detectable by the test (mosaicism).
- This test does not have the ability to detect all of the long-term medical risks that I/my child might experience. The result of this test does not guarantee my health or the health of my child/fetus. Other diagnostic tests may still need to be done, especially when only a genetic screening test has been performed previously.
- Occasionally, an additional sample may be needed if the initial specimen is not adequate.

### Patient Confidentiality and Genetic Counseling

It is recommended that I receive genetic counseling before and after having this genetic test. I can find a genetic counselor in my area here: [www.nsgc.org](http://www.nsgc.org). Further testing or additional consultations with a health care provider may be necessary.

To maintain confidentiality, the test results will only be released to the referring health care provider, to the ordering laboratory, to me, to other health care providers involved in my/my child's diagnosis and treatment, or to others as entitled by law. The United States Federal Government has enacted several laws that prohibit discrimination based on genetic test results by health insurance companies and employers. In addition, these laws prohibit unauthorized disclosure of this information. For more information, I understand that I can visit [www.genome.gov/10002077](http://www.genome.gov/10002077).

### International Specimens

If I/my child reside outside the United States, I attest that by providing a sample for testing, I am not knowingly violating any export ban or other legal restriction in the country of my/my child's residence.

**Additional information about the specific test being ordered is available from my health care provider or I can go to the GeneDx website, [www.genedx.com](http://www.genedx.com). This information includes the complete gene lists, the specific types of genetic disorders that can be identified by the genetic test, the likelihood of a positive result, the limitations of genetic testing, as well as information about how specimens and information are stored and used.**