

CLIENT NAME/ID (FOR LAB USE): \*Mandatory fields PATIENT INFORMATION ORDERING PROVIDER INSTITUTION/PRACTICE NAME LAST NAME\* FIRST NAME\* SEX ASSIGNED AT BIRTH\* DATE OF BIRTH (MM/DD/YYYY)\* INSTITUTION PHONE / FAX INSTITUTION EMAIL MRN ETHNICITY ORDERING PROVIDER(S) ADDRESS NPI (USA)/MINC (Canada) PROVIDER TITLE (MD, DO, GC) CITY COUNTRY STATE POSTAL CODE PROVIDER ADDRESS PHONE FMAII CITY STATE POSTAL CODE COUNTRY SAMPLE COLLECTION DATE\* (MM/DD/YYYY) PROVIDER PHONE FAX/EMAIL REPORT TO SAMPLE TYPE\*: O Blood O Buccal/Saliva GC/PRIMARY CONTACT NAME Extracted DNA & DNA Source Other: (Blood, Buccal/Saliva, Tissue, Fibroblast): GC/PRIMARY CONTACT PHONE/FAX GC/PRIMARY CONTACT EMAIL PATIENT ACKNOWLEDGEMENT I have read the Informed Consent document and I give permission to Fulgent Genetics and STATEMENT OF INFORMED CONSENT its entities to perform genetic testing as described. I also give permission for my specimer By signing below, I, the ordering Medical Provider, confirm that testing is medically necessary and clinical information to be used in de-identified studies at Fulgent and for publication, if and that test results may impact medical management for the patient. appropriate. My name or other personal identifying information will not be used in or I attest that the patient has received and read the Fulgent Informed Consent document, or linked to the results of any studies and publications. More information is available at has had it read to them, and that I have fully informed the patient about the purpose. www.fulgentgenetics.com/policies/privacy-policy. capabilities, and limitations of the ordered test. The patient has voluntarily given his or her full consent for the ordered test and a signed copy of this consent is available on file. Any Check this box if you are a New York state resident and give permission for Fulgent to retain any remaining sample longer than 60 days after sample collection. Fulgent Informed Consent that the patient agrees to at a later date will supersede and replace this Informed Consent. Opt out of research Patient Signature (Required for billing purposes) Date (MM/DD/YYYY) Ordering Provider Signature (Required)\* Date (MM/DD/YYYY)\* PATIENT COMMUNICATION CONSENT By checking this box, you acknowledge and agree that: a. By executing this agreement, you are providing express written consent for Fulgent, CSI, and Inform Diagnostics, their affiliates and subsidiaries, and parties making contact on their behalf to call and text you using automatic telephone dialing systems and artificial or pre-recorded messages at the telephone number you have provided, about your out-of-pocket estimation, even if your telephone number is currently listed on any state, federal, local or corporate Do Not Call list. b. Your Consent to be contacted through the use of automatic telephone dialing systems and artificial or pre-recorded messages is not required in order to purchase any property, goods, or services. If you would like to speak with our Benefits Investigation team, you can reach them at +1888.FULGENT (+1888.385.4368), opt 3. **TEST OPTIONS** Step 1\*: Select Test(s) Step 2\*: Select Test Option(s) Singleton (Proband) (FT-TP01963) Rapid FulGenome - Whole Genome Analysis ☐ **Duo** (FT-TP01962)<sup>‡</sup> ☐ **Trio** (FT-TP01871)<sup>‡</sup> Add RISE Analysis Check this box if you wish to receive ACMG secondary findings Specimen Requirements: for the proband 1 x 4 mL EDTA TUBE \* All testing is started immediately upon arrival at Fulgent. Parental samples MUST arrive at the same time as the proband or they will not be processed as part of the For pediatric patients, including infants, we can accept a minimum of FulGenome. The test order will automatically update to reflect which family member 1 - 2 mL. Insufficient volume can cause delays or test failure. Saliva/buccal swabs will be accepted for family member testing (Duo/Trio); Additional panel or test option requests however, they are not recommended and will result in testing delays (~1 day). Step 3\*: Include Clinical Indications/Suspected Diagnosis Clinical information is REQUIRED for testing. Please attach medical records and/or

complete Page 3.

Exome/Genome Sequencing is a patient-centric, phenotype-driven analysis designed to report only variants that are of plausible clinical relevance. Please provide relevant

clinical history, suspected diagnosis, or family history.



FAMILY SAMPLES FO	OR DUG	O/TRI	OTESTING -									
COMPLETE THIS SECTION IF FAMILY SAMPLES HAVE BEEN SUBMITTED FOR TESTING						The lab may perform confirmation of parental relationships for quality control or other purposes.  Check here to opt-out See informed consent for more details.						
LAST NAME*	FIRST NAME*			LAST	NAME*	FIRS		FIRST NAME*				
SEX ASSIGNED AT BIRTH*	RTH* DATE OF BIRTH (MM/DD/YYYY)*				SEX	ASSIGNED AT BIRTH*		DATE OF BIRTH (MM/DD/YYYY)*		YYYY)*		
LATION TO PRIMARY PATIENT*  AFFECTED/UNAFFECTED STATUS*					RELA	ATION TO PRIMARY PATI	ENT*	AFFECTED/UNAFFECTED STATUS*				
SAMPLE COLLECTION DATE*	(MM/DD/	/YYYY)			SAMI	PLE COLLECTION DATE	* (MM/DD/	YYYY)				
SAMPLE TYPE*: O Blood O Buccal/Saliva					SAME	SAMPLE TYPE*: O Blood O Buccal/Saliva						
Extracted DNA & DNA Source (Blood, Buccal/Saliva, Tissue, Fibroblast):					O Extracted DNA & DNA Source (Blood, Buccal/Saliva, Tissue, Fibroblast):							
PATIENT ACKNOWLEDGEMENT  I have read the Informed Consent document and I give permission to Fulgent Genetics and its entities to perform genetic testing as described. I also give permission for my specimen and clinical information to be used in de-identified studies at Fulgent and for publication, if appropriate. My name or other personal identifying information will not be used in or linked to the results of any studies and publications. More information is available at www.fulgentgenetics.com/policies/privacy-policy.					l have entiti clinic appro linke	PATIENT ACKNOWLEDGEMENT  I have read the Informed Consent document and I give permission to Fulgent Genetics and its entities to perform genetic testing as described. I also give permission for my specimen and clinical information to be used in de-identified studies at Fulgent and for publication, if appropriate. My name or other personal identifying information will not be used in or linked to the results of any studies and publications. More information is available at www.fulgentgenetics.com/policies/privacy-policy.						
Check this box if you are a New York state resident and give permission for Fulgent to retain any remaining sample longer than 60 days after sample collection.						Check this box if you are a New York state resident and give permission for Fulgent to retain any remaining sample longer than 60 days after sample collection.						
Opt out of research						Opt out of research						
x					X	X						
Patient Signature (Required in BILLING OPTIONS  Please select a billing opt	ion and c	complet	e the relevant fie		Insuran		Self-Pay	By sig Fulge (inclu	pring above, the patient of ent and its entities to cont ding via text), and author es to release medical info	act them directly izes Fulgent and its		
INSURANCE/BILLING I ICD-10 VALID CODE	REFERRA			HOSPITAL ST		ards, ABN, medical criteria	rorm al Inpatient		est to the assigned insura on-Hospital Outrea			
				COLLECTED,			l Outpatier	_				
RIMARY INSURANCE ID		INS	INSURANCE NAME		STATE	GROUP		INSURANCE PHONE #				
INSURANCE PLAN		NA	NAME OF INSURED			RELATION TO PATIENT		DATE OF BIRTH (MM/DD/YYYY)				
SECONDARY INSURANCE ID		INS	INSURANCE NAME			GROUP		INSURANCE PHONE #				
INSURANCE ID			NAME OF INSURED			RELATION TO PATIENT		DATE OF BIRTH (MM/DD/YYYY)				
INSTITUTIONAL BILLIN  Use institution information  Use information below for	above for	billing			O U	F-PAY se patient information ab se information below for		ng the	signing above, the patier Igent and its entities to c em directly, and use the p tructions to bill the indic	ontact to contact provided billing		
INSTITUTION/PRACTICE NAM	ME	AT.	TENTION TO		PAYO	OR LAST NAME		PAY	OR FIRST NAME			
ADDRESS					ADD	RESS		•				
CITY	STATE		POSTAL CODE	COUNTRY	CITY	CITY			POSTAL CODE	COUNTRY		
PHONE					PHO	PHONE						



CLINICAL INFORMATION	ON —						
CLINICAL DETAILS			Th f-	-4			
☐ Mosaicism ☐ Consanguinity	details of any selected indications:  Organ Transplant  Known Chromosomal Gain/Los	s	There are many factors which may affect genetic diagnostic testing: such as gene-gene interactions, high-risk ethnicity groups, and transplants. Please list any that may apply.				
☐ Bone Marrow Transplant	☐ Known Gene Gain/Loss						
CLINICAL PRESENTATIO	DN		There are many no	recentations which may not seem like a			
Please indicate any clinical preservel relevant to genetic testing:	entations and/or findings that may be		There are many presentations which may not seem like a direct association for disease. Please list the most suspected presentations and attach detailed medical				
- Behavior - Conditions - Pedigree/Family History	<ul><li>Phenotypes</li><li>Physical</li><li>Symptoms</li></ul>	records and/or pedigree.					
CLINICAL TESTING							
	entations and/or findings that may be re	elevant to	Please also include tests that had a negative result.  These tests help our clinical staff process the results of your testing.				
<ul><li>Karyotype</li><li>Previous Genetic Testing</li><li>Vision</li><li>Hearing</li></ul>	<ul><li>Growth Measurements</li><li>Biochemical Testing</li><li>Imaging</li><li>Pathology Reports</li></ul>						
FAMILY HISTORY —							
FAMILY MEMBER 1		FAMILY MEMB	FAMILY MEMBER 2				
LAST NAME   FIRST NAME		LAST NAME		FIRST NAME			
DEL ATION TO DATIENT		RELATION TO PATI	- NIT				
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SEX ASSIGNED AT BIRTH			SEX ASSIGNED AT BIRTH				
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DIAGNOSIS AND/OR STMPTOMS			W 211,1E I O 1,19				
AGE OF ONSET DOB (MM/DD/YYYY)		AGE OF ONSET		DOB (MM/DD/YYYY)			



#### **INSTRUCTIONS** -

- 1. Complete the patient and provider information section.
- Read and sign the informed consent policy statement. The complete
  patient informed consent form for genetic testing can be found on
  FulgentGenetics.com. Signature from the provider on Page 1 of the TRF is
  required for all testing. Signature from the patient is required for billing
  purposes and communication consent.
- 3. Select or write in the test name and indicate any relevant test options. Please call us if you have any questions.
- 4. Visit our website for our most updated list of available genes.
- For Duo/Trio testing, please complete the Family Samples section or submit a separate TRF for each sample.
- 6. Please visit FulgentGenetics.com for specimen requirements.
- Extracted DNA must be extracted from a CLIA-certified laboratory or a laboratory meeting equivalent requirements as determined by CAP and/ or CMS.

### REQUIRED FOR INSURANCE CHECKLIST

☐ Detailed medical record (pedigree if available)
☐ ICD-10 code(s)
☐ Physician, patient, and insured signatures
☐ Copy of insurance card(s) - front/back
☐ Insurer specific forms (i.e. ABN)
☐ Insurance authorization, if available
☐ For medicare, medicare criteria form is required

For the most updated information and limitations on our products and services, please visit FulgentGenetics.com.