

**LG-OrT-FR-026**

## MOLECULAR GENETICS AND CYTOGENETICS TESTS REQUEST FORM

Karyotyping and Rapid Aneuploidy (QF PCR) Analyses	FISH Panels
<p><input type="checkbox"/> <b>C8392391</b> Karyotyping (Chromosome Analysis); Abortion Material+Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</p> <p><input type="checkbox"/> <b>C8390710</b> Karyotyping (Chromosome Analysis); Amniotic Fluid (Twins)+Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</p> <p><input type="checkbox"/> <b>C8390788</b> Karyotyping (Chromosome Analysis); Amniotic Fluid+Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</p> <p><input type="checkbox"/> <b>C8390716</b> Karyotyping (Chromosome Analysis); Chorionic Villus Sampling (CVS)(Twins)+Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</p> <p><input type="checkbox"/> <b>C8390715</b> Karyotyping (Chromosome Analysis); Chorionic Villus Sampling (CVS)+Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</p> <p><input type="checkbox"/> <b>C8390703</b> Karyotyping (Chromosome Analysis); Peripheral Blood</p> <p><input type="checkbox"/> <b>C8392374</b> Karyotyping (Chromosome Analysis); Fetal Blood (Cordocentesis)+Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</p> <p><input type="checkbox"/> <b>C8390702</b> Karyotyping (Chromosome Analysis); Bone marrow</p> <p><input type="checkbox"/> <b>C8390711</b> Karyotyping (Chromosome Analysis); Hemato-oncologic (peripheral blood)</p> <p><input type="checkbox"/> <b>C8392322</b> Chromosomal Instability Syndromes, DEB Test</p> <p><input type="checkbox"/> <b>C8391003</b> Chromosomal Instability Syndromes; MMC Test</p> <p><input type="checkbox"/> <b>C8392309</b> Chromosomal Instability Syndromes, SCE (Sister Chromosomes Exchange) Analysis</p> <p><input type="checkbox"/> <b>C8392297</b> Chromosomal Instability Syndromes, Spontaneous Breakage Analysis</p> <p><input type="checkbox"/> <b>C8392035</b> Karyotyping (Chromosome Analysis) Solid Tissue Sampling (Skin Biopsy)</p> <p><input type="checkbox"/> <b>C8392249</b> Karyotyping ( Subtelomeric Screening); Infertility, Bad Obstetric History Panels</p>	<p><input type="checkbox"/> <b>C8389920</b> <b>FISH_ALL Panel</b>; 8q24 cMYC, 9p21/CEP9 P16 (CDKN2A), t(9;22) BCR/ABL, 11q23 MLL, t(12;21) TEL/AML1, 14q32 IGH, 17p13 P53, E2A/PBX1 t(1;19), Trisomy 10, Trisomy 4, Monosomy 3</p> <p><input type="checkbox"/> <b>C8389926</b> <b>FISH_NHL (Non Hodgkin Lymphoma) Panel</b>; 3q27.3 BCL6, t(8;14) IGH/MYC, t(11;14) CCND1/IGH, t(14;18) IGH/BCL2, 14q32 IGH, 17p13 P53, 18q21 MALT1</p> <p><input type="checkbox"/> <b>C8389980</b> <b>FISH_Multiple Myeloma (MM) Panel</b>; t(4;14) IGH/FGFR3, t(11;14) IGH/CCND1, del 13 (RB1)/D13S25 (13q14), del D134S319/13q34/Trisomy 12, t(14;16) IGH/MAF, 14q32 IGH, 17p13.1 p53</p> <p><input type="checkbox"/> <b>C8389922</b> <b>FISH CLL Panel</b>; MYB (6q23.3)/D6Z1, ATM (11q22.3), IGH/CCND1, 13q14/13q34/CEP12, RB1/13q14), IGH (14q32), IGH/BCL2, TP53 (17p13.1)/CEP17</p> <p><input type="checkbox"/> <b>C8389976</b> <b>FISH_CML Panel</b>; t(9;22) BCR/ABL, CEP8, t(15;17) PML/RARA</p> <p><input type="checkbox"/> <b>C8392621</b> <b>FISH_AML Panels</b>; 5p15.31/5q31.1 EGR1, 7q22.1/7q31, t(8;21) AML1/ETO, 11q23 MLL, 16q22 (inv16), t(15;17) PML/RARA, 17p13 P53 t(6;9)(p22;q34) DEK/NUP214, t(7;12)(q36;p13) MNX1/ETV6, t(9;22) inv(3)</p> <p><input type="checkbox"/> <b>C8392622</b> <b>FISH_MDS Panels</b>; 5p15.31/5q31.1 EGR1, 7q22.1/7q31, 17p13 P53, 20q12/20q13.12, Trisomy 8</p> <p><input type="checkbox"/> <b>C8392629</b> <b>FISH_MPN Panels</b>; FIP1L1/CHIC2/PDGFR, t(9;22) BCR/ABL, 7q22/del 7q36, 13q34, +8 (trisomy 8), 17p13.1 (p53), 20q12, Trisomy 21</p> <p><input type="checkbox"/> <b>C8392644</b> <b>FISH_Neuroblastoma Panels</b>; (1p, 11q, 17q and N-MYC)</p>
Microarray (Molecular Karyotyping) Analyses	FISH Analyses
<p><input type="checkbox"/> <b>C8392124</b> <b>Microarray (Abortion Material)</b></p> <p><input type="checkbox"/> <b>C8392094</b> <b>Microarray (Postnatal Diagnostic Array)</b></p> <p><input type="checkbox"/> <b>C8392100</b> <b>Microarray TRIO (Prenatal Screening Array, Parent-Child), Amniotic Fluid</b></p> <p><b>Note:</b> Parent-fetal analyses done together. EDTA and heparinized blood samples of parents are required. Rapid aneuploidy analysis (QF PCR), long-term cell-tissue culture, and chromosome analysis are included</p> <p><input type="checkbox"/> <b>C8392163</b> <b>Microarray TRIO (Prenatal Screening Array, Parent-Child), CVS material</b></p> <p><b>Note:</b> Parent-fetal analyses done together. EDTA and heparinized blood samples of parents are required. Rapid aneuploidy analysis (QF PCR), long-term cell-tissue culture, and chromosome analysis are included</p> <p><input type="checkbox"/> <b>C8392274</b> <b>Microarray, Prenatal Array, CVS</b></p> <p><b>Note:</b> Rapid aneuploidy analysis (QF PCR), long-term cell-tissue culture, and chromosome analysis are included</p> <p><input type="checkbox"/> <b>C8392275</b> <b>Microarray, Prenatal Array, Amniotic Fluid</b></p> <p><b>Note:</b> Rapid aneuploidy analysis (QF PCR), long-term cell-tissue culture, and chromosome analysis are included (</p> <p><input type="checkbox"/> <b>C8392409</b> <b>Microarray, Prenatal Array, Cord Blood</b></p> <p><b>Note:</b> Rapid aneuploidy analysis (QF PCR), long-term cell-tissue culture, and chromosome analysis are included</p> <p><input type="checkbox"/> <b>C8392278</b> <b>Microarray, Prenatal Array, Parents</b></p>	<p><input type="checkbox"/> <b>C8392072</b> FISH_t(11;19)(MLL/MLLT1)</p> <p><input type="checkbox"/> <b>C8392073</b> FISH_t(9;11) MLLT3/MLL</p> <p><input type="checkbox"/> <b>C8392232</b> FISH_t(4;11) MLL/AFF1</p> <p><input type="checkbox"/> <b>C8389913</b> FISH_t(11;14) IGH/CCND1</p> <p><input type="checkbox"/> <b>C8389910</b> FISH_t(14;18) IGH/BCL2</p> <p><input type="checkbox"/> <b>C8389966</b> FISH_t(15;17) PML/RARA</p> <p><input type="checkbox"/> <b>C8389914</b> FISH_(3q27) BCL6</p> <p><input type="checkbox"/> <b>C8389911</b> FISH_t(8;14) IGH/MYC</p> <p><input type="checkbox"/> <b>C8389965</b> FISH_t(9;22) BCR/ABL</p> <p><input type="checkbox"/> <b>C8389967</b> FISH_t(8;21) AML1/ETO</p> <p><input type="checkbox"/> <b>C8392617</b> FISH_t(6;9) DEK NUP214</p> <p><input type="checkbox"/> <b>C8392238</b> FISH_IKZF1 (7p12.2)</p> <p><input type="checkbox"/> <b>C8392411</b> FISH_IKZF3</p> <p><input type="checkbox"/> <b>C8392237</b> FISH_CRLF2 Rearrangement</p> <p><input type="checkbox"/> <b>C8389948</b> FISH_+12 (Trisomy 12)</p> <p><input type="checkbox"/> <b>C8389950</b> FISH_+8 (Trisomy 8)</p> <p><input type="checkbox"/> <b>C8389982</b> FISH_18q21 (MALT1)</p> <p><input type="checkbox"/> <b>C8389957</b> FISH_Del 11q22.3 (ATM)</p> <p><input type="checkbox"/> <b>C8389921</b> FISH_Del 13q14 (D13S25)</p> <p><input type="checkbox"/> <b>C8389975</b> FISH_Del 13q14 (D13S319)</p> <p><input type="checkbox"/> <b>C8389969</b> FISH_Del 13q14 (RB1)</p> <p><input type="checkbox"/> <b>C8389959</b> FISH_Del 13q34</p> <p><input type="checkbox"/> <b>C8389923</b> FISH_Del 20q12</p> <p><input type="checkbox"/> <b>C8389964</b> FISH_Del 5q31 (EGR1)</p> <p><input type="checkbox"/> <b>C8389963</b> FISH_Del 5q33-q34 (CSF1R)</p> <p><input type="checkbox"/> <b>C8389962</b> FISH_Del 7q22/del 7q35</p> <p><input type="checkbox"/> <b>C8389925</b> FISH_Del 9p21 (p16/CDKN2A)</p> <p><input type="checkbox"/> <b>C8389952</b> FISH_Del/amp 17p13.1 (p53)</p> <p><input type="checkbox"/> <b>C8389970</b> FISH_Del/t(11q23) MLL</p> <p><input type="checkbox"/> <b>C8392080</b> FISH_E2A (TCF3)</p> <p><input type="checkbox"/> <b>C8389927</b> FISH_IGH (14q32)</p> <p><input type="checkbox"/> <b>C8389960</b> FISH_inv (16)(p13q22)</p> <p><input type="checkbox"/> <b>C8392059</b> FISH_Monosomy 5</p> <p><input type="checkbox"/> <b>C8389862</b> FISH_Chimerism, X/Y Analysis with FISH</p> <p><input type="checkbox"/> <b>C8392062</b> FISH_Monosomy/Trisomy 7</p> <p><input type="checkbox"/> <b>C8389979</b> FISH_8q24 MYC</p> <p><input type="checkbox"/> <b>C8392043</b> FISH_PDGFRB</p> <p><input type="checkbox"/> <b>C8392063</b> FISH_t(1;19)(q23;p13) TCF3/PBX1</p>

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FISH Analyses		Molecular Genetic Tests	
<input type="checkbox"/> C8389916	FISH_t(14;16) IGH/MAF	<input type="checkbox"/> C8392149	Alpha Thalassemia HBA1, HBA2 Gene Deletion Duplication Analyses
<input type="checkbox"/> C8389917	FISH_t(17q21;?)/i(17q) RARA	<input type="checkbox"/> C8389815	Alpha Thalassemia HBA1, HBA2 Gene Analysis
<input type="checkbox"/> C8389981	FISH_t(4;14) IGH/FGFR3	<input type="checkbox"/> C8389842	Angiotensin ACE ins/del (ID) Polymorphism Analysis
<input type="checkbox"/> C8391012	FISH_t(12;21)(p13;q22) TEL/AML1	<input type="checkbox"/> C8392174	ATP1A3 Gene Analysis
<input type="checkbox"/> C8392152	FISH_Monosomy/Trisomy 10	<input type="checkbox"/> C8389605	Ankylosing spondylitis, HLA Analysis (HLA B*27)
<input type="checkbox"/> C8392151	FISH_Monosomy/Trisomy 4	<input type="checkbox"/> C8389837	Apo B Gene Genotype Analysis
<input type="checkbox"/> C8392410	FISH_Monozomi 1p36	<input type="checkbox"/> C8389834	Apo E Gene Genotype Analysis
<input type="checkbox"/> C8392363	FISH_Centromeric Y Chromosome	<input type="checkbox"/> C8392433	APC Gene Analysis
<input type="checkbox"/> C8392091	FISH_Subtelomeric Probe Two Chromosomes	<input type="checkbox"/> C8392095	Ataxia Telangiectasia, ATM Gene Sequence Analysis
<input type="checkbox"/> C8392090	FISH_Subtelomeric Probe Single Chromosome	<input type="checkbox"/> C8392359	Bartter Syndrome Tip 4A, BSND Gene Analysis
<input type="checkbox"/> C8391011	FISH_Subtelomeric Screening (All Chromosomes)	<input type="checkbox"/> C8389607	Behçet's disease, HLA Analysis B5 (B*51 and B*52)
<input type="checkbox"/> C8392339	FISH_Alagille Syndrome 20p12 Microdeletion	<input type="checkbox"/> C8390110	B-Raf (BRAF) p.V600E Mutation Analysis
<input type="checkbox"/> C8390202	FISH_DiGeorge 22q11.2 Microdeletion	<input type="checkbox"/> C8389820	Beta Thalassemia HBB Gene Analysis
<input type="checkbox"/> C8390203	FISH_Cri du chat 5p Microdeletion	<input type="checkbox"/> C8392380	Beta Thalassemia HBB Gene Deletion Duplication Analysis
<input type="checkbox"/> C8390208	FISH_Kallmann Xp22 Microdeletion	<input type="checkbox"/> C8389888	Beta Thalassemia HBB Gene Analysis, Prenatal
<input type="checkbox"/> C8390205	FISH_Miller-Dieker 17p13 Microdeletion	<input type="checkbox"/> C8389848	Beta Fibrinogen FGB Gene Polymorphism Analysis
<input type="checkbox"/> C8392064	FISH_Rubinstein-Taybi 16p13 Microdeletion	<input type="checkbox"/> C8392398	Biotinidase deficiency, BTD Sequence Analysis
<input type="checkbox"/> C8390206	FISH_Smith-Magenis 17p11 Microdeletion	<input type="checkbox"/> C8392230	BRCA1/BRCA2 Gene Deletion Duplication Analysis
<input type="checkbox"/> C8390204	FISH_Williams 7q11.23 Microdeletion	<input type="checkbox"/> C8392190	BRCA1-BRCA2 Gene Analysis
<input type="checkbox"/> C8390207	FISH_Wolf-Hirschhorn 4p16.3 Microdeletion	<input type="checkbox"/> C8389900	CADASIL Disease NOTCH 3 Gene Common Mutation Analysis
<input type="checkbox"/> C8392058	FISH_CHARGE Syndrome(8q12.2) Microdeletion	<input type="checkbox"/> C8389247	Canavan Disease ASPA Gene Analysis
<input type="checkbox"/> C8392060	FISH_Prader-Willi/Angelman (SNRPN)	<input type="checkbox"/> C8392614	Carrier Screening (SMA, Fragile X and Cystic Fibrosis 18 Mutations)
<input type="checkbox"/> C8392065	FISH_SHOX Microdeletions	<input type="checkbox"/> C8392348	CD40 Gene Analysis
<input type="checkbox"/> C8392061	FISH_SOTOS Syndrome (5q35.3)	<input type="checkbox"/> C8389200	Charcot Marie Tooth Type 1A PMP22 Deletion Duplication Analysis
<input type="checkbox"/> C8392164	FISH_Monosomy, Trisomy X	<input type="checkbox"/> C8389202	Charcot Marie Tooth Type 1B MPZ Gene Analysis
<input type="checkbox"/> C8392081	FISH_Trisomy 21	<input type="checkbox"/> C8389208	Charcot Marie Tooth Type 1E PMP22 Gene Analysis
<input type="checkbox"/> C8392076	FISH_SRY Analysis	<input type="checkbox"/> C8389211	Charcot Marie Tooth X-Linked Dominant Type GJB1 Gene Analysis
<input type="checkbox"/> C8392071	FISH_19q13/19p13	<input type="checkbox"/> C8392024	Coeliac Disease, HLA Analysis (HLA-DQ2 and HLA-DQ8)
<input type="checkbox"/> C8392014	FISH_2p24.3 N-MYC	<input type="checkbox"/> C8389861	Chimerism, PCR-based STR Fragment Analysis
<input type="checkbox"/> C8392074	FISH_PTEN	<input type="checkbox"/> C8392030	Chimerism, PCR-based STR Fragment Analysis, Preliminary
<input type="checkbox"/> C8392012	FISH_ALK Rearrangements	<input type="checkbox"/> C8392360	Chimerism, T Cell PCR-based STR Fragment Analysis
<input type="checkbox"/> C8392045	FISH_EWSR1	<input type="checkbox"/> C8392361	Chimerism, B Cell PCR-based STR Fragment Analysis
<input type="checkbox"/> C8399905	FISH_ROS1	<input type="checkbox"/> C8389125	Cystic Fibrosis CFTR Gene Analysis
<input type="checkbox"/> C8392070	FISH_1p36/1q25	<input type="checkbox"/> C8392216	Cystic Fibrosis, CFTR Gene, Deletion Duplication Analysis
<input type="checkbox"/> C8392264	FISH_PDGFR4_4q12 Rearrangements	<input type="checkbox"/> C8389191	Cystic Fibrosis CFTR Gene Analysis, prenatal
<input type="checkbox"/> C8392499	FISH_FIP1L1/CHIC2/PDGFR4 Deletion, Rearrangement	<input type="checkbox"/> C8392610	Cystic Fibrosis Carrier Screening
<input type="checkbox"/> C8392618	FISH_3q26.2 EVI1 (MECOM) Rearrangements	<input type="checkbox"/> C8392611	Cystic Fibrosis Carrier Screening, Prenatal
<input type="checkbox"/> C8392290	FISH_PAX5 Rearrangement	<input type="checkbox"/> C8389906	Clonality B Cell Test
<input type="checkbox"/> C8392296	FISH_Seathre-Chatzen Syndrome 7q11.23 Microdeletion	<input type="checkbox"/> C8389907	Clonality T Cell Test
<input type="checkbox"/> C8392147	Locus-Specific FISH Analysis	<input type="checkbox"/> C8392150	Congenital Adrenal Hyperplasia CYP21A2 Gene Deletion Duplication Analysis
<input type="checkbox"/> C8392654	FISH_t(7;12) (q36;p13) MNX1/ETV6	<input type="checkbox"/> C8392033	Congenital Adrenal Hyperplasia CYP21A2 Gene Analysis
<b>Thrombophilia Panels</b>		<input type="checkbox"/> C8392126	Congenital Deafness Connexin 26 (GJB2) Gene Analysis
<input type="checkbox"/> C8389838	<b>Thrombophilia Panel;</b> (Factor II, Factor V Leiden, Factor XIII, MTHFR 677, MTHFR 1298, PAI, Factor V Cambridge)	<input type="checkbox"/> C8392140	Congenital Neutropenia ELANE (ELA 2) Gene Analysis
<input type="checkbox"/> C8389838	<b>Cardiovascular Risk Panels;</b> (Factor V Leiden, Factor V (H1299R) (R2), Factor II G20210A), MTHFR 677T, MTHFR A1298C, PAI-1 4G/5G, ACE gene I/D, Factor XIII V34L, $\beta$ -Fibrinogen -455 G>A, GPIIIa L33P, Apo B, ApoE)	<input type="checkbox"/> C8392079	Congenital Neutropenia Type 3 HAX1 Gene Analysis
<b>Molecular Genetic Tests</b>		<input type="checkbox"/> C8392210	CDAN1 Gene Analysis
<input type="checkbox"/> C8392295	5 Alpha Reductase Deficiency, SRD5A2 Gene Analysis	<input type="checkbox"/> C8392214	Ceruloplasmin, CP Gene Analysis
<input type="checkbox"/> C8392265	5-Fluorouracil Sensitivity (DPD Mutation Test)	<input type="checkbox"/> C8392243	Costello Syndrome, HRAS Gene Analysis
<input type="checkbox"/> C8389610	Autosomal Dominant Periodic Fever (TRAPS) TNFR1 Gene Analysis	<input type="checkbox"/> C8392266	DNAF2 Gene Analysis
<input type="checkbox"/> C8389616	Achondroplasia Disease, FGFR3 Gene Common Mutation Analysis	<input type="checkbox"/> C8392251	Diabetes Mellitus Type 1, INS Gene Analysis
<input type="checkbox"/> C8392159	Achondroplasia Disease FGFR3 Gene Common Mutation Analysis, Prenatal	<input type="checkbox"/> C8389207	Duchenne Muscular Dystrophy (DMD) Dystrophin Deletion Duplication Analysis
<input type="checkbox"/> C8392142	Achondroplasia Disease, FGFR3 Gene Analysis	<input type="checkbox"/> C8389285	Duchenne Muscular Dystrophy (DMD) Deletion Duplication Analysis, Prenatal
<input type="checkbox"/> C8392161	Achondroplasia Disease FGFR3 Gene Analysis, Prenatal	<input type="checkbox"/> C8392125	Duchenne Muscular Dystrophy (DMD) Dystrophin Gene Analysis
<input type="checkbox"/> C8392343	Acute Intermittent Porphyria, HMBS Gene Analysis	<input type="checkbox"/> C8392013	E-CADHERIN CDH1 Gene Analysis
<input type="checkbox"/> C8389988	Acute Myeloid Leukemia (AML) CEBPA Gene Analysis	<input type="checkbox"/> C8390103	EGFR Gene Common Mutation Analysis
<input type="checkbox"/> C8389902	Alpha 1-Antitrypsin Genotyping	<input type="checkbox"/> C8392619	ETFDH Gene Sequencing Analysis
<input type="checkbox"/> C8392129	Alport syndrome COL4A3, COL4A4, COL4A5 Gene Analyses	<input type="checkbox"/> C8392280	Ethylmalonic Encephalopathy, ETHE1 Gene Analysis
		<input type="checkbox"/> C8392381	EXT1 Gene Sequencing Analysis

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## MOLECULAR GENETICS AND CYTOGENETICS TESTS REQUEST FORM

Molecular Genetic Tests	Molecular Genetic Tests
<input type="checkbox"/> C8392382 EXT2 Gene Sequencing Analysis	<input type="checkbox"/> C8392020 Interleukin 28B (Interferon lambda 3) IFNL3 Gene Analysis
<input type="checkbox"/> C8392306 Fabry Disease, GLA Gene Analysis	<input type="checkbox"/> C8389945 Imatinib (Gleevec) Resistance ABL1 Gene Mutation Analysis
<input type="checkbox"/> C8389827 Factor II (Prothrombin) Polymorphism Analysis	<input type="checkbox"/> C8392261 Kaposi Sarcoma, IL6 Gene Analysis
<input type="checkbox"/> C8392006 Factor V H1299R Polymorphism Analysis	<input type="checkbox"/> C8390117 KIT (Mast cell growth factor receptor) Gene D816V Mutation Analysis
<input type="checkbox"/> C8391007 Factor V Cambridge Polymorphism Analysis	<input type="checkbox"/> C8389901 KIT (Mast cell growth factor receptor) Gene Exon 9, 11, 13, 17 Mutation Analyses
<input type="checkbox"/> C8389831 Factor V_Leiden Polymorphism Analysis	<input type="checkbox"/> C8390112 KRAS Exon 2, 3, 4 Gene Mutation Analysis
<input type="checkbox"/> C8389993 Factor XIII Polymorphism Analysis	<input type="checkbox"/> C8392498 KRT5 Gene Sequence Analysis
<input type="checkbox"/> C8392143 Fanconi Anemia Deletion Duplication Analysis	<input type="checkbox"/> C8389939 JAK2 Gene V617F Mutation Analysis
<input type="checkbox"/> C8389909 Familial Fructose Intolerance Aldolase B (ALDOB) Gene Analysis	<input type="checkbox"/> C8389974 JAK2 Gene Mutation Analysis (Exon 12)
<input type="checkbox"/> C8392133 Familial Hypercholesterolemia (ADH) Panel	<input type="checkbox"/> C8392325 JAK3 Gene Analysis
<input type="checkbox"/> C8392132 Familial Hypertrophic Cardiomyopathy Panel	<input type="checkbox"/> C8389501 Leber's Optic Atrophy MTND1 Gene Mutation Analysis
<input type="checkbox"/> C8389282 Familial Transthyretin Amyloidosis, TTR Gene Analysis	<input type="checkbox"/> C8392245 Leptin Gene Analysis
<input type="checkbox"/> C8392130 Familial Adenomatous Polyposis (FAP) APC and MUTYH Gene Analyses	<input type="checkbox"/> C8392349 Lesch Nyhan Syndrome - HPRT1 Gene Analysis
<input type="checkbox"/> C8389600 Familial Mediterranean Fever (FMF) MEFV Gene Analysis	<input type="checkbox"/> C8392273 MGMT Gene Analyses
<input type="checkbox"/> C8392093 FGFR2 Gene Analysis	<input type="checkbox"/> C8392128 Marfan Syndrome FBN1 Gene Analysis
<input type="checkbox"/> C8392138 FLT3 Gene ITD (Internal tandem duplication) Analysis	<input type="checkbox"/> C8392641 Maple Syrup Urine Disease, Type IA, BCKDHA Gene Sequencing
<input type="checkbox"/> C8389930 FLT3 Gene p.D835 Mutation Analysis	<input type="checkbox"/> C8389903 MELAS MT-ND5 Gene Common Mutation Analysis
<input type="checkbox"/> C8392645 FLT3 Mutations Analysis (IDT and D835)	<input type="checkbox"/> C8389904 MELAS MT-TL1 Gene Common Mutation Analysis
<input type="checkbox"/> C8392493 FXN Gene Sequence Analysis	<input type="checkbox"/> C8389293 Metachromatic Leukodystrophy ARSA Gene Analysis
<input type="checkbox"/> C8389206 Fragile X FMR1 Gene CGG Triplet Repeat Status Analysis, Postnatal	<input type="checkbox"/> C8392455 MDS NGS Panel (Cytogenetics and FISH_DeI 5q)
<input type="checkbox"/> C8389210 Friedreich's Ataxia FXN Gene GAA Triplet Repeat Status Analysis	<input type="checkbox"/> C8390108 Microsatellite instability
<input type="checkbox"/> C8389250 Fructose 1,6 Bisphosphatase Deficiency FBP1 Gene Analysis	<input type="checkbox"/> C8392134 MODY Panel
<input type="checkbox"/> C8392440 G6PC Gene Analysis	<input type="checkbox"/> C8389845 MTHFR Gene c.A1298C Polymorphism Analysis
<input type="checkbox"/> C8392347 G6PC3 Gene Analysis	<input type="checkbox"/> C8389841 MTHFR Gene c.C677T Polymorphism Analysis
<input type="checkbox"/> C8389730 Galactosemia GALT Gene Common Mutation Analysis	<input type="checkbox"/> C8389829 MTHFR Gene c.C677T and c.A1298C Polymorphism Analyses
<input type="checkbox"/> C8389701 Gaucher Disease GBA Gene Analysis	<input type="checkbox"/> C8392609 MPN Screening Test (CALR, MLP, JAK2 (Ekzon 12 and V617F), CSF3R)
<input type="checkbox"/> C8392437 GATA2 Gene Analysis	<input type="checkbox"/> C8392148 Multiple Endocrine Neoplasia Type 1 MEN1 Gene Analysis
<input type="checkbox"/> C8392004 GATA1 Gene Common Mutation Analysis	<input type="checkbox"/> C8390130 Multiple Endocrine Neoplasia Type 2A RET Gene Sequence Analysis
<input type="checkbox"/> C8392239 Galactokinase Deficiency, GALK1 Gene Analysis	<input type="checkbox"/> C8399906 Myeloproliferative Disorders CALR Gene Common Mutation Analysis
<input type="checkbox"/> C8392240 GJB6 Gene Analysis	<input type="checkbox"/> C8389989 Myeloproliferative Leukemia Protein, MPL Gene Common Mutation Analysis
<input type="checkbox"/> C8392135 Genetic Arrhythmia Panel	<input type="checkbox"/> C8392623 Myeloid NGS Panels
<input type="checkbox"/> C8392287 Gilbert Syndrome, UGT1A1 Gene Analysis	<input type="checkbox"/> C8389218 Myotonic Dystrophy Type 1 DMPK Gene CTG Triplet Repeat Status Analysis
<input type="checkbox"/> C8389280 Glycogen Storage Disease Type V PYGM Gene Common Mutation Analysis	<input type="checkbox"/> C8392000 Neuronal Ceroid-Lipofuscinoses 1 (NCL Type1) Common Mutation Analysis
<input type="checkbox"/> C8392268 Glucose 6 Phosphate Dehydrogenase Deficiency, G6PD Gene Analysis	<input type="checkbox"/> C8392001 Neuronal Ceroid-Lipofuscinoses 2 (NCL Type2) Common Mutation Analysis
<input type="checkbox"/> C8392362 Glycogen Storage Disease Type IV, GBE1 Gene Analysis	<input type="checkbox"/> C8392002 Neuronal Ceroid-Lipofuscinoses 3 (NCL Type3) Common Mutation Analysis
<input type="checkbox"/> C8389844 Glycoprotein IIIA ITGB3 Gene Polymorphism Analysis	<input type="checkbox"/> C8392092 Noonan Syndrome PTPN11 Gene Analysis
<input type="checkbox"/> C8392136 Glioma H3F3A (H3.3) Gene p.K27 and p.G34 Mutation Analysis	<input type="checkbox"/> C8392162 Noonan Syndrome PTPN11 Gene Analysis, Prenatal
<input type="checkbox"/> C8392040 Glioma; IDH1, IDH2 and hTERT Genes Common Mutation Analyses	<input type="checkbox"/> C8392145 Neurofibromatosis Type 1 NF1 Gene Deletion Duplication Analysis
<input type="checkbox"/> C8389244 GM1 Gangliosidosis GLB1 Gene c.1594A>G (S532G) Mutation Analysis	<input type="checkbox"/> C8389219 Neurofibromatosis Type 1 NF1 Gene Analysis
<input type="checkbox"/> C8392099 Gonadal Cell Freezing Preparation (STR Analysis)	<input type="checkbox"/> C8392173 Neurofibromatosis Type 2 NF2 Gene Analysis
<input type="checkbox"/> C8389870 Griscelli Syndrome Type 1 MYO5A Gene Common Mutation Analysis	<input type="checkbox"/> C8392324 Neurofibromatosis Type 2 NF2 Gene Deletion Duplication Analysis
<input type="checkbox"/> C8392139 Griscelli Syndrome Type 2 RAB27A Gene Analysis	<input type="checkbox"/> C8392358 Neutrophil Specific Antigen 1, PRV-1 (CD 177) Gene Analysis
<input type="checkbox"/> C8392352 HADHA Gene Analysis	<input type="checkbox"/> C8392338 NPHP1 Gene Deletion Duplication Analysis
<input type="checkbox"/> C8389836 Hemochromatosis HFE Gene Common Mutation Analysis	<input type="checkbox"/> C8390123 Nucleophosmin (NPM1) Gene Exon 12 Mutation Analysis
<input type="checkbox"/> C8390107 Hereditary Cancer Panel (BRCA1-BRCA2 and 24 Gene Analysis)	<input type="checkbox"/> C8392213 NRAS Exon 2, 3, 4 Gene Mutation Analysis
<input type="checkbox"/> C8392353 Hereditary Fructose Intolerance Aldolase B (ALDOB) Deletion/Duplication Analysis	<input type="checkbox"/> C8392007 Opitz-Kaveggia Syndrome MED12 Gene Common Mutation Analysis
<input type="checkbox"/> C8392288 HMX1 Gene Analysis	<input type="checkbox"/> C8389620 Osteogenesis Imperfecta COL1A1 Gene Analysis
<input type="checkbox"/> C8392131 HNPCC (Hereditary Nonpolyposis Colorectal Cancer) Panel	<input type="checkbox"/> C8389614 Osteogenesis Imperfecta COL1A1 Gene c.G1245T Polymorphism Analysis
<input type="checkbox"/> C8389216 Huntington's Disease Huntingtin (IT15) CAG Triplet Repeat Status Analysis	
<input type="checkbox"/> C8392177 Hypophosphatasia, ALPL Gene Analysis	
<input type="checkbox"/> C8392272 Hypoplastic Left Heart Syndrome Panel (NKX2-5, GJA1)	
<input type="checkbox"/> C8392158 iAMP 21 Amplification Analysis	
<input type="checkbox"/> C8392366 IL21 Gene Analysis	
<input type="checkbox"/> C8392326 IL2RG (IL2 Receptor Gamma) Gene Analysis	

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**LG-OrT-FR-026**

## MOLECULAR GENETICS AND CYTOGENETICS TESTS REQUEST FORM

Molecular Genetic Tests	Molecular Genetic Tests
<ul style="list-style-type: none"> <li><input type="checkbox"/> C8392160 Osteogenesis Imperfecta COL1A2 Gene Analysis</li> <li><input type="checkbox"/> C8389259 Oligodendroglioma 1p/19q Deletions</li> <li><input type="checkbox"/> C8392291 P53 Gene Common Mutation (Exon 5, 6, 7, 8, 9) Analysis</li> <li><input type="checkbox"/> C8392313 Paraganglioma Panels; SDHB, SDHC, SDHD, VHL Gene Analysis</li> <li><input type="checkbox"/> C8389839 PAI1 Gene -675 4G/5G Polymorphism Analysis</li> <li><input type="checkbox"/> C8392146 Pelizaeus Merzbacher PLP 1 Gene Deletion Duplication</li> <li><input type="checkbox"/> C8392344 PAX3 Gene Analysis</li> <li><input type="checkbox"/> C8389283 Pelizaeus Merzbacher PLP 1 Gene Analysis</li> <li><input type="checkbox"/> C8392345 Perforin 1, PRF1 Gene Analysis</li> <li><input type="checkbox"/> C8390122 PDGFRA Gene Exon 12, 14, 18 Mutation Analysis</li> <li><input type="checkbox"/> C8389214 Prader-Willi-Angelman Syndrome SNRPN Gene Methylation Pattern Analysis</li> <li><input type="checkbox"/> C8392236 Primary Hyperoxaluria Types 1, AGXT Gene Analysis</li> <li><input type="checkbox"/> C8389241 Primary Torsion Dystonia, TOR1A Exon 5 Mutation Analysis</li> <li><input type="checkbox"/> C8389703 Phenylketonuria PAH Gene Analysis</li> <li><input type="checkbox"/> C8392356 Phenylketonuria PAH Gene Deletion/Duplication Analysis</li> <li><input type="checkbox"/> C8392178 PITX1 Gene Analysis</li> <li><input type="checkbox"/> C8392226 PFIC 1, ATP8B1 Gene Analysis</li> <li><input type="checkbox"/> C8392227 PFIC 2, ABCB11 Gene Analysis</li> <li><input type="checkbox"/> C8392228 PFIC 3, ABCB4 Gene Analysis</li> <li><input type="checkbox"/> C8392235 PABPN1 Gene Analysis</li> <li><input type="checkbox"/> C8392258 PNPLA6 Gene Analysis</li> <li><input type="checkbox"/> C8392332 Porphyria Cutanea Tarda, UROD Gene Analysis</li> <li><input type="checkbox"/> C8392330 PRSS1 Gene Analysis</li> <li><input type="checkbox"/> C8392354 PTEN Deletion/Duplication Analysis</li> <li><input type="checkbox"/> C8392292 Pycnodysostosis Syndrome, CTSK Gene Analysis</li> <li><input type="checkbox"/> C8389222 Rett Syndrome MECP2 Gene Analysis</li> <li><input type="checkbox"/> C8392355 Rett Syndrome MECP2 Gene Deletion/Duplication Analysis</li> <li><input type="checkbox"/> C8390785 Rapid Aneuploidy Analysis (X, Y, 13, 18, 21)</li> <li><input type="checkbox"/> C8392175 RPE65 Gene Analysis</li> <li><input type="checkbox"/> C8389810 Sickle Cell Anemia HBB Gene p.E7V Mutation Analysis Postnatal</li> <li><input type="checkbox"/> C8389890 Sickle Cell Anemia HBB Gene p.E7V Mutation Analysis Prenatal</li> <li><input type="checkbox"/> C8392446 SHOX Gene Sequencing Analysis</li> <li><input type="checkbox"/> C8392293 Smith Lemli Opitz Syndrome, DHCR7 Gene Analysis</li> <li><input type="checkbox"/> C8392252 Somatic BRCA1-BRCA2 Gene Analysis</li> <li><input type="checkbox"/> C8389287 Spinal Muscular Atrophy (SMA) Deletion/Duplication Analysis Prenatal</li> <li><input type="checkbox"/> C8389217 Spinal Muscular Atrophy (SMA) Deletion/Duplication Analysis Postnatal</li> <li><input type="checkbox"/> C8389231 Spinocerebellar Ataxia (SCA) Type 1 ATXN1 CAG Triplet Repeat Status Analysis</li> <li><input type="checkbox"/> C8389232 Spinocerebellar Ataxia (SCA) Type 2 ATXN2 CAG Triplet Repeat Status Analysis</li> <li><input type="checkbox"/> C8389233 Spinocerebellar Ataxia (SCA) Type 3 ATXN3 CAG Triplet Repeat Status Analysis</li> <li><input type="checkbox"/> C8389634 Spinocerebellar Ataxia (SCA) Type 6 CACNA1A CAG Triplet Repeat Status Analysis</li> <li><input type="checkbox"/> C8389635 Spinocerebellar Ataxia (SCA) Type 7 ATXN7 CAG Triplet Repeat Status Analysis</li> <li><input type="checkbox"/> C8389230 Spinocerebellar Ataxia (SCA) Type 1,2,3,6,7 Triplet Repeat Status Analysis</li> <li><input type="checkbox"/> C8392331 SPINK1 Gene Analysis</li> <li><input type="checkbox"/> C8392211 SEC23B Gene Analysis</li> <li><input type="checkbox"/> C8392242 SH2D1A Gene Analysis</li> <li><input type="checkbox"/> C8392097 Targeted Mutation Analysis (Single Mutation)</li> <li><input type="checkbox"/> C8392166 Targeted Mutation Analysis, Prenatal (Single Mutation)</li> <li><input type="checkbox"/> C8392491 Targeted Mutations Analysis (Two Mutations)</li> <li><input type="checkbox"/> C8392492 Targeted Mutations Analysis (Tree Mutations)</li> <li><input type="checkbox"/> C8392267 Thyroxine-binding Globulin Deficiency, TBG (SERPINA7) Gene Analysis</li> <li><input type="checkbox"/> C8392336 Thyroid Hormone Resistance THRB Gene Analysis</li> <li><input type="checkbox"/> C8389990 TPMT Genotyping</li> </ul>	<ul style="list-style-type: none"> <li><input type="checkbox"/> C8392357 TPO Gene Analysis</li> <li><input type="checkbox"/> C8392215 TSC2 Gene, Deletion Duplication Analysis</li> <li><input type="checkbox"/> C8392289 TSH Receptor Gene (TSHR) Analysis</li> <li><input type="checkbox"/> C8392659 TWIST1 Gene Analysis</li> <li><input type="checkbox"/> C8389410 Y Chromosome Microdeletions Screening</li> <li><input type="checkbox"/> C8392016 Von Hippel Lindau VHL Gene Analysis</li> <li><input type="checkbox"/> C8389840 Warfarin Resistance VKORC1 Polymorphism Analysis</li> <li><input type="checkbox"/> C8389843 Warfarin Resistance CYP2C9 Gene Polymorphism Analysis</li> <li><input type="checkbox"/> C8389835 Wiskott-Aldrich Syndrome WAS Gene Analysis</li> <li><input type="checkbox"/> C8392036 Wilm's Tumor_1p/16q deletions</li> <li><input type="checkbox"/> C8392003 Wilson's Disease ATP7B Gene Common Mutation Analysis</li> <li><input type="checkbox"/> C8392294 X-linked ALD (ABCD1) Gene Analysis</li> </ul> <div style="text-align: center; border-top: 1px solid black; border-bottom: 1px solid black; padding: 2px;"><b>Fusion Transcripts</b></div> <ul style="list-style-type: none"> <li><input type="checkbox"/> C8389931 t(8;21) AML1/ETO Fusion Transcripts</li> <li><input type="checkbox"/> C8389938 inv 16(p13;q22) CBFβ-MYH11 Fusion Transcripts</li> <li><input type="checkbox"/> C8389943 t(4;11)(q21;q23) MLL-AF4 Fusion Transcripts</li> <li><input type="checkbox"/> C8389937 t(15;17) PML/RARA Fusion Transcripts</li> <li><input type="checkbox"/> C8391013 t(1;19) E2A-PBX Fusion Transcripts</li> <li><input type="checkbox"/> C8389955 t(12;21) TEL/AML1 Fusion Transcripts</li> <li><input type="checkbox"/> C8389935 t(9;22) BCR/ABL Fusion Transcript</li> </ul> <div style="text-align: center; border-top: 1px solid black; border-bottom: 1px solid black; padding: 2px;"><b>Carrier Screening Test</b></div> <ul style="list-style-type: none"> <li><input type="checkbox"/> C8392614 Carrier Screening (SMA, Fragile X and Cystic Fibrosis 18 Mutations)</li> </ul>

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