

PLACE 1 BARCODE ON FORM AND 1 ON SAMPLE (REOUIRED: NAME/DOB)

# **Cardiac Test Requisition Form**

First Name	Last I	Vame		Middle Initial	Clinic Name	
Social Security #	Date of Birth Sex □ F	Ethnicity □ M □ Africo		□ Ashkenazi lewish	Caucasian Hispanic	□ Other
Address				City	State Zip	Phone
INSURANCE: Please provide a legible copy o Name of Insured	of the front and back of the patient's in Relationship to Pa		O INSURANCE: Insurance Compan	Self Pay WC,	/Auto (Date of Injury) Member/ID Number	Other Group Number
Collector Name (Print)		Date Colle	ected	Time Collected		<i>Fasting</i> □ Yes □ No
Specimen Type  OCD-100 (Buccal)			Specimen Storage	iture 🗌 Refrigerated	Specimen Shipping	ure 🔲 Cooling/Ice Pack
MOLECI LAP DIAGNOSTICS TESTING OPTIONS						

Comprehensive Cardiovascular NGS

TNNC1, TNNT2, TPM1, TRDN, TTN, TTR, TXNRD2, VCL (137 genes)

A2ML1, ABCC9, ACADVL, ACTA2, ACTC1, ACTN2, ADA2, AGL, AKAP9, ALMS1, ANK2, ANKRD1, BAG3, BRAF, CACNA1C, CACNB2, CALR3, CASQ2, CAV3, CAVIN4, CBL, CHRM2, COL3A1, CPT2, CRYAB, CSRP3, CTF1, CTN-NA3, DES, DMD, DNAJC19, DOLK, DSC2, DSG2, DSP, DTNA, ELAC2, EMD, EYA4, F9, FBN1, FHL1, FHL2, FKRP,

NAS, DES, DMD, DNAJCT9, DUCK, DSC2, DSG2, DSP, DTINA, ELAC2, EMID, ETAA, PS, FENT, FHLT, FHLZ, FKRF, FKTN, FLNC, FXN, GAA, GATA4, GATA6, GATAD1, GLA, GPD1L, HCN4, HRAS, ILK, JPH2, JUP, KCNE1, KCNE2, KCNE3, KCNH2, KCNJ2, KCNJ3, KCNJ8, KCNQ1, KRAS, LAMA4, LAMP2, LDB3, LMNA, MAPZK1, MAPZK2, MIB1, MTO1, MYBPC3, MYH11, MYH6, MYH7, MYL2, MYL3, MYLK, MYLK2, MYOM1, MYOZ2, MYPN, NDUFB11, NEBL, NEXN, NF1, NKX2-5, NPPA, NRAS, PDLIM3, PKP2, PLN, PRDM16, PRKAG2, PTPN11, RAF1, RANGRF, RSA1, RBM20, RIT1, RRAS, RYR2, SCN1B, SCN3B, SCN4B, SCN5A, SDHA, SGCD, SHOC2, SLC22A5, SLC2A10, SMAD3, SNTA1, SOS1, SPRED1, TAZ, TBX20, TCAP, TGFB81, TGFBR2, TMEM43, TMEM70, TMPO, TNN13, NNC1, TNNC1, TNNT2, TDM1, TDN1, TTN TTNIPD, VCI (127, Capac)

Cardiac Genomics Test Please select the Panel to be tested. Please attach patient Medication List.

## □ Comprehensive Cardiomyopathy NGS

ABCC9, ACADVL, ACTC1, ACTN2, ADA2, AGL, AKAP9, ALMS1, ANK2, ANKRD1, BAG3, BRAF, CACNA1C, CACNB2, CALR3, CASQ2, CAV3, CAVIN4, CBL, CHRM2, CPT2, CRYAB, CSRP3, CTF1, CTNNA3, DES, DMD, DNAIC19, DOLK, DSC2, DSC2, DSP, DTNA, ELAC2, EMD, EYA4, FHL1, FHL2, FKTN, FLNC, FXN, GAA, GATA4, GATA6, GATAD1, GLA, GPD1L, HCN4, HRAS, ILK, JPH2, JUP, KCNE1, KCNE2, KCNE3, KCNH2, KCNJ2, KCNJ5, KCNJB, KCNQT, KRAS, LADVA, LAMPZ, LDB3, LMNA, MAP2KI, MAP2K2, MB1, MCO1, MYBPG3, MYH6, MYH7, MYL2, MYLK2, MYOM1, MYOZ 2, MYPN, NDUFB11, NEBL, NEXN, NF1, NKX2-5, NPPA, NRAS, PKP2, PLN, PRDM16, PRKAG2, RAF1, RANGRF, RASA1, RBM20, RIT1, RRAS, RYR2, SCN1B, SCN3B, SCN4B, SCN5A, SDHA, SGCD, SLC 22A5, SNTA1, SPRED1, TAZ, TCAP, TGFB3, TMEM43, TMEM70, TMPO, TNNC1, TNNT2, TPM1, TRDN, TTN, TTR, TXNRD2, VCL (118 GENES)

### Comprehensive Arrthymia NGS

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В СС9, АСТС1, АСТN2, АGL, АКАР9, ALMS1, ANK2, BAG3, CACNA1C, CACNB2, CASQ2, CRYAB, CSRP3, DES, DMD, DOLK, DSC2, DSG2, DSP, EMD, EYA4, FHL1, FKRP, FKTN, FLNC, GAA, GATAD1, GLA, GPD1L, HCN4, JPH2, JUP, KCNE1, KCNE2, KCNE3, KCNH2, KCNJ2, KCNJ5, KCNQ1, LAMA4, LAMP2, LDB3, LMNA, MYBPC3, MYH6, MYH7, MYL2, MYL3, MYLK2, MYOZ2, MYPN, NEXN, NKX2-5, PKP2, PLN, PRKAG2, PTPN11, NEXPONDED (2010) (201 RAF1, RBM20, RIT1, RYR2, SCN1B, SCN3B, SCN4B, SCN5A, SGCD, SLC22A5, SNTA1, TAZ, TBX20, TGFB3, TMEM70, TNN13, TNNT2, TTR, TRDN, TTN (77GENES)

### Personal/Family History Questionnaire Please complete Questionnaire FAMILY HISTORY ATIENT'S PERSONAL HISTORY (Hx Clinical Details Personal Hx Age at Dx Relationship Maternal Paternal Cancer Site(s) Age at Dx 🗆 Yes 🗆 No Mosaicism Consanguinty 🗆 Yes 🗆 No Bone Marrow Transplant 🗆 Yes 🗆 No Organ Transplant 🗆 Yes 🗆 No Known Chromosomal Gain/Loss □ Yes □ No Known Gene Gain/Loss 🗆 Yes 🗆 No Clinical Presentation Please indicate any clinical presentations and/or Clinical Testing Please indicate any clinical testing results and/or findings that may be relevant to aenetic testina: findings that may be relevant to genetic testing: □ Karyotype□ Vision□ Previous Genetic Testing□ Hearing □ Behavior □ Conditions □ Pedigree/Family History □ Growth Measurements □ Imaging □ Phenotypes □ Physical □ Symptoms Biochemical Testing □ Pathology Results ICD-10 DIAGNOSIS CODES: Additional documentation supporting Medical Necessity may be attached. □ |42.7 Cardiomyopathy due to drug and external agent Hypertensive heart disease with heart failure STEMI) myocardial infarction involving left circumflex coronary artery □ I50.21 Acute systolic (congestive) heart failure □ I21.21 □ I50.22 Chronic systolic (congestive) heart failure □ I21.4 □ I22.2 NSTEMI) myocardial infarction □ I50.23 Acute on chronic systolic (congestive) heart failure NSTEMI) myocardial infarction □ I50.31 Acute diastolic (congestive) heart failure □ I25.110 □ I25.730 Atherosclerotic heart disease of native coronary artery with unstable angina pectoris □ I50.32 Chronic diastolic (congestive) heart failure Atherosclerosis of nonautologous biological coronary artery bypass graft(s) with I50.33 Acute on chronic diastolic (congestive) heart failure Autoroscerosis of non-decomposed biological coloniary arcely bypass grands, with unstable angina pectoris Atherosclerosis of native coronary artery of transplanted heart with unstable angina □ I50.41 Acute combined systolic (congestive) and diastolic (congestive) heart □ I25.750 failure □ I25.760 Atherosclerosis of bypass graft of coronary artery of transplanted heart with □ I50.42 Chronic combined systolic (congestive) and diastolic (congestive) heart unstable angina failure □ I25.790 Atherosclerosis of other coronary artery bypass graft(s) with unstable angina □ I50.43 Acute on chronic combined systolic (congestive) and diastolic (congestive) pectoris heart failure Dilated cardiomyopathy □ 142.0 □ R06.01 Orthopnea □ I42.1 Obstructive hypertrophic cardiomyopathy □ R06.02 Shortness of breath □ I42.2 □ I42.3 Other hypertrophic cardiomyopathy Endomyocardial (eosinophilic) disease □ R06.2 Wheezing □ R06.82 Tachypnea, not elsewhere classified Endocardial fibroelastosis 🗆 R06.89 Other abnormalities of breathing □ I42.6 Alcoholic cardiomyopathy □ Other I, the provider, attest that I am the ordering physician or am authorized under applicable laws and regulations to order genetic testing for the patient. I attest that this test is medically necessary for the diagnosis or detection of a disease or disorder, and that the results will be used in medical management and care decisions for the patient. I further attest that any Medical Necessity Required for insurance information entered on this Test Requisition Form, or otherwise provided by me on behalf of the patient, is true and correct to the best of my knowledge, and that the patient has consented to eceive communications about his/her genetic test from Mainstream Diagnostic Laboratory I, the patient, voluntarily consent to the collection and testing of my sample. I certify that the specimen is fresh and has not been adulterated in any manner. I authorize the laboratory to release the results of this testing to the ordering provider. I further authorize my insurance benefits to be paid directly to Mainstream Diagnostic Laboratory for services rendered. I acknowledge that the lab may be treated as an out-of network provider. In the event I receive payment for laboratory services from my insurer, I will remit said payment to the lab within 14 days of receipt. I will either en-dorse the original check, or produce a personal check for the entire payment amount, and forward it to the lab. When selecting Self Pay above, I acknowledge financial responsibility for all lab charges associated with the processing of this test requisition. All rights to the samples will belong to the laboratory conducting the testing. There will be no compensation in the event of an invention resulting from research and development using this sample. I agree to allow my provided samples to be used for the purpose of (diagnosis/ research) (development/quality control). I understand that if I agree, any information identifying me will be kept confidential so that it will not be possible to determine from whom the sample was drawn. Your signature on this form indicates that you understand to your satisfaction the information about Mainstream Diagnostic Laboratory and agree to have the test done. In no way does this waive your legal rights or release anyone from their legal and professional responsibilities. If you have further questions concerning matters related to this consent, you may wish to seek professional genetic counseling prior to signing this form. Consultation with a medical genetics, genetic counselor, or your referring healthcare provider also may be warranted after the test has been completed. Patient Informed Consent Patient must consent test has been completed. I give permission for my sample and clinical information to be used in de-identified studies at Mainstream Diagnostic Laboratory and for publication, if Mainstream Diagnostic Laboratory deems it appropriate. I understand that my name and/or other identifying information will NOT be used in or linked to the results of any studies and publications. More information is available at www.mainstreamlab.com. Opt In for Research Clinic Phone/Fax Provider Name (Print) Provider NPI # Clinic Address Provider Signature Date Date Patient Signature (or Legal Guardian)

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