OMNIHEALTH DIAGNOSTICS, LLC CLIA: 45D2089485 Lab Director: Akhar Afshan Ali Address: 1840 N Greenville Ave, Suite 176 Richardson, TX 75081 Phone: 972.887.3444 | Fax: 972.887.3443 PLEASE SUBMIT THE FOLLOWING WITH REQUISITION FORM

□ Statement of Medical Necessity (Signed by Physician)

□ Informed Consent Form (Signed by Pt & Physician)

□ SOAP & Progress Note (Signed by Physician)

CARDIO-PULMONARY TESTING REQUISITION FORM								
PATIENT INFORMATION								
Patient First Name		Patier	nt Last Name		Biological Sex 🔲 F 🗌 M			
Date of Birth (MM/DD/YYYY)	Phone Numb			l Address				
Address			City		State	Zip		
Ethnicity: African American Asian Caucasian Hispanic Jewish(Ashkenazi) Portuguese Other								
PATIENT INSU	RANCE INFORMATION	1		SPECI	MEN INFO	RMATION		
🗌 Insurance 🗌 Self-Pay 🗌	Client Bill		Date Sample	Date Sample Collected (mm/dd/yy) (required)				
Name of the insurance	Secondary Insurance, I	Medical Reco	ord#					
Insurance Policy/ID number	Name of the insured			Buccal Swab Other (specify source)				
Insurance Group number	Date of Birth of Insured	d Other (specify source)						
ORDERING PHYSICIAN/SENDING FACILITY (Each Listed person will receive a copy of the report)								
Facility Name (Facility Code): Addre			255:	is: City:				
State/Country : 2					Phone:	hone:		
Ordering Licensed Provider Name (Last, First)(Code)		NPI#		Phone		Fax/Email		
Additional Results Recipients								
Genetic Counselor or Other Medical Provider Name (Last, First)(Code)				Phone/Fax/Email				
Signature Required for Processing Medical Professional Signature:       Date :								
STATEMENT OF MEDICAL NECESSITY								
By submission of this test requisition and accompanying sample(s), I: (i) authorize and direct to perform the testing indicated; (ii) certify that the person listed as the ordering provider is authorized by law to order the test(s) requested; (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 and relatives', when applicable, 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:				Date:				
INDICATIONS FOR TESTING (CHECK ALL THAT APPLY)								
Diagnostic 🗌 Family history 🗌 Positive or normal control 🔲 Other								

Will Patient management be changed depending on the test results?

CLINICAL HISTORY (PLEASE SUPPLY CLINIC NOTES AND PEDIGREE)					
		Types (s) of Arrhythmia:			
□ No personal history of Cardiopulmonary disease		☐ Family or personal history of chronic bronchitis? ☐Yes ☐No ☐ Pulmonary edema? ☐Yes ☐No			
Sudden Lungs Failur 🔲 Y 🗌 N (if yes): # Episodes: Age first incident:		Family history of heart failure? Yes No			
History of Cardiopulmonary $\Box$ Y $\Box$ N Age at dx:		Family or personal history of a COPD? Yes No Cardiac arrhythmias? Yes No			
Type(s) of Cardiopulmnary:		☐ History of right-sided heart failure? ☐ Yes ☐ No ☐ Collection of fluid in legs or belly area? ☐ Yes ☐ No			
		Diagnosed with Emphysema? 🗌 Yes 🗌 No			
History of Arrhythmia 🛛 Y 🗋 N		Family history show the segregation of pulmonary emphysema? Yes No Diagnosed with alpha 1-antitrypsin deficiency-related pulmonary emphysema or early			
Age at dx:		onset pulmonary emphysema?  Yes No			
		Diagnosed with cor pulmonale? Yes No			
CLINICAL	INFORMATION (DETAILEI	D MEDICAL RECORDS MUST BE ATTACHED)			
Reason for testing: O Diagnosis O Presymptomatic diagr		esting			
Please check all that apply. This is not a substitute for submi	ting clinical records.				
Diagnosis	Marfan/TAAD/HDCT	Abnormal heart morphology			
	Aortic/Arterial aneurysm     Aertic/Arterial dispection	<ul> <li>Bicuspid aortic valve</li> <li>Coarctation of aorta</li> </ul>			
O ARVC O Brugada syndrome	<ul> <li>Aortic/Arterial dissection</li> <li>Aortic root dilation</li> </ul>	$\bigcirc$ Heart murmur			
○ Brugada syndrome ○ CPVT	Aortic root dilation     Arachnodactyly	⊖ Heterotaxy			
ODCM	<ul> <li>Arterial tortuosity/ectasia</li> </ul>	O Hypoplastic left heart			
O Ehlers-Danlos syndrome	$\bigcirc$ Arthralgia	$\bigcirc$ Mitral valve prolapse			
O HCM	<ul> <li>Atypical scarring of skin</li> </ul>	<ul> <li>Patent ductus arteriosus</li> </ul>			
O HHT	<ul> <li>Beighton score</li> </ul>	O Patent foramen ovale			
○ Hypertension	O Bifid uvula	<ul> <li>O Tetralogy of Fallot</li> <li>O Ventricular septal defect</li> </ul>			
○ Loeys-Dietz syndrome	O Blue sclerae	<ul> <li>Atrial septal defect</li> </ul>			
○ LQT syndrome	O Bruising susceptibility	O Other:			
<ul> <li>Noncompaction Cardiopulmonary (LVNC)</li> </ul>	<ul> <li>Cleft lip</li> </ul>	РАН			
O Marfan syndrome	O Cleft palate	<ul> <li>Pulmonary hypertension</li> </ul>			
O PAH	○ Craniosynostosis	Cardiopulmonary			
	○ Cutis laxa	O Chronic bronchitis			
<ul> <li>○ SQT syndrome</li> <li>○ Sudden Cardiac Arrest</li> </ul>	<ul> <li>Dental crowding</li> </ul>	O Chronic obstructive pulmonary disease (COPD)			
$\bigcirc$ Sudden Death	○ Dural ectasia	Congestive heart failure			
Echocardiogram	<ul> <li>Ectopia lentis</li> </ul>	O Emphysema			
O Aortic root dimensions:	<ul> <li>Flexion contracture</li> </ul>	Other O Abnormality of the periventricular white matter			
O Z-score:	○ High palate	$\bigcirc$ Angiokeratomas			
○ EF%:	○ Hollow organ rupture:	○ Anhydrosis			
O LVEDD:	•	ntestinal perforation O Café-Au-Lait Macules			
O Z-score:		O Hearing impairment:			
O Max LV wall thickness:	○ Hypertelorism	○ Sensorineural ○ Conductive			
○ Normal	<ul> <li>Joint contractures</li> </ul>	<ul> <li>Craniosynostosis</li> <li>Curtis transmission</li> </ul>			
O Report Included	<ul> <li>Joint dislocations</li> </ul>	<ul> <li>Cystic hygroma</li> <li>Downslanted palpebral fissures</li> </ul>			
ECG	<ul> <li>Joint hypermobility</li> <li>Masta Object evitation</li> </ul>	<ul> <li>Downstanted palpebra insores</li> <li>Dysmorphic features:</li> </ul>			
O Prolonged QTc interval: Max QTc:	<ul> <li>Meets Ghent criteria</li> <li>Micrographia / Betrographia</li> </ul>				
Nax Grc:	<ul> <li>Micrognathia / Retrognathia</li> <li>Midface retrusion</li> </ul>	© Elevated CPK			
O Report Included	<ul> <li>Mitral valve prolapse</li> </ul>	O Hypotonia			
Arrhythmia/Cardiopulmonary	<ul> <li>Mitrai valve prolapse</li> <li>Myopia</li> </ul>	<ul> <li>Increase nuchal translucency</li> </ul>			
O Abnormal atrioventricular conduction	$\bigcirc$ Osteoarthritis	O Intellectual disability			
○ Atrial fibrillation	<ul> <li>O Pectus carinatum</li> </ul>	○ Keratoconus			
⊖ Bradycardia	<ul> <li>Pectus excavatum</li> </ul>	Muscle weakness     Muscathy			
○ Fatty replacement of ventricular myocardial tissue	O Pes Planus	<ul> <li>Myopathy</li> <li>Renal insufficiency</li> </ul>			
O Heart transplant	○ Pneumothorax	O Short neck			
○ Syncope	O Recurrent fractures	O Thromboembolism			
O Torsades de pointe	Retinal detachment     Secliaria (Kurbasia (airela un)	O Type:			
O Ventricular tachycardia	<ul> <li>Scoliosis/Kyphosis (circle w</li> <li>Skin findinge, Specify</li> </ul>	vnat applies) :			
	<ul> <li>Skin findings, Specify:</li> <li>Stroke</li> </ul>				
<ul> <li>Arteriovenous malformation</li> <li>Epistaxis</li> <li>Tall stature</li> </ul>					
O Telangiectasia	O Velvety skin				
Dislipidemias	2 -				
○ Atherosclerosis					
○ Corneal Arcus					
○ LDL-C levels					
○ Xanthomatosis					

○ Other:\_

<ul> <li>ACTR2</li> <li>DSC</li> <li>PRN</li> <li>DSC</li> <li>PRN</li> <li>COLDA</li> <li>DSC</li> <li>PRN</li> <l< th=""><th colspan="5">Custom Cardio-Pulmonary NGS Testing (Select the genes below) or 📃 Comprehensive Cardio-Pulmonary NGS Testing Panel (Test All Genes)</th></l<></ul>	Custom Cardio-Pulmonary NGS Testing (Select the genes below) or 📃 Comprehensive Cardio-Pulmonary NGS Testing Panel (Test All Genes)								
AAC12         BAC         COL11AL         DOLX         PR8P         AMP2         MYH6         PLN         SCNB         OS1         TRPMA           AAC1C1         BRAF         COL11AL         DSC2         PKIN         LDRA         MYH2         PR0D1         SCNB         TAX         TWN           ACTNL         BRAFR         COL11AL         DSC2         PF         NHA         LDRA         MYH2         PPA2         SCNB         TAX         TWN           ACMNL         BRAFR         COL11AL         DSC2         PF         LDRA         MYH3         PPA2         SCNB         TRX2         TWN           ADMTS2         CAXNAC         COL12AL         DTW         GAA         LDX         MYH3         PPR01         SCNB         TGF83           ADMTS2         CAXNAD1         COL3AL         EFEMP2         CATAA         LBRC10         MYHX         PPRK30         SCNBA         TGF83           ALMK3         CAND2         COL3AL         EFEMP2         CATAA         LBRC10         MYHX         PPRK30         SCNAA         TGF83           ALMK3         CAMB01         CALM3         CAMAA         PARC2         SCMA1         SCANAA         TGF83		CardioGe	enomics Gen	nes					
Pulmonary Genes         CCDC39       COLQ       DNAH1       ELMOD2       ITGA3       PHOX2B       RTEL1       SFTPB       TERC       FBLNS         CCDC40       CSF2RA       DNAH1       FLCN       MECP2       PIH1D3       SCNN4       SFTPC       TERT       ELN         CFTR       CSF2RA       DNAH1       GASS       NF1       NAF1       RAPSN       SCNN14       SICA42       TINF2       LTBP4         CHAT       DKC1       DNAI1       GASS       NF1       NRE       SCNN15       SICA5A2       TSC1         CHRNA1       OHAF1       DNAI1       GASS       NK2-1       RSPH3       SERPINA1       SICA5A2       TSC1         CHRNB1       DNAAF2       EDN3       HP54       PARN       RSPH4A       SFTPA1       SMPD1       ZEB2         CHRNE       DNAAF2       EDN3       HP54       PARN       RSPH4A       SFTPA2       STAT3       EFEMP2         E78.4 - Other Hyperlipidemia, unspecified       I35.9       Nontheumatic aortic valve disorder, unspecified       R60.2       -Fontenss of breath         IB37.1 - Hyperispidemia, unspecified       I42.0       -Other specified cardioxacular       R66.02       -Shontenss of dyspnea         IB35.9	ACTA2       BAG3       COL11A1       DOLK         ACTC1       BGN       COL11A2       DSC2         ACTN2       BMPR2       COL12A1       DSE         ACVRL1       BRAF       COL1A1       DSG2         ADAMTS2       CACNA1C       COL1A2       DSP         AKAP9       CACNA2D1       COL3A1       EFEM         ALDH18A1       B3GALT6       COL5A1       EIF2A         ALMS1       CACNB2       COL5A1       EIF2A         ANK2       CALM1       COL9A1       EMD         ANK2       CALM3       COL9A1       END         APOB       CASQ2       COL9A3       EYA4         ATP6V0A2       CAV1       CRYAB       FBLN.	K <ul> <li>FKRP</li> <li>FKTN</li> <li>FLNA</li> <li>FINC</li> <li>GAA</li> <li>GATA4</li> <li>GATA5</li> <li>GATA6</li> <li>GATA1</li> <li>GDF2</li> <li>GJA5</li> <li>KCNJ8</li> <li>KCNK3</li> </ul>	<ul> <li>LAMP2</li> <li>LDB3</li> <li>LDLR</li> <li>LDLRAP1</li> <li>LMNA</li> <li>LOX</li> <li>LRRC10</li> <li>LTBP4</li> <li>MAP2K1</li> <li>MAP2K2</li> <li>MAT2A</li> <li>MED12</li> <li>MIB1</li> </ul>	<ul> <li>MYH6</li> <li>MYH7</li> <li>MYL2</li> <li>MYL3</li> <li>MYL4</li> <li>MYLK</li> <li>MYLK2</li> <li>MYOZ2</li> <li>MYOZ2</li> <li>MYPN</li> <li>NEBL</li> <li>NEXN</li> <li>NKX2-5</li> <li>NOTCH1</li> <li>NRAS</li> </ul>	<ul> <li>PLN</li> <li>PLOD</li> <li>PPA2</li> <li>PRDM</li> <li>PRKAI</li> <li>PRKG</li> <li>PTPN</li> <li>PYCR</li> <li>RAF1</li> <li>RANG</li> <li>RASA</li> <li>RBM2</li> <li>RIN2</li> </ul>	SCN1B           SCN2B           SCN3B           SCN4B           SCN3B           SCN4B           SCN4B           SCN4B           SCN4B           SCN4D           SMAD3	<ul> <li>SOS1</li> <li>TAZ</li> <li>TBX20</li> <li>TCAP</li> <li>TECRL</li> <li>TGFB3</li> <li>TGFB2</li> <li>TGFBR2</li> <li>TGFBR2</li> <li>TGFBR2</li> <li>TGFBR2</li> <li>TMP0</li> <li>VTNNC1</li> <li>TNNT2</li> <li>TNXB</li> </ul>	TRPM4 TTN TXNRD2	
CCDC39       COLQ       DNAH1       ELMOD2       ITGA3       PHOX2B       RTEL1       SFTPB       TERC       FBLNS         CCDC40       CSF2RA       DNAH1       FLCN       MECP2       PHIND3       SCNA4       SFTPC       TERC       FBLNS         CFR       CSF2RB       DNAH1       FOXF1       NAF1       RAPSN       SCNA4       SFTPC       TERC       TERC       FBLNS         CHRNA1       CHRND       DNAL2       GRA1       NKX2-1       RSPH3       SCNN1A       SLC7A7       TSC2         CHRNB1       DNAAF2       EDN3       HP54       PARN       RSPH3       SFTPA2       STAT3       EFEMP2         CardioGenomics Disease         CHNE       DNAAF2       EDN3       HP54       PARN       RSPH9       SFTPA2       STAT3       EFEMP2         CardioGenomics Disease         E37.1 +Nperlipidemia       135.9 - Nontheumatic aortic valve disorder, unspecified       R06.09 - Other forms of dyspnea       R06.02 - Shortness of breath         B92.9 - Other Chronic Pain       142.0 - dilated Cardiovascular       R06.09 - Other forms of dyspnea       R06.3 - Periodic breathing         125.5 - Sthemic Cardiovascular       143.9 - Nonrheumatic darial fbrillation       R06.3 - Periodic breathing	B3GAT3 CHRM2 DES FHL1				C RYR2	SMAD9	TPM1		
CCDC40       CSF2RA       DNAH11       FLCN       MECP2       PIH1D3       SCNA4       SFTPC       TERT       ELN         CFTR       CSF2RA       DNAH5       FOXF1       NAF1       RAPSN       SCNN1A       SLC6A5       TSC1       LTBP4         CHAN1       CHRND       DNAL1       GAS8       NF1       RFT       SCNN1B       SLC6A7       TSC2         CHRNA1       CHRND       DNAL1       HPS1       NK2-1       RSPH3       SERPINA1       SLC7A7       TSC2         CHRNE       DNAAF1       DNAL1       HPS1       NK88       RSPH4A       SFTPA1       SMPD1       ZEB2         CHRNE       DNAAF2       EDN3       HPS4       PARN       RSPH3       SFTPA2       STAT3       EFEM22         E78.4 - Other Hyperlipidemia       IS5.9 - Nonrheumatic aortic valve disorder, unspecified       R60.9 - Edema, unspecified       R06.00       R06.00       SF002       Shortness of breath         G58.29 - Other Chronic Pain       I35.9 - Nonrheumatic aortic valve disorder, unspecified       R06.00       S60.00       S60.00       Spenea, unspecified         I10 - Essential (Primary) Hypertension       I35.9 - Nonrheumatic aortic valve disorder, unspecified       R06.00       S60.00       S60.00       Spenea, unspecified		Pulmo	onary Genes						
CardioGenomics Disease         E78.4 - Other Hyperlipidemia       I35.9 - Nonrheumatic aortic valve disorder, unspecified       R60.9 - Edema, unspecified         E78.5 - Hyperlipidemia, unspecified       I42.0 - dilated Cardiovascular       R06.02 - Palpitations         E87.1 - Hypo - osmolality and / or hypernatremia       I42.5 - Other restrictive Cardiovascular       R06.02 - Shortness of breath         G89.29 - Other Chronic Pain       I42.9 - Supraventricular tachycardia       R06.00 - Other forms of dyspnea         I10 - Essential (Primary) Hypertension       I42.9 - Junctional premature depolarization       R06.09 - Other forms of dyspnea         coronary artery without angina pectoris       I48.2 - Chronic atrial fibrillation       R06.3 - Periodic breathing         I25.5 - Ischemic Cardiovascular       I49.9 - Unspecified atrial fibrillation       R06.83 - Snoring         I25.5 - Schemic Cardiovascular       I49.9 - Unspecified atrial fibrillation       R06.89 - Other apnormalities of breathing         I25.5 - Schemic Cardiovascular       I49.9 - Unspecified atrial fibrillation       R06.89 - Other apnormalities of breathing         I25.5 - Schemic Cardiovascular       I49.8 - Other specified atrial fibrillation       R06.89 - Other apnormalities of breathing         I25.9 - Other forms of chronic ischemic heart disease       R00.1 - Bradycardia, unspecified       R07.2 - Precordial pain         I35.9 - Chonnic ischemic heart disease       I50.21 - A	CCDC40       CSF2RA       DNAH1I       F         CFTR       CSF2RB       DNAH5       F         CHAT       DKC1       DNAI1       C         CHRNA1       CHRND       DNAI2       C         CHRNB1       DNAAF1       DNAL1       F	ELCN         Image: Meconstruction           FOXF1         NAF*           GAS8         NF1           GLRA1         NKX:           HPS1         NME	P2 PI 1 R/ 2-1 R5 38 R5	H1D3 APSN ET 5PH3 5PH4A	<ul> <li>SCNA4</li> <li>SCNN1A</li> <li>SCNN1B</li> <li>SERPINA1</li> <li>SFTPA1</li> </ul>	<ul> <li>SFTPC</li> <li>SLC34A2</li> <li>SLC6A5</li> <li>SLC7A7</li> <li>SMPD1</li> </ul>	<ul> <li>TERT</li> <li>TINF2</li> <li>TSC1</li> <li>TSC2</li> <li>ZEB2</li> </ul>	🗆 ELN	
E78.4 - Other Hyperlipidemia       I35.9 - Nonrheumatic aortic valve disorder, unspecified       R60.9 - Edema, unspecified         E78.5 - Hyperlipidemia, unspecified       I42.0 - dilated Cardiovascular       R06.02 - Palpitations         E87.1 - Hypo - osmolality and / or hypernatremia       I42.5 - Other restrictive Cardiovascular       R06.02 - Shortness of breath         G89.29 - Other Chronic Pain       I42.9 - Supraventricular tachycardia       R06.00 - Dyspnea, unspecified         I10 - Essential (Primary) Hypertension       I48.0 - Paroxysmal atrial fibrillation       R06.3 - Periodic breathing         coronary artery without angina pectoris       I48.2 - Chronic atrial fibrillation       R06.89 - Other abnormalities of breathing         I25.5 - Ischemic Cardiovascular       I49.91 - Unspecified atrial fibrillation       R06.89 - Other abnormalities of breathing         I25.9 - Other forms of chronic ischemic heart disease       R00.1 - Bradycardia, unspecified       R07.2 - Precordial pain         I34.1 - Nonrheumatic mitral (valve) insufficiency       I50.21 - Acute systolic (congestive) heart failure       R07.89 - Other chest pain         I34.2 - Nonrheumatic mitral valve disorder, unspecified       I50.32 - Chronic diastolic (congestive) heart failure       R07.43 - Nonspecific datoradises, unspecified         I34.2 - Nonrheumatic mitral valve disorder, unspecified       I50.32 - Chronic diastolic (congestive) heart failure       R07.43 - Nonspecific abnormal electrocardiogram (ECG)(EKG)	ICD-10 DIAGNOSIS CODES WITH DESCRIPTION								
<ul> <li>E78.5 - Hyperlipidemia, unspecified</li> <li>E78.5 - Hyperlipidemia, unspecified</li> <li>I42.0 - dilated Cardiovascular</li> <li>R00.2 - Palpitations</li> <li>R06.02 - Shortness of breath</li> <li>R06.02 - Shortness of breath</li> <li>R06.00 - Dyspnea, unspecified</li> <li>I10 - Essential (Primary) Hypertension</li> <li>I42.9 - Supraventricular tachycardia</li> <li>R06.09 - Other forms of dyspnea</li> <li>I25.10 - Atherosclerotic heart disease of native coronary artery without angina pectoris</li> <li>I48.2 - Chronic atrial fibrillation</li> <li>R06.3 - Periodic breathing</li> <li>I25.5 - Ischemic Cardiovascular</li> <li>I49.2 - Junctional premature depolarization</li> <li>R06.3 - Periodic breathing</li> <li>I25.6 - Silent Myocardial Ischemia</li> <li>I49.9 - Unspecified atrial fibrillation</li> <li>R06.89 - Other abnormalities of breathing</li> <li>I25.9 - Chronic ischemic heart disease</li> <li>R00.1 - Bradycardia, unspecified</li> <li>I35.9 - Other nonreheumatic mitral (valve) insufficiency</li> <li>I34.2 - Nonrheumatic mitral (valve) stenosis</li> <li>I35.8 - Other nonreheumatic antiral valve disorder, unspecified</li> <li>I35.1 - Nonrheumatic antiral valve disorder, unspecified</li> <li>I35.2 - Nonrheumatic aortic (Valve) stenosis</li> <li>I35.2 - Other heumatic aortic (Valve) stenosis</li> <li>I52.3 - Other heumatic aortic (Valve) stenosis</li> <li>I52.5 - Other heart disease classified elsewhere</li> <li>I35.1 - Nonrheumatic aortic (Valve) stenosis with</li> <li>R60.0 - Localized edema</li> <li>Z01.812 - Encounter for preprocedural</li> </ul>		CardioGer	nomics Dise	ease					
□ I35.8 - Other Nonrheumatic aortic (valve) disorders □ R60.1 - Generalized edema □ Z01.818 - Encounter for other preprocedural examination	<ul> <li>E78.5 - Hyperlipidemia, unspecified</li> <li>E87.1 - Hypo - osmolality and / or hypernatremia</li> <li>G89.29 - Other Chronic Pain</li> <li>I10 - Essential (Primary) Hypertension</li> <li>I25.10 - Atherosclerotic heart disease of native coronary artery without angina pectoris</li> <li>I25.5 - Ischemic Cardiovascular</li> <li>I25.6 - Silent Myocardial Ischemia</li> <li>I25.9 - Other forms of chronic ischemic heart disease</li> <li>I25.9 - Other forms of chronic ischemic heart disease</li> <li>I25.9 - Chronic ischemic heart disease, unspecified</li> <li>I34.1 - Nonrheumatic mitral (valve) prolapse</li> <li>I34.2 - Nonrheumatic mitral (valve) prolapse</li> <li>I35.8 - Other nonreheumatic mitral valve disorders</li> <li>I35.0 - Nonrheumatic aortic (Valve) Insufficiency</li> <li>I35.1 - Nonrheumatic aortic (Valve) Insufficiency</li> <li>I35.2 - Nonrheumatic aortic (valve) stenosis with insufficiency</li> <li>I35.8 - Other Nonrheumatic aortic (valve) disorders</li> </ul>			rdiovascular rictive Cardiovascular ricular tachycardia I premature depolarization al atrial fibrillation rial fibrillation ed atrial fibrillation cified cardiac arrhythmias dia, unspecified tolic (congestive) heart failure ystolic(congestive) heart failure liastolic (congestive) heart failure chronic diastolic (congestive) heart failure chronic diastolic (congestive) heart failure ase, unspecified t diseases classified elsewhere and Collapse edema hypercholesterolemia ed edema			<ul> <li>R00.2 - Palpitations</li> <li>R06.02 - Shortness of breath</li> <li>R06.00 - Dyspnea, unspecified</li> <li>R06.09 - Other forms of dyspnea</li> <li>R06.3 - Periodic breathing</li> <li>R06.83 - Snoring</li> <li>R06.89 - Other abnormalities of breathing</li> <li>R07.9 - Chest pain, unspecified</li> <li>R07.2 - Precordial pain</li> <li>R07.89 - Other chest pain</li> <li>R07.89 - Other chest pain</li> <li>R07.89 - Other chest pain</li> <li>R07.81 - Nonspecific abnormal electrocardiogram (ECG)(EKG)</li> <li>Z79.01 - Long term (current) use of anticoagulants</li> <li>Z01.810 - Encounter for preprocedural cardiovascular examination</li> <li>Z01.812 - Encounter for preprocedural laboratory examination</li> <li>Z01.818 - Encounter for other</li> </ul>		
Pulmonary Disease									
C34.12-Malignant Neoplasm of upper lobe, left bronchus or lungJ20.6-acute bronchitis due to rhinovirusC34.2-Malignant Neoplasm of Middle lobe, bronchus or lungJ20.7-Acute bronchitis due to ochovirusC34.31-Malignant Neoplasm of lower lobe, left bronchus or lungJ20.8-Acute bronchitis due to other specified organismsC34.32-Malignant Neoplasm of lower lobe, left bronchus or lungJ20.9-Acute bronchitis, unspecifiedE84.0-Cystic Fibrosis with pulmonary manifestationsJ16.8-Pneumonia due to other specified infectious organismsG47.33-Obstructive sleep apneaJ18.9-Pneumonia, unspecified organismJ27.0-Primary Pulmonary HypertensionJ44.1-Obstructive chronic bronchitis, with (acute) exacerbationJ44.1-Chronic Obstructive Pulmonary Disease with acute exacerbationJ44.1-Obstructive chronic bronchitis, with (acute) exacerbationJ44.1-Chronic Obstructive Pulmonary Disease NOSJ45.20-Mild Intermittent AsthmaJ20.3-Acute bronchitis due to Mycoplasma pneumoniaeJ45.23-Mild Intermittent Asthma with acute exacerbationJ20.4-Acute bronchitis due to ParainfluenzaJ45.40-Moderate persistent Asthma with acute exacerbationJ20.5-Acute bronchitis due to Parainfluenza virusJ45.23-Mild Persistent Asthma with acute exacerbationJ20.6-Acute bronchitis due to other specified organismsJ45.23-Mild Persistent Asthma with acute exacerbationJ20.6-Acute bronchitis due to other specified organismsJ45.23-Mild Persistent Asthma with acute exacerbationJ20.6-Acute bronchitis due to respiratory syncytial virusJ45.23-Mild Persistent AsthmaJ20.6-Acute bronchitis due to other specified organismsJ45.23-Mild Persistent AsthmaJ20.6-Ac	<ul> <li>C34.2-Malignant Neoplasm of Middle lobe, bronchus or lung</li> <li>C34.31-Malignant Neoplasm of lower lobe, right bronchus or lung</li> <li>C34.32-Malignant Neoplasm of lower lobe, left bronchus or lung</li> <li>E84.0-Cystic Fibrosis with pulmonary manifestations</li> <li>G47.33-Obstructive sleep apnea</li> <li>I27.0-Primary Pulmonary Hypertension</li> <li>J44.1-Chronic Obstructive Pulmonary Disease with acute exacerbation</li> <li>J44.1- Chronic Obstructive Pulmonary disease NOS</li> <li>J20.0- Acute bronchitis due to Mycoplasma pneumoniae</li> <li>J20.1-Acute bronchitis due to coxsackievirus</li> <li>J20.3-Acute bronchitis due to respiratory syncytial virus</li> <li>J20.5-Acute bronchitis due to rhinovirus</li> <li>J20.7-Acute bronchitis due to other specified organisms</li> <li>J20.9-Acute bronchitis due to other specified organisms</li> <li>J20.9-Acute pulmonary Edema</li> </ul>			<ul> <li>J20.7-Acute bronchitis due to echovirus</li> <li>J20.8-Acute bronchitis due to other specified organisms</li> <li>J20.9-Acute bronchitis, unspecified</li> <li>J16.8-Pneumonia due to other specified infectious organisms</li> <li>J18.9-Pneumonia, unspecified organism</li> <li>J40-Bronchitis, not specified as acute or chronic</li> <li>J44.1-Obstructive chronic bronchitis, with (acute) exacerbation</li> <li>J45.20-Mild Intermittent Asthma</li> <li>J45.23-Mild Intermittent Asthma with status asthmaticus</li> <li>J45.40-Moderate persistent Asthma</li> <li>J45.42-Moderate persistent Asthma with acute exacerbation</li> <li>J45.21-Mild Intermittent Asthma with acute exacerbation</li> <li>J45.21-Mild Intermittent Asthma with acute exacerbation</li> <li>J45.23-Mild Intermittent Asthma with acute exacerbation</li> <li>J45.42-Moderate persistent Asthma with acute exacerbation</li> <li>J45.21-Mild Intermittent Asthma with acute exacerbation</li> <li>J45.23-Mild Persistent Asthma with acute exacerbation</li> <li>J45.23-Mild Persistent Asthma with acute exacerbation</li> <li>J45.24-Moderate persistent Asthma with acute exacerbation</li> <li>J45.32-Mild Persistent Asthma with acute exacerbation</li> <li>J45.32-Mild Persistent Asthma with acute exacerbation</li> <li>J45.32-Mild Persistent Asthma with acute exacerbation</li> <li>J45.32-Severe persistent Asthma with acute exacerbation</li> <li>J45.30-Severe persistent Asthma with acute exacerbation</li> </ul>					

R07.81-Pleurodynia	J90-Pleural effusion, not elsewhere classified
□ J45.20 Mild Intermittent Asthma	J98.11-Atelectasis
J45.23-Mild Intermittent Asthma with status asthmaticus	J98.19-Other pulmonary collapse
J45.31-Mild Persistent Asthma with acute exacerbation	J98.2-Interstitial emphysema
J45.40-Moderate persistent Asthma	□ J81.0-Acute pulmonary edema
J45.42-Moderate persistent Asthma with status asthmaticus	J95.84-Transfusion related acute lung injury (TRALI)
J45.21-Mild Intermittent Asthma with acute exacerbation	<b>J96.00</b> -Acute respiratory failure, unspecified whether with hypoxia or hypercapnia
J45.30-Mild Persistent Asthma	□ J96.0-Acute respiratory failure
J45.32-Mild Persistent Asthma with status asthmaticus	□ J96.02-Acute respiratory failure with hypercapnia
J45.41-Moderate persistent Asthma with acute exacerbation	J98.4-Other disorders of lung
J45.52-Servere persistent Asthma with status asthmaticus	J96.10- Chronic respiratory failure, unspecified whether with hypoxia or hypercapnia
J45.50-Servere persistent Asthma	□ J96.11- Chronic respiratory failure with hypoxia
J45.51-Servere persistent Asthma with acute exacerbation	□ J96.12-Chronic respiratory failure with hypercapnia
R22.2-Localized swelling, mass and lump, trunk	<b>J96.20</b> - Acute/Chronic respiratory failure, unspecified whether with hypoxia or hypercapnia
R09.02 Hypoxemia	□ J96.21-Acute/Chronic respiratory failure with hypoxia
R91.8-Nonspecific abnormal finding of lung field in diagnostic imaging	J96.22-Acute/Chronic respiratory failure with hypercapnia
R94.2-Abnormal results of pulmonary function studies	J98.4-Other disorders of lung
A41.9-Sepsis, unspecified organism Malignant neoplasm of trachea, bronchus, lung	N17.9-Acute kidney failure, unspecified
C33-Trachea	R06.02-Shortness of breath
<b>C34.00</b> -Unspecified main bronchus	□ R06.2-Wheezing
<b>C34.10</b> -Upper lobe unspecified bronchus or lung	<b>R09.89-</b> Other specified symptoms and signs involving the circulatory and respiratory systems
<b>C34.2</b> -Middle lobe bronchus or lung	□ R05-Cough
<b>C34.30</b> -Lower lobe bronchus or lung	R07.1-Chest pain on breathing
<b>C34.80-</b> Overlapping sites of unspecified bronchus or lung	R07.81-Pleurodynia
<b>E84.0-</b> Cystic fibrosis with pulmonary manifestation	R22.2-Localized swelling, mass and lump, trunk (chest mass)(localized swelling of chest)
<b>G47.33-</b> Obstructive sleep apnea (adult) (pediatric)	R91.8-Other nonspecific abnormal finding of lung field(lung mass)
126.99-Other pulmonary embolism without acute corpulmonale	R91.1-Solitary pulmonary nodule
127.0-Primary pulmonary hypertension	R91.8-Other nonspecific abnormal finding of lung field
□ <b>195.9</b> -Hypotension, unspecified	R94.2-Abnormal results of pulmonary function studies
<b>J20.0-</b> Acute bronchitis due to Mycoplasma pneumoniae	R09.02-Hypoxemia
<b>J20.0-</b> Acute bronchitis due to Mycoplasma pneumoniae	J98.4-Other disorders of lung
<b>J20.1</b> -Acute bronchitis due to Hemophilius influenzae	□ <b>R65.20</b> -Severe sepsis without septic shock (sequence the underlying infection first)
<b>J20.2</b> -Acute bronchitis due to streptococcus	Z85.118-Personal history of malignant neoplasm of bronchus and lung
<b>J20.3</b> -Acute bronchitis due to coxsackievirus	<b>Z79.01-</b> Long-term (current) use of anticoagulants
J20.4-Acute bronchitis due to parainfluenza virus	

#### Additional ICD10 codes:

# **INFORMED CONSENT**

For the purposes of this consent, "I", "my", and "your" will refer to me or to my child, including my unborn child, if my child is the person for whom the healthcare provider has ordered testing.

**PURPOSE OF THIS TEST** - The purpose of this test is (a) to see if I may have a genetic variant or chromosome rearrangement causing a genetic disorder; or (b) to evaluate the chance that I will develop or passon a genetic disorder in the future. If I already know the specific gene variant(s) or chromosome rearrangement that causes the genetic disorder in my family, I agree to inform the laboratory of this information.

#### WHAT TYPE OF TEST RESULTS CAN I EXPECT FROM GENETIC TESTING?

1. Positive: A change in your DNA was found, which is very likely the cause of your features/symptoms. This is the most straightforward test result, which can be used as the basis to test other family members to determine their chances of having either the disease or a child with the disease.

2. Negative: No variants were found to explain your symptoms. This does not mean that you do not have a genetic condition. It is still possible that there is a genetic variant not found by the test that was ordered. Your healthcare provider or genetic counselor may discuss more testing either now or in the future.

3. Variant of Uncertain Significance (VUS): A change in a gene was found. However, we are not sure whether this variant is the cause of your symptoms/features. More information is needed. We may suggest testing other family members to help figure out the meaning of the test result.

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 find you are at risk for another genetic condition I am not aware of or it may indicate differences in the number or rearrangement of sex chromosomes. We may disclose this information to the ordering healthcare provider if it likely affects medical care. Because medical and scientific knowledge is constantly changing, new information that becomes available may supplement the information **OmniHealth Diagnostics, LLC** used to interpret my results.

Healthcare providers can contact OmniHealth Diagnostics, LLC at any time to discuss the classification of an identified variant.

#### WHAT IS TRIO/DUO-BASED GENETIC TESTING?

For some genetic tests, including samples from the biological parents and/or other biological relatives along with the patient's sample can help with the interpretation of the test results. These tests are often referred to as "trio tests" since they typically include samples from the patient and both parents. Samples from relatives should be submitted with the patient's sample.

I understand that **OmniHealth Diagnostics, LLC** will use the relative sample(s) when needed for the interpretation of my test results and that my test report may include clinical and genetic information about relative when it is relevant to the interpretation of the test results. I further understand that relatives will not receive an independent analysis of data nor a separate report.

## RISKS AND LIMITATIONS OF GENETIC TESTING

1. 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.

2. Accurate interpretation of test results may require knowing the true biological relationships in a family. I understand that if I fail to accurately state the biological relationships in my family, it could lead to incorrect interpretation of the test results, incorrect diagnoses, and/or inconclusive test results. If genetic testing reveals that the true biological relationships in a family are not as I reported them, including non-paternity (the reported father is not the biological father) and consanguinity (the parents are related by blood), I agree to have these findings reported to the healthcare provider who ordered the test.

3. Although genetic testing is highly accurate, inaccurate results may occur. These reasons include, but are not limited to mislabeled samples, inaccurate reporting of clinical/medical information, rare technical errors, or other reasons.

4. I understand that this test may not detect all of the long-term medical risks that I might experience. The result of this test does not guarantee my health and that additional diagnostic tests may still need to be done.

5. I agree to provide an additional sample if the initial sample 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 at www.nsgc.org. Further testing or additional consultations with a healthcare provider may be necessary.

To maintain confidentiality, test results will only be released to the referring healthcare provider, the ordering laboratory, to me, to other healthcare providers involved in my care, diagnosis and treatment, or to others with my consent or as permitted or required by law. Federal laws prohibit unauthorized disclosure of this information. More information can be found at: www.genome.gov/10002077

#### **INTERNATIONAL SAMPLES**

If I 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 residence.

#### SAMPLE RETENTION

After testing is complete, my sample may be de-identified and be used for test development and improvement, internal validation, quality assurance, and training purposes. **OmniHealth Diagnostics, LLC** will not return DNA samples to you or to referring healthcare providers, unless specific prior arrangements have been made. I understand that samples from residents of New York State will not be included in the de-identified research studies described in this authorization and will not retain them for more than 60 days after test completion, unless specifically authorized by my selection. The authorization is optional, and testing will be unaffected if I do not check the box for the New York authorization language. **OmniHealth Diagnostics, LLC** will not perform any tests on the biological sample other than those specifically authorized.

### DATABASE PARTICIPATION

De-identified health history and genetic information can help healthcare providers and scientists understand how genes affect human health. Sharing this de-identified information helps healthcare providers to provide better care for their patients and researchers to make new discoveries. **OmniHealth Diagnostics**, **LLC** shares this type of information with healthcare providers, scientists, and healthcare databases. **OmniHealth Diagnostics**, **LLC** will not share any personally identifying information and will replace the identifying information with a unique code not derived from any personally identifying information. Even with a unique code, there is a risk that I could be identified based on the genetic and health information that is shared. **OmniHealth Diagnostics**, **LLC** believes that this is unlikely, though the risk is greater if I have already shared my genetic or health information with public resources, such as genealogy websites.

## **INFORMED CONSENT**

#### EXOME/GENOME SEQUENCING SECONDARY FINDINGS

Applicable Only for Full Exome Sequencing and Genome Sequencing Tests.
 Oees not pertain to Xpanded® or Slice tests

As many different genes and conditions are analyzed in an exome or genome sequencing test, these tests may reveal some findings not directly related to the reason for ordering the test. Such findings are called "incidental" or "secondary" and can provide information that was not anticipated.

Secondary findings are variants, identified by an exome or genome sequencing test, in genes that are unrelated to the individual's reported clinical features.

The American College of Medical Genetics and Genomics (ACMG) has recommended that secondary findings identified in a specific subset of medically actionable genes associated with various inherited disorders be reported for all probands undergoing exome or genome sequencing. Please refer to the latest version of the ACMG recommendations for reporting of secondary findings in clinical exome and genome sequencing for complete details of the genes and associated genetic disorders. Reportable secondary findings will be confirmed by an alternate test method when needed.

WHAT WILL BE REPORTED FOR THE PATIENT? - All pathogenic and likely pathogenic variants associated with specific genotypes identified in the genes (for which a minimum of 10X coverage was achieved by exome sequencing), as recommended by the ACMG. WHAT WILL BE REPORTED FOR RELATIVES? - The presence or absence of any secondary finding(s) reported for the proband will be provided for all relatives analyzed

by an exome or genome sequencing test.

**LIMITATIONS** - Pathogenic and/or likely pathogenic variants may be present in a portion of the gene not covered by this test and therefore are not reported. The absence of reportable secondary findings for any particular gene does not mean there are no pathogenic and/or likely pathogenic variants in that gene. Pathogenic variants and/or likely pathogenic variants that may be present in a relative, but are not present in the proband, will not be identified, or reported. Only changes at the sequence level will be reported in the secondary findings report. Larger deletions/duplications, abnormal methylation, triplet repeat or other expansion variants, or other variants not routinely identified by clinical exome and genome sequencing will not be reported.

**FINANCIAL AGREEMENT AND GUARANTEE** - For insurance billing, I understand and authorize **OmniHealth Diagnostics**, **LLC** to bill my health insurance plan on my behalf, to release any information required for billing, and to be my designated representative for purposes of appealing any denial of benefits. I irrevocably assign to and direct that payment be made directly to I understand that my out-of-pocket costs may be different than the estimated amount indicated to me by **OmniHealth Diagnostics**, **LLC** as part of a benefit investigation. I agree to be financially responsible for any and all amounts as indicated on the explanation of benefits issued by my health insurance plan. If my insurance provider sends a payment directly to me for services performed by **OmniHealth Diagnostics**, **LLC** on my behalf, I agree to endorse the insurance check and forward it to **OmniHealth Diagnostics**, **LLC** within 30 days of receipt as payment towards **OmniHealth Diagnostics**, **LLC** claim for services rendered. **MEDICARE** 

A completed Advance Beneficiary Notice (ABN) is required for Medicare patients.

#### DIGITAL PATIENT LETTER CONSENT

Applicable Only for Commercial Insurance

• Estimate is provided by your health insurance company and therefore NO estimate will be sent for any orders placed with federal or state-funded insurance plans (e.g. Medicare, Medicaid, Tricare, etc.), institutional bill, or patient bill (self-pay).

To provide you with the estimated out-of-pocket expenses related to your test, **OmniHealth Diagnostics, LLC** will send you an email and/or text with the link to access your personalized Digital Patient Letter.

In order to send this information, we need your consent and agreement to the following items:

1. can use your email address or mobile phone number solely for the purpose of **OmniHealth Diagnostics**, LLC sending your estimated financial obligation. Text message data rates may apply. is not responsible for undelivered messages due to incorrect or illegible contact information.

2. will send you an email and/or text message containing a link to view your personalized Patient Letter that includes the test out-of-pocket estimate. The link is time-sensitive and will only be available for 72 hours from the time the message is sent. In order to view the estimate, you must click the link in the message.

3. If you take no action, **OmniHealth Diagnostics, LLC** will assume that you agree to move ahead with testing and will bill your health insurance. You can approve testing with insurance, switch to self-pay, or cancel the test via the link within the given 72-hour window. In turn, **OmniHealth Diagnostics, LLC** if receives your sample(s) and the billing method hasn't been changed, or the test hasn't been cancelled, we will move ahead with testing as ordered, and you will be responsible for any out-of-pocket costs for the completion of the test(s).

## **Patient Signature**

I hereby assign all rights and benefits under my health plan and all rights and obligations that I and my dependents have under my health plan to **OmniHealth Diagnostics, LLC** its assigned affiliates and authorized representatives for laboratory services furnished to me by **OmniHealth Diagnostics, LLC** irrevocably designate, authorize and appoint **OmniHealth Diagnostics, LLC** or its assigned affiliates and their authorized representatives as my true and lawful attorney-in-fact for the purpose of submitting my claims, obtain a copy of my health plan document, Summary Plan Description, disclosure, appeal, litigation or other remedies in accordance with the benefits and rights under my health plan and in accordance with federal or state laws. If my health plan fails to abide by my authorization and makes payment directly to me, l agree to endorse the insurance check and forward it to **OmniHealth Diagnostics, LLC** inmediately upon receipt. I hereby authorize **OmniHealth Diagnostics, LLC** its assigned affiliates and authorized representatives to contact me or my health Plan/administrator for billing or payment purposes by phone, text message, or email with the contact information that I have provided to **OmniHealth Diagnostics, LLC**, in compliance with federal and state laws. **OmniHealth Diagnostics, LLC**, its assigned affiliates and their authorized representatives may release to my health plan administrator, my employer, and my authorized representative my personal health information for the purpose of procuring payment of **OmniHealth Diagnostics, LLC** and for all the laboratory services. I understand the acceptance of insurance does not relieve me from any responsibility concerning payment for laboratory services and that I am financially responsible for all charges whether or not they are covered by my insurance.

Signature of Patient or Patient Representative / Relationship to Patient

## ORDERING PHYSICIAN SIGN HERE

#### Physician must only order tests that are medically necessory for the diagnosis or treatment of a 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. Furthermore, all information on this Requisition Form is true to the best of my knowledge. I agree to provide the Care Plan notes and Letter of Intent for this order if the insurance requests the lab to gather the medical necessity for any reason

Date: