COLLAGEN DIAGNOSTIC LABORATORY

UW LABORATORY FOR PRECISION DIAGNOSTICS

Peter H. Byers, MD, University of Washington, Lab H-561 Health Sciences, 1959 NE Pacific St, Seattle, WA 98195 Main Phone (206) 543-0459; Genetic Counselors (206) 543-5464; Fax (206) 616-1899; www.uwcpdx.org

LABORATORY TEST REQUISITION FORM

PATIENT INFORMATION	REPORTING RESULTS	
NAME:	REFERRING PHYSICIAN (REQUIRED):	
DOB:	NAME:	
SEX ASSIGNED AT BIRTH: MALE FEMALE	NPI #: PHYSICIAN SPECIALTY:	
YOUR PATIENT ID#:	GENETIC COUNSELOR:	
ADDRESS:	INSTITUTION:	
CITY: STATE: ZIP:	ADDRESS:	
PHONE:	CITY: STATE: ZIP:	
[]	COUNTRY:	
SAMPLE TYPE:	PHONE:	
	FAX:	
Blood (PREFERRED) 5-7cc, purple top EDTA	EMAIL:	
DNA (minimum $1\mu g$; must be extracted in		
<i>CLIA accredited or other suitable lab</i>)	REFERRING LAB:	
☐ Saliva	SEND OUT COORDINATOR:	
Amniocytes (2 T25-flasks, cultured)	ADDRESS:	
CVS Cells (2 T-25 flasks, cultured)	CITY: STATE: ZIP:	
•	COUNTRY:	
Other:	PHONE:	
COLLECTION DATE:	FAX:	

data registries and/or for publication. Individuals may refuse to allow CDL to use their information and specimen by signing below. A description of this testing policy and research approach is available at *www.uwcdl.org*

OPT OUT SIGNATURE

DATE

CLINICAL INFORMATION – PLEASE ATTACH CLINIC NOTE SUSPECTED DIAGNOSIS: Osteogenesis Imperfecta Ehlers-Danlos Syndrome Marfan Syndrome Loeys-Dietz Syndrome ☐ Familial Aneurysm Syndrome ☐ Other: CLINICAL DESCRIPTION - PLEASE ATTACH CLINIC NOTE: Blue sclerae Lens dislocation Dentinogenesis Imperfecta Hearing Loss High arched palate □ Translucent skin □ Easy bruising Atrophic scarring Bifid Uvula Dural ectasia Scoliosis Bone deformity Short Stature ☐ Joint hypermobility ☐ Joint dislocations Congenital contractures Fractures (age and location): Vascular Event: dissection, aneurysm, rupture (age and location): Other findings/history: Positive family history (please include pedigree) Special Instructions and Additional Information (if prenatal, include EDD):

	TEST REQUESTED (See website for current costs and CPT codes)						
	Mutation/Variant Testing (Please provide positive control and copy of report if testing done at another laboratory) of Relative (Proband): CDL#: Relationship to Proband: Mutation/Variant to be tested: Mutation/Variant to be tested:						
Panels an	id single gene tests include both sequencing and deletion/duplication analysis by next-generation sequencing						
Osteogenesis Imperfecta (OI) and genetic bone disorders							
	Autosomal Dominant OI Panel (COL1A1, COL1A2, IFITM5)						
	OI and Genetic Bone Disorders Panel (ALPL, ANO5, B3GAT3, B4GALT7, BMP1, CASR, CCDC134, COL1A1, COL1A2, COL2A1, CREB3L1, CRTAP, FGFR3, FKBP10, GORAB, IFITM5, KDELR2, LRP5, MBTPS2, MESD, NBAS, P3H1, P4HB, PHLDB1, PLOD2, PLOD3, PLS3, PPIB, RUNX2, SEC24D, SERPINF1, SERPINH1, SLC26A2, SOX9, SP7, SPARC, TAPT1, TENT5A, TMEM38B, TNFRSF11B, TRIP11, WNT1, XYLT2)						
Ind	lividual Genes:						
	COL1A1 and COL1A2						
	IFITM5						
	Caffey Disease (COL1A1 c.3040C>T), Sanger sequencing						
	Hypophosphatasia (ALPL)						
	X-Linked Osteoporosis (PLS3)						
Osteop	petrosis Panel						
	(AMER1, CA2, CLCN7, CTSK, FAM20C, FERMT3, LEMD3, LRP5, OSTM1, PLEKHM1, SNX10, TCIRG1, TNFRSF11A, TNFSF11)						
Ehlers-	-Danlos Syndromes (EDS)						
	Comprehensive EDS Panel (ADAMTS2, AEBP1, ATP7A, C1R, C1S, CHST14, COL1A1, COL1A2, COL3A1, COL5A1, COL5A2, DSE, FKBP14, FLNA, PLOD1, SLC39A13, THBS2)						
	Classical and Vascular EDS Panel (COL5A1, COL5A2, COL3A1)						
	Classical EDS (COL5A1 and COL5A2)						
	Vascular EDS (COL3A1)						
	Arthrochalasia EDS (Exon 6 COL1A1/COL1A2), Sanger sequencing						
	Dermatosparaxis EDS (ADAMTS2)						
	Kyphoscoliotic EDS (<i>PLOD1</i> and <i>FKBP14</i>)						
	Periodontal EDS (C1S and C1R)						
Comp	lex EDS-like Disorders						
	(B3GALT6, B4GALT7, B3GAT3, CHST14, CHST3, XYLT1)						
	lar Genetic Disorders						
	Arterial Aneurysm Panel (ACTA2, BGN, CBS, COL1A1, COL3A1, FBN1, FBN2, FOXE3, IPO8, LOX, LTBP3, MAT2A, MFAP5, MYH11, MYLK, NOTCH1, PLOD3, PRKG1, SKI, SLC2A10, SMAD2, SMAD3, SMAD4, SMAD6, TGFB2, TGFB3, TGFBR1, TGFBR2)						
	Marfan Syndrome and Loeys-Dietz Syndrome Panel (FBN1, SMAD2, SMAD3, SMAD4, TGFB2, TGFB3, TGFBR1, TGFBR2)						
	Marfan syndrome (FBN1)						
	COL4A1 and COL4A2 Related Disorders (COL4A1 and COL4A2)						
Cutis	Laxa Panel						
	(ALDH18A1, ATP6V0A2, ATP6V1A, ATP6V1E1, ATP7A, EFEMP2, ELN, FBLN5, GORAB, LTBP4, PYCR1, RIN2, SLC2A10)						
Ectop	ia Lentis Panel						
	(ADAMTS10, ADAMTS17, ADAMTSL4, CBS, FBN1)						
Alpoi	rt Syndrome						
	Alport Syndrome Panel (COL4A3, COL4A4, COL4A5)						
	X-Linked Alport Syndrome (COL4A5)						
Stickler Syndrome							
	Stickler Panel (COL2A1, COL9A1, COL9A2, COL9A3, COL11A1, COL11A2)						
Othe	r Tests						
	Maternal Cell Contamination Studies (fragment analysis)						
	Targeted pre-mRNA Splicing Analysis (cultured fibroblasts required)						

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BILLING INFORMATION

Requested laboratory testing will be initiated only AFTER accurate billing information is provided. Canadian samples: Providers must acquire approval of all tests, including reflex testing, before submitting samples.							
BILL INSTITUTION: INSTITUTIONAL BILLING							
Institution Name:							
Tax ID Number:	Ι	PO#:					
City:	State:	ZIP:					
Phone:	Fax:						
Send Result Report:							
BILL PATIENT: SELF-PAY or INTERNATIONAL SAMPLES							
Check (payable to UW Physicians) or Money Order Amount (USD):							
Credit Card: Please contact CDL billing at (206) 685-5007 to provide credit card information over a secure line.							
Electronic Funds Transfer (EFT) (See website for details) EFT Amount: USD							
BILL INSURANCE: <u>PATIENT INSURANCE</u> *Patient needs to sign billing consent below* Preauthorization is <u>required</u> for all insurance billing over \$600.00 (note testing is not started until pre-authorization complete) Do insurance preauthorization PROCEED if approved; include PreAuth Form. Clinic notes are required.							
Preauthorization alre	Preauthorization already obtained; PreAuth approval# Attach copy of P.A.						
No Preauthorization needed (for tests less than \$600.00; confirm patient signs below) Please attach a copy of card for all insurance billing Name on Policy:							
Relationship to Patient: S	Self Parei	nt Spouse	Other (please specify):				
Cardholder DOB: Dates of Coverage:							
Patient Policy ID#:							
Group Name:		Group #:					
Insurance Co. Name:							
Claims Billing Address:							
my health care provider necessary for r reimburse my medical genetic services	ssignated insurance eimbursement. I as in full due to usual tot paid in full, co-p	sign and authorize in and customary rate li ayments, and policy	surance payments to the CDL. I understand my mits, benefit exclusions, coverage limits, lack o deductibles except where my liability is limited	his form and any other information provided by insurance carrier may not approve and of authorization, medical necessity, or otherwise. by contract or State or Federal law. A duplicate			
Patient Signature:			Date:				

BILL MEDICARE: Medicare does not pre-authorize under any circumstances.

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

SHIP SAMPLE OVERNIGHT TO:

Peter H. Byers, MD UW Laboratory for Precision Diagnostics H-561, Health Science Bldg. 1959 NE Pacific Street Seattle, WA 98195