# COLLAGEN DIAGNOSTIC LABORATORY

### UW MEDICINE CENTER 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; <a href="https://www.uwcpdx.org">www.uwcpdx.org</a>

## LABORATORY TEST REQUISITION FORM

PATIENT INFORMATION	REPORTING RESULTS
NAME:	REFERRING PHYSICIAN (REQUIRED):
DOB:	NAME:
SEX: ☐ MALE ☐ FEMALE	NPI #:
YOUR PATIENT ID#:	PHYSICIAN SPECIALTY:
ADDRESS:	GENETIC COUNSELOR:
	INSTITUTION:
CITY: STATE: ZIP:	ADDRESS:
PHONE:	CITY: STATE: ZIP:
	COUNTRY:
SAMPLE TYPE:	PHONE:
	FAX:
☐ Blood ( <b>PREFERRED</b> )5-7cc, purple top EDTA	EMAIL:
□ DNA (minimum 5µg)	
☐ Saliva	REFERRING LAB:
	SEND OUT COORDINATOR:
	ADDRESS:
CVS Cells (2 T-25 flasks, cultured)	CITY: STATE: ZIP:
Cultured fibroblasts	COUNTRY:
☐ Stored cells (CDL Repository)	PHONE:
	FAX:
data registries and/or for publication. Individua	fied data from tested individuals may be submitted to clinical genetic als may refuse to allow CDL to use their information and specimen by ng policy and research approach is available at <a href="https://www.uwcdl.org">www.uwcdl.org</a>
CLINICAL INFORMA	ATION – PLEASE ATTACH CLINIC NOTE
SUSPECTED DIAGNOSIS:	
☐ Osteogenesis Imperfecta ☐ Ehlers-Danlos Synd	drome 🔲 Marfan Syndrome 🔲 Loeys-Dietz Syndrome
☐ Familial Aneurysm Syndrome ☐ Other:	
CLINICAL DESCRIPTION – PLEASE ATTACH CLINIC	NOTE:
☐ Blue sclerae ☐ Hearing Loss ☐ High archeo	
☐ Translucent skin ☐ Easy bruising ☐ Atrophic sc	
☐ Bone deformity ☐ Short Stature ☐ Joint hypern	
	John distribution — Congestian continuentes
<ul><li>☐ Fractures (age and location):</li><li>☐ Vascular Event: dissection, aneurysm, rupture (age and</li></ul>	Alocation):
vascular Everit. dissection, afteurysin, rupture (age and	a location).
Other findings/history:	
Positive family history (please include pedigree)	
Special Instructions and Additional Information (if pren	natal, include EDC):

### **TEST REQUESTED** (See website for current costs and CPT codes)

Name of Relative (Proband): (Please provide copy of report	t if testing done at a CDL#:	nother laboratory)	
Relationship to Proband:	Gene: Mutation/Variant:		riant:
	Sequencing 1	Deletion/Duplication	Seq reflex to Del/Dup
Osteogenesis Imperfecta (OI) and genetic bone disorders			
Autosomal Dominant OI Panel (COL1A1, COL1A2, IFITM5)			
OI and Genetic Bone Disorders Panel			
(ALPL, B3GAT3, B4GALT7, BMP1, COL1A1, COL1A2, CREB3L1, CR P4HB, PLOD2, PLOD3, PLS3, PPIB, SEC24D, SERPINF1, SERPINH1, S			
Individual Gene(s):			
COL1A1 and COL1A2			
IFITM5			
Caffey Disease (COL1A1 c.3040C>T)			
Hypophosphatasia (ALPL)			
X-Linked Osteoporosis (PLS3)			
Osteopetrosis Panel			
(AMER1, CA2, CLCN7, CTSK, FAM20C, FERMT3, LEMD3, LRP5, OSTN	M1, PLEKHM1, SN	IX10, TCIRG1, TNFRSF	11A, TNFSF11)
Ehlers-Danlos Syndromes (EDS)			
Comprehensive EDS Panel (COL5A1, COL5A2, COL3A1, FLNA, PLOD1, COL1A1, COL1A2, AD.	☐ AMTS2, C1S, C1R	□ R, ATP7A, CHST14, FKP	☐ PBP14, SLC39A13)
Classical and Vascular EDS Panel (COL5A1, COL5A2, COL3A1)			
Classical EDS, types I, II (COL5A1 and COL5A2)			
Vascular EDS, type IV (COL3A1)			
Ocular-scoliotic EDS, type VI (PLOD1)			
Arthrochalasia, type VIIA/B (Exon 6 COL1A1/COL1A2)			
Dermatosparaxis, type VIIC (ADAMTS2)			
Periodontal EDS, type VIII (C1S and C1R)			
FKBP14-related EDS			
Complex EDS-like Disorders			
(B3GALT6, B4GALT7, B3GAT3, CHST14, CHST3, XYLT1)			
Vascular Genetic Disorders			
Arterial Aneurysm Panel	П	П	П
(ACTA2, BGN, CBS, COL3A1, FBN1, FBN2, FOXE3, LOX, MAT2, MF. SMAD2, SMAD3, SMAD4, TGFB2, TGFB3, TGFBR1, TGFBR2)	— AP5, MYH11, MY	LK, PLOD3, PRKG1, SK	
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, EL	LN, FBLN5, GORA	AB, LTBP4, PYCR1, RIN	(2, SLC2A10)
Ectopia Lentis Panel			
(ADAMTS10, ADAMTS17, ADAMTSL4, CBS, FBN1)			
Alport Syndrome			
Alport Syndrome Panel (COL4A3, COL4A4, COL4A5)			
X-Linked Alport Syndrome (COL4A5)			
Maternal Cell Contamination Studies			

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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	N: <u>INSTITUT</u>	<u>IONAL BILLING</u>	
Institution Name:			
Tax ID Number:		PO#:	
City:	State:	ZIP:	
Phone:	Fax:		
Send Result Report:			
BILL PATIENT: SE	I F-PAY or IN	TERNATIONAL !	SAMPLES
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Electronic Funds T			•
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BILL INSURANCE	: PATIENT I	NSURANCE	
Preauthorization is requ	ired for all insu	rance billing	
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•		• •	ER before proceeding; include PreAuth Form
<del>-</del>		d; PreAuth approva	
Please attach a copy	•		
Name on Policy:	oj cara jor an	insurance offing	3
Relationship to Patient:	Self Pa	arent Spouse	Other (please specify):
Cardholder DOB:		of Coverage:	Other (picase specify).
	Dates	n Coverage.	
Patient Policy ID#:		C #	
Group Name:		Group #:	
Insurance Co. Name:			
Claims Billing Address:			
<b>Patient Insurance Billin</b>	ng Consent:		
			third party administrator the information on this form and any other information provided by
, ,		C	urance payments to the CDL. I understand my insurance carrier may not approve and mits, benefit exclusions, coverage limits, lack of authorization, medical necessity, or otherwise.
			leductibles except where my liability is limited by contract or State or Federal law. A duplicate
or faxed copy of this authorization	is considered the sa	me as the original docume	ent.
Patient Signature:			Date:
RILL MEDICAID:	For Medicaid R	illing the following	g is required: Medicaid claims address, copy of card, and Letter of
Medical Necessity from			5 is required. The decide claims address, copy of card, and better of
•			Notice (ABN) is required for Medicare patients (see website).
	. I completed 11	a , and beneficially	1.5 des (1251), 15 required for intedicate patients (see Webbite).

### SHIP SAMPLE OVERNIGHT TO:

Peter H. Byers, MD

UW Medicine Center for Precision Diagnostics
Lab H-561, Health Science Bldg.

1959 NE Pacific Street
Seattle, WA 98195