

Supplementary Table 18. Rare variants (CADD > 20) identified in EDS patients of differing clinical EDS subtypes, in genes not currently associated with human disease or variants in genes not currently associated with an EDS phenotype. These variants have high in silico pathogenicity scores and some published evidence of biological plausibility.

Patient ID	Variant ID	Age	Sex	Clinical Diagnosis	Beighton score	Villefranche Major/Minor	Aortic & Other Vascular Involvement	Auto. Dom. Family History	Skin Biopsy	Gene NM	Current Gene annotation	Protein Domain	Rs ID ClinVar	gnomAD allele frequency	CADD DANN	ACMG classification (See footnote) criteria
34	50	30-39	F	HDCT	3	A, C, E, H, I dj	Carotid artery dissection	-	normal	PTGER4 NM_000958.3 c.644G>T	a)	p.Arg215Leu - helical transmembrane (3AA)	-	0 0.998	29.2 0.998	
404	51	40-49	M	hEDS	9	A, C, H, I a, d, f, l, u	-	+	Occasional irregular collagen fibril	MMP25 NM_022468.5 c.580C>T	a)	p.His194Tyr -	rs1004972120 -	0 -	28.9 -	
446	52	40-49	M	HDCT	4	A, C, E, I d, i, f, u	Carotid artery dissection	+	irregular collagen fibril size	ADAMTS5 NM_007038.5 c.2314A>G	a)	p.Thr772Ala - spacer domain	-	0 0.998	22.6 0.998	
446	53	40-49	M	HDCT	4	A, C, E, I d, i, f, u	Carotid artery dissection	+	irregular collagen fibril size	ADAMTS16 NM_139056.4 c.2459G>A	a)	p.Arg820Gln - spacer domain	rs748937514 -	0.0000281 -	32 0.999	
446	54	40-49	M	HDCT	4	A, C, E, I d, i, f, u	Carotid artery dissection	+	irregular collagen fibril size	NFAT5 NM_138713.4 c.3446T>A	a)	p.Val1149Asp -	-	0 0.981	25.8 0.981	
505	55	10-19	F	HDCT	-	H e, l, u	-	+	-	ROBO2 NM_002942.5 c.2018G>A	c)	p.Arg679His Fibronectin III2 346696 (LB)	rs376737394 -	0.000121 -	34 0.999	VUS PM2, PP3 (Supp) BP6 (S)
566	56	60-69	M	hEDS	5	A, C, E, H, I, J x, y, aa	-	biparental	Collagen fibril size variability	SYAP1 NM_032796.4 c.37C>T	a)	p.Gln13Ter -	0 -	36 0.998		
703	57	10-19	F	hEDS	-	C, H t, u	-	-	-	LZTS1 NM_021020.5 c.1483G>A	a)	p.Glu495Lys -	rs150225368 -	0.0005212 0.997	22.8 0.997	
761	58	20-29	M	hEDS	6	B, C, H, I, J d, f, t, u, v	-	+	-	C9 NM_001737.5 c.1052C>G	c)	p.Ser351Cys - Transmembrane	rs1999424520 -	0.0000318 0.991	25.5 0.991	VUS PM2
1396	59	0-9	M	kEDS	7	C, H, J e, f, u, w	-	+	-	INO80D NM_017759.5 c.1822-1823del eAAC	a)	p.Thr608Ter -	0 -	35 -		
1450	60	30-39	F	hEDS	-	B, C, H, I a, t, u premature rupture of membranes	-	+	Collagen fibril size variability	MMP8 NM_002424.3 c.679C>T	a)	p.His227Tyr -	rs769627751 -	0.00000518 0.995	23.6 0.995	
1491	61	20-29	F	hEDS	6	C, H d, f, t, y	-	-	-	FBN3 NM_032447.5 c.6988C>T	a)	p.Arg2330Trp - TB9 domain	rs372443838 -	0.0000678 0.999	34 0.999	
1620	62	20-29	M	hEDS	6	C, H, I d, f, t, u	-	+	-	ITGA2 NM_002203.4 c.1027A>G	c)	p.Asn343Asp -	0 -	28.4 0.998	VUS PM2	
1625	63	60-69	F	HDCT	-	- B, t megacolon	ApR	-	-	TGFBI1, NM001042454.3 c.199C>T	a)	p.Arg671rp - Nr Phosphoserine	-	0 0.999	35 0.999	
1695	64	20-29	F	hEDS	8	C, H, I f, u	-	+	-	NOTCH4 NM_004557.4 c.3203C>A	a)	p.Pro1068His -	rs765636311 -	0 0.994	22.4 0.994	
1717	65	40-49	F	hEDS	7	C, H d, t	-	-	-	C3 NM_000064.3 c.910C>T	c)	p.Arg304Trp - Neighbours phosphoserine	rs1189452748 -	0.00000399 0.999	24.4 0.999	VUS PM2

ACMG criteria as per Richards et al. (9): P = pathogenic, LP = likely pathogenic, VUS/LP = variant of uncertain significance close to criteria for LP classification, VUS = variant of uncertain significance, LB = likely benign, B = benign. Individual criteria (9), Table 3)

VUS* are defined here as including VUS that according to ACGS criteria are "hot", "warm" or "tepid" Variants of Uncertain Significance (Figure 6 of <https://www.acgs.uk.com/media/11631/uk-practice-guidelines-for-variant-classification-v4-01-2020.pdf>).

Segregation analysis, re-evaluation for specific phenotypic features and/or further functional analysis may enable variant reclassification, using ACMG criteria.

EDS Diagnostic Criteria as per list in Supplementary Table 1.