## **RESEARCH LETTER**



# Enrichment of Rare Variants in Loeys– Dietz Syndrome Genes in Spontaneous Coronary Artery Dissection but Not in Severe Fibromuscular Dysplasia

**S** pontaneous coronary artery dissection (SCAD) is a prime cause of acute myocardial infarction in women < 50 years. Fibromuscular dysplasia (FMD) is present in 17% to 86% of patients with SCAD.<sup>1</sup> In contrast, the prevalence of SCAD in FMD cohorts is <3%.<sup>1</sup> Although the association between these 2 conditions has gained significant interest over the past years, the convergent underlying molecular mechanisms remain elusive.

Familial recurrence studies pointed to a genetic contribution for SCAD and FMD.<sup>2</sup> In case reports and a small-scale exome sequencing-based cohort study, SCAD has occasionally been linked to FBN1, COL3A1, and SMAD3 mutations.<sup>2</sup> For FMD, a low yield of molecular testing for syndromic thoracic aortic aneurysm and dissection genes has been reported.<sup>2,3</sup> We aimed to investigate the contribution of rare genetic variants in 25 known thoracic aortic aneurysm and dissection genes to the cause of SCAD and FMD because of the frequent association between the 2 entities. Haloplex-based gene panel sequencing was performed in a large multicenter unrelated patient cohort comprising 179 patients with SCAD±FMD and 102 patients with severe FMD only. The study has full United Kingdom Health Research Authority ethical clearance, and Institutional Review Board approval was obtained from the University of Antwerp and Université Catholique de Louvain. All patients signed informed consent. Variant filtering involved selection of heterozygous nonsynonymous coding or splice site (±2bp and +5bp) variants that are absent in gnomAD v2.1.1, have a minor allele frequency <0.01%, or between 0.01% and 0.1% combined with a CADD score (Combined Annotation Dependent Depletion) >20. After Sanger sequencing validation, selected variants were classified according to the American College of Medical Genetics guidelines. As an independent "control" cohort, gnomAD (genome aggregation database) v2.1.1 (approximately representing the general population) was filtered identically.<sup>4</sup> Two case-control burden analyses were subsequently performed using Fisher exact (expected values <5) or chi-square statistics (expected values  $\geq$ 5): SCAD±FMD versus gnomAD and FMD only versus gnomAD. To address the multiple testing burden, the significance of the association tests was evaluated using false discovery rate analysis by means of the q-value method. A q value <0.05 was considered significant. Segregation analysis was carried out for all filtered variants in the associated genes and (likely) pathogenic variants in the other genes if DNA of relatives was available.

Of the patients with SCAD, 38.5% also showed FMD. Connective tissue disease features (including joint hypermobility or dislocations, thin or smooth or hyperelastic skin, varicose veins, pes planus or talipes, hallux valgus, pectus asymmetry or excavatum/carinatum, inguinal hernia, scoliosis, cutis marmorata, joint pain, easy bruising, striae, atrophic scars, tendon rupture, osteoarthritis) were displayed in 26.3% and extracoronary arterial involvement in 26.3%, and 21.8% had a positive

Aline Verstraeten, PhD : Alexandre Persu<sup>®</sup>, MD, PhD\* David Adlam<sup>®</sup>, DPhil\* Bart Loeys<sup>®</sup>, MD, PhD\*

\*Drs Persu, Adlam, and Loeys contributed equally.

The full author list is available on page 1023.

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#### Table. Overview of Rare Variants in LDS Genes and (Likely) Pathogenic Variants in Other Thoracic Aortic Aneurysm and Dissection Genes Extracoronary MAF ACMG Arterial **Gene Variant** qnomAD Classification M/F CTD ΗT SCAD FMD Involvement **Family History** SMAD2 c.619C>G VUS (PP2; PM2) F Cx/RCA 0 Ν Y (p.Pro207Ala) SMAD2 c.1082A>C 0 LP (PP2-3; Μ Υ Υ LAD Tortuosity carotid/ (p.Asn361Thr) PM1-2) iliac artery SMAD2 c.1108C>T 0 LP (PP2-3; F Ν Ν RCA R/CC/other Arterial dissection (p.Pro370Ser) PM1-2) SMAD3 c.871+1G>A 0 LP (PP3; PM1-F Υ Ν LAD Infrarenal Sister: incidentally identified 2.4)aneurysm small brain aneurysm (–) Brother: splenic aneurysm requiring surgery (+) Mother: abdominal aortic rupture (?) *TGFB2* c.440C>T VUS (PM1) 4.1E-4 F Ν Υ LAD R (p.Pro147Leu) TGFB2 c.440C>T 4.1E-4 VUS (PM1) F LAD Ν Ν (p.Pro147Leu) VUS (PP3; PM1) F TGFB2 c 631C>T 3.2E-5 Ν Ν LAD R/CC Vertebral artery (p.Arg211Cys) dissection TGFB2 c.740G>T VUS (PP3: F LAD 0 Ν Ν (p.Gly247Val) PM1-2)

<i>TGFB3</i> c.454C>T (p.Arg152Trp)	1.4E-5	VUS (PP3; PM1)	F	Y	N	LAD	R	Common iliac artery aneurysm	
<i>TGFB3</i> c.463C>T (p.Arg155Trp)	0	VUS (PP3; PM1-2)	F	Y	N	OM2		Tortuosity vertebral arteries	
<i>TGFB3</i> c.813G>C (p.Lys271Asn)	7.4E-5	VUS (PM1)	F	?	Y	Cx	?		Brother: thoracic aortic aneurysm (?)
<i>TGFB3</i> c.1202T>C (p.Leu401Pro)	0	LP (PP3,5; PM1-2)	F	N	N		NA		Father: abdominal and thoracic aortic aneurysm (+)
<i>TGFBR1</i> c.739G>A (p.Glu247Lys)	0	VUS (PP3; PM1-2)	Μ	N	N	LAD			
<i>TGFBR1</i> c.1499G>C (p.Gly500Ala)	4.0E-6	VUS (-)	F	N	Y	LAD	R/CC	Arterial dissection	
<i>TGFBR2</i> c.1082A>G (p.Tyr361Cys)	0	VUS (PP3; PM1-2)	F	N	Y	NA		CC aneurysm and arterial dissection	
<i>TGFBR2</i> c.1718C>T (p.Ser573Leu)	1.6E-5	VUS (PP3)	F	N	N	RDP	R		
<i>TGFBR2</i> c.1396+5G>T	0	LP (PP3; PM2,4,6)	F	Y	N	Cx			(de novo)
<i>LOX</i> c.893T>G (p.Met298Arg)	0	P (PP3; PM1-2; PS1,3)	F	Y	Y	Cx	R		Mother: brain aneurysm (?)
<i>FLNA</i> c.3806G>A (p.Gly1269Asp)	0	LP (PP2-3; PM2,4)	F	N	Y	LAD		Mild aortic dilation	Mother: thoracic aortic aneurysm and rupture (?)
									Daughter: BAV with mild ascending aortic aneurysm (?)
COL3A1 c.737delG (p.Gly246Aspfs*17)	0	P (PP3; PM2; PVS1)	F	N	?		R	Arterial aneurysm	
<i>COL3A1</i> c.2177G>T (p.Gly726Val)	0	LP (PP2-3; PM1- 2,5)	F	N	Y	LAD/Cx			

Extracoronary arterial involvement includes tortuosity, aneurysm, or dissection of noncoronary arteries. Family history is deemed positive on documentation of at least 1 first- or second-degree relative with SCAD, FMD, or arterial aneurysm/dissection. ? indicates unknown variant status; (–), variant is absent; (+), variant is present; ACMG, American College of Medical Genetics; BAV, bicuspid aortic valve; CC, cervico-cephalic; CTD, connective tissue disease findings; Cx, circumflex coronary artery; F, female; FMD, fibromuscular dysplasia; gnomAD, genome aggregation database; HT, hypertension; LAD, left anterior descending coronary artery; LDS, Loeys–Dietz syndrome; LP, likely pathogenic; M, male; MAF, minor allele frequency; N, no; NA, yes, but the location is unknown; OM2, obtuse marginal 2; P, pathogenic; R, renal; RCA, right coronary artery; RDP, right descending posterior coronary artery; SCAD, spontaneous coronary artery dissection; TAAD, thoracic aortic aneurysm and dissection; VUS, variant of uncertain significance; and Y, yes. family history. In the FMD-only cohort, these frequencies were 0%, 84.3%, and 2.9%, respectively. Multifocal FMD was observed in 84.3% of cases, and in 46.1% of patients, FMD was observed in multiple vascular beds.

In the single-gene burden analyses, only SMAD2 reached significant enrichment for rare variants in patients with SCAD±FMD compared with the general population after false discovery rate correction (3/358) [0.8%] versus 172/241932 [0.1%], P=1E-3, g=1.7E-2). However, we did observe a higher frequency of variants in all 6 Loeys–Dietz syndrome (LDS) genes (ie, TGFBR1/2, SMAD2/3, TGFB2/3; Table). A combined LDS gene burden analysis confirmed a highly significant *P* value surviving false discovery rate correction (16/358 [4.5%] versus 3485/235972 [1.5%]; P=3E-6, q=1E-4). The patient cohort with variants in the LDS genes was not statistically significantly different from the whole SCAD±FMD cohort with respect to sex, age at SCAD event, family history, SCAD location, FMD co-occurrence, connective tissue disease findings, history of hypertension, and concurrence with extracoronary tortuosity or aneurysm/ dissection. None of the variant carriers was clinically diagnosed with LDS or presented with the typical LDS triad, suggesting that these variants can cause a milder phenotype. Segregation analysis in the limited number of relatives from which DNA was available hints to variable expressivity (Table). Although no significance was reached for the other analyzed thoracic aortic aneurysm and dissection genes, we also identified likely pathogenic variants in COL3A1, FLNA, and LOX in patients with SCAD±FMD with a positive family history of arteriopathy (Table). Whereas pathogenic COL3A1 variants were described in patients with SCAD previously,<sup>2</sup> FLNA and LOX variants have not. Altogether, thoracic aortic aneurysm and dissection panel analysis of the SCAD±FMD cohort reached a rare variant uptake of 10.6%.

Despite previous identification of elevated TGFB1 and TGFB2 plasma levels in patients with mostly severe FMD,<sup>2</sup> neither the single-gene analyses nor combined LDS gene analysis (1/204 [0.5%] versus 3485/235972 [1.5%]; P=3.8E-1) yielded significant P values in our FMD-only series. However, we did identify 1 multifocal FMD patient with a *TGFB3* variant (p.Leu401Pro) that was previously reported in LDS,<sup>5</sup> and for the first time identified a patient with FMD (multifocal) with a pathogenic *COL3A1* variant (p.Gly246Aspfs\*17).

In conclusion, we demonstrated that rare variants in the known LDS genes impinge on SCAD risk, implying a pathophysiological role for dysregulated transforming growth factor  $\beta$  signaling and, hence, opening new avenues for SCAD research and future prevention and therapy. Although validation in other SCAD cohorts is warranted, our results advocate for routine molecular diagnostic screening of LDS genes in patients with SCAD, even in those without

connective tissue disease manifestations. We showed that pathogenic *FLNA* and *LOX* variants are occasionally found in SCAD±FMD cases and revealed the presence of pathogenic *COL3A1* variants in both patients with SCAD±FMD and patients with FMD only.

Gene panel sequencing was done for ACTA2, BGN, COL3A1, ELN, EMILIN1, FBN1/2, FLNA, FOXE3, LOX, MAT2A, MFAP5, MYH11, MYLK, NOTCH1, PRKG1, SKI, SMAD2/3/4/6, TGFB2/3, and TGFBR1/2. For all samples, at least 99% of the target region was covered at  $\geq$ 30×. The following transcripts were used to annotate all variants in the associated genes or (likely) pathogenic variants in any of the other genes: ENST00000402690, ENST00000327367, ENST00000366929, ENST00000 238682, ENST00000374994, ENST00000359013, ENST00000304636, ENST00000369850, and ENST00 000231004.

#### **ARTICLE INFORMATION**

Data sharing: The data that support the findings of this study are available from the corresponding author upon reasonable request.

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#### **Authors**

Aline Verstraeten, PhD; Melanie H.A.M. Perik, MSc; Anna A. Baranowska<sup>®</sup>, MSc; Josephina A.N. Meester, PhD; Lotte Van Den Heuvel, MSc; Jarl Bastianen, MSc; Marlies Kempers, MD, PhD; Ingrid P.C. Krapels, MD, PhD; Angela Maas, MD, PhD; Andrea Rideout, MSc; Anthony Vandersteen, MD, PhD; Glenda Sobey, MD; Diana Johnson, MSc; Erik Fransen, PhD; Neeti Ghali<sup>®</sup>, MD; Tom Webb<sup>®</sup>, PhD; Abtehale Al-Hussaini, MBBS; Peter de Leeuw<sup>®</sup>, MD, PhD; Philippe Delmotte, MD; Marilucy Lopez-Sublet, MD; Marco Pappaccogli<sup>®</sup>, MD; Muriel Sprynger<sup>®</sup>, MD; Laurent Toubiana, PhD; European/International Fibromuscular Dysplasia Registry and Initiative (FEIRI); Lut Van Laer, PhD; Fleur S. Van Dijk, MD, PhD; Mikka Vikkula<sup>®</sup>, MD, PhD; Nilesh J. Samani, MD; Alexandre Persu<sup>®</sup>, MD, PhD; David Adlam<sup>®</sup>, DPhi; Bart Loeys<sup>®</sup>, MD, PhD

#### Correspondence

Correspondence to Bart Loeys, MD, PhD, Center of Medical Genetics, Faculty of Medicine and Health Sciences, University of Antwerp and Antwerp University Hospital Prins Boudewijnlaan 43, 2650 Antwerp, Belgium; or David Adlam, MD, PhD, Department of Cardiovascular Sciences, NIHR Leicester Biomedical Research Centre, University of Leicester, Glenfield Hospital, Groby Road, Leicester LE3 9QP, United Kingdom. Email bart.loeys@uantwerpen.be or da134@le.ac.uk

### Affiliations

Center of Medical Genetics, University of Antwerp and Antwerp University Hospital, Belgium (A. Verstraeten, M.H.A.M.P., J.A.N.M., L.V.D.H., J.B., E.F., L.V.L., B.L.). Department of Cardiovascular Sciences and National Institute for Health Research Leicester Biomedical Research Centre, Glenfield Hospital, United Kingdom (A.A.B., T.W., A.A.-H., N.J.S., D.A.). Department of Human Genetics (M.K., B.L.), Department of Cardiology (A.M.), Radboud University Nijmegen Medical Center, The Netherlands. Department of Clinical Genetics (I.P.C.K.), Department of Medicine (P.D.L.), Maastricht University Medical Centre, The Netherlands. Maritime Medical Genetics Service, Izaak Walton Killam (IWK) Health Centre, Halifax, Canada (A.R., A. Vandersteen). Division of Medical Genetics, Department of Pediatrics, Dalhousie University, Halifax, Canada (A. Vandersteen). Ehlers Danlos Syndrome National Diagnostic Service, Sheffield Clinical Genetics Department, Northern General Hospital, United Kingdom (G.S., D.J.). StatUa Center for Statistics, University of Antwerp, Belgium (E.F.). Ehlers-Danlos Syndrome, National Diagnostic Service, Northwick Park and St. Mark's Hospitals, Harrow, United Kingdom (N.G., F.S.V.D.). Division of Cardiology, Centre Hospitalier Universitaire Ambroise

Paré, Mons, Belgium (P.D.). Department of Internal Medicine, European Society of Hypertension Excellence Centre, Centre Hospitalier Universitaire (CHU) Avicenne, assistance publique hôpitaux de paris (AP-HP), Bobigny, France (M.L.-S.). Pole of Cardiovascular Research, Institut de Recherche Expérimentale et Clinique, Division of Cardiology, Cliniques Universitaires Saint-Luc (M.P., A.P.), Human Molecular Genetics, de Duve Institute (M.V.), Université Catholique de Louvain, Brussels, Belgium. Division of Internal Medicine and Hypertension Unit, Department of Medical Sciences, University of Turin, Italy (M.P.). Cardiology Department, Universitý of Liège Hospital, Belgium (M.S.). Sorbonne Université, Université Paris 13, Sorbonne Paris Cité, Institut national de la santé et de la recherche médicale (INSERM), UMR\_S 1142, Laboratoire d'Informatique Médicale et d'Ingénieurie des Connaissances en e-Santé (LIMICS), Institut de recherche pour la valorisation des données de santé (IRSAN), France (L.T.).

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#### **APPENDIX**

#### Collaborators of the European/International Fibromuscular Dysplasia Registry and Initiative (FEIRI)

Alexandre Persu<sup>®</sup>, Marco Pappaccogli<sup>®</sup>, Christophe Beauloye, Patrick Chenu, Frank Hammer, Pierre Goffette, Parla Astarci, André Peeters, Robert Verhelst, Miikka Vikkula<sup>®</sup>, Patricia Van der Niepen, Frank Van Tussenbroek, Tine De Backer, Sofie Gevaert, Dimitri Hemelsoet, Luc Defreyne, Hilde Heuten, Laetitia Yperzeele, Thijs Van der Zijden, Jean-Philippe Lengelé, Jean-Marie Krzesinski, Muriel Sprynger<sup>®</sup>, Philippe Delmotte, Peter Verhamme, Thomas Vanassche, Pasquale Scoppettuolo, Jean-Claude Wautrecht, Bojan Jelaković, Zivka Dika, Rosa Maria Bruno, Stefano Taddei, Caterina Romanini, Ilaria Petrucci, Franco Rabbia, Marco Pappaccogli<sup>®</sup>, Silvia Di Monaco, Gian Paolo Rossi, Silvia Lerco, Pietro Minuz, Giancarlo Mansueto, Sergio De Marchi, Denise Marcon, Bram Kroon, Peter de Leeuw<sup>®</sup>, Wilko Spiering, Bert-Jan van den Born, Esteban Poch, Enrique Montagud-Marrahi, Alicia Molina, Elena Guillen, Marta Burrel, Gregor Wuerzner, Lucia Mazzolai, Giacomo Buso

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