

Cardiovascular manifestations of hypermobile Ehlers–Danlos syndrome and hypermobility spectrum disorders

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
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Abstract

Introduction: Mitral valve prolapse and aortic root dilatation are reported in association with hypermobile Ehlers–Danlos syndrome (hEDS), but the full phenotypic spectrum of cardiovascular complications in this condition has not been studied in the aftermath of updated nosology and diagnostic criteria. **Methods:** We performed a retrospective review of 258 patients (> 94% adults) referred to a multidisciplinary clinic for evaluation of joint hypermobility between January 2017 and December 2020 and diagnosed with hEDS or a hypermobility spectrum disorder (HSD) to determine the incidence and spectrum of cardiovascular involvement. **Results:** Mitral valve prolapse was present in 7.5% and thoracic aortic dilatation in 15.2%. Aortic dilatation was more frequent in individuals with hEDS (20.7%) than with HSD (7.7%) and similarly prevalent between males and females, although was mild in > 90% of females and moderate-to-severe in 50% of males. Five individuals (1.9%) with hEDS/HSD had extra-aortic arterial involvement, including cervical artery dissection (CeAD, $n = 2$), spontaneous coronary artery dissection (SCAD, $n = 2$), and SCAD plus celiac artery pseudoaneurysm ($n = 1$). This is the first series to report the prevalence of CeAD and SCAD in hEDS/HSD. **Conclusions:** Cardiovascular manifestations in adults with hEDS/HSD, especially females, are typically mild and readily assessed by echocardiography. Since the risk of progression has not yet been defined, adults with hEDS/HSD who are found to have aortic dilatation at baseline should continue ongoing surveillance to monitor for progressive dilatation. Cardiovascular medicine specialists, neurologists, and neurosurgeons should consider hEDS/HSD on the differential for patients with CeAD or SCAD who also have joint hypermobility.

Keywords

aortic disease, cervical artery dissection (CeAD), Ehlers–Danlos syndrome, genetics, spontaneous coronary artery dissection

Introduction

The Ehlers–Danlos syndromes (EDS) are a heterogeneous group of hereditary connective tissue disorders affecting collagen production, structure, and function, characterized by a range of clinical features, from joint hypermobility and tissue fragility to arterial aneurysms, dissections, and organ rupture.¹ Recently, updated nosology delineates 13 EDS subtypes, highlighting associated co-morbid conditions. Within these guidelines, the commonest form, hypermobile EDS (hEDS), is further distinguished from the related hypermobility spectrum disorders (HSD), but with a higher expected incidence of systemic complications.²

Potentially serious cardiovascular manifestations are well cited in the vascular, classical, and cardiac-valvular forms of EDS.^{3–5} In contrast, only mild, nonprogressive thoracic aortic dilatation and mitral valve prolapse (MVP) have historically been recognized as cardiovascular features of hEDS.^{6,7} Incidence of these disorders were previously

reported between 6% and 21%,^{8,9} prompting echocardiography to be routinely included as part of the evaluation of joint hypermobility syndromes. Although these initial reports

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Table 1. Definition of hypermobile Ehlers–Danlos syndrome and hypermobility spectrum disorder diagnoses.

Diagnosis	Main diagnostic features
Hypermobile Ehlers-Danlos syndrome (hEDS)	Generalized joint hypermobility (positive Beighton score), positive family history, and/or specific systemic manifestations
Generalized hypermobility spectrum disorder (G-HSD)	Generalized joint hypermobility (positive Beighton score) with musculoskeletal involvement
Localized hypermobility spectrum disorder (L-HSD)	Localized joint hypermobility (negative Beighton score) with musculoskeletal involvement
Historical hypermobility spectrum disorder (H-HSD)	Historical presence of joint hypermobility (e.g., positive 5-point questionnaire) but negative Beighton score, musculoskeletal involvement
Asymptomatic generalized joint hypermobility (A-GJH)	Generalized joint hypermobility, no musculoskeletal involvement

Diagnostic criteria are based on Malfait et al., 2017² and Castori et al. 2017.¹⁵ All diagnoses are based on exclusion of other acquired and hereditary connective tissue disorders or genetic conditions that may have overlapping features, on either clinical or molecular diagnostic grounds.

were based on small cohorts with mixed ages that included other forms of EDS and predated the updated nosology, both aortic root dilatation and MVP are included as part of the 2017 hEDS diagnostic criteria.² In the contemporary diagnostic era, echocardiography is performed in up to 90% of patients under evaluation for EDS¹⁰ but, at least in children and young adults,^{10–12} has not been found to reveal significant cardiac pathology,¹¹ prompting some to suggest that echocardiographic screening may not be necessary as part of a routine hEDS evaluation. Additionally, extra-thoracic arterial complications, such as spontaneous cervical artery dissection (CeAD),^{13,14} have been rarely reported prior to 2017 in patients with hEDS as part of broader vascular disease series. Notably, a recent comprehensive review of vascular phenotypes in nonvascular subtypes of EDS did not include hEDS or HSDs.⁴ The incidence and range of severity of cardiovascular complications in older adults with hEDS or HSD have not been systematically evaluated in the era of the updated 2017 nosology.

Methods

The study population included patients referred to our multidisciplinary Cardiovascular Genetics Program for evaluation of a heritable connective tissue disorder between January 2017 and December 2020 for suspected joint hypermobility-related symptoms by cardiologists, primary care providers, rheumatologists, orthopedists, pain management specialists, neurologists, vascular medicine specialists or self-referred, and evaluated by a study investigator (ARK, BDG or LM). Electronic health records (EHR) were reviewed for demographic data, diagnoses of hEDS or HSD (based on updated 2017 criteria^{2,15}; see Table 1 for definitions) documented by a study investigator, other cardiovascular diagnoses, and clinical genetic testing results where available. The study was conducted under the approval of Mount Sinai's Institutional Review Board (GCO # 19-0883 ISMMS) utilizing a waiver of informed consent.

Echocardiographic reports were reviewed for all subjects where available and data were systematically collected, including presence or absence of MVP, mitral valve insufficiency presence/severity, and end-diastolic aortic diameters at the sinuses of Valsalva, tubular ascending

segment and arch measured from parasternal long-axis views perpendicular to the long axis of the aorta, using the leading edge technique. Ascertainment of aortic dilatation was determined by evaluating the documented maximal thoracic aortic diameters within established aortic nomograms for age, sex, and height¹⁶; measurements ≥ 2 SDs above the mean were deemed dilated.

Descriptive statistics for categorical variables were reported as frequency and percentages. Statistical analysis was performed using Microsoft Excel. Fisher's exact test (two-tailed) was used to compare rates of aortic dilatation between sexes and in hEDS versus alternative hypermobility diagnosis (HSD or hypermobility disorder not otherwise specified). A $p \leq 0.05$ was considered statistically significant.

Results

During the study period, 258 patients met the inclusion criteria and were diagnosed with either hEDS or an HSD (Table 1). The median age was 31 years (range 4–73; 94.2% ≥ 18 years) and 90% were female. The diagnosis of hEDS was made in 116 (45%) subjects (Table 2).

Valvular assessment

Of the 254 subjects (98.4%) with echocardiographic data available, 19 (7.5%) had MVP and five (2.0%) had a structural mitral valve abnormality such as borderline prolapse, bowing or buckling. The incidence of MVP was similar in individuals with hEDS and HSDs (8, 42% vs 11, 58%, $p = 0.17$). Mild mitral regurgitation was seen in 14 individuals and mild-to-moderate in two (Table 3).

Thoracic aortic assessment

Among 230 echocardiograms reporting aortic dimensions, the thoracic aorta was dilated in 35 (15.2%), upper limit of normal in 12 (5.2%), and normal in 183 (79.6%) individuals. The prevalence of aortic dilatation was higher in patients diagnosed with hEDS (24/116, 20.7%) compared to the pooled HSD cohort (11/142, 7.7%) ($p = 0.003$). There were similar proportions of males (4/25, 16.0%) and

Table 2. Subject demographics.

Diagnosis	Total n (%)	Total Male/Female/NB-TG
hEDS	116 (45.0)	14/100/2
G-HSD	79 (30.6)	6/73/0
L-HSD	33 (12.8)	2/31/0
H-HSD	17 (6.6)	2/15/0
A-GJH	9 (3.5)	0/9/0
HSD not otherwise specified	4 (1.6)	1/3/0
Total	258	25/231/2

Mean age, years (range): 31 (4–73).

A-GJH, asymptomatic generalized joint hypermobility; G-HSD, generalized HSD; hEDS, hypermobile Ehlers–Danlos syndrome; H-HSD, historical HSD; HSD, hypermobility spectrum disorder; L-HSD, localized HSD; NB-TG, nonbinary/transgender (female-to-male).

Table 3. Mitral valve anomalies and severity in 258 subjects with hypermobile Ehlers–Danlos syndrome or hypermobility spectrum disorder with available echocardiograms.

Anomaly	Severity	Total n (%)
Mitral valve prolapse	Mild	5 (1.9)
	Mild-to-moderate	19 (7.4)
	Total	24 (9.3)
Mitral regurgitation	Mild	14 (5.4)
	Mild-to-moderate	2 (0.8)
	Total	16 (6.2)

females (31/231, 13.4%) with hEDS/HSD who had aortic dilatation ($p = 0.76$). However, males with dilatation had higher maximal thoracic aortic diameters than females (average 4.3 vs 3.5 cm, $p < 0.001$), despite the male subcohort being slightly younger (median age 33.5 vs 44 years, $p = 0.31$) and of similar body surface area (BSA) (1.87 vs 1.78 mm², $p = 0.052$). Among all individuals with hEDS/HSD, there was a trend in increasing prevalence of aortic dilatation by age (Figure 1). Dilatation was mild in > 90% of females and did not show dramatic rise by decade of life (Table 4). However, two of the four males had moderate-to-severe dilatation (4.6 cm and 4.9 cm), which was confirmed by magnetic resonance angiography.

Extra-aortic arterial features

Extra-aortic arterial involvement was present in five patients, four of whom also had aortic dilatation (Table 5). Two, both also with aortic root dilatation, had CeAD: one (hEDS) had bilateral spontaneous carotid artery dissection and the other (H-HSD) had dissection of the distal left internal carotid artery with a dissecting pseudoaneurysm of the left vertebral artery. Three individuals had a diagnosis of spontaneous coronary artery dissection (SCAD) or angiographic evidence of this confirmed by a study investigator (DKD or JWO); two of these (both female) had aortic root dilatation and one (male, G-HSD) did not. One female with hEDS and both SCAD and aortic root dilatation also had a

focal outpouching of the proximal celiac artery (5 mm width × 5 mm depth) suggestive of a saccular aneurysm. All five individuals were assessed and ruled out for fibromuscular dysplasia (FMD) by a study investigator (DKD or JWO). Extra-aortic findings were similarly distributed between cases of hEDS ($n = 2$) and HSDs ($n = 3$).

Genetic assessment

Genetic testing through Clinical Laboratory Improvement Amendments (CLIA)-approved commercial laboratories was performed as part of the clinical evaluation (indicated on the basis of either personal history of aortic dilatation or extra-aortic arterial findings or family history of aneurysm or dissection) or available from outside evaluations on 41/258 (15.9%) subjects. Targeted gene sequencing for vascular ± classical EDS (either *COL3A1* alone or with *COL5A1* and *COL5A2*) was performed in 16 individuals, and either a comprehensive aortopathy/arteriopathy gene panel (including the genes related to Marfan, *FBNI*; Loeys–Dietz, *TGFBR1*, *TGFBR2*, *SMAD3*, *TGFB2*; and vascular EDS, *COL3A1* syndromes, along with additional nonsyndromic heritable thoracic aneurysm and dissection genes) or a heritable connective tissue disorders panel was used in 25 cases. Of 22 patients with aortic dilatation and five patients with extra-aortic manifestations who underwent genetic testing, results were either negative or identified a variant of uncertain significance (VUS) (Table 6). One additional subject, a 35-year-old female, had been referred after carrier screening during pregnancy revealed a likely pathogenic partial deletion in *TNXXB* (121-bp deletion in exon 35); this subject was diagnosed with localized HSD (L-HSD) and had no cardiovascular phenotype. Heterozygous pathogenic variants in *TNXXB*, which encodes tenascin-X, an extracellular protein that regulates elastic fiber integrity, have been described in case series as causing features of EDS (joint hypermobility, dislocations, and chronic pain).¹⁷ Interestingly, this patient had multiple affected relatives with hypermobility symptoms, but none was found to carry the *TNXXB* variant, suggesting that it may not have been causative or there may be an oligogenic etiology to her L-HSD. Two individuals, neither with cardiovascular phenotypes, carried pathogenic variants that were interpreted to be unassociated with their presentations. One was a 33-year-old female with hEDS who had prior genetic testing on a heritable connective tissue diseases panel and was found to be heterozygous for a pathogenic variant in *FKBP14*. Since this gene is associated with autosomal recessive kyphoscoliotic EDS, the result was interpreted to be clinically insignificant. Another individual, a 61-year-old female with L-HSD, proceeded to genetic evaluation for a possible metabolic condition and was found on whole exome sequencing to harbor a heterozygous pathogenic variant in *FLG*; this gene is associated with ichthyosis vulgaris and was not relevant to HSD. Following completion of genetic testing, no alternative genetic arteriopathy was apparent in the patients with arterial findings, other than clinically diagnosed hEDS/HSD.

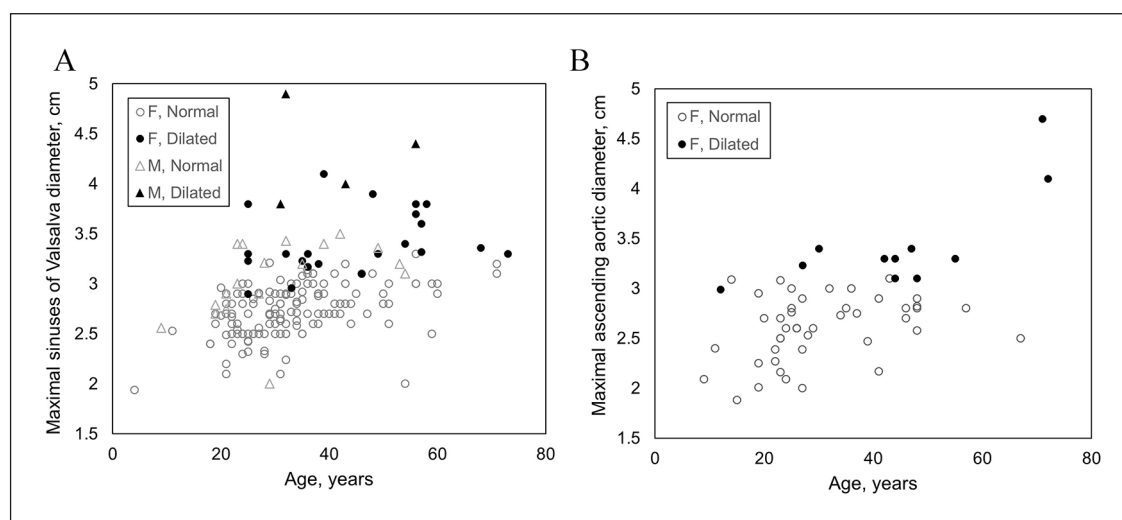


Figure 1. Distribution of thoracic aortic diameters by age measured in individuals with hypermobile Ehlers–Danlos syndrome or hypermobility spectrum disorder with available echocardiographic data. Measurements are represented according to anatomic location where aortic diameter was maximal, either (A) sinuses of Valsalva or (B) tubular ascending aorta.

Table 4. Age distribution of hypermobile Ehlers–Danlos syndrome or hypermobility spectrum disorder diagnoses and within-group average maximal thoracic aortic diameters among females and males with noted aortic dilatation.

Age group (years)	Female		Male	
	<i>n</i> hEDS/HSD	Maximal aortic diameter (average, cm)	<i>n</i> hEDS/HSD	Maximal aortic diameter (average, cm)
< 20	0/2	2.9	–	–
20–29	4/0	3.4	–	–
30–39	7/0	3.4	3/0	4.2
40–49	3/4	3.3	1/0	4.6
50–59	5/2	3.6	–	–
60–69	1/0	3.4	–	–
70–79	1/2	4	–	–

hEDS, hypermobile Ehlers–Danlos syndrome; HSD, hypermobility spectrum disorder or hypermobility disorder not otherwise specified.

Table 5. Arterial phenotypes in five subjects with hypermobile Ehlers–Danlos syndrome or hypermobility spectrum disorder.

Subject	Hypermobility diagnosis	Aortic dilatation	CeAD	SCAD	Celiac artery PA
1	hEDS	x	x		
2	hEDS	x		x	x
3	H-HSD	x	x		
4	G-HSD	x		x	
5	G-HSD			x	

CeAD, cervical artery dissection; G-HSD, generalized hypermobility spectrum disorder; hEDS, hypermobile Ehlers–Danlos syndrome; H-HSD, historical hypermobility spectrum disorder; PA, pseudoaneurysm; SCAD, spontaneous coronary artery dissection.

Discussion

A contemporary description of the full spectrum of cardiovascular manifestations in patients with the commonest form of EDS, hEDS, and related HSDs across all age groups was previously lacking. We, therefore, retrospectively assessed 258 subjects with hEDS or HSD diagnosed according to the updated 2017 clinical criteria. The two cardiovascular conditions that have historically been cited in association with hypermobility disorders, MVP^{2,18} and aortic root dilatation,⁶ were seen in 7.5% and 15.2% of the

subjects with available echocardiograms, respectively, with severity typically mild. MVP was enriched compared to the expected population prevalence of 0.7%.¹⁹ Since hEDS and HSD, as defined in the updated 2017 guidelines, occur on a spectrum and share many systemic associations,²⁰ it is not surprising that we found a similar incidence of MVP among both diagnoses.

Previous studies have documented an association between thoracic aortic dilation and joint hypermobility syndromes, but this is emerging as an age-dependent phenomenon. Although pre-2017 pediatric series cited the rate

Table 6. Reportable genetic variants in hypermobile Ehlers–Danlos syndrome or hypermobility spectrum disorder subjects.

Age (years)	Sex	Clinical diagnosis	CV phenotype	Test type	Gene	Zygosity	cDNA variant	AA change	Classification of variant
33	F	hEDS	None	CTD	<i>FKBP14</i>	het	c.362dupC	p.E122RfsX7	Pathogenic variant for autosomal recessive disorders
35	F	L-HSD	None	PGT	<i>TNXB</i>	het	121-bp deletion in exon 35		Likely pathogenic
61	F	L-HSD	None	WES	<i>FLG</i>	het	c.1501C>T	p.Arg501X	Pathogenic for ichthyosis vulgaris
44	F	G-HSD	Aortic root dilatation + SCAD	hTAAD panel	<i>COL5A1</i>	het	c.3260G>C	p.Gly1087Ala	VUS
54	F	H-HSD	Aortic root dilatation + CeAD	hTAAD panel	<i>COL5A2</i> <i>MYLK</i>	het het	c.379A>C c.1802A>G	p.Ile127Leu p.His601Arg	VUS VUS
48	F	hEDS	Aortic root dilatation + CeAD	hTAAD + CTD panels	<i>TNXB</i> <i>ADAMTS2</i>	het het	c.2083C>T c.1793G>A	p.Arg69Trp p.Arg598His	VUS VUS
71	F	H-HSD	Aortic root dilatation	hTAAD panel	<i>COL5A1</i> <i>FLNA</i>	het het	c.2708C>T c.7604G>A	p.Pro903Leu p.Cys2535Tyr	VUS VUS
58	F	hEDS	Aortic root dilatation	hTAAD panel	<i>SKI</i>	het	c.1255G>A	p.Val419Met	VUS
53	M	EDS-NOS	MVP	COL3A1	<i>COL3A1</i>	het	c.3418-5C>A	intronic	VUS
26	F	hEDS	None	CTD	<i>COL1A1</i> <i>B3GAT3</i>	het het	c.1018G>A c.554G>A	p.Ala340Thr p.Gly185Glu	VUS VUS

AA, protein-level amino acid; CV, cardiovascular; cDNA, coding DNA; CeAD, cervical artery dissection; CTD, connective tissue disorder panel; G-HSD, generalized hypermobility spectrum disorder; hEDS, hypermobile Ehlers–Danlos syndrome; H-HSD, historical hypermobility spectrum disorder; hTAAD, heritable thoracic aortic aneurysm and dissection; L-HSD, localized hypermobility spectrum disorder; MVP, mitral valve prolapse; NOS, not otherwise specified; PGT, prenatal genetic testing; het, heterozygous; SCAD, spontaneous coronary artery dissection; VUS, variant of uncertain significance; WES, whole exome sequencing.

of aortic root dilatation in children with hEDS at 14%,¹² recent studies including children meeting the more stringent 2017 hEDS criteria observed this in less than 2%.^{10,11} Based on this lower prevalence, some have advocated that echocardiographic evaluations should not be routinely performed for hEDS in the absence of clinical findings or positive family history. To our knowledge, the present study is the first to report on echocardiographic findings in an older adult population with all hEDS/HSD diagnoses made according to the updated 2017 guidelines. Here, we found aortic dilatation in 15.2% of patients, reaffirming this important association. The prevalence trended higher across increasing age brackets. In the context of previous reports, we conclude that aortic dilatation in hEDS/HSD emerges later in adulthood; echocardiographic assessments may therefore not be necessary in children, but should be included as part of baseline testing for patients ≥ 21 years. Our data indicate that aortic dimensions in hEDS/HSD remain generally within the mildly dilated range, especially for females, consistent with previous reports. Although interpretation is limited by the retrospective study design, it is unlikely that aortic dilatation in the context of hEDS/HSD poses exaggerated risk of aortic emergency because we included adults over a wide age range and did not find any with aortic dissection/rupture. Of note, two males with hEDS had moderate-to-severe dilatation; aortopathy gene panel sequencing was normal in both cases, with no clear alternative genetic diagnosis. Until the

natural history of aortic dilatation can be clarified through large longitudinal studies, patients with hEDS/HSD and aortic dilatation should have ongoing surveillance according to established society guidelines.²¹ Of note, we found aortic root dilatation more frequently in patients with hEDS compared to HSDs; because this feature confers a ‘point’ on the hEDS diagnostic checklist, it is challenging to determine whether the risk of aortic pathology is truly higher in hEDS than HSDs.

Extra-aortic arterial involvement (aneurysms, dissections) is frequently cited in the vascular and classical forms of EDS and has not previously been categorically linked to hEDS. We identified systemic arterial complications in five patients with hEDS/HSD: two with CeAD and three with SCAD, one of whom also had a celiac artery pseudoaneurysm. Although the population prevalence of CeAD is unknown, community-based stroke studies have estimated it at one to three per 100,000.^{22,23} Similarly, the prevalence of SCAD is not known but is estimated at around 0.6–0.24%.²⁴ Therefore, we report the first description of incidence of CeAD (0.7%) and SCAD (1.8%) in hEDS/HSD, and interpret these phenotypes to be enriched in this cohort. Genetic panel testing was performed based on these clinical indications and did not reclassify the hEDS/HSD diagnoses to an alternative genetic arteriopathy. Vascular medicine specialists should consider hEDS/HSD as part of the differential for these conditions, especially when other

underlying conditions such as FMD and genetically defined arteriopathies (i.e., vascular EDS) have been ruled out. Of note, the prevalence of joint hypermobility in FMD is reportedly low,²⁵ so it should be feasible to clinically differentiate these potential underlying causes. Furthermore, since CeAD may present with neurologic features including headache/migraine or acute neurologic changes, neurologists and headache medicine specialists should be attuned to this association, especially in patients with known hEDS/HSD.²⁰

The genetic basis of hEDS/HSD is not yet known, but efforts are underway to molecularly characterize these disorders. Through the current clinical classification system and advances in genetic analyses, future investigations may illuminate genetic mediators of aortic and arterial risk in patients with hEDS/HSD. Further study is needed in patients with SCAD and CeAD to determine whether there is a robust association with hEDS/HSD.

Of the 258 individuals included in this study, 245 (95%) were referred to our Cardiovascular Genetics program based on a patient complaint of joint pain or hypermobility or a family history of EDS. Only eight patients were referred based on known arterial abnormalities (aortic dilatation, $n = 4$; suspected or confirmed SCAD, $n = 3$; or cervical artery dissection, $n = 1$). Following the evaluation, 31 individuals received a new diagnosis of thoracic aortic dilatation. Since the vast majority of individuals did not have known cardiovascular abnormalities prior to referral and diagnosis of hEDS/HSD, and because CeAD, SCAD, and peripheral arterial aneurysm/pseudoaneurysm are not part of the diagnostic criteria, we feel confident that this study was not biased towards ascertainment of cardiovascular manifestations in hEDS/HSD. Still, this retrospective study has several limitations inherent to the study design. Although this study contributes important information on the prevalence of aortic dilatation in older adults with hEDS/HSD, there were no longitudinal assessments made and therefore it cannot be used to describe the natural history of this finding. Prospective studies are needed to determine whether the aortic dilatation seen in hEDS/HSD confers an increased risk for future aneurysmal growth and/or dissection. In subjects with extra-aortic findings, diagnoses of SCAD and CeAD were made prior to the time of the hEDS/HSD assessment. Pan-vascular imaging is not routine or recommended in hEDS/HSD, so subclinical incidence of these manifestations could not be robustly ascertained through this study and may not be fully represented. Genetic testing to exclude other genetic connective tissue disorders relied on targeted genes or panels, not genome-wide studies. Lastly, due to lack of objective tests, diagnoses of hEDS/HSD remain clinically challenging.

Conclusion

Cardiovascular manifestations are present in up to 15% of subjects with hEDS and HSD and mainly include mild aortic dilatation and mild MVP. We report, for the first time, that SCAD and CeAD occur, respectively, in 1.8% and 0.7% of patients with hEDS/HSD. Until the natural history

of aortic dilatation is fully elucidated, a baseline echocardiographic study in adults under evaluation for hEDS is reasonable and those for whom dilatation is noted should have ongoing surveillance. Patients with extra-thoracic arteriopathy and joint hypermobility should be screened for hEDS/HSD, as these underlying diagnoses are likely under-recognized.

Declaration of conflicting interests

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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Data availability

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

References

1. Tinkle B, Castori M, Berglund B, et al. Hypermobile Ehlers-Danlos syndrome (a.k.a. Ehlers-Danlos Syndrome Type III and Ehlers-Danlos syndrome hypermobility type): Clinical description and natural history. *Am J Med Genet C Semin Med Genet* 2017; 175: 48–69.
2. Malfait F, Francomano C, Byers P, et al. The 2017 international classification of the Ehlers-Danlos syndromes. *Am J Med Genet C Semin Med Genet* 2017; 175: 8–26.
3. Eagleton MJ. Arterial complications of vascular Ehlers-Danlos syndrome. *J Vasc Surg* 2016; 64: 1869–1880.
4. D'hondt S, Van Damme T, Malfait F. Vascular phenotypes in nonvascular subtypes of the Ehlers-danlos syndrome: A systematic review. *Genet Med* 2017; 20: 562–573.
5. Chu LC, Johnson PT, Dietz HC, et al. Vascular complications of Ehlers-Danlos syndrome: CT findings. *Am J Roentgenol* 2012; 198: 482–487.
6. Wenstrup RJ, Meyer RA, Lyle JS, et al. Prevalence of aortic root dilation in the Ehlers-Danlos syndrome. *Genet Med* 2002; 4: 112–117.
7. Atzinger CL, Meyer RA, Khoury PR, et al. Cross-sectional and longitudinal assessment of aortic root dilation and valvular anomalies in hypermobile and classic Ehlers-Danlos syndrome. *J Pediatr* 2011; 158: 826–830.e1.
8. Asher SB, Chen R, Kallish S. Mitral valve prolapse and aortic root dilation in adults with hypermobile Ehlers-Danlos syndrome and related disorders. *Am J Med Genet A* 2018; 176: 1838–1844.
9. Meester JA, Verstraeten A, Schepers D, et al. Differences in manifestations of Marfan syndrome, Ehlers-Danlos syndrome, and Loeys-Dietz syndrome. *Ann Cardiothorac Surg* 2017; 6: 582–594.
10. Rauser-Foltz KK, Starr LJ, Yetman AT. Utilization of echocardiography in Ehlers-Danlos syndrome. *Congenit Heart Dis* 2019; 14: 864–867.

11. Paige SL, Lechich KM, Tierney ES, Collins RT 2nd. Cardiac involvement in classical or hypermobile Ehlers–Danlos syndrome is uncommon. *Genet Med* 2020; 22: 1583–1588.
12. Ritter A, Atzinger C, Hays B, et al. Natural history of aortic root dilation through young adulthood in a hypermobile Ehlers–Danlos syndrome cohort. *Am J Med Genet A* 2017; 173: 1467–1472.
13. Debette S, Goeggel Simonetti B, Schilling S, et al. Familial occurrence and heritable connective tissue disorders in cervical artery dissection. *Neurology* 2014; 83: 2023–2031.
14. Hermanns-Le T, Manicourt D, E. Pierard G. Familial expression of spontaneous cervical artery dissections and Ehlers–Danlos syndrome hypermobile type. *J Skin Stem Cell* 2014; 1: e27023.
15. Castori M, Tinkle B, Levy H, et al. A framework for the classification of joint hypermobility and related conditions. *Am J Med Genet C* 2017; 175: 148–157.
16. Saura D, Dulgheru R, Caballero L, et al. Two-dimensional transthoracic echocardiographic normal reference ranges for proximal aorta dimensions: Results from the EACVI norre study. *Eur Heart J Cardiovasc Imaging* 2016; 18: 167–179.
17. Zweers MC, Bristow J, Steijlen PM, et al. Haploinsufficiency of TNXB is associated with hypermobility type of Ehlers–Danlos syndrome. *Am J Hum Genet* 2003; 73: 214–217.
18. Gensemer C, Burks R, Kautz S, et al. Hypermobile Ehlers–Danlos syndromes: Complex phenotypes, challenging diagnoses, and poorly understood causes. *Dev Dyn* 2021; 250: 318–344.
19. Sattur S, Bates S, Movahed MR. Prevalence of mitral valve prolapse and associated valvular regurgitations in healthy teenagers undergoing screening echocardiography. *Exp Clin Cardiol* 2010; 15: e13–e15.
20. Malhotra A, Pace A, Ruiz Maya T, et al. Headaches in hypermobility syndromes: A pain in the neck? *Am J Med Genet A* 2020; 182: 2902–2908.
21. Hiratzka LF, Bakris GL, Beckman JA, et al. 2010 ACCF/AHA/AATS/ACR/ASA/SCA/SCAI/SIR/STS/SVM guidelines for the diagnosis and management of patients with thoracic aortic disease. A Report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines, American Association for Thoracic Surgery, American College of Radiology, American Stroke Association, Society of Cardiovascular Anesthesiologists, Society for Cardiovascular Angiography and Interventions, Society of Interventional Radiology, Society of Thoracic Surgeons, and Society for Vascular Medicine. *J Am Coll Cardiol* 2010; 55: e27–e129.
22. Schievink WI, Mokri B, Whisnant JP. Internal carotid artery dissection in a community. Rochester, Minnesota, 1987–1992. *Stroke* 1993; 24: 1678–1680.
23. Béjot Y, Daubail B, Debette S, et al. Incidence and outcome of cerebrovascular events related to cervical artery dissection: The Dijon Stroke Registry. *Int J Stroke* 2014; 9: 879–882.
24. Hayes SN, Kim ESH, Saw J, et al. Spontaneous coronary artery dissection: Current state of the science: A scientific statement from the American Heart Association. *Circulation* 2018; 137: e523–e557.
25. O’Connor S, Kim ESH, Brinza E, et al. Systemic connective tissue features in women with fibromuscular dysplasia. *Vasc Med* 2015; 20: 454–462.