

Diagnostic Criteria for Hypermobile Ehlers-Danlos Syndrome (hEDS)

This diagnostic checklist is for doctors across all disciplines to be able to diagnose EDS



Patient name:	DOB:	DOB: DOV: Evaluator:		
The clinical diagnosis of hypermobile	EDS needs the simultaneou	s presence of all	criteria, 1 and 2 and 3.	
CRITERION 1 – Generalized Joint	Hypermobility			
One of the following selected: □ ≥6 pre-pubertal children and adoles □ ≥5 pubertal men and woman to age □ ≥4 men and women over the age of	50 Beighton Sco	re:/9		
If Beighton Score is one point below age ☐ Can you now (or could you ever) pla ☐ Can you now (or could you ever) ber ☐ As a child, did you amuse your frien ☐ As a child or teenager, did your shou ☐ Do you consider yourself "double jo	ce your hands flat on the floor nd your thumb to touch your fo ds by contorting your body into ulder or kneecap dislocate on m	without bending y orearm? o strange shapes o	or could you do the splits?	
CRITERION 2 – Two or more of th	e following features (A,	B, or C) must b	e present	
without a history of significant gain Bilateral piezogenic papules of the h Recurrent or multiple abdominal he Atrophic scarring involving at least t	or loss of body fat or weight neel rnia(s) wo sites and without the forma- rolapse in children, men or nulli w palate or more of the following: on both sides, (ii) positive thumb	tion of truly papyra iparous women wit b sign (Steinberg s		
Feature A total:/12 Feature B				
☐ Positive family history; one or more	first-degree relatives indepen	dently meeting the	e current criteria for hEDS	
Feature C (must have at least one) ☐ Musculoskeletal pain in two or more ☐ Chronic, widespread pain for ≥3 more ☐ Recurrent joint dislocations or frank	nths			

CRITERION 3 - All of the following prerequisites MUST be met

- 1. Absence of unusual skin fragility, which should prompt consideration of other types of EDS
- 2. Exclusion of other heritable and acquired connective tissue disorders, including autoimmune rheumatologic conditions. In patients with an acquired CTD (e.g. Lupus, Rheumatoid Arthritis, etc.), additional diagnosis of hEDS requires meeting both Features A and B of Criterion 2. Feature C of Criterion 2 (chronic pain and/or instability) cannot be counted toward a diagnosis of hEDS in this situation.
- 3. Exclusion of alternative diagnoses that may also include joint hypermobility by means of hypotonia and/or connective tissue laxity. Alternative diagnoses and diagnostic categories include, but are not limited to, neuromuscular disorders (e.g. Bethlem myopathy), other hereditary disorders of the connective tissue (e.g. other types of EDS, Loeys-Dietz syndrome, Marfan syndrome), and skeletal dysplasias (e.g. osteogenesis imperfecta). Exclusion of these considerations may be based upon history, physical examination, and/or molecular genetic testing, as indicated.

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