

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:	DOV:	Evaluator:	
The clinical diagnosis of hypermobile EDS needs the simultaneous presence of all criteria, 1 and 2 and 3.				
CRITERION 1 – Generalized Joint Hy	permobility			
One of the following selected: □ ≥6 pre-pubertal children and adolescen □ ≥5 pubertal men* and women* to age □ ≥4 men* and women* over the age of	50 Beighton Sco	ore:/9		
If Beighton Score is one point below age- and sex-specific cut off, two or more of the following must also be selected to meet criterion: □ Can you now (or could you ever) place your hands flat on the floor without bending your knees? □ Can you now (or could you ever) bend your thumb to touch your forearm? □ As a child, did you amuse your friends by contorting your body into strange shapes or could you do the splits? □ As a child or teenager, did your shoulder or kneecap dislocate on more than one occasion? □ Do you consider yourself "double jointed"?				
CRITERION 2 — Two or more of the	following features (A _,	B, or C) must be	present	
without a history of significant gain or Bilateral piezogenic papules of the hee Recurrent or multiple abdominal herni Atrophic scarring involving at least two	loss of body fat or weight I a(s) sites and without the forma pse in children, men or null alate nore of the following: both sides, (ii) positive thum	ition of truly papyrace iparous women witho b sign (Steinberg sigi	nen in adolescents, men or pre-pubertal we eous and/or hemosideric scars as seen in cla out a history of morbid obesity or other kr	assical EDS
Feature A total:/12				
Feature B ☐ Positive family history; one or more fir	st-degree relatives indeper	dently meeting the c	urrent criteria for hEDS	
Feature C (must have at least one) ☐ Musculoskeletal pain in two or more lin ☐ Chronic, widespread pain for ≥3 month	S			

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.

