Connecting the Issues by Understanding Our Connective Tissues



Brought to you by Chiari Bridges



Connective tissues provide strength and flexibility to structures such as ...

- Ligaments, Cartilage, Bursa
- Cutaneous Tissue (skin)
- Adipose tissue (fat)
- Muscles
- Organs

- Blood
- Lymph nodes
- Bones
- Blood vessels
- Nerves

What are Heritable Disorders of Connective Tissues (HDCTs)?



There are over 200 identified Heritable Disorders of Connective Tissue (HDCTs), conditions such as:

- Ehlers-Danlos Syndrome (pronounced: ee-lers dan-low)
- Marfan Syndrome
- Loeys-Dietz Syndrome
- Stickler Syndrome
- Osteogenesis Imperfecta
- Pseudoxanthoma Elasticum
- Epidermolysis bullosa

All involve one of the many **proteins** that make up our body's connective tissue.

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Ehlers-Danlos Syndromes (EDS) are a group of hereditary connective tissue disorders involving defects in our collagen.

There are thirteen known subtypes of Ehlers-Danlos syndromes (EDS).

- Classical EDS (cEDS)
- Classical-like EDS (clEDS)
- Cardiac-valvular EDS (cvEDS)
- Vascular EDS (vEDS)
- Arthrochalasia EDS (aEDS)
- Dermatosparaxis EDS (dEDS)
- Kyphoscoliotic EDS (kEDS)
- Brittle Cornea Syndrome (BCS)

- Spondylodysplastic EDS (spEDS)
- Musculocontractural EDS (mcEDS)
- Myopathic EDS (mEDS)
- Periodontal EDS (pEDS)
- Hypermobile EDS (hEDS)
- New Unnamed sub-type involving the AEPB1



Crossover Symptoms



While each sub-type of the Ehlers-Danlos Syndromes (EDS) has its own characteristics and symptoms, it's important to remember that EDS symptoms are known to crossover from one sub-type to another.

So, while symptomology can help narrow down a diagnosis, it is not unusual for someone with one sub-type, to have symptoms or complications that seem more consistent with another subtype.

Hereditary

Ehlers-Danlos Syndromes (EDS) are known to be hereditary. Some are known to be autosomal recessive (where both parents must have the mutated gene to pass it down to a child) and others are autosomal dominant (where only one parent need have the mutated gene to pass it down to the child). It's important to remember that you can't control what you pass down to your children genetically, so it should NEVER be allowed to become a blame game!



Collagen is a Structural Protein that Makes Up 30% of the Proteins in the Human Body

Collagen is often likened to the "**fibrous glue** that holds us together at a cellular level," providing structure to our organs, muscles, bones, skin, blood vessels, and connective tissues.

Imagine the difference between something bonded together with **Gorilla Glue** versus watered down **Elmer's Glue**. The creation using Elmer's Glue wouldn't be as strong. The consistency of Elmer's Glue could cause anything depending on it to lose its form and then gravity will cause it to prolapse/herniate.



Structure Gone Awry



- Ehlers-Danlos Syndromes (EDS) are known to involve a mutation in our body's collagen, but it's not as simple as it sounds. EDS doesn't just affect our skin and joints, but everything in our bodies that was designed to interact with that collagen. That is why we tend to have so many problems throughout our bodies that you would think couldn't be related. It really is our connective tissues that connect the issues.
- Our Intervertebral Discs are comprised of collagen, therefore the discs of the Ehlers-Danlos Syndromes (EDS) patient, tend to dehydrate, thin, bulge, slip, and herniate into the spinal cord. The average person might have 1-2 disc problems in their lifetime, EDS patients tend to have many discs bulging and herniated at any given time, often in different levels of the spine. This condition is known as Degenerative Disc Disease (DDD).



Connecting EDS & Chiari Malformations

The same theory can be used to explain why many of us with EDS have prolapsed cerebellar tonsils.

- > EDS is known for causing organ prolapses and your cerebellum really is just a complex organ.
- Therefore, if you have EDS and Chiari, it should be considered an Acquired Chiari Malformation until all possible EDS related comorbidities are ruled out!
 - Even if posterior fossa hypoplasia is confirmed, it is irresponsible not to give adequate attention to all comorbid conditions that can push/pull the cerebellar tonsils down, creating an Acquired (Secondary) Chiari Malformation.

Studies have shown that Ehlers-Danlos Syndromes (EDS) can have <u>many</u> neurological and spinal manifestations, to include:



- Intracranial Hypertension (IH/IIH)
- Chiari Malformations (CM)
- Atlantoaxial & Craniocervical Instability (AAI/CCI)
- Segmental Kyphosis & Instability
- Tethered Cord Syndrome (TCS)
- CSF Leaks causing Spontaneous Intracranial Hypotension (SIH)
- Dystonia & Other Movement Disorders
- Neuromuscular Complications
- Acquired Chiari Malformations

Chiari Malformation Type 1 (CM1) has been reported as a comorbid condition in hypermobile EDS (hEDS). The precise incidence of the CM1 and EDS association is unknown, but the female to male ratio is higher in the CM1 and EDS subgroup (9:1) than in the general CM1 population alone (3:1).

-Dr. Fraser Henderson

- It is a matter of concern that so many surgeons find herniated cerebellar tonsils, diagnose it as a Chiari Malformation and assume it to be congenital.
- They offer relief from decompression surgery, without checking for any pathological conditions that might be causing this hernia of the cerebellar tonsils.
- Most of the literature simply says that an Acquired Chiari is rare without ever defining what would constitute an Acquired Chiari Malformation.
- If you have EDS and Chiari, it should be considered an Acquired Chiari Malformation until all possible EDS related comorbidities are ruled out!
 - Even if posterior fossa hypoplasia is confirmed, it is irresponsible not to give adequate attention to all the other comorbid conditions that are considered pathological to Acquired Chiari Malformations.



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Hypermobility

• It may seem counter-intuitive, but many hypermobile people also suffer from stiffness and tightness in their joints and muscles. Muscle stiffness and tightness is caused by the extra work muscles need to do to try and keep a hypermobile person's joints stable. We affectionately refer to them as *Stiff Zebras*.

"When you hear hoofbeats, think horses, not zebras!"



In medical school, students are taught to not neglect the obvious, with the analogy of, "when you hear hoofbeats, think horses, not zebras."

This philosophy poses a problem for those with Ehlers-Danlos syndromes, as it's not that our condition is rare, but that it's rare that doctors know anything about it; and patients often go years or decades pursuing answers to our numerous symptoms.



Multisystemic Manifestations

Because collagen is so abundant in the human bodies, it often affects multiple systems and a multidisciplinary approach to is often needed:

- Coordinate care
- Maximize treatment
- Reduce medical visits.

Sleep Apnea



Studies show that those with Ehlers-Danlos Syndromes (EDS) have a higher prevalence of symptomatic <u>Obstructive</u> Sleep Apnea (32% vs 6% in the general population).

For this reason, we recommended that **EVERYONE** with EDS is tested for Sleep Apnea and use their CPAP **RELIGIOUSLY!**

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Statistics

Recent statistics are proving Ehlers-Danlos to be far more common than initially believed (just a few years ago the prevalence was believed to be 1 in 5,000 to 1 in 20,000 depending on the subtype). The most recent numbers published on its prevalence is 1 in 2,500 to 1 in 5,000 people depending on the subtype (1.5-3 million people worldwide).

A Cure for Ehlers-Danlos Syndromes

There are no known cures for Ehlers-Danlos syndromes (EDS). There is no way to change/repair our mutated collagen.

We deal with one symptom at a time as our bodies try to interact with our faulty collagen as if it's normal. We typically have a multitude of medical specialists (for each manifestation) and a box of braces to try and help keep our joints in place.



"At this point in time, I put EDS in the category of being in the top three or four most severe pain problems. A lot of people for example think that cancer pain is the worst pain, but let me assure you that many EDS patients have pain far beyond any cancer patient I've ever seen. And so it's one of the pain problems that is severe, has been very troublesome, many physicians are afraid of the disease and of the kind of pain that EDS patients have."

-Dr. Forest Tennant