ATTR-CM (transthyretin amyloid cardiomyopathy) is a serious, underrecognized, and underdiagnosed type of amyloidosis that affects the heart and is associated with heart failure.

YOUR SYMPTOMS COULD ADD UP TO SOMETHING MORE SERIOUS THAN YOU REALIZE.

ATTR-CM SYMPTOMS MAY INCLUDE:

- Heart Failure with Preserved Ejection Fraction
- Irregular Heartbeat
- Shortness of Breath
- GI Problems
- Pain or Numbness in Lower Back or Legs
- Diagnosed Carpal Tunnel Syndrome

Learn more about ATTR-CM throughout this brochure or visit SignsofATTR-CM.com
WHAT IS ATTR-CM?

Amyloidosis is a group of diseases in which certain proteins change shape, or “misfold,” and can build up in different parts of the body. When these misfolded transthyretin proteins build up in your heart, it may lead to ATTR-CM, a serious and often underdiagnosed condition that is associated with heart failure.

HERE’S WHAT HAPPENS IN THE BODY WITH ATTR-CM

Imagine your body as a factory, composed of many systems working together to maintain your health. The liver is just one part of that system, but it carries out many important jobs that can affect your entire body, including the heart.

The liver produces transthyretin, a transport protein that carries the hormone thyroxine and vitamin A (retinol) throughout the bloodstream.

When someone has ATTR-CM, either due to aging (wild-type) or an inherited genetic variant (hereditary), the protein becomes unstable and misfolds.

Over time, the misfolded proteins join together and build up in the body, including in the heart (causing the heart muscle to thicken and stiffen, eventually leading to heart failure).

WANT TO LEARN MORE?
Click here or scan the QR code to watch a video about how ATTR-CM affects the body. Using the camera on your smartphone, hold your device so that the QR code is visible on your screen. Your device will recognize the code and provide a notification that links to the video.
TYPES OF ATTR-CM
There are 2 types of ATTR-CM—wild-type and hereditary.

WILD-TYPE ATTR-CM (wtATTR)

- Associated with aging
- Most often affects white men over the age of 60
- May be the most common form of ATTR-CM

V122I almost exclusively affects African Americans

In the United States, the V122I mutation is found almost exclusively in individuals of African ancestry. Approximately 3-4% of African Americans in the U.S. are thought to be carriers of the mutation. However, not all individuals with the V122I mutation develop symptoms of hereditary ATTR-CM.

HEREDITARY ATTR-CM (hATTR)

- Caused by a change (or “mutation”) in one of your genes
- Passed down from a relative
- Affects both men and women, with symptoms beginning as early as 50 to 60 years old
- There are more than 120 known mutations that cause hATTR; the most common mutation in the United States is V122I
UNDERSTAND THE SIGNS AND SYMPTOMS OF ATTR-CM

Did you know that some seemingly unrelated signs and symptoms could be caused by ATTR-CM? Your body may be sending you a message.

SYMPTOMS OF ATTR-CM MAY INCLUDE:

- **FATIGUE** (excessive tiredness)
- **TENDON RUPTURE** (eg, biceps, Achilles tendon)
- **GASTROINTESTINAL ISSUES** (eg, diarrhea, constipation, nausea, or feeling full quickly)
- **SHOULDER, HIP, AND/OR KNEE PAIN**
- **SWELLING IN LOWER LEGS** (known as peripheral edema)
- **EYE DISORDERS** (eg, glaucoma)
- **SHORTNESS OF BREATH**
- **IRREGULAR HEARTBEAT** (known as cardiac arrhythmia)
- **BILATERAL CARPAL TUNNEL SYNDROME** (numbness, tingling, or pain in your fingers)
- **PAIN OR NUMBNESS IN LOWER BACK/LEG DUE TO NARROWING OF LOWER SPINE** (known as lumbar spinal stenosis)
- **DECREASED OR STRANGE TINGLING SENSATION/PAIN IN TOES/FEET** (known as peripheral neuropathy)

These examples are for illustrative purposes only. Signs and symptoms may vary from patient to patient.

While these signs and symptoms don’t necessarily indicate that you have ATTR-CM or another condition, any one or combination should be mentioned to your doctor.
If you have unresolved heart failure symptoms, ask your doctor about ATTR-CM as soon as possible.

TALKING TO YOUR DOCTOR
Advocating for yourself or a loved one with the disease can help you get many of the answers you need.

THE ROAD TO AN ATTR-CM DIAGNOSIS CAN BE COMPLEX AND FRUSTRATING

Awareness among patients, and even physicians, remains low, which results in ATTR-CM being underdiagnosed.

• Underdiagnosis and delayed diagnosis occur because the symptoms of ATTR-CM mimic those of other more common causes of heart failure
• Some ATTR-CM patients report visiting up to 5 different doctors before receiving the correct diagnosis

Ready to talk to your doctor about ATTR-CM but aren’t sure how to start the conversation? Click here or scan the QR code (using your phone’s camera) to create a custom doctor discussion guide—a useful tool to help get the conversation started.

“I was diagnosed with ATTR-CM 11 years after the presentation of my first symptom of carpal tunnel and 10 years after my second symptom of heart failure.”

— WALT, ATTR-CM patient, age 71
HOW ATTR-CM IS SUSPECTED

If your doctor suspects ATTR-CM based on clinical clues, they may conduct certain diagnostic tests for further evaluation.

INITIAL TESTS

Your doctor may first order tests to assess how your heart is working and look for signs of ATTR-CM. While none of these tests are typically used to confirm an ATTR-CM diagnosis, they can help your doctor learn more about your heart and determine the need for additional diagnostic testing.

**Electrocardiogram (ECG)**
- Reads electrical signals from your heart
- Can reveal conditions like irregular heartbeat (ie, atrial fibrillation), among other findings, that may be related to ATTR-CM

**Echocardiogram (ECHO)**
- Uses sound waves to create images of your heart
- Findings help determine the speed and direction of blood flow in the heart
- Findings associated with ATTR-CM include, but are not limited to, heart failure with preserved ejection fraction (HFpEF), which relates to the amount of blood that passes through the heart with each beat

**Cardiac magnetic resonance imaging (cardiac MRI)**
- Uses radio waves, magnets, and a computer to create images of your heart to look for abnormalities

Your doctor must also rule out another form of cardiac amyloidosis, known as light-chain amyloidosis (AL), using blood and urine tests. This is an important step, as AL amyloidosis and ATTR-CM are managed in different ways.
DIAGNOSING ATTR-CM

Once AL amyloidosis is ruled out, your doctor may order additional tests to help diagnose ATTR-CM.

DIAGNOSTIC TESTS

There are several tests that can help confirm a diagnosis of ATTR-CM or identify whether you or a loved one are at risk.

Nuclear scintigraphy—a noninvasive imaging test
- A small amount of dye is injected into your body
- After 1-3 hours, a special camera takes images of your body
- These images can help your doctor understand if transthyretin (TTR) amyloid fibrils are present in your heart
- Also referred to as a PYP (pyrophosphate) scan

Cardiac biopsy—samples taken from your heart muscle tissue
- Your cardiologist conducts the biopsy while you are awake
- If amyloid fibrils are found in the removed tissue sample, it is sent out to a lab
- The lab can help determine if they are TTR amyloid strands or not
- Your doctor may also do biopsies from other parts of your body—but a cardiac biopsy is more accurate to detect ATTR-CM

If you are diagnosed with ATTR-CM, genetic testing and counseling are recommended to determine whether you have the wild-type or hereditary form

Genetic Testing
- Will help confirm or rule out the hereditary form of ATTR-CM (hATTR)
- Determines whether family members are at risk, as the mutation that causes hATTR can be inherited
- Usually performed using blood or saliva samples
YOU ARE NOT ALONE

Resources are available to help if you or a loved one have been diagnosed with ATTR-CM.

Amyloidosis Support Groups
• Provides education through support meetings, live in 30 cities
• Provides education through virtual webinars, as well as support groups on Facebook

Amyloidosis Research Consortium
• Provides comprehensive support and information for patients
• Accelerates development of and access to new and innovative treatments
• Drives research that will have the greatest impact on patients

Amyloidosis Foundation
• Supports research for an earlier diagnosis
• Educates medical professionals
• Provides patients with a comprehensive range of services

amyloidossupport.org  arcl.org  amyloidosis.org

Don’t miss out on important updates!
Visit yourheartsmessage.com/resources to join our mailing list and stay informed about ATTR-CM.

Visit us on Facebook! Hear firsthand stories and connect with others impacted by ATTR-CM at Facebook.com/YourHeartsMessage.

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