#### Hereditary transthyretin amyloidosis (hATTR)

Hereditary transthyretin amyloidosis (hATTR) is an autosomal dominant disease. It is caused by one or more pathogenic variations in the TTR gene.

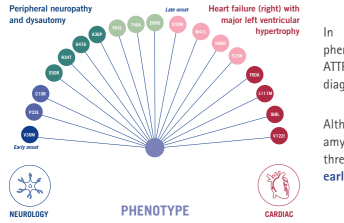
People with one or more pathogenic variations in the  $\Pi R$  gene produce an «abnormal» TTR protein throughout their lives. Amyloid deposits begin to form and then accumulate until they cause a chronic disease, mainly affecting the nerves and/or heart, and sometimes the kidneys, eyes and synovial tissues (tendons and ligaments).

Symptoms can appear at any time in early adulthood.

Phénotype de l'amylose à TTR Phenotype of TTR amy Ocular manifestation Conjunctival vesse Ð (JK) rinheral sensory-motor infection (due to neuropathy Length-dependent

#### To date, there are more than 120 known pathogenic variants.

Some favour neuropathy, others purely cardiac disease, or mixed forms, in some cases with renal and/or ocular involvement.



In clinical practice, the phenotypic variability of ATTR amyloidosis can make diagnosis complex.

Progressive progressio

Although ATTR rare. amvloidosis is a lifethreatening disease, and early diagnosis is vital.

# The most common forms of hATTR amyloidosis

Hereditary transthyretin amyloidosis is often referred to as hereditary amyloid neuropathy when the disease primarily affects the nerves - or familial amyloid cardiomyopathy when the disease primarily affects the heart.

#### Hereditary amyloid neuropathy

J.C Hereditary amyloid transthyretin neuropathy is a rare (500 cases recorded in France), systemic, autosomal dominant disease resulting from pathogenic variants in the transthyretin (TTR) protein gene. It affects adults only.

The most frequent variation observed in France in hereditary amyloid neuropathies is V30M (p.Val50Met). It is found in two-thirds of cases. However, other rarer variants such as S77Y and 1107V may be encountered.



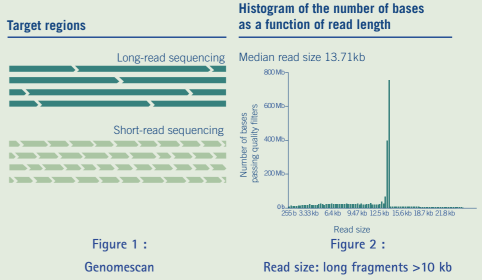
#### Familial amyloid cardiomyopathy

The prevalence of transthyretin-mutated cardiac amyloidosis is difficult to assess because the diversity of symptoms it causes makes it difficult to diagnose.

Certain transthyretin variants, which preferentially cause cardiac damage, are found in 0.4% to 3% of the population, depending on ethnic origin. The V122I variant (p.Val142IIe) is the most common worldwide.

The diagnosis of these transthyretin pathologies is based on two main steps:

- the identification of amyloid deposits :
  - for cardiomyopathy: cardiac ultrasound (to look for MHC), ECG and NTproBNP/Tn assay (prognostic value) and Tc99m bone scan (specific for TTR amyloidosis).
  - **b** for neuropathy: on nerve biopsy, immunostained for transthyretin
- the identification of an amyloidogenic variation in the TTR gene by genetic analysis.











# Long-read sequencing in the diagnosis of hATTR amyloidosis

Long-read sequencing or third-generation sequencing makes it possible to produce significantly longer reads than second-generation sequencing (short-read), i.e. several thousand (or even millions) of base pairs vs. a few hundred (Fig.1 and 2).

In hATTR amyloidosis, long-read sequencing offers a number of advantages, as it enables:

Sequencing of the entire TTR gene in a single analysis.

Save time and reduce diagnostic costs compared with a succession of targeted sequencing operations (exon by exon).

A diagnostic response within 4 weeks.

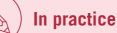
#### **Clinical benefits**

Identifying the responsible variation(s) makes it possible to:

- Confirm or rule out the diagnosis of hATTR amyloidosis:
  - Other conditions also present neuropathic symptoms similar to hereditary amyloid neuropathy, such as diabetic and alcoholic neuropathy, Charcot-Marie-Tooth disease and light chain amyloidosis.
  - From a cardiological point of view too, misdiagnosis is not unthinkable. hATTR should be excluded in patients with hypertrophic cardiomyopathy secondary to haemochromatosis, Fabry disease or cardiac sarcoidosis.
- Provide prognostic information on the pathology with regard to the variation detected.
- Initiate therapeutic management as early as possible: anti-amyloid treatment, symptomatic treatments if necessary, prevention and management of organ failure
- Offer genetic counselling and family screening (for adults):
  - ▶ to detect subjects at risk of developing the disease early and offer them regular monitoring and effective treatment as soon as symptoms start,
- or to reassure non-carriers once and for all.



- Over 120 known pathogenic variants in TTR amyloidosis.
- The age of onset of the disease, the type of symptoms, the prognosis and treatment depend on the variant responsible.
- Analysis of the TTR gene using long-read sequencing enables the entire gene to be studied in a single analysis, with results available within 4 weeks.
- The earlier the treatment is delivered, the more the progression of the disease can be delayed.



Test	Molecular study of the $TTR$ gene using next-generation sequencing
Test code	TTR
Délai	4 weeks
Sample	5 mL EDTA whole blood or extracted DNA
Storage and	Refrigerated

transport	Refrigerated
Technique	Next generation sequencing
Price	Contact us
Required documents	<ul> <li>B12-INTGB test request form</li> <li>Specific information sheet R76-INTGB available on www.eurofins-biomnis.com &gt; Test guide &gt; Test code: TTR</li> </ul>

# References

French National Protocol for Diagnosis and Care (Protocole National de Diagnostic et de Soins (PNDS)) - Cardiac amyloidosis. 2020-2021

French National Protocol for Diagnosis and Care (Protocole National de Diagnostic et de Soins (PNDS)) - Familial amyloid neuropathy. July 2022

Figure 1: https://www.genomescan.nl/long-read-sequencing-2/

Figure 2: Molecular study of the MEFV gene using Nanopore sequencing. Eurofins Biomnis.

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# For more information

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# Hereditary transthyretin amyloidosis

The contribution of long-read sequencingof the *TTR* gene for rapid and accurate diagnosis