Clinical Case Report

Late onset cardiomyopathy as presenting sign of ATTR A45G amyloidosis caused by a novel TTR mutation (p.A65G)

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Objective: The clinical description of a novel TTR gene mutation characterized by a late onset amyloid cardiomyopathy.

Methods and Results: A 78-year-old man of Dutch origin with recent surgery for bilateral carpal tunnel syndrome (CTS) was admitted to our hospital because of heart failure with preserved ejection fraction (55%). Cardiac ultrasound showed thickened biventricular walls, and cardiac magnetic resonance imaging also showed late gadolinium enhancement. Early signs of a polyneuropathy were found by neurophysiological testing. A few months later, his 72-year-old sister was admitted to an affiliated hospital because of heart failure caused by a restrictive cardiomyopathy. In both patients, a subcutaneous abdominal fat aspirate was stained with Congo red and DNA was analyzed by direct sequencing of exons 1 to 4 of the transthyretin (TTR) gene. Both fat aspirates revealed transthyretin-derived (ATTR) amyloid. 99mTc-diphosphonate scintigraphy further confirmed cardiac ATTR amyloidosis in the male patient. DNA analysis of both patients showed a novel TTR mutation c.194C > G that encodes for the gene product TTR A45G. The 56-year-old daughter of the male patient had the same mutation.

Conclusions: A novel amyloidogenic TTR mutation was found in a Dutch family. The clinical presentation of ATTR A45G amyloidosis in the affected family members was heart failure due to a late-onset cardiomyopathy. The systemic nature of this disease was reflected by bilateral CTS and by early signs of a polyneuropathy in the index patient.

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1. Introduction

The etiology of hereditary transthyretin-derived (ATTR) amyloidosis is the deposition of amyloid fibrils in the extracellular matrix due to mutations of the gene encoding transthyretin (TTR). More than 110 TTR mutations have been associated with hereditary ATTR amyloidosis [1,2]. Disease presentation, severity, and clinical course of ATTR amyloidosis are variable. Usually the disease starts with the clinical picture of a polyneuropathy accompanied by autonomic neuropathy, but sometimes, a cardiomyopathy is the presenting manifestation. In addition, albeit less frequent, other organs can be affected leading to carpal tunnel syndrome (CTS), vitreous opacities, or loss of renal function [1–3]. Clinical course and disease severity vary among different mutations [4]. Despite an initial monosymptomatic presentation, other organ involvement usually develops during disease progression. In the current report, we describe a Dutch family with a novel TTR mutation with late-onset cardiomyopathy, that is, not symptomatic before the sixth decade [5]. Although cardiac presentation was the prominent feature, a recent history of bilateral CTS and preclinical involvement of the peripheral nervous system were recognized in one of the two cases reflecting the systemic nature of the disease.

2. Patients and methods

A 78-year-old man and his 72-year-old sister, both known with atrial fibrillation (AF), presented with symptoms of heart failure at a hospital a couple of months after each other. Amyloidosis was considered a possible cause; consequently, a subcutaneous abdominal fat tissue aspirate was obtained that showed the presence of amyloid in both cases,
and subsequently, both cases were referred to our Amyloidosis Center of Expertise. A 56-year-old daughter of the male patient was referred later for clinical investigation.

Ample (more than 100 mg) subcutaneous abdominal fat tissue was aspirated using a 16-Gauge needle. Amyloid was extracted in 6-M guanidine hydrochloride and further analyzed as described [6]. The TTR concentration in fat tissue of the extracted amyloid was assessed immunochromatically using an indirect enzyme-linked immunosorbent assay [7] and also chemically by proteomics using a selected reaction-monitoring-mass spectrometry assay (SRM-MS; kindly provided by Paula Picotti and Paul Boersema, Zürich, Switzerland). The amount of Congo red-stained birefringent material was assessed semiquantitatively [8].

DNA was analyzed in all cases by direct sequencing of exons 1 to 4 of the TTR gene. The primary translation product is 20 amino acids longer than the mature TTR protein because the signal peptide and propeptides are included. This leads to amino acid position numbers of the translation product that are 20 higher than those of the corresponding mature protein [2].

A full clinical investigation to detect other organs affected by amyloid was offered to all patients. Due to the observational nature of this study, according to Dutch law, informed consent and permission of the local institutional review board were not required.

3. Results

3.1. Case 1

A 78-year-old man, known with AF for the last 2 years, presented with symptoms of heart failure at our hospital. Plasma levels of NT-proBNP and hs-Troponin T were 1877 ng/l (N=125) and 48 ng/l (N=14), respectively. One year before, he suffered from CTS in both hands, which was successfully treated by surgery. His electrocardiogram showed AF, low-voltage QRS complexes, and a left anterior fascicular block. A 24-h rhythm monitoring did not reveal other arrhythmias. Echocardiography showed concentric left ventricular wall hypertrophy and a left ventricular ejection fraction (EF) of 55%. Hypertrophy of the right ventricular wall (7 mm) was also observed. A multigated acquisition scan showed a left ventricular EF of 65% and a right ventricular EF of 44%. Cardiac magnetic resonance (CMR) imaging showed late gadolinium enhancement and a thickened interventricular septum (18 mm). Based on these clinical findings, amyloidosis was considered to be present. Easily accessible subcutaneous abdominal fat tissue was aspirated as screening biopsy for amyloidosis. The tissue was stained with Congo red, which indeed showed the presence (abundant, visual grade 4+) of amyloid (Fig. 1) [8]. No signs of immunoglobulin free light chain overproduction (serum kappa free light chain 13.1 mg/l (N=20) and lambda free light chain 13.0 mg/l (N=32)) nor other signs of a plasma cell dyscrasia were found in serum, urine, or bone marrow.

These findings made light chain-derived systemic AL amyloidosis very unlikely. The creatinine clearance was 24 ml/min, and no proteinuria was observed. 123I-MIBG scintigraphy showed a normal washout (34%) and low heart–mediastinum ratio (late uptake 1.16) indicative of cardiac denervation [10]. A fine needle subcutaneous abdominal fat aspirate yielded a TTR concentration of amyloid in fat of 658-ng/mg fat tissue (N=7.5 ng/mg) [6,7]. These findings indicated ATTR amyloidosis to be present. Wild-type ATTR amyloidosis was excluded because subsequent TTR gene analysis showed a novel mutation c.194C>G that encodes the primary translation product (p.A65G) and the mature protein TTR A45G [11]. SRM-MS proteomics confirmed the specific TTR mutation in the amyloid extracted from fat tissue. The patient was therefore diagnosed with hereditary ATTR A45G amyloidosis with prominent cardiac involvement and CTS. He had not noticed any symptoms of peripheral or autonomic neuropathy. Neurophysiological testing, however, revealed an early-stage length-dependent axonal sensory polyneuropathy in the feet of the patient. Quantitative sensory testing of the ankle also showed thin fiber neuropathy. He was treated with the TTR tetramer-stabilizing drug tafamidis for his polyneuropathy [12], and the clinical condition of both his heart and nerves seemed to stabilize during the next 2 years of follow-up.

3.2. Case 2

Family history of Case 1 revealed that the 72-year-old sister of the patient suffered from paroxysmal AF. She was admitted to an affiliated hospital because of heart failure caused by restrictive cardiomyopathy a few months after presentation of her brother. Upon further clinical investigation, she had no clear signs of CTS, polyneuropathy, or autonomic dysfunction. However, fat tissue analysis also showed amyloid (moderate, 3+) [8]. The TTR concentration of amyloid in fat was 122-ng/mg fat tissue (N=7.5 ng/mg) [6]. TTR gene analysis identified the same TTR mutation. Therefore, the sister was also diagnosed with hereditary ATTR A45G amyloidosis. She was not interested in further clinical investigations nor in disease monitoring during follow-up.

3.3. Case 3

The daughter of Case 1 presented at the age of 56 at the outpatient clinic with nonspecific complaints. TTR gene analysis showed the same genotype as her father and aunt. A full diagnostic work-up for amyloidosis was initiated: symptoms and signs of polyneuropathy and autonomic neuropathy were lacking, NT-proBNP was marginally elevated (145 ng/L) with a normal hs-Troponin T (10 ng/l). No cardiac uptake was visible on the 99mTc-diphosphonate bone scintigraphy, and fat...
tissue analysis did not show any amyloid deposits. Therefore, at that
time, amyloidosis could not be diagnosed in the daughter, and she
was scheduled for regular follow-up as a carrier of the mutation.

4. Discussion

A novel amyloidogenic TTR mutation, c.194C > G that encodes for the
primary translation product (p.A65G) and the mature protein TTR
A45G, was found in a Dutch family. This mutation was presented during
the International Symposium on Amyloidosis in Indianapolis in 2014
and subsequently included in the TTR mutations registry [2]. The clinical
picture of ATTR amyloidosis in the two affected family members was
foremost a late-onset cardiomyopathy [4,5,13]. 99mTc-diphosphonate
scintigraphy is currently considered useful in clinical practice and
more specific than ultrasound and CMR for noninvasive detection of
an ATTR amyloid cardiomyopathy [9].

Late-onset cardiomyopathy, frequently preceded by CTS, is typically
caused by wild-type ATTR amyloidosis [14]. Initially, this diagnosis was
considered in our index patient. However, some mutations causing he-
ereditary ATTR amyloidosis are characterized by late-onset cardiomyop-
athy, such as in Afro-Caribbean patients with ATTR V122I amyloidosis
who present with cardiac failure at median age 75 (range, 50–90)
years [15]. In the Italian population with different mutations, the medi-
an age at onset of exclusive cardiomyopathy in hereditary ATTR amy-
loidosis is 70 (62–75) years [4]. A German focus with ATTR V20I
amyloidosis with exclusive cardiomyopathy and occasionally CTS has
a median age at onset of 60 years [5]. Therefore, in an elderly man
with ATTR amyloid cardiomyopathy and CTS a search for a TTR mutation
is mandatory.

The TTR c.194C > G mutation is located at the end of exon 2 on the
TTR gene. A single base change, a cytosine nucleotide replaced by a gua-
nine nucleotide, is predicted to result in an amino acid change from ala-
nine to glycine. Our group reported in 2014 this novel mutation to the
Website concerning TTR mutations in hereditary amyloidosis [2]. Key
features of both cases were biventricular hypertrophy and AF. Neu-
rologic manifestations were also present, albeit less prominent. In 2016,
another case with this TTR (p.A65G) mutation and amyloid cardiomy-
opathy was reported brieﬂy [13]. These observations are in line with re-
ports on all three other mutations of the same amino acid 65 of the
TTR gene: (p.A65S), (p.A65T), and (p.A65D), in which the amino acid ala-
nine (A) has been replaced by serine (S), threonine (T), and aspartic
acid (D), respectively; therefore, all four mutations on this position 65
present as a cardiomyopathy [13,16–18]. However, neurologic sequelae
can be present and should be sought for. In particular, a history of CTS
— especially bilateral — in patients presenting with restrictive cardiomy-
opathy is a strong indicator that amyloidosis may be present [19].

By using the American College of Medical Genetics criteria for the interpre-
tation of sequence variants the current TTR c.194C > G mutation can be
classified as pathogenic [20].

Obvious limitations of this study are the small number of patients in
this family that were studied and the short follow-up period. Another
limitation is the absence of proof of actual presence of ATTR amyloid
in cardiac tissue. Only one case history of a patient with both AL and
ATTR amyloid simultaneously present in the heart has convincingly
been documented [21]. Besides, a clearly elevated serum lambda free
light chain was found in this case. The specificity of intense cardiac uptake on diphosphate scintigraphy is currently accepted as noninvasive evidence of the presence of ATTR amyloid in the myocardium in patients without an elevated free light chain in the blood [9]. Exclusion of AL amyloid is necessary but not sufficient for a diagnosis of ATTR amyloid. The conformation of both TTR and its particular mutation in ATTR amyloid by direct typing using SRM-MS is requisite for typing this amyloid as ATTR. Although the heart was not directly sampled, the imaging, the clinical scenario, and proof of ATTR amyloid in the fat tissue are sufficiently compelling.

Diuretic treatment is currently the only rational treatment option of heart failure with preserved ejection fraction (HFpEF) [22]. However, in ATTR amyloidosis the progression of polyneuropathy slows down after tetramer-stabilizing treatment by tafamidis and diffusinual in which the circulating TTR, a tetramer consisting of four identical molecules surrounding in the middle a tunnel for the transport of thyroxin, is protected against dissociation into highly amyloidogenic TTR dimers and monomers [12,23]. The possible effect of tafamidis on progression of ATTR amyloid cardiomyopathy is currently studied in a randomized clinical trial, the ATTR-ACT study [24]. A second trial in which the effect of siRNA on TTR production by the liver and thereby on progression of hereditary ATTR polyneuropathy is currently studied in the APOLLO trial [25]. A comparable ENDEAVOUR trial on hereditary ATTR cardiomyopathy has been stopped recently because of unexplained increased mortality in the treatment arm. The results of the ongoing two trials are expected in 2018. A favorable result might have big consequences for patients with systemic amyloidosis. In: Picken MM, Dogan A, Marmot MG, editors. Amyloid in Transthyretin Amyloidosis. Circulation 2016;133:2404–12.

5. Conclusion

The novel TTR c.194C > G mutation found in this family is clinically characterized by late-onset cardiomyopathy, CTS, and less prominent neurological involvement.

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References