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Mild clinical presentation in KLHL40-related Nemaline Myopathy (NEM 8)

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Abstract

Nemaline myopathies (NM) are clinically and genetically heterogeneous muscle diseases characterized by the presence of nemaline bodies (rods) in muscle fibers. Mutations in the *KLHL40* (kelch-like family member 40) gene (NEM 8) are common cause of severe/lethal nemaline myopathy. We report on an 8-year-old girl born to consanguineous Moroccan parents, who presented with hypotonia and poor sucking at birth, delayed motor development, and further mild difficulties in walking and fatigability. A muscle biopsy revealed the presence of nemaline bodies. *KLHL40* gene Sanger sequencing disclosed a never reported pathogenic homozygous mutation who resulted in absent KLHL40 protein expression in muscle. This further expand the phenotypical spectrum of KLHL40 related nemaline myopathy.

**Keywords:** Nemaline Myopathy; *KLHL40*
1. Introduction

Nemaline myopathy (NM) is a rare congenital muscle disorder associating hypotonia, muscle weakness, and often skeletal deformities with the presence of nemaline bodies (rods) in muscle fibers [1]. This disorder has a marked clinical variability, ranging from neonatal lethal to mild non-progressive forms with onset in childhood and adulthood. NM has been classified into six clinical categories according to the severity of the disease, the age of onset and the pattern of muscle weakness [2].

To date, eleven genes have been identified for NM: α-skeletal actin (ACTA1, MIM#161800) [3], muscle specific coflin (CFL2, MIM#610687) [4], nebulin (NEB, MIM#256030) [5], slow troponin T (TNNT1, MIM#605355) [6], β-tropomyosin (TPM2, MIM#609285) [7], slow α-trompomyosine (TPM3, MIM#609284) [8], kelch-repeat and BTB-[POZ]-domain containing 13 (KBTBD13, MIM#609273) [9] and kelch-like family member 40 (KLHL40, MIM#615340) [10] and member 41 (KLHL41, MIM#607701) [11], leiomodin 3 (LMOD3, MIM#616112) [12] and very recently myosin XVIIIB (MYO18B) [13].

Mutations in the KLHL40 gene (NEM 8) have been associated with severe/lethal NM [10]. KLHL40 belongs to the superfamily of kelch-repeat-containing proteins [14] and is crucial for myogenesis through regulation of E2F1-DP1 [15] and for skeletal muscle maintenance, binding to NEB and LMOD3.

Almost all (96%) of 32 affected individuals described in the seminal paper by Ravenscroft et al. had acute respiratory failure, swallowing problems. Foetal akinesia, hypokinesia and contractures were present at birth in more the three quarters of the cases. The average age at death was 5 months (n=14) [10]. Later on, Kawase et al. described a KLHL40 mutated patient with congenital locked-in state, who died at 4 years [17]. More recently, mutations in KLHL40 have been described in a 9-year old
female with severe NM patient and myastenic features, who surprisingly responded to acetylcolinesterase inhibitors [18].

Here we describe a mild NM myopathy case presenting a novel KLHL40 homozygous mutation.

2. Case Report

We present the case of a 8 year old girl third born to Moroccan consanguineous (the parents were first degree cousins) parents with no familial history for neuromuscular conditions. Of note congenital cataract was found in one of her two older sisters. Pregnancy was uneventful and she was born at 39 gestational weeks with a weight of 3200 grams (25th percentile), a length of 50 cm (50th percentile) and a cranial circumference of 39 cm (97th percentile). At three days of age she was admitted in a neonatal unit due to axial hypotonia and poor sucking. She was on nasogastric feeding until the 15th day, after which she was released on oral nutrition. Her evolution slowly improved. At 7 months, hypomimia and equinus deformity of the foot were noticed. The patient was able to support her head at 6 months. Walking was achieved at around 20 months.

At 3 years, she was able to run and ride a bike but had difficulties climbing the stairs. She had a waddling gait, deficit of the orbicularis oculi and drooling.

At five years, she had a good function of upper and lower limbs but all the activities were performed very slowly. The neck flexors and extensors were weak. At six years and a half, the Gowers sign was negative. The neck extensors were stronger than the flexors and inversely the knee extensors weaker than the flexors. From the respiratory point of view, multiple ear infections led to a mild hearing loss in the left
ear. At 7 years, the patient was complaining a lot about tiredness and she was only able to walk for 10 minutes. She was using a manual wheel chair for long distances. She had predominant axial weakness. Coughing was rather inefficient and the sniff nasal inspiratory pressure was low.

Last seen at 8 years and 2 months, she weighed 20 kg (5th percentile), measured 129 cm (50th percentile), and was of normal intelligence (no formal IQ test was performed) (Fig. 1). She walked 599 meters at the 6-minute walking test. She was able to climb some steps without the rail. The Gowers sign was negative. She had been under Tyrosine treatment 500mg b.i.d. for one year and a half, with mild and non-objective improvement; her mother found that she was doing “slightly better” with the treatment. A complete muscular testing was performed using the Medical Research Council 6-point score (MRC). Flexors and extensor of the hip, knee and foot, abductors and adductors of the hip on both sides were graded all at 4; with the exception of the right knee extensor (graded 5) and the hip adductors (graded 3) on both sides. The right upper limb strength was slightly higher for the elbow flexors, finger and wrist flexors and extensors (graded 5°) compared to the left side (graded 4). The neck extensors were graded 3 compared to the flexors (graded 2).

During the first months of life, an intensive research of inborn errors of metabolism was performed (serum amino acid chromatography, carnitine, acyl-carnitine, lactic acid, vacuolated peripheral blood lymphocytes, CDG syndrome, urinary amino acid and organic acid chromatography) and no specific abnormality was detected. The cerebral neonatal MRI showed T2 hyper intensity of the periventricular white matter with nonspecific high diffusion. The karyotype was XY and methylation analysis by Southern blotting for Prader Willi syndrome and CTG sequence repeats analysis in DMPK gene by PCR and TP-PCR on the two strands for Steinert myotonic dystrophy...
were negative. The whole body 3T MRI was performed at the age of 9. It showed no particular changes of the muscles. The T2 cartography of the muscular tissue of the tights and calves showed normal values.

A deltoid muscle biopsy performed at 3 years of age showed the presence of clusters of elongated protein inclusions found both in cytoplasmic and subsarcolemmal areas of around 30% of fibres, staining red with the Gomori thricrome (Fig. 2 A,B). The inclusions corresponded to nemaline bodies. Ultrastructural analysis confirmed the presence of nemaline bodies presenting the typical Z-disk electrondensity and net like organization. (Fig. 2 C,D).

NM gene panel Sanger sequencing of ACTA1, TPM2, TPM3, CFL2, TNNT1, KBTBD13, KLHL40, KLHL41 and LMOD3 revealed a homozygous KLHL40 gene c.1498C>T missense mutation leading to a p.Arg500Cys change at the protein level. The mutation was found only one time in the DIVAS database at an heterozygous level [19]. As the North African population was underrepresented in genetics databases, we screened an Arabic cohort present in the laboratory (100 individuals). The c.1498C>T transition was not found.

The p.Arg500Cys occurs in a conserved domain in the fourth Kelch Repeat (Fig 3 E) and is predicted to be deleterious according to prediction software (Polyphen,SIFT, CADD, Fathmm, PROVEAN) [20].

Based on KLHL40 PDB structure 4ASC, the Arg500 residue interacted with the Glu528 inside a beta sheet by ionic interaction (Fig 3 B). The foldX software predicts an important variation of free energy [21].
The segregation was confirmed in both parents who carried the c.1498C>T at heterozygous state. One sister was heterozygous carrier of the mutation. (Fig 3 A, C).

Western blot analysis performed using a rabbit polyclonal KLHL40 (KBTBD5) antibody from sigma (HPA024463) on protein extracted from skeletal muscle revealed absence of KLHL40 protein (Fig.2 E). RT-PCR analysis revealed that KLHL40 transcript was found overexpressed in the patient biopsy compared to a same age control biopsy (fig 3 D).

3. Discussion

We report the case of 8 year old girl with KLHL40 gene-related nemaline myopathy, carrying a new mutation, c.1498C>T. Her clinical phenotype is milder compared to those described in the literature.

Nemaline myopathy related to KLHL40 varies very much in severity from neonatal death to very rare cases of survival into adolescence.

The most frequent mutation in patients with KLHL40 NEM, c.1582G>A has been found in Japanese, Kurdish and Turkish persons. Ravenscroft et al. showed that c.1582G>A mutation suggests a milder phenotype compared to the cases where this mutation is absent. However, regardless of the genotype, the severity of the disease differs greatly, from death at 20 days to survival at adolescence. One explication could be given by modifying factors [10].
The screening of 129 probands with a milder NEM phenotype did not identify any 
*KLHL40* mutation, confirming that the mutations found by Ravenscroft et al. are 
probably restricted to severe cases [10].

Regarding the present mutation, it is noteworthy that it leads to increased transcript, 
but absent protein on the western blot. An explication could reside in a deep 
restructuration of the protein that expresses a new epitope not recognized by the 
antibody used in the Western blot. Another possibility is that the protein is either not 
produced, or it is rapidly degraded. We could assume that other proteins might 
possibly take over *KLHL40* function to compensate for its loss but this is yet to be 
determined. Indeed, complete absence of the protein is otherwise difficult to 
conciliate with the milder phenotype.

At the current time, there are insufficient MRI data to comment upon whether *KLHL40* 
mutations give rise to specific patterns of muscle involvement, as for *ACTA1* 
(proximal leg involvement) or *NEB* (predominant distal leg atrophy) [22].

A limitation of the present report is that it concerns a single case. However, *KLHL40* 
is a recognized cause of nemaline myopathy and other nemaline-myopathy genes 
have been ruled out. The mutation is predicted to be pathogenic, as it is absent from 
the healthy population. Finally, western blot analysis revealed the absence of 
*KLHL40* protein expression supporting the pathogenicity of c.1498C>T.

A recent report [18] demonstrated the dramatic effect of pyridostigmine in a patient 
with *KLHL40* severe phenotype. Pyridostigmine or other neuromuscular junction-
acting molecules were not tried in the present case for several reasons: the clinical 
features of our patient (myopathic facies with no ophthalmoplegia and lack of 
variability of muscular signs) did not determine us to start an anticholinergic
treatment; another reason was that our patient was already on Tyrosine and we tried to avoid multiple therapy in order to assess Tyrosine efficacy. It is likely that other missense mutations in the KLHL40 will be associated with such milder phenotype in the coming years. Phenotype-genotype relation are important to recognize, not only to avoid diagnosis misleading, but also to adequately inform the parents at the time of diagnosis, or in case of prenatal counselling.

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**References**


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**Figure legends**

**Figure 1** Clinical presentation at 8 years: slender phenotype, hypomimia (A), scapular winging (B), facial weakness (C)

**Figure 2** Muscle biopsy from biceps brachii showing rod clusters (nemaline bodies) in the sub sarcolemmal region on trichrome Gomori (A); scarce rods in the centre of the muscular fibre on trichrome Gomori (B); electronic microscopy confirmed the presence of nemaline bodies in the sub sarcolemmal region (C-D). Western blot performed on skeletal muscle protein extracts by routine methods using the anti-KLHL40 (KBTBD5) form Sigma. A 69–53 KDa band corresponding to KLHL40 in protein extracts from an age-matched control was revealed, while no signal was detectable in the patient’s biopsy. Myosin heavy chain was used as a loading control (E).

**Figure 3** (A) Pedigree of the proband’s family and homozygous for the missense mutation (T/T). Unaffected parents and one sibling were both carriers (C/T) and one sibling wasn’t bearer of the mutation (C/C); (B, upper part) Structural modelling of the p.Arg500Cys mutation. The crystal structures of the B kelch domain of KLHL40. The fourth Kelch repeat is in blue. The mutation location is represented by a yellow star. (B, lower part) Focus of the Arg500 residu. The Arginine 500 (Arg500) interacts with the Glutamine 528. Hydrogen bonds are represented in yellow doted lines; (C) Sequence chromatographs of KLHL40 fourth exon of the different family genotype. The mutation location is framed. +/+ means homozygous for the C (or reference nucleotide), +/- means heterozygous for the mutation (T) and -/- homozygous for the mutation; (D) Quantitative expression of the KLHL40 transcript extracted from muscle.
biopsy. CB is a control biopsy from a control patient and PB the patient biopsy. In PB, KLHL40 is overexpressed compared to CB KLHL40 expression; (E) Alignment of the corresponding region of KLHL40 containing the index case’s mutation. The Arginine 500 is conserved between various species.