Neuroacanthocytosis Syndromes in an Italian Cohort: Clinical Spectrum, High Genetic Variability and Muscle Involvement

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Abstract: Neuroacanthocytosis (NA) syndromes are a group of genetically defined diseases characterized by the association of red blood cell acanthocytosis, progressive degeneration of the basal ganglia and neuromuscular features with characteristic persistent hyperCKemia. The main NA syndromes include autosomal recessive chorea-acanthocytosis (ChAc) and X-linked McLeod syndrome (MLS). A series of Italian patients selected through a multicenter study for these specific neurological phenotypes underwent DNA sequencing of the VPS13A and XK genes to search for causative mutations. Where it has been possible, muscle biopsies were obtained and thoroughly investigated with histochemical assays. A total of nine patients from five different families were diagnosed with ChAc and had mostly biallelic changes in the VPS13A gene (three nonsense, two frameshift, three splicing), while three patients from a single X-linked family were diagnosed with McLeod syndrome and had a deletion in the XK gene. Despite a very low incidence (only one thousand cases of ChAc and a few hundred MLS cases reported worldwide), none of the 8 VPS13A variants identified in our patients is shared by two families, suggesting the high genetic variability of ChAc in the Italian population. In our series, in line with epidemiological data, McLeod syndrome occurs less frequently than ChAc, although it can be easily suspected because of its X-linked mode of inheritance. Finally, histochemical studies strongly suggest that muscle pathology is not simply secondary to the axonal neuropathy, frequently seen in these patients, but primary myopathic alterations can be detected in both NA syndromes.
Keywords: neuroacanthocytosis syndromes; chorea-acanthocytosis; McLeod syndrome; VPS13A gene; XK gene

1. Introduction

Chorea-acanthocytosis (ChAc, OMIM #200150) is a rare hereditary neurodegenerative condition, characterized by the young adult-onset of involuntary choreiform movements and red blood cells (RBCs) acanthocytosis. The phenotypic spectrum is relatively wide and variable, encompassing psychiatric symptoms, cognitive impairment, peripheral neuropathy, myopathy and epilepsy [1]. Besides chorea, other motor manifestations may include facial and oromandibular dystonia, tics, parkinsonism and postural tremor. Although the misshaped RBCs are considered the diagnostic hallmark, the acanthocytes can sometimes be absent or even make a late appearance in the course of the disease. Other common laboratory findings in ChAc include increased levels of creatine kinase and liver enzymes. Together with the X-linked McLeod syndrome (MLS, OMIM #300842), due to loss-of-function (LOF) variants of the XK gene (OMIM *314850), ChAc accounts for the “core” neuroacanthocytosis (NA) syndromes, defined by the combination of RBC acanthocytosis and basal ganglia-related neurological disorders [1,2]. ChAc represents one of the most common differential diagnoses of Huntington’s disease (HD). The inheritance pattern of ChAc is autosomal recessive and patients are either homozygous or compound heterozygous for LOF variants in the VPS13A gene (OMIM *605978) encoding chorein, a large protein involved in the intracellular vesicle trafficking [1]. Mammalian VPS13A belongs to a small gene family that codes for ubiquitously expressed proteins derived by the duplication of an ancestral yeast gene [3]; pathogenic variants in the other genes of this family underlie other neurologic conditions such as Cohen syndrome (VPS13B), early-onset autosomal recessive Parkinson’s disease (VPS13C) and spinocerebellar ataxia with saccadic intrusions (VPS13D). Although ChAc inheritance is generally accepted as autosomal recessive, patients with possible autosomal dominant ChAc have occasionally been reported [4–7] but remain questionable in the pre-exome sequencing era. Indeed, a second VPS13A variant may initially go unnoticed, as in the affected siblings reported by Saiki et al. [4] where a second VPS13A variant was subsequently identified [8].

We describe here a series of Italian patients with NA syndromes recruited as part of a multicenter study, that were screened for pathogenic variants in the VPS13A and XK genes. In a total of six families, we identified eight different VPS13A variants, mostly unreported, and a single exon deletion of the XK gene. The clinical description, including detailed muscular pathology, laboratory findings and imaging features are reported.

2. Materials and Methods

2.1. Patient Series

The patients with NA syndromes described here come from clinical–diagnostic evaluations carried out in the neurology departments of three different University hospitals in Italy. All examiners used commonly adopted scales according to shared criteria to assess the core phenotype and disability score i.e. the Unified Huntington’s Disease Rating Scale (UHDRS), the Montreal Cognitive Assessment (MoCA), the Mini Mental State Evaluation (MMSE), and the Manual Muscle Test (MMT)-Medical Research Council (MRC) grading scale. With the exception of 2 patients, all other cases described here were not previously reported. This study was conducted according to the Declaration of Helsinki and written informed consent was obtained from all participants.

2.2. Molecular Analysis

The molecular diagnostics of probands, and other affected or unaffected family members, were performed between 2016 and 2018 in the same laboratory. Total DNA was isolated from peripheral blood using a MagPurix automatic extractor. All exons and
flanking regions of the \textit{VPS13A} and \textit{XK} genes were amplified from genomic DNA and customized primers (listed in the Supplementary Material). Amplicons were purified using Exo-Sap, labeled using a BigDye Terminator v3.1 Cycle sequencing kit (Applied Biosystem, Foster City, CA, USA) and directly sequenced on an ABI 3500 Genetic Analyzer (Applied Biosystem, Foster City, CA, USA). Total RNA was purified from blood and was reversely transcribed using the First Strand cDNA Synthesis Kit (Roche, Hamburg, Germany) according to the manufacturer’s random primer protocol. The effect of the \textit{VPS13A} intronic variant on splicing in family B was examined by RT-PCR using gene specific primers and Sanger sequencing. Primer sequences have been reported in the Supplementary Material and PCR conditions are available upon request. The identification of exon 1 deletion in the \textit{XK} gene in family F was obtained by quantitative real-time PCR (qPCR) in a MicroAmp® optical 96-well plate using exon specific oligonucleotide primers and SYBR Green mastermix (ThermoFisher, Waltham, MA, USA). The relative quantification of the copy number was performed according to the comparative method (2$^{-\Delta\Delta Ct}$) [9]. The mean result of three independent experiments run in triplicate was compared with that of normal controls whose expression was arbitrarily attributed the value of 1. Western blotting was performed in some patients on blood in order to detect the VPS13A protein using the anti-chor1 antibody as described previously [10].

2.3. Muscle Studies  
2.3.1. Chorea-Acanthocytosis  
Open muscle biopsies on patients from two different families with ChAC were performed at the Department of Neurology of the University of Campania “Luigi Vanvitelli”: patient II:4 from family A had a biopsy on the vastus lateralis of the right quadriceps, while patients II:2 and II:3 from family D were biopsied on the left deltoid. Muscle tissues were snap-frozen in isopentane, pre-cooled in liquid nitrogen and stored at $-260\,^\circ\text{C}$ until sectioning. 7 and 10-µm-thick serial cryo-sections were used for histology and histochemistry and stainings included: hematoxylin eosin (H&E), Gomori trichrome, periodic acid Schiff (PAS), Sudan III, adenosine triphosphatase (ATPase) at pH 9.6, 4.6 and 4.3, AP (acid phosphatase) and oxidative enzymes such as nicotinamide adenine dinucleotide-tetrazolium reductase (NADH-TR), succinate dehydrogenase (SDH) and cytochrome oxidase (COX). Qualitative and quantitative light microscopy studies were carried out on muscle slices using a Nikon Eclipse Ni® photomicroscope equipped with the NIS Elements F4.30.00 image analyses system.

2.3.2. McLeod Syndrome  
Following the same protocol described above, muscle studies were also performed on three individuals affected by McLeod syndrome: F-III:4 had biopsy on the right peroneus brevis at the Neurology Division of the “Federico II” University of Naples, while patients F-III:5 and F-III:9 were biopsied on the right quadriceps, respectively, at the Department of Neurology of University of Campania “Luigi Vanvitelli” and at the Neurology Unit of Padua University.

3. Results  
Table 1 summarizes the clinical data of all 12 patients belonging to the six families (A–F). Individual patients’ histories are described in detail in the Supplementary Material, while muscle studies performed on patients A-II:4, D-II:2, D-II:3, F-III:4, F-III:5 and F-III:9 are reported in a dedicated section.
Table 1. Main clinical, laboratory and histological features. Acanthocytes: percentage of acanthocytes on peripheral blood smear; Serum CK level: normal range 60–190 U/L; OCD: obsessive-compulsive disorder; CN: caudate nuclei; LL: lower limbs; MUP: motor unit potential; ChAc: chorea-acanthocytosis; MLS: X-linked McLeod syndrome; n.a: not available. * previously reported by Peluso et al. [11].

|   | A-II:4 (ChAc) | A-II:5 (ChAc) | B-II:4 (ChAc) | B-II:5 (ChAc) | C-II:4 (ChAc) | C-II:5 (ChAc) | D-II:2 (ChAc) | D-II:3 * (ChAc) | E-II:2 * (ChAc) | F-III:4 (MLS) | F-III:5 (MLS) | F-III:9 (MLS) |
|---|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|
| Age at onset | 6 24 27 25 11 20 40 30 28 20 | 39 25 36 30 48 41 52 (dead) 46 53 59 (dead) 63 45 (dead) |   |   |   |   |   |   |   |   |   |   |
| Onset Symptoms | Behavioral disorder | Anxiety disorder | Orofacial tics | Behavioral Disorder | Tics, seizures | Psychosis, OCD | Feeding dystonia, muscle weakness | Seizures | Mood and behavioral disorders | Muscle weakness | Seizures | Behavioral disorder |
| Acanthocytes (%) | 7.3–19.2 | 8.3 | 8.6 | 3.1 | 16.4 | 28 | 17 | 13 | 40 | 9–10 | 18 | 5–7 |
| Serum CK level (U/L) | 600 | 1971 | 489–948 | 5564 | 872–3435 | 1180 | 3000 | 1522 | 1500 | 2500 | 1000 | 1000–8000 |
| Choreic movements (onset age) | YES (15) | No | YES (29) | No | YES (30) | YES (30) | YES (42) | No | No | YES (50) | YES (59) | YES (45) |
| other movement disorders | Feeding dystonia, Akathisia, Tics | Tics | Tics | Facial dyskinesia, dystonia | Dystonia, bradykinesia, postural tremor | Feeding dystonia | No | Bradykinesia, tremor, motor slowing, dystonia | No | Buccal stereotypes | Oro-facial dystonia |
| Seizures (onset age) | YES (23) | No | No | YES (25) | YES (11) | No | No | YES (30) | No | No | YES (38) | YES (40) |
| Psychiatric symptoms (onset age) | YES (6) | YES (23) | No | YES (26) | No | YES (20) | No | No | YES (28) | No | No | YES (20) |
| Brain MRI anomalies | Atrophy of CN and hippocampus | No | No | Left lateral ventricular enlargement | Ventricular enlargement, calcific meningioma | Ventricular enlargement, arachnoid cyst | Cerebral cortex and CN atrophy | Atrophy of CN and hippocampus | Atrophy of CN | No | No | Atrophy of CN |
| Axonal polyneuropathy | Sensorimotor | Sensory | Sensory | Sensory | n.a. | Sensory | Sensorimotor | Sensory | No | Sensorimotor | Sensorimotor | Sensorimotor |
| EMG pattern | A-II:4 (ChAc) | A-II:5 (ChAc) | B-II:3 (ChAc) | B-II:5 (ChAc) | C-II:3 (ChAc) | C-II:4 (ChAc) | D-II:2 (ChAc) | D-II:3* (ChAc) | E-II:2* (ChAc) | F-III:4 (MLS) | F-III:5 (MLS) | F-III:9 (MLS) |
|-------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|
| Chronic denervation with myopathic MUP | Normal | Myopathic MUP | n.a. | n.a. | n.a. | Active and chronic denervation | Normal | Myopathic MUP with chronic denervation | Normal | Myopathic MUP with chronic denervation | Myopathic MUP with chronic denervation |
| Muscle weakness (onset age) | Lower limb-girdle (35) | No | Lower limb-girdle (29) | No | Lower limb-girdle (37) | Lower limb-girdle (38) | Lower limb-girdle (39) | Upper and lower limb girdle (40) | Upper and lower limb girdle (45) | Upper and lower limb girdle (40) |
| other muscular signs and symptoms | No | No | n.a. | n.a. | n.a. | Fasciculations, cramps, muscle atrophy | Fasciculations, cramps muscle atrophy | No | Cramps at LL | Cramps at LL |
| Muscle biopsy | Type-grouping, multinicrome fibers | n.a. | n.a. | n.a. | n.a. | Bizarre fibers, atrophic fascicles | Type-grouping, core fibers | n.a. | Fiber size variability, rimmed vacuoles | Fiber size variability, inflammatory infiltrates | Fiber size variability, increased central nuclei |
| Cardiac anomalies (onset age) | No | No | paroxysmic tachycardia (7) | No | paroxysmic atrial fibrillation (36) | mild conductive anomalies (39) | No | No | No | YES (57) | No | No |
Table 2 lists the VPS13A variants identified in the first five families (A–E) next to their frequency in the Genome Aggregation Database (gnomAD) and the expected protein changes.

| Family | CDNA Variant | Exon | Allele Frequency (gnomAD) | Variant Type | Protein Change | Previously Reported in |
|--------|--------------|------|---------------------------|--------------|----------------|------------------------|
| A      | c.3817C>T (pat) | 34   | 1 / 251198                | Nonsense     | p.R1273*       | NR                     |
|        | c.1114_1115del (mat) | 13   | 1 / 250630               | Frameshift deletion | p.K372Vfs*4 | [8]                     |
| B      | c.3339+4_3339+10delinsTATAGCTGTATATAAATTATTTAA | IVS31 | NF | Splice-site indel | ? | NR                     |
| C      | c.1078C>T     | 13   | 3 / 25076                | Nonsense     | p.Q360* | [8,12–14] |
|        | c.7867C>T     | 56   | 1 / 251190               | Nonsense     | p.R2623* | [8,12–14] |
| D      | c.2512+2T>G   | IVS24 | NF | Splice-site substitution | ? | NR                     |
| E      | c.7736_7739del | 55   | 5 / 251030               | Frameshift deletion | p.R2579Nfs*26 | NR                     |
|        | c.2825-10T>G  | IVS26 | NF | Splice-site substitution | ? | NR                     |

3.1. Family A (Chorea-Acanthocytosis)

This pedigree (Figure 1A), originating from two Vesuvian territories, includes two affected individuals. Molecular analysis of the VPS13A gene in the proband A-II:4 confirmed the ChAc diagnosis and revealed the presence of three variants: c.3817C>T, c.1114_1115delAA and c.3592A>C; the last two were found to be in cis on the maternal copy of the gene (Supplementary Figure S1). The first variant (inherited from the father) introduces a stop codon in position 1273 (p.R1273*) and the second (inherited from the mother) is a frameshift variant disrupting the protein after residue 372 (p.K372Vfs*4). While these two variants are both pathogenic and clearly account for the patients’ phenotype, a third variant (c.3592A>C) has been co-inherited with the maternal frameshift, causing a likely benign missense change (p.K1198Q). The same variants have been detected in the younger brother (A-II:5) while their older sister (A-II:3) carries only the paternal c.3817C>T (p.R1273*) nonsense variant.

3.2. Family B (Chorea-Acanthocytosis)

This consanguineous pedigree (Figure 1B) originates from a town near Brindisi, in the south-eastern tip of Italy, and the parents are second cousins. Molecular analysis eventually revealed a homozygous VPS13A variant in both affected brothers B-II:3 and B-II:5 (c.3339+4_3339+10delinsTATAGCTGTATATAAATTATTTAA). As shown in Supplementary Figure S2, this insertion–deletion, close to the splice donor site in intron 31, alters the canonical splicing of VPS13A mRNA, likely resulting in protein loss-of-function. The two other brothers (II:2 and II:4) are both heterozygous carriers, like their mother I:2. The older sister II:1 turned out to be homozygous for the wild-type allele, while the father I:1 (an obligate carrier) declined genetic testing.

3.3. Family C (Chorea-Acanthocytosis)

This non-consanguineous family has two probands, C-II:3 and C-II:4 (Figure 1C). Only in 2017, at the age of 41, molecular analysis of VPS13A was eventually performed on C-II:4, revealing a compound heterozygosity for two nonsense variants c.1078C>T (p.Q360*) and c.7867C>T (p.R2623*). Sequences of the two variants are shown in Supplementary.
Figure S3. Shortly after his younger brother, C-II:3 also received molecular confirmation of his ChAc diagnosis and was found to carry the same two nonsense variants. Two other unaffected brothers (II:2 and II:5) have not been tested yet.

Figure 1. Pedigrees of the five choreo-acanthocytosis families (A–E), all with a clear recessive mode of inheritance. However, consanguinity was confirmed only for pedigree B (parents are second-degree cousins) and suspected for pedigree D (both parents carry the same VPS13A variant). Affected individuals are shaded in black. Family members that were sequenced are indicated by a dash on top of their symbol.

3.4. Family D (Chorea-Acanthocytosis)

This family has two affected brothers (II:2 and II:3) and originates from Herculaneum, south-east of Naples, close to Mount Vesuvius (Figure 1D). Only the two affected brothers have been tested and the proband (D-II:2) has recently passed away. Although no parental consanguinity had been reported, VPS13A sequencing revealed a homozygous splicing variant (c.2512+2T>G) that abolishes the donor site of intron 24 Supplementary Figure S4). Genetic testing of the younger brother D-II:3 confirmed the presence of the same homozygous VPS13A variant.

3.5. Family E (Chorea-Acanthocytosis)

The proband (E-II:2) is the second son of a non-consanguineous marriage (Figure 1E). This patient, along with D-II:3, was previously reported by Peluso et al. [11], who made a biochemical diagnosis of ChAc since Western blotting had proven the absence of chorein, the VPS13A protein. Eventually VPS13A sequencing revealed that patient E-II:2 is a compound heterozygote for two different variants, namely, a 4-bp deletion (c.7736_7739del) causing a frameshift in exon 55 and an intronic substitution close to the acceptor site of intron 26 (c.2825-10T>G). Disruption of normal mRNA splicing has been confirmed with cDNA sequencing of the relevant VPS13A exons (Supplementary Figure S5).
3.6. Family F (McLeod Syndrome)

This large pedigree (Figure 2) originates from the metropolitan area of Venice and all patients but the proband F-III:5 are now dead. The familial history clearly suggested X-linked inheritance and, in fact, the absence of the Kell antigen on red blood cells confirmed this suspicion. Genetic testing eventually proved that proband III:5 was positive for a hemizygous deletion of exon 1 of the \( XK \) gene. As shown in the pedigree, four other individuals have been reported to be affected by family members and died before DNA analysis was available. However, III:4 and III:9 were subjected to muscle biopsy as described in the following section.

![Figure 2. Pedigree of family F is compatible with X-linked inheritance. The molecular test of III:5 indirectly confirmed the diagnosis of McLeod syndrome for the other patients (I:1, I:2, III:4, and III:9) who had already died. Obligate carriers are indicated with a small black circle within their symbol.](image)

3.7. Muscle Studies

Muscle biopsies were performed in three patients with ChAc and in three individuals of the MLS family; the main histological findings are illustrated in Figure 3.

3.7.1. Chorea-Acanthocytosis

In patient A-II:4 (Figure 3A–C) the muscle biopsy disclosed predominantly neurogenic alterations, with fiber caliber variability due to small groups of atrophic angulated fibers, next to some hypertrophic type I and type II fibers at myosinATPase pH 9.4; rare unstructured core fibers were also detected, suggesting chronic denervation. Muscle specimens obtained from the two affected siblings of family D showed a significantly different degree of muscle neurogenic degeneration. In patient D-II:2 (Figure 3D–F) a severe variability of fiber size was observed due to large groups of atrophic fibers (\( \bar{d} < 20 \mu m \)) intermingled with several giant fibers (\( \bar{d} > 120 \mu m \)). Splitting phenomena and the severe disruption of myofibrillar architecture in the form of whorled and bizarre-shaped fibers were also seen with oxidative enzymes staining (Figure 3E–F). The biopsy of Patient D-II:3 (Figure 3G–I) showed a histological picture similar to A-II:4, characterized by a slight variability in fiber size, atrophy of both type I and II fibers, and fiber-type grouping phenomena.
Figure 3. Histological findings on muscle biopsy. The first column shows images of muscle stained with haematoxylin-eosin (H&E), the second with succinate dehydrogenase (SDH) and the third with myosin ATPase at pH 9.4. (A–C) Case A-II:4 affected by ChAc: (A) moderate fiber size variability with some hypo-atrophic angulated fibers (black arrow) next to large hypertrophic fibers; (B) the great majority of type 1 fibers display multi-minicores and some poorly structured cores (white head-arrow); (C) atrophic fibers are organized in small-medium groups (type-grouping phenomena). (D–F) Case D-II:2 affected by ChAc: (D) very severe fiber size variability with numerous atrophic angulated fibers and atrophic fascicles, next to giant hypertrophic round fibers (black arrow); (E) severe myofibrillar architecture disruption with bizarre fiber shape, vortex fibers and presence of unstructured cores in most fibers (black arrow); (F) disruption of muscle architecture where fiber types are hardly distinguishable. (G–I) Case D-II:3 affected by ChAc: (G) poor fiber size variability with some hypotrophic fibers (black arrow); (H) rare structured core fibers (white arrowhead); (I) fibers grouping according to their histochemical type thus suggesting central denervation and reinnervation. (L–N) Case F-III:5 affected by McLeod syndrome: (L) fiber size variability with numerous clumps of naked nuclei (black arrow) and an infiltrate of inflammatory cells (white head-arrow); (M) numerous moth-eaten fibers (black stars); (N) some type grouping phenomena.

3.7.2. McLeod Syndrome

The two brothers (F-III:4 and F-III:5) displayed diffuse neuropathic muscle changes with a modest increase in endomysial connective tissue. Moreover, in case F-III:5 (Figure 3L–N), diffuse inflammatory infiltrates were evident along with some moth-eaten and core-like
fibers, while in patient F-III:4, a single fiber with a “rimmed vacuole” was detected (pictures not available). Finally, in the biopsy of patient F-III:9 (pictures not available), myopathic changes with a moderate variability in muscle fiber size were detected, consisting in prevalence of type I and IIC fibers and increased central nuclei.

4. Discussion

Here, we report 12 Italian patients with a molecular diagnosis of NA, nine belonging to five ChAC families (carrying bi-allelic variants of the VPS13A gene) and three belonging to the same family with McLeod syndrome (harboring a deletion of exon 1 of the X-linked XK gene). The clinical history of these patients, summarized in Table 1 and detailed in Supplementary Material (Table S1 and Table S2), allowed us to highlight some aspects of the NA syndromes. Due to the small number of patients with MLS (Family F), a clinical comparison with ChAc patients (Families A–E) does not add much to what has already been published [15,16]. As already reported, also in our series the ChAc phenotype is variable among patients, even within the same family: due to the high variability and low prevalence, NA is underreported and diagnosed with a clinically significant delay [17,18]. Among the core symptoms, tics and choreic movements, with a prevalence of orofacial dyskinesia, stand out. This is a first major phenotypic difference with the most frequent form of hereditary chorea, i.e., Huntington’s disease (HD), in which choreic movements mainly affect limbs at onset; another distinguishing feature is the occurrence of epilepsy in ChAc (even at an early stage), while epilepsy is rare and late in typical adult-onset HD patients. On the other hand, psychiatric symptoms are as frequent as those found in other hereditary choreas.

Muscle pathology of the NA syndromes is poorly understood, however, a distinction between ChAc and MLS must be made as they are two separate conditions. In fact, a neurogenic pattern at muscle biopsy was reported in most ChAc patients, though minor myopathic changes were observed in few cases [19,20]. Conversely, a primary myopathic pattern was reported in the MLS series [20]. Actually, the pathogenesis of muscle alterations in ChAc is still a matter of debate. Indeed, bizarre fibers, such as nucleus fibers, can be found in both dystrophic process and chronic denervation [21]. Whereas some studies focused on neurogenic muscle wasting in NA, little is known about the mechanisms underlying primary myopathy. Conformational and functional anomalies of tTGase products were proposed to alter the deformability of both erythrocytes and muscle membrane in ChAc [22,23]. Saiki et al. [24] proved the loss of chorein expression in the skeletal muscle of ChAc patients and suggested that this protein plays a role in mitochondrial activity. Since VPS13A is ubiquitously expressed, mutated chorein could be detected in all tissues of ChAc patients, including the skeletal muscle [25]. Undoubtedly, the pattern of fascicular atrophy found in patient D-II:2 (Figure 3D–F) allowed us to hypothesize that the second motor neuron body or the motor nerve roots may be the primary site of pathogenic events in ChAc, at least in some patients. However, the detection of whorled and bizarre shaped fibers (Figure 3E–F) suggests a concomitant primary myopathic process: as far as we know, these peculiar alterations have not been previously reported in ChAc patients. Furthermore, we note that muscle changes were very severe in the biopsy of patient D-II:2 (Figure 3D–F) compared to his brother D-II:3 (Figure 3G–I); although the two brothers share the same VPS13A splicing variant, this phenotypic heterogeneity cannot be explained only by disease duration, but genetic and/or epigenetic factors might be involved. The mixed neurogenic and myopathic pattern of muscle pathology in ChAc, already described in the literature, is further confirmed by our findings in A-II:2 (Figure 3A–C).

Let us now consider the other NA form, i.e., MLS, which was initially defined as a benign X-linked myopathy with acanthocytes [26], although Hewer et al. [27] claimed that it is neither benign nor a pure myopathy. Since a dystrophic pattern of myopathy was reported in some rare cases [28], an involvement of dystrophin (whose gene is 4 Mb upstream of XK on the X chromosome) was sought but not confirmed [29,30]. Immunocytochemical studies suggested that lack of XK expression in MLS may destabilize the normal muscle
structure and function, but the protein role remains unclear [31]. The 444-amino acid long XK protein has 10 transmembrane domains, it belongs to the 4.1R multiprotein complex of the red blood cell membranes and is joined to the Kell protein by a disulphide bond [32]. The deletion of exon 1 of the XK gene identified in our family F (Figure 2), which had been reported at least once before [33], is expected to completely abolish the presence of the protein in all cells of hemizygous males. Finally, the clinical and muscular phenotype (Figure 3L–N) of family F confirms that MLS is a mixed neuropathic and myopathic condition, with variable severity of CNS involvement. Based on our findings and those of the literature, histological signs of primary myopathy along with less severe neurogenic changes should be considered more suggestive of MLS rather than of ChAc.

We should now focus on the high genetic variability of VPS13A variants identified in our ChAc families (Figure 1), all of Italian origin: it is worth noting that, differently to what has been reported in the Japanese population [8,34], our patients do not present with recurrent variants (Table 2). Furthermore, the two most frequent Japanese variants (p.R1471* and p.V2738Afs*5), present in more than half of the patients [8,34], are absent in our series. In agreement with previous reports [34], where missense pathogenic variants are exceedingly rare, we also found three nonsense, two frameshift and three splice-site variants in our nine ChAc patients (Table 2). Only one missense variant (p.K1198Q) has been detected in Family A, but it is predicted to be likely benign and, most importantly, it was found in cis with the pathogenic frameshift variant (p.K372Vfs*4).

As shown in Table 2, only two of the listed VPS13A variants were previously reported in ChAc patients; the first such variant (c.7867C>T) introduces a stop codon truncating chor ein at arginine 2623 (p.R2623*). This variant was first identified by Dobson-Stone et al. [12] in the original Kentucky family, clinically described by Critchley et al. [35] in 1968 and was afterwards reported in several independent patients [8,13,14]. The other previously reported variant is a 2-bp deletion (c.1114_1115del) causing an early frameshift (p.K372Vfs*4), reported just once in the extensive series of Tomiyasu et al. [8]; interestingly, although they analyzed patients from 11 countries, the only patient carrying this variant (in homozygosity) was Italian. The c.1114_1115del frameshift might be therefore an Italian variant, also found in our family A from Naples where it associates in compound heterozygosity with the c.3817C>T nonsense variant (Supplementary Figure S1). The other six variants found in our nine ChAc patients are apparently novel, and three of them are not even found in the Genome Aggregation Database (gnomad.broadinstitute.org/, last accessed on 30 December 2020) [36]; this unexpected high genetic variability suggests that many undetected VPS13A variants are actually present in the Italian and possibly other European populations. Although Italy has a very diverse genetic structure due to its central position in the Mediterranean and the numerous migratory waves that has mixed its population over the millennia, we conclude from our study that VPS13A variant screening should be considered in patients with chorea (after HD has been excluded) even in the absence of acanthocyte studies.

5. Conclusions

To summarize, we reported a relatively large series of NA patients proving that ChAc and MLS, although rare, should be always considered in the differential diagnosis of suspected hereditary chorea. Depending on the laboratory techniques available, chorein dosage with Western blotting and Kell blood group expression may be tested first or DNA sequencing of the VPS13A and XK genes will be performed. Moreover, since a primary myopathy is now suspected not only in MLS but also in ChAc patients, it would be useful to perform a muscle biopsy and protein expression studies in all confirmed NA patients in order to thoroughly study skeletal muscle involvement in these conditions.
Supplementary Materials: The following are available online at https://www.mdpi.com/2073-4254/12/3/344/s1, Figure S1: (A) c.1114_1115del; (C) c.3817C>T, Figure S2: (A) c.3339+4_3339+10delinTTAGCTGTATTTAAAAATTA; (B) exon 30-33 cDNA amplicon, Figure S3: (A) c.1078C>T; (B) c.7867C>T, Figure S4: c.2512+2T>G (homozygous), Figure S5: (A) c.7736_7739del; (B) c.2825-10T>G; (C) effect of the c.2825-10T>G splice-site variant on cDNA sequence, Table S1. Sequencing primers employed for genetic testing of VPS13A., Table S2. Sequencing primers employed for genetic testing of XK.

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