Multiomic elucidation of a coding 99-mer repeat-expansion skeletal muscle disease

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Twenty-two individuals across four generations suffer a chromosome 19p13.3-linked autosomal dominant progressive myopathy with distinctive pathology including rimmed ubiquitin-positive autophagic vacuolation [6] (Fig. 1 and Supplementary data). A recombination in the newest (youngest) affected patient (V:13) and repeat linkage analysis on six patients (IV:10, IV:17, III:18, IV:3, IV:23, V:13) refined the disease haplotype to 5.12 Mb containing 164 genes. Sanger (24 genes), whole-exome, whole-genome (Supplementary Table 1) and whole skeletal muscle RNA sequencing proved unrevealing.

Immunohistochemical workup showed that patients’ vacuoles and subsarcolemmal regions stained positive for FK2 and p62/SQSTM1 markers, respectively, of ubiquitinated proteins and autophagy (Fig. 1d). The number of stained fibers correlated with clinical severity (Supplementary data). To query the FK2 target(s), we microdissected vacuoles and unaffected myofiber parts for quantitative mass spectrometry. Among the more than 700 identified proteins (Supplementary Table 2), perilipin-4 was the most highly (almost 20-fold) over-represented in vacuoles versus control myofiber regions (Supplementary Fig. 2). Perilipins coat the phospholipid monolayer surrounding lipid droplets and regulate the latter [2]. All five perilipins share an amphipathic domain composed of an 11-amino acid (aa) sequence, which is particularly extensive in perilipin-4, the 11-mer being repeated three times to generate a 33-mer, in turn repeated 29 or 31 times [11] (Supplementary Fig. 3). Perilipin-4 is the most abundantly expressed perilipin in muscle, notwithstanding which, aside from reduced cardiac triacylglycerol proteins and autophagy (Fig. 1d). The number of stained fibers correlated with clinical severity (Supplementary data). To query the FK2 target(s), we microdissected vacuoles and unaffected myofiber parts for quantitative mass spectrometry. Among the more than 700 identified proteins (Supplementary Table 2), perilipin-4 was the most highly (almost 20-fold) over-represented in vacuoles versus control myofiber regions (Supplementary Fig. 2). Perilipins coat the phospholipid monolayer surrounding lipid droplets and regulate the latter [2]. All five perilipins share an amphipathic domain composed of an 11-amino acid (aa) sequence, which is particularly extensive in perilipin-4, the 11-mer being repeated three times to generate a 33-mer, in turn repeated 29 or 31 times [11] (Supplementary Fig. 3). Perilipin-4 is the most abundantly expressed perilipin in muscle, notwithstanding which, aside from reduced cardiac triacylglycerol
levels, its absence in mouse results in no cardiac, skeletal muscle or other impairment [10, 11].

Since PLIN4 maps to our linked region, we revisited the genomic and transcriptomic patient data and noticed an unusually high coverage in PLIN4 exon 3 (Supplementary Fig. 4). PCR amplification of this exon in patient genomic DNA and muscle RNA revealed the wild-type band, and a second ~1000 bp higher band (Fig. 2a) not present in unaffected relatives or in 60 ethnic controls. The 31 × 33-aa amphipathic domain of perilipin-4 is encoded by 31 × 99 repetitive sequences in exon 3 [4], which poses a computational challenge for aligning short sequencing reads. We amplified cDNA from patient muscle RNA and obtained Oxford Nanopore long-read sequencing, which confirmed that the higher band is an expansion of the normal 31 × 99-nucleotide sequence to 40 × 99 bases, resulting in 297 (9 × 33) extra amino acids (Supplementary Fig. 5). Muscle extract Western blotting with a perilipin-4 antibody showed the presence of a second band consistent in size with the genetic expansion in patients, and absent from controls (Fig. 2b). Immunohistochemistry with the same antibody showed a major increase in perilipin-4 positivity in subsarcolemmal regions and vacuoles of patients compared to controls. The perilipin-4 signal most exactly reproduced the staining with the FK2 (Fig. 2c) and p62/SQSTM1 antibodies (Supplementary Fig. 6). These staining correlated with the diseased muscle fiber type, namely, slow-twitch Type I fibers, known to contain the highest amounts of intramyocellular lipids. Oil Red O staining showed normal lipid content and distribution (Supplementary Fig. 7 and data). Aggrephagy pathway components beyond FK2 and p62/SQSTM1, namely, NBR1 and WDFY3, were upregulated, the former (Fig. 2d) colocalizing with perilipin-4, FK2 and p62/ SQSTM1, the latter (Fig. 2e) increased in subsarcolemmata near perilipin-4 positivity but without co-localization. In aggrephagy, p62/SQSTM1 interacts with NBR1, and the two, as an autophagy receptor complex and through their shared LC3-interacting regions, bridge the aggregating ubiquitinated proteins with LC3. Meanwhile, WDFY3 shuttles from the nucleus to the cytoplasm to scaffold the overall structure with PtdIns3P-containing membranes and encapsulate the aggregates in autophagosomes for degradation [8]. The present disease is characterized by dominantly inherited progressively increasing mobilization of aggrephagy at sites of progressive accumulation of a mutated protein, suggesting that the mutation is leading to aggregation, likely through misfolding, exceeding aggrephagic capacity. Continuous formation and fusion of failing aggrephagic vesicles possibly leads to ever larger vacuoles, which disrupt the organization of myofibers and alter their contractile abilities, resulting in atrophy.

Many cases of Inclusion Body Myopathy, the most common of the myopathies, exhibit aggrephagic activation, including NBR1 deposition, not dissimilar to the present patients [5]. The proportion of cases that are due to misfolded proteins, potentially including perilipin-4, remains to be determined.

Perilipin-4 shares its amphipathic domain structure with α-synuclein and exchangeable lipoproteins (ApoA, ApoC and ApoE) [3]. All known mutations (all missense) of α-synuclein and ApoA1 in familial Parkinson disease and amyloidosis, respectively, localize to these proteins’ amphipathic regions and transition the repeating helices of these domains to amyloidogenic β pleats [1, 7]. Genomic repeat sequences predispose to expansion [9]. To our knowledge, ours is the first report of an amphipathic domain repeat expansion in disease, and identification of the expansion was only possible with long-read sequencing. The possible occurrence of germline or somatic pathogenic amphipathic region repeat expansions in proteins possessing these domains in their related diseases should be explored.
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Author contributions AR, MM and BAM conceived and designed the study and wrote the manuscript with input from all authors. MM and BAM supervised and provided critical discussion of the data. AR, SN, MAS and KB performed genetic experiments and/or interpreted data. MV, KM and RAK preformed proteomic experiments and/or interpreted data. AR, EI, FB, BC and GS performed molecular, immunohistochemistry, immunoblot, confocal and electron microscopy experiments. LM and RM acquired and analyzed clinical data. PW and JJD provided experimental support and advice.

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