# Dystroglycanopathies: Genetic Bases of Muscular Dystrophies Due to Alteration in the O-Glycosylation of α-Dystroglycan

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#### **Abstract**

Congenital muscular dystrophies (CMDs) are inherited, progressive and heterogeneous muscle disorders. A group of CMDs are dystroglycanopathies, also called  $\alpha$ -dystroglycanopathies, where there is an abnormal glycosylation of protein  $\alpha$ -dystroglycan. Hypoglycosylation of  $\alpha$ -DG results in different severities of congenital muscular dystrophies and they present with progressive muscle weakness and loss of motor functions. This article first focuses on the CMDs, their classification according to the observed symptoms or the protein involved in the resulting phenotype. We then focus on dystroglycanopathies, the importance of its correct O-glycosylation of the  $\alpha$ -dystroglycan given its important structural function, considering the enzymes involved in said glycosylation and the phenotypes that can result, to finally address current therapeutics for these diseases with the aim of increasing current knowledge.

### **Keywords:**

Congenital muscular dystrophies (CMDs), a-dystroglycan protein (aDG), glycosylation, muscle disorders, dystroglycanopathies.

### Introduction

Congenital muscular dystrophies (CMDs) are inherited, progressive and heterogeneous muscle disorders [1]. They affect children at birth or appear during the first months of life with a predominant autosomal recessive mode of inheritance, except for laminopathies (L-CMD) and some Ullrich congenital muscular dystrophies (U-CMDs) [2]. These disorders are characterized by congenital hypotonia, delayed motor development and early onset of progressive muscle weakness, and they can comprise the involvement of brain and eyes [1].

In principle, the CMDs were classified by histological and clinical criteria.

The classification of these diseases is complex [2], and in general it is based on phenotype characteristics defined by clinical criteria and histochemical analysis considering the following criteria:

- Involvement or non-involvement of the Central Nervous System.
- 2. Increased muscle enzymes € eatine Kinase (CK)).
- Neuro-radiological abnormalities.
- 4. Protein expression (Laminin α2, Dytroglycan, collagen).

Clinical features, immunohistochemical staining, Western blot, brain and muscle's Magnetic Resonance Imaging (MRI) are all useful tools to direct genetic testing.

They can be also classified according to the subcellular location of the mutated protein and/or the consequences at the skeletal muscle level (extracellular matrix, sarcolemma, basal lamina, endoplasmic reticulum, and nuclear envelope). The CK levels are low or remain unchanged [1,3].

In 2018, Quijano-Roy *et al* set out a global algorithm that facilitates the classification into five main forms of CMDs according to the affected gene [1] but there are several genes that have been discovered in the last years [1,4] (Figure 1).

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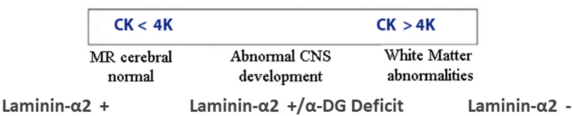


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# **Congenital Muscular Dystrophy**

### **Central Nervous System affected**

## **Central Nervous System Not affected**



CMD ULLRICH (U-CMD) (Col6A1,Col6A2,Col6A3) RIGID SPINE (RS-CMD1) (SEPN1) LAMINOPATHIES (LMNA)

MUSCLE EYE BRAIN

MEROSINOPATHIES (CMD1A)
(Lama2)

WALKER WARBURG

### CMD FUKUYAMA TYPE

Figure 1. Classification of Congenital Muscular Dystrophies. Adapted from Quijano et al 2018.

A summary proposal by Zambon and Mutoni considers the subcellular location of the mutated protein and they classified CMDs as follows [4]:

 Defective structural proteins of basal membrane or extracellular matrix of muscle fibers

**LAMA2-RD:** The laminin  $\alpha$ 2–related muscular dystrophies (LAMA2-RDs), previously known as merosin-deficient congenital muscular dystrophy type 1A (MDC1A), are autosomal recessive disorders caused by pathogenic variants in the LAMA2 gene [5–6]. LAMA2 encodes for the alpha-2 subunit of heterotrimeric laminin-2 protein (made up of  $\alpha 2$ ,  $\beta 1$ , and  $\gamma 1$  subunits) with the α2 subunit called laminin-211 or merosin serving as a tissuespecific component of the extracellular matrix with a key role in myotubes stability and apoptosis [7]. The spectrum of LAMA2-RD ranges from severe CMD (90% of cases) to milder, later childhood-onset LGMD largely due to the amount of residual Lm-211 protein in muscle (complete vs partial deficiency, CD vs PD) [4]. To date, there are no effective treatments for LAMA2-RDs [7]. With an estimated prevalence of 0.6-0.7/100,000, LAMA2-RD is one of the most common types of CMDs. It is clinically manifested by hypotonia and weakness at birth, the development of contractures of large joints, and progressive respiratory involvement. Muscle atrophy and severe weakness typically prevent independent ambulation [7–9].

**COL6-RD**: Collagen VI-related myopathies are hereditary myopathies caused by mutations in either COL6A1, COL6A2

or COL6A gene, each encoding a subunit of collagen VI [10–11]. The clinicopathological hallmarks include distal hyperlaxity, proximal joint contractures, protruding calcanei, skin hyperkeratosis, scoliosis and respiratory insufficiency [12]. Approximately three quarters of individuals with congenital/severe COL6-RD acquire the ability to walk independently, but subsequently their motor skills decline, with loss of ambulation occurring around 9–11 years of age [4].

Integrin  $\alpha$ -7 deficient CMD: This CMD is caused by mutation in integrin  $\alpha$ 7 (ITGA7) and it is a rare subtype. This gene has 28 exons, among which 26 code for protein [13]. Only a few patients diagnosed with CMD were found to have ITGA7 mutation [6,13–16] and it is not clear whether all variants are pathogenic mutations. Variants of ITGA7 present microcephaly, agenesis of the corpus callosum, cerebellar hypoplasia, seizures, scoliosis, hemivertebrae, asymmetric extremities, and hypopigmented skin macules [16]. These had various symptoms and different grades of severity. The consistent clinical features were muscle weakness and increased CK level [13].

Integrin  $\alpha$ -9 deficient CMD: The  $\alpha$  retegrin is one of the younger evolutionary additions to the integrin family of receptors. This integrin subunit is expressed in a variety of cell types and binds to a plethora of ligands, some of which are restricted to specific tissues, or are upregulated during development or in pathophysiological conditions. There have not been many reports describing mutations in integrin  $\alpha$ -9 [17]. This could indicate that mutations in  $\alpha$ -9 is a rare event, and that it is the  $\alpha$ -9 expression level (or a lack of it) that is important in

developing pathophysiological conditions, which was already suggested when ITGA9 was first cloned [18].

Dystroglycanopathies (DGP)/defects of α-DG glycosylation: They are a group of muscular dystrophies where there is an abnormal glycosylation of  $\alpha$ -dystroglycan protein ( $\alpha$ -DG). This protein is a fundamental component of the Dystrophin glycoprotein complex (DGC), which is essential to link the extracellular matrix (ECM) to the intracellular actin cytoskeleton [4]. The clinical symptoms are diverse, ranging from severe congenital to adult-onset limb-girdle types [19-20]. Primary dystroglycanopathy is due to defects in the coding sequence of a-DG [21-22]. Around 20 genes have been associated with secondary dystroglycanopathies, including glycosyltransferases for O-mannitol type glycosylation, a kinase, enzymes involved in nucleotide sugar synthesis (dolichol—P-Mannose), proteins necessary for joining the α-DG-linked core glycan structure with distal ligand-binding region and Golgi membrane trafficking proteins [23-24].

### -Endoplasmic reticulum protein

**SEPN1-related Myopathy:** SEPN1 or SELENON is a ubiquitous endoplasmic/sarcoplasmic reticulum (ER/SR) protein encoded by the SELENON (or SEPN1) gene. It protects cells against oxidative stress or ER-stress and defends calcium homeostasis by counteracting ERO1-mediated oxidation of the ATPdependent Ca2+ pump SERCA [25]. Mutations of the SEPN1 gene are characterized by muscle weakness and fatigue leading to scoliosis and life-threatening respiratory failure. Core lesions and focal areas of mitochondria depletion in skeletal muscle fibers are the most common histopathological lesions [26]. This myopathy comprises a spectrum of pathological conditions encompassing rigid spine muscular dystrophy, multi-minicore disease, congenital fiber-type disproportion and desmin-related myopathy with Mallory body-like inclusions [4]. This st reveals SEPN1-RM as a more severe and progressive disease than previously thought. While motor abilities were reported to be stable, we und loss of ambulation in 10% of the cases with full follow-up data. Muscle functional performance and respiratory function (particularly diaphragmatic fatigue) declined systematically from the end of the third decade, even in mild cases. But lifespan was reduced even in 2 mild cases with optimum respiratory support [27].

### Nuclear envelope proteins

LMNA-Related CMD: A-type lamins (lamin A and C) are intermediate filament proteins expressed in differentiated cells [28]. Together with B-type lamins, they form the nuclear lamina, an organized meshwork found under neath the inner nuclear envelope [29]. Laminopathies are a heterogeneous group of disorders caused by mutations in the LMNA gene encoding lamin A/C. This includes striated skeletal and cardiac muscles, and it includes Emery–Dreifuss muscular dystrophy (EDMD), limb-girdle muscular dystrophy type 1B (previously

known as LGMD1B) and isolated dilated cardiomyopathy with conduction system defects and arrhythmias (DCMCD). LMNA-related congenital muscular dystrophy (L-CMD) has been described as an autosomal dominant muscle disorder related to a dominant de novo mutation in LMNA, so far, the most severe form of striated muscle laminopathies [30]. L-CMD group can operationally be defined as having an onset of skeletal muscle manifestations within the first two years of life, when early motor development, that includes walking and running, should be typically attained. It is of great medical importance to distinguish this group of patients due to the potential of early life-threatening complications involving nutritional, respiratory and cardiac compromise [31].

SYNE-Related CMD (nesprin): EDMD is associated with at least seven gene mutations, of which SYNE1 mutation is relatively less common [32]. The SYNE1 (Spectrin Repeat Containing Nuclear Envelope Protein 1) gene has an autosomal dominant inheritance pattern, and its mutations might result in defects in the expression product nesprin-1 [33], a protein characterized by the presence of multiple spectrin repeats which is highly expressed in striated muscles [34]. Besides, the mutations in the SYNE1 gene cause spinocerebellar ataxia type 8, myogenic multiplex arthrogryposis congenital with features of EDMD, intellectual disability with spastic paraplegia, and axonal neuropathy [35–37].

### -Proteins involved in ER to Golgi apparatus trafficking

There are two genes that represent the first membrane trafficking proteins implicated in  $\alpha$ -DG hypoglycosylation. TRAPPC11 and GOSR2, that each have a role in membrane trafficking in the biosynthetic pathway, have been implicated as candidate dystroglycanopathy genes [4].

TRAPPC11: Transport protein particle (TRAPP) is a supramolecular protein complex that functions in the localization of proteins to the Golgi compartment. The TRAPPC11 subunit has been implicated in muscle disease through the identification of homozygous and compound heterozygous deleterious mutations in individuals with limb girdle muscular dystrophy and congenital muscular dystrophy. These individuals also display membrane trafficking defects in cultured fibroblasts; this gene should be considered in the diagnostic evaluation of patients with CMD [24,38].

GOSR2: Pathogenic variants in the Golgi SNAP receptor complex 2 gene (GOSR2, also known as Membrin) are well-known to be associated with autosomal recessive progressive myoclonic epilepsy (PME) [39], but some new compound heterozygous variants in the GOSR2 gene have expanded the clinical spectrum [24,40]. This variant has progressive muscle weakness and areflexia, and it developed seizures in early childhood. Muscle biopsy showed an active dystrophic process and hypoglycosylation of alpha-dystroglycan. This is suggestive of a dystroglycanopathy, which is a known cause of congenital muscular dystrophy [41].

There are another condition (called **CMD PLUS**) that share clinical features with CMD, for example, Megaconial CMD or Marinesco-Sjogren Syndrome (MSS) [4]. These conditions have overlapping pathological and clinical changes that often lead to diagnostic difficulties with CMD. Some of them are listed in Table 1.

This review focuses on dystroglycanopathies, also called  $\alpha$ -dystroglycanopathies (CMD- $\alpha$ DG).

### **Dystroglycanopathies**

These pathologies owe their etiology to mutations in genes involved in the O-glycosylation of  $\alpha$ -DG protein [42]. At least 20 genes are involved in the correct O-glycosylation of α-DG. They are classified into primary and secondary dystroglycanopathies [20] (Table 2). The first ones are due to mutations of the DAG1 gene that alter the state of the DG core protein with potential repercussions on the glycosylation state of  $\alpha$ -DG. The second ones are due to mutations in the genes that participate in the O-glycosylation of α-DG including enzymes involved in nucleotide sugar synthesis and Golgi membrane trafficking proteins [23–24]. Primary dystroglycanopathies are comparatively less studied due to the small number of cases identified so far. DAG1 mutations are rare, recessive mutations that are found in consanguineous families. Dystroglycanopathies exhibit a broad clinical spectrum [43], because of the mutation on the activity of the protein involved. One of the characteristics of these dystrophies is the involvement of central nervous system disorders, such as malformation of the brain (type II lissencephaly) and mental retardation (although there are times that this occurs without structural abnormalities). Cases with heart failure and eye symptoms have also been observed [44-48].

The glycosylation status of  $\alpha$ -DG is strictly regulated with respect to both developmental stage and tissue (in brain, heart, skeletal muscle, and kidney,  $\alpha$ -DG is modified in a way that it can function as an ECM-receptor) [49].

# pnctional Importance and Genetic Bases of α-Dystroglycan

The dystrophin glycoprotein complex (DGC), which is essential to link the extracellular matrix (ECM) to the intracellular actin cytoskeleton, takes center stage in several physiological and pathological contexts, playing a particularly important role in skeletal muscle. This gives stability to a big number of tissues, such as skeletal and smooth muscles, the brain and peripheral nervous system, the neuromuscular junction, the interface between endothelial cells and the surrounding astrocytes endfeet at the blood-brain barrier, the kidney glomeruli basement membrane, and the lungs at the epithelia-connective border [20,50–51].

DAG1 gene has a highly conserved sequence between species and has been mapped to human chromosome 3p21 [52]. Coding sequence is organized into two exons, separated by a large intron. The derived 5.8 kb transcript contains an 895-residue open reading frame. This propeptide is post-translationally cleaved by an unknown protease at residue 653 (P) to yield  $\alpha$ -and  $\beta$ -dystrogly ( $\alpha$ -DG and  $\beta$ -DG) [51,53–54].  $\beta$ -DG protein contains a single contains and a 121-residue C-terminal cytoplasmic tail that is enriched in proline.  $\alpha$ -DG is an extracellular protein that contains three potential N-linked glycosylation sites. DGC complexes associated with various proteins via  $\alpha$ -DG y  $\beta$ -DG.

**Table 1.** Genes and Protein implicated in CMDs called CMD plus.

OtherCMDs/CMD spectrum	Gene symbol	OMIN ref	Protein/Function
Mitochondrial CMD	СНКВ	602,541	Choline kinase
MD withcerebellarinvolmente	MSTO1	617,675	Mitocondrial fusion
MD with extrapiramidal signs	MICUI	615,673	Mitochondrial Ca2+uptake/ mitochondria mediated sarcolema repair
CMD with cataracts and intellectual disability	INPP5K	617,404	Inositol polyphosphate-5-phosphatase K
Marinesco-Sjogren syndrome	SIL-1	248,800	Protein Folding
Mucolipidosis type IV	MCOLN1	252,650	Nonselective cation chanel in lysosomal endosomal trafficking
MD/hearing Loss/Ovarian insufficiency Syndrme	GGPSI	606,982	Mevalonat/isoprenoid pathway

**Table 2.** Genes associated to dystroglycanopathies.

Gene	OMIM ref	Protein Functions
Primary dystroglycanopathy		
DAG1	616,538	Dystroglycan
Secondary dystroglycanopathy		
POMT1	607,423	Protein O-mannosyltransferase
POMT2	607,439	Protein O-mannosyltransferase
POMGnT1	606,822	Protein O-mannose $\beta$ 1,2-N-acetylglucosaminyltransferase; Core M1 synthesis
POMGnT2	614,828	Protein O-mannose $\beta$ 1,2-N-acetylglucosaminyltransferase; Core M3 synthesis
B3GALT2	610,194	$\beta$ -1,3-N-acetylgalactosaminyltransferase; Comre M3 synthesis
POMK	615,247	Protein O-mannose kinase; Phosphorylation of Core M3
FKTN	607,440	Ribitol phosphate transferase, tandem ribitol synthesis
FKRP	606,596	Ribitol phosphate transferase, tandem ribitol synthesis
ISPD/CRPPA	614,631	CDP-ribitolpyrophosphorylase; synthesis of CDP-ribitol (donor substrate of FKTN/FKRP)
TMEM5/RXYLT1	605,862	β-1,4-xylosyltrnasferase; synthesis of linker structure between tandem ribitol and matriglycan
B4GAT1	615,287	β-1,4-Glucuronyltransferase; synthesis of linker structure between tandem ribitol and matriglycan
LARGE1	603,590	$\alpha 3$ -Xylosyl and $\beta 3$ -glucuronyltransferase; matriglycan synthesis
GMPPB	615,320	GDP-mannose pyrophosphorylase required for the formation of GDP-Man; Dolichol-phosphate-mannose synthesis
DPM1	603,503	Dolichol-phosphate-mannose synthase; Dolichol-phosphate-mannose synthesis
DPM2	603,564	Dolichol-phosphate-mannose synthase; Dolichol-phosphate-mannose synthesis
DPM3	605,951	Dolichol-phosphate-mannose synthase; Dolichol-phosphate-mannose synthesis
DOLK	610,768	Dolichol kinase required for formation of dolichol -phosphate; Dolichol-phosphate-mannose synthesis

β-DG (Figure 2) subunit binds intracellularly to dystrophin, that joins to the intracellular actin cytoskeleton; and it extracellularly binds to α-DG, which in turn binds via Laminin 2 to the extracellular matrix [20,54] (Figure 3). The core skeletal muscle DGC also contains the sarcoglycans (α-SG, β-SG, δ-SG and γ-SG [50–51]), the sarcospan [55] and the syntrophins [56]. In addition, several extra- and intracellular proteins are less tightly associated with the DGC, such as nitric oxide synthase (nNOS) and dystrobrevin [54,57–59].

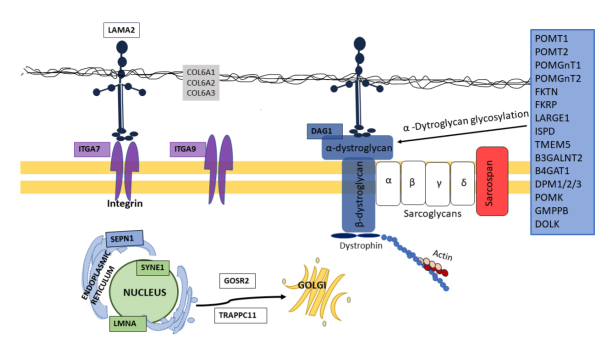
# Spgar Chain Structure of α-DG

α-DG has a mucin-type O-glycosylation site in the central region of the molecule and it contains more than 40 Ser/Thr residues that form an O-glycan cluster [53]. O-glycosylation is an even more complex process [60].  $\alpha$  -DG is heavily N- and O-glycosylated, as well as O-mannosylated [61–62]. The first 18 amino acids of the mucin-like domain have been shown to be important for this O-mannosylation of  $\alpha$ -DG to be carried

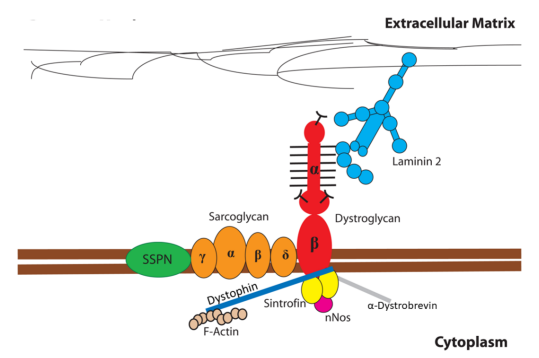
out correctly [21]. The sequential O-glycosylation process of  $\alpha$ -DG forms different structures called nuclei or Core (Core 1 to 3). Core 1 is crucial for the binding of the extracellular components [50].

# re M1 and Core M2

The O-mannosylation initiates in the endoplasmic reticulum where POMT1 and POMT2 (O-mannosyl-transferases) form a complex that transfer a O-Man in an alpha linkage to serine and threonine residues of  $\alpha$ -DG [63]. POMT1-POMT2 complex uses dolichol phosphate mannose (Dol-P-Man) as a donor substrate. The synthesis of Dol-P-Man is carried out by GMPPB (GDP-mannose pyrophosphorylase), DPM1/2/3 (Dolichol-phosphate-mannos synthase), and DOLK (Dolichol Kinase) [43]. POMGnT1 acts in the Golgi, where it transfers  $\beta$ 2-linked GlcNAc residues to the mannose residues added by POMT1 and POMT2 during synthesis of the core M1 and M2 glycans [64]. Core M2 structure is found specifically in the brain [65].



**Figure 2.** Schematic representation of the Dystrophin glycoprotein complex (CDGP) in skeletal muscle. The two dystroglycan subunits ( $\alpha$ -DG and  $\beta$ -DG) interact non-covalently to form a bridge between the extracellular matrix and the actin cytoskeleton. The cytosolic domain of  $\beta$ -DG is anchored to actin through interaction with dystrophin.  $\alpha$ -DG interacts with the ectodomain of  $\beta$ -DG on the extracellular side of the plasma membrane. DG acts as a receptor for extracellular matrix proteins such as laminins. Adapted from Brancaccio 2019.



**Figure 3.** Scheme of genes implicated in CMDs and their localization in skeletal muscle cell. Adapted from Zambon and Mutoni 2021. LAMA2, Laminin-2 protein; COL6A1, COLA2 and COL6A3, α-chains of Callagen VI monomers and tetramers; ITGA7, integrin α-9; ITGA9, integrin α-9; SEPN1, Selenoprotein 1; SYNE, Spectrin Repeat Containing Nuclear Envelope Protein 1; LMNA, Laminin a/c; TRAPPC11, Transport protein particle 11; GOSR2, Golgi SNAP receptor complex 2; DAG1, Dytroglycan; POMT1, Protein O-mannosyl-transferase 1; POMT2, Protein O-mannosyl- transferase 2; POMGNT1, Protein O-mannose β1,2-N-acetylglucosminytranferase 1; POMGNT2, Protein O-mannose β1,4-N-acetylglucosminytranferase 2; TMEM 5, transmembrane protein 5; RXYLT1, ribitol-5-phosphate xylosyltransferase 1; B4GAT1, β1,4-glucuronyltransferase; B3GALNT2, β1,3-N\_acetylgalactosaminyltransferase; POMK, Protein O-mannose Kinase; FKTN, Fukutin; FKRP, Fukutin related protein; LARGE1, like-acetylglucosaminyltransferase/LARGE xylosyl-and glucoronyltransferase 1; ISPD, isoprenoid synthase domain-containing protein; CDP-ribitopyrophosphorylse A.

# re M3 and Matriglycan Importance

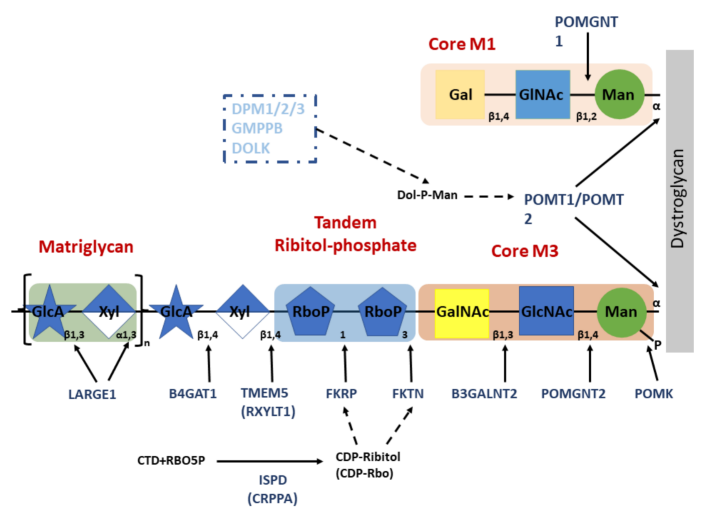
A small subset of O-Man modified sites, apparently exclusively on  $\alpha\text{-DG},$  are extended in the endoplasmic reticulum by a GlcNAc in a beta-1,4 linkage by POMGNT2 to generate the Core M3 glycans [49]. This is further elaborated into a trisaccharide by the action of a beta-1,3-N-acetylgalactosamine (GalNAc) transferase, B3GALNT2 [53,66]. Core M3 has a phosphorylation at the C6 position of O-mannose, being POMK [67] the one that transfers a phosphate group from adenosine 5'-triphosphate (ATP), forming the phospho-nucleus structure [68].

The  $\beta$ 1,4 branch is subsequently modified by a series of Golgi enzymes [69] that include B3GALNT2 ( $\beta$ 1,3-N-acetylgalactosaminyl transferase), the ribitol transferases FKTN (Fukutin) and FKRP (fukutin-related protein), TMEM5 (Ribitol-

5-phosphate xylosylransferase), and B4GAT1 ( $\beta$ 1,4-glucoronic acid transferase). Finally, the LARGE1 lengthens the previous linear carbohydrate branch by synthesizing a unique repeating disaccharide structure [-3-xylose- $\alpha$ 1,3-glucuronic acid- $\beta$ 1-]n, called matriglycan [53] (Figure 4). This complex mediates  $\alpha$ -DG interactions with LG domains of extracellular matrix (ECM) proteins such as laminins, agrin and perlecan and pikachurin [50–51,70].

This unique heteropolysaccharide is expressed in most tissues and it plays diverse roles, from acting as viral receptor to neuronal development [71–76].

Multiple studies, carried out in cultured cells and mouse skeletal muscles, have demonstrated that forced expression of LARGE increases the MW of  $\alpha$ -DG and its binding ability to LG domains [77–78].



**Figure 4.** Representation of the sequence of genes involved in the O-Glycosylation of α-DG and matriglycan formation. Man, mannose; GlcNAc, N-acetylglucosamine; GalNac, N-acetylglactosamine: RboP, ribitol phosphate; Xyl, xylose; GlcA, glucronic acid; Gal, galactose; Rbo5P, ribitol-5-phosphate; POMT1, Protein O-mannosyl-transferase 1; POMT2, Protein O-mannosyl-transferase 2; POMGNT1, Protein O-mannose β1,2-N-acetylglucosminytranferase 1; POMGNT2, Protein O-mannose β1,4-N-acetylglucosminytranferase 2; TMEM 5, transmembrane protein 5; RXYLT1, ribitol-5-phosphate xylosyltransferase 1; B4GAT1, β1,4-glucuronyltransferase; B3GALNT2, β1,3-N\_acetylgalactosaminyltransferase; POMK, Protein O-mannose Kinase; FKTN, Fukutin; FKRP, Fukutin related protein; LARGE1, like-acetylglucosaminyltransferase/LARGE xylosyl-and glucoronyltransferase 1; ISPD, isoprenoid synthase domain-containing protein; CDP-ribitopyrophosphorylse A.

LARGE is a type II transmembrane protein that contains two distinct domains: one with homology to  $\beta 3GNT1$ , and another with homology to proteins belonging to glycosyltransferase family 8. LARGE has two glycosyltransferase activities: a  $\alpha 3$ -xylosyltransferase activity and a  $\beta 3$ -glucuronyltransferase activity. The LARGE paralog, LARGE2, possesses the same enzymatic function, although its optimal pH and pattern of expression differ from those of LARGE [79–80].

# Dystroglycanopathies & Congenital Disorders of Glycosylation

Congenital Disorders of Glycosylation (CDG) are a group of rare inborn errors of metabolism and include some forms of dystroglycanopathies [60].

These disorders (CDG) are genetic diseases caused by deficient glycoprotein and glycolipid glycan synthesis and attachment [81–82]. Most are multisystem disorders with variable phenotype severity and neurological involvement. CMDs due to CDG are clinically and genetically heterogeneous diseases and present progressive muscle weakness and loss of motor functions [1–2]. These present a stereotyped phenotype with early-onset muscle deficit with or without central nervous system involvement [83–85].

The glycosylation is a complex biological process involving many pathways. CDG are sub-grouped into defects of protein N-glycosylation, protein O-glycosylation, lipid glycosylation, GDP-anchor glycosylation and multiple glycosylation defects [81,86]. O-glycosylation is a common covalent modification of serine and threonine residues of mammalian glycoproteins [53].

Many of the described mutations for muscular dystrophy affect the O-glycosylation pathways for the biosynthesis of  $\alpha$ -dystroglycan. It has been described that new types of CDG result in defects in the O-mannosyl glycosylation pathway [87–92].

The multitude of clinical phenotypes resulting from defective O-mannosylation highlights the biomedical significance of this unique modification [47,93]. The understanding of this modification is important for the development of novel therapeutics [91,94–96].

Elucidation of the molecular pathological mechanisms of CDG associated with DG glycosylation abnormalities will be an important issue in understanding the mechanisms of dystroglycanopathies. Characterizing a glycoproteome profile of patients prior to and on treatment will help to better understand the changes of a plethora of glycoproteins and related clinical observations in dystroglycanopathies [81–82,86,97].

# Penotypes Resulting from Defects in α-Dg Glycosylation

When the enzymes that participate in the O-glycosylation pathway of the  $\alpha$ -DG subunit show failures [98–99] or a

pathogenic variant is observed in DAGI gene, a decrease in the binding capacity of  $\alpha$ -DG to the other proteins of the CDGP complex is observed, including in laminin. This generates a severe clinical phenotype of muscular dystrophy, brain abnormalities, and often optic abnormalities. DMCs such as Walker-Warburg Syndrome (WWS), Muscle-Eye-Brain disease (SEM) and Fukuyama Congenital Muscular Dystrophy (FC-CMD) [100] are assumed to be  $\alpha$ -dystroglycanopathies. They all present a severe degree of muscular dystrophy with deterioration in the muscles, brain, and eye, and they have overlapping phenotypes that make their specific diagnosis difficult.

Dystroglycanopathies present a broad phenotypic spectrum that can even overlap, making diagnosis difficult. This is probably because of mutations involving the functions of gene products (enzyme activity), rather than variation in the causative genes [43]. In addition to the changes in the DAG1 gene, changes in the enzymes involved in the glycosylation of  $\alpha$ -DG must be considered. Moreover, it is important not to forget the effect of variations in genes (such as ISPD or GMPPB) and their corresponding enzymes responsible for the manufacture of carbohydrate building blocks in the cytosol, indirectly modifying  $\alpha$ -DG glycosylation [20,24,91,101].

There is a clinical-radiological imaging technology classification that divides dystroglycanopathies into seven groups, including CMDs and limb girdle dystrophies [22,102–103,106].

- Walker-Warburg Syndrome (WWS): onset prenatally or at birth. With eye abnormalities and severe structural brain abnormalities, including complete agyria or severe lissencephaly with only rudimentary cortical folding, it marked hydrocephalus, severe cerebellar involvement and complete or partial absence of the corpus callous. Patients assigned to this category are incompatible with life or have very limited survival.
- CMDs muscle-eye-brain/Fukuyama (MEB/CMD-FC) type: multiple malformations of the CNS (brain, cerebellum, or trunk) of less severity than the previous group.
- CMDs with mental retardation and localized structural CNS involvement (CRB-CMD).
- CMDs with intellectual disability and without CNS morphological abnormality (MR-CMD), possible microcephaly and/or moderate involvement of the cerebral white matter.
- 5. CMDs without intellectual disability (no MR-CMD): this category includes CMD-1C.
- Girdle dystrophies with intellectual disability (LGMD-MR). It may include microcephaly and/or moderate brain and white matter abnormalities.
- Limb Girdle muscular dystrophies without intellectual impairment (LGMD-not MR). It includes LGMD2I (FKRP), 2L (ISPD [107–112]) and 2M (FKTN).

# alker-Warburg Syndrome: WWS

It is the most severe phenotype, and clinical manifestations appear early and can be detected in the prenatal stage by imaging techniques [113]. The clinical manifestations include congenital muscular dystrophy, brain, and eye anomalies: hydrocephalus, abnormal migration, retinal dysplasia and encephalocele [45,114]. The eye anomalies involve both anterior and posterior chambers with retinal detachment and blindness. Microphthalmia, thalamus, optic nerve hypoplasia, colobomas and iris malformation, cataract, and cornea defects may also be found. Mutations in B3GLNT2, B4GAT1, DAG1, FKRP, FKTN, GMPPB, ISPD, or LARGE are associated to Walker-Warburg Syndrome with brain and eye abnormalities [48]. The clinical features may also be associated, even if not frequently, to facial dysmorphism and cleft lip or palate [19]. It has been shown that mutations in POMT1 [115], FKTN, FKRP [116] and B3GALNT2 [117–118] can each give rise to the clinical WWS phenotype, even this phenotype should also be suspected when mutations in POMGnT1 and LARGE1[45] are present [119]. The most common known causes of WWS are mutations in glycosyltransferases protein O-mannosyltransferase 1 and 2 (POMT1 and POMT2 [120]) [45,63,121]; these mutations have been observed in a few patients [114,122].

# scle-Eye-Brain Syndrome: MEB

This disease is characterized by structural ophthalmological abnormalities and brain malformation. It was first reported in Finnish patients with poor psychomotor development, hypotonia and loss of reflexes at the age of 1 [123]. Patients are unable to sit still and they have seizures, and high levels of CK are detected [124].

Ophthalmologic abnormalities are very common. The EMG usually shows myopia, corneal opacity, cataracts and dysgenesis of the anterior chamber, congenital glaucoma, hypoplastic choroid and optic atrophy, among other alterations. Brain malformations such as argyric hemispheres, polymicrogyria in various cortical segments, and severe cortical disorganization are observed on MRI [125–127]. Muscle Eye Brain disease involves genetic mutations in B3GLNT2, B4GAT1, DAG1, FKRP, FKTN, GMPPB, ISPD, or LARGE. There are frequent mutations [103–106,128–129] in genes that have been associated with this pathology [116,130].

# 1D Fukuyama (CMD-F)

This pathology is caused by mutations in the FKTN [131] gene. This disease is highly prevalent in Japan, where the founder mutation was detected, a 3 kb insertion in 3'UTR [127], although there are others associated with this phenotype. This homozygous mutation is present in 80% of patients [131]. Other mutations in non-Japanese patients have been detected and involve a loss of localization in the Golgi or missense mutations [132–133].

Some mutations involve total or partial loss of function. This depends on how involved it is its glycosyltransferase enzymatic domain, which is essential in the binding of the first Rob 5 to the M3 core of  $\alpha$ -DG [93].

This phenotype is characterized by the appearance of congenital muscle weakness, severe mental retardation and delayed motor development with severe epilepsy associated with significant muscle involvement of the facial muscles and marked hypotonia in the extremities [134].

# uscular Dystrophy 1C (CMD-1C)

This subtype of CMD was first reported by Brockington *et al*, who identified a new member of the fukutin family of proteins, FKRP [135–136]. The clinical manifestations appear in the first weeks of life with a pronounced increase in CK. Some patients present normal brain structure, cardiac anomalies and respiratory failure, severe weakness in the muscles of the shoulder girdle, calves, and thighs in young age with null ambulation [19].

Although there is a decrease in  $\alpha$ -DG levels, there would be a mechanism by which this deficiency would be compensated in the brain, so there is no significant impact on the CNS, although there would be a subgroup that would present neurological abnormalities resulting from mutations in FKRP [19].

# uscular Dystrophy 1D (CMD-1D)

It is caused by mutations in the LARGE1 gene [137–138], which encodes the enzyme acetyl-glusosaminyl transferase. In this pathology, there is profound mental retardation and sometimes retinopathy [19,77]. The first human mutation reported for LARGE1 gave rise to the new classification of (DMC-1D) [19]. It was reported in a 17-year-old girl with congenital muscular dystrophy, profound mental retardation, white matter changes, and very subtle structural abnormalities on MRI of the brain. A significant reduction in  $\alpha$ -DG was observed at biopsy [139]. A missense mutation (c.G1525A) and a 1 bp insert (c.1999insT) in the human LARGE1 gene homologous to mouse LARGE were described in this patient [48,139–140].

# nb Girdle Muscular Dystrophy LGMD2I

It is caused by mutations in the FKRP gene [141], which also implies a secondary deficit of  $\alpha$ -DG and merosin [139]. In patients with LGMD2I, symptoms appear between the ages of 6 and 13 [142]. Frequent muscular pseudohypertrophy (calves, tongue) is observed. There are cases of severe respiratory failure that can precede loss of gait with an elevated risk of cardiomyopathy. Near half of the patients develop weakness in the ventricles [143].

In Table 3 we list some disease-causing variants associated to phenotypes more frequent to dystroglycanopathies.

**Table 3.** DISEASE-CAUSING VARIANTS associated to dystroglycanopathies Phenotypes. These variants are interpreted as pathogenic in ClinVar (Genomic variation as it relates to human health). NM\_ correspond to NCBI Reference Sequence used in the construction of this table.

Gene	Phenotype	Disease-causing variants	Reference
DAG1 NM_004393.6	WWS MEB CMD-1C CMD-1D	c.15G>A (p.Val5=); c.41C>A(p.Ser14Ter); c.235C>T (p.Arg79Ter); c.285+1G>A c.330G>A (p.Trp110Ter); c.440del(p.Gln147fs); c.454_467del (p.Phe152fs) c.556G>T (p.Glu186Ter) c.721_722del (p.Phe241fs) c.743del (p.Ala248fs) c.839del (p.Pro280fs)	[164] [165] [166] [167]
POMT2 NM_013382.7	WWS CMD-MR LGMD 2N	c.49_50delinsA (p.Arg18fs) c.248+1G>C; c.248+2T>C; c.248+5G>C c.311A>T (p.Asp104Val) c.431T>G (p.Met144Arg) c.462G>A (p.Trp154Ter) c.648C>A (p.Cys216Ter) c.673del (p.Trp225fs) c.678del (p.Trp226fs) c.737G>A (p.Gly246Asp) c.791del (p.Leu264fs) c.879_880del (p.Thr295fs) c.881A>G (p.Tyr294Cys) c.924-2A>C; c.958C>T (p.Gln320Ter) c.1006+1G>A c.1034_1035del (p.Val373Phe) c.1123_1124dup (p.Tyr376fs) c.1237C>T (p.Arg413Ter) c.1253+1G>A c.1261del (p.Arg421fs) c.1293dup (p.Met432fs) c.1417C>T (p.Arg473Ter) c.1445G>T (p.Gly3482Val) c.1555G>T (p.Glu519Ter) c.1597A>G (p.Tyr666Cys) c.1917C>A (p.Gly726Glu)	[120] [168] [169] [170] [171] [172] [173] [174] [175]

Table 3. Cont.

Gene	Phenotype	Disease-causing variants	Reference
		c.58dup (p.Val20fs) c.72del (p.Met25fs)	
		c.97C>T (p.Arg33Ter)	
		c.130G>A (p.Glu44Lys)	
		c.193G>A (p.Gly65Arg)	
		c.264G>A (p.Crp88Ter)	
		c.270_280delAATTGGAGCAG (p.Gly92fs)	
		c.280+1G>T	
		c.414del (p.Leu138_Leu139insTer)	
		c.418_420del (p.Met140del)	
		c.430A>G (p.Asn144Asp)	
		c.443C>A (p.Thr148Asn)	
		c.579_580del (p.Val195fs)	
		c.598G>C (p.Ala200Pro)	
		c.606del (p.lle203fs)	
		c.699+62del; c.699+67G>A	
		c.841C>T (p.Gln281Ter)	
		c.859_871del (p.Gly287fs)	F223
		c.978C>A (p.Tyr326Ter)	[22]
		c.990T>A (p.Tyr330Ter)	[45]
		c.1087C>T (p.Gln363Ter)	[60]
POMT1	WWS	c.1091del (p.Leu364fs)	[96] [97]
NM_001077365.2	LGMD	c.1093_1094insGGAGCACGGTGTGGAACGTGGG (p.Val365fs)	[104]
		c.1175C>T (p.Thr392Met)	[112]
		c.1175+3del	[116]
		c.1195_1196del (p.Leu399fs)	[98]
		c.1204dup (p.His402fs)	[]
		c.1272+1G>A	
		c.1361T>G (p.Leu454Ter)	
		c.1364del (p.Lys455fs)	
		c.1391G>C (p.Trp464Ser)	
		c.1417G>C (p.Gly473Arg) c.1457G>A (p.Trp486Ter)	
		c.1474C>T (p.Arg492Ter)	
		c.1474C21 (p.Nig4721e1)	
		c.1680G>C (p.Trp560Cys)	
		c.1798C>T (p.Arg600Ter)	
		c.1837_1852dup (p.Gly618fs)	
		c.1892C>T (p.Pro631Leu)	
		c.1921C>T (p.116651254)	
		c.1939G>A (p.Ala647Thr)	
		c.2097C>A (p.Tyr699Ter)	
		c.2101dup (p.Asp701fs)	
		c.2144_2147dup (p.Asp716fs)	

Table 3. Cont.

Gene	Phenotype	Disease-causing variants	Reference
POMGNT1 NM_017739.4	MEB LGMD	c.187C>T (p.Arg63Ter)     c.593del (p.Ser198fs)     c.595C>T (p.Gln199Ter)     c.643C>T (p.Arg215Ter)     c.931C>T (p.Arg311Ter)     c.932G>A (p.Arg311Gln)     c.1282C>T (p.Gln428Ter)     c.1324C>T (p.Arg442Cys)     c.1325G>A (p.Arg442His)     c.1350_1354del (p.Trp451fs)     c.1469G>A (p.Cys490Tyr)     c.1478C>G (p.Pro493Arg)     c.1694_1695del (p.Ser565fs)     c.1738C>T (p.Arg580Ter)     c.1719del (p.His573fs)     c.1814G>A (p.Arg605His)     c.1895C>G (p.Ser632Ter)     c.1895C>G (p.Ser632Ter)     c.1876del (p.Val626fs)	[47] [102] [103] [104] [105] [106] [128]
POMGNT2 NM_032806.6	LGMD	c.118C>T (p.Arg40Ter) c.410_411delinsG (p.Ala137fs) c.494T>C (p.Met165Thr) c.503T>C (p.Phe168Ser) c.509Gel (p.Asp170fs) c.590G>A (p.Trp197Ter) c.745C>T (p.Gln249Ter) c.758C>T (p.Pro253Leu) c.820_821del (p.Lys274fs) c.1000_1003del (p.Leu334fs) c.1232_1233del (p.Gln411fs) c.1333C>T (p.Arg445Ter) c.494T>C; c.758C>T	[9 <del>4</del> ] [167] [168]
B3GALNT2 NM_152490.5	WWS MEB LGMD	c.51_73dup (p.Ser25fs) c.199C>T (p.Arg67Ter) c.308_309del (p.Val103fs) c.448C>T (p.Arg150Ter) c.753del (p.Val252fs) c.755T>G (p.Val252Gly) c.824_825dup (p.lle276fs) c.875G>C (p.Arg292Pro) c.1066_1067del (p.Thr355_Asp356insTer) c.1423C>T (p.Gln475Ter)	[178] [117] [179] [180] [181] [118]

Table 3. Cont.

Gene	Phenotype	Disease-causing variants	Reference
POMK NM_032237.5	WWS LGMD	c.10C>T (p.Gln4Ter) c.43dup (p.Arg15fs) c.152del (p.Asp51fs) c.238_239del (p.Glu80fs) c.288del (p.Leu97fs) c.325C>T (p.Gln109Ter) c.386_387del (p.Leu129fs) c.410T>G (p.Leu137Arg) c.452_455dup (p.His152fs) c.907C>T (p.Arg303Ter) c.917dup (p.Leu306fs)	[67] [182] [183] [184]
Fukutin (FKTN) NM_001079802.2	WWS CMD-F	c.42del (p.Thr14_Leu15insTer)	[45] [131] [93] [134] [132] [133]

Table 3. Cont.

Gene	Phenotype	Disease-causing variants	Reference
FKRP NM_024301.5	WWS MEB CMD-1C LGMD 2I	c.77G>A (p.Trp26Ter) c.151G>T (p.Val51Phe) c.158_162dup (p.Glu55fs) c.162_165dup (p.Phe56fs) c.224del (p.Pro75fs) c.266C>T (p.Pro89Leu) c.313C>T (p.Gln105Ter) c.511_523del (p.Leu171fs) c.515dup (p.Asn172fs) c.515dup (p.Asn172fs) c.526C>T (p.Arg176Ter) c.540_570dup (p.Cys191fs) c.650dup (p.Val218fs) c.826C>A (p.Leu276lle) c.919del (p.Tyr307fs) c.919T>A (p.Tyr307Asn) c.928G>T (p.Glu310Ter) c.939G>A (p.Trp313Ter) c.948del (p.Cys317fs) c.970G>T (p.Glu324Ter) c.1075del (p.Trp359fs) c.1174C>A (p.Ser385Ter) c.1170_1171del (p.Gly391fs) c.1213G>T (p.Val405Leu) c.1256_1257del (p.Pro419fs) c.1387A>G (p.Asn463Asp) c.1394A>C (p.Tyr465Ser) c.258-2A>G	[22] [45] [69] [141] [116] [185] [127] [143] [144]
ISPD(CRPPA) NM_001101426.4	LGMD WWS	c.364G>C (p.Ala122Pro) c.466G>A (p.Asp156Asn) c.550C>T (p.Arg184Ter) c.638T>G (p.Met213Arg) c.643C>T (p.Gln215Ter) c.647C>A (p.Ala216Asp) c.704_705del (p.Glu235fs) c.773C>A (p.Ser258Ter) c.789+2T>G c.802C>T (p.Arg268Ter) c.835+2T>C c.(534+1_535-1)_(933+1_934-1)del c.1120-1G>T c.1123_1126del (p.His375fs) c.1354T>A (p.Ter452Arg)	[107] [122] [108] [110] [111] [112] [109]
TMEM5/ RXYLT1 NM_014254.3	Cobblestome Lissencephaly	c.169+2T>C c.279del (p.Gly94fs) c.429-2A>G c.649del (p.Arg217fs) c.795del (p.Arg266fs) c.1018C>T (p.Arg340Ter) c.1064_1091del (p.Asp355fs)	[154] [186] [187]
B4GAT1 NM_006876.3 NM_006876.2	WWS	c.1207G>T (p.Glu403Ter) c.1168A>G; c.1217C>T c.864T>A (p.Tyr288Ter) c.821_822insTT (p.Glu274fs)	[188] [189]

Table 3. Cont.

Gene	Phenotype	Disease-causing variants	Reference
LARGE1 NM_133642.5	WWS MEB CMD-1D	c.265C>T (p.Arg89Ter) c.283C>T (p.Arg95Ter) c.334G>T (p.Glu112Ter) c.620_621del (p.Glu207fs) c.871del (p.Gly292fs) c.992C>T (p.Ser331Phe) c.1102C>T (p.Gln368Ter) c.1209del (p.Phe404fs) c.1483T>C (p.Trp495Arg) c.1525G>A (p.Glu509Lys) c.1699del (p.Leu567fs) c.1811del (p.Leu604fs) c.1999dup (p.Cys667fs) c.2089G>T (p.Val697Leu)	[129] [148] [149] [138] [137] [130]
GMPPB NM_021971.4	LGMD	c.395C>G (p.S132C) c.64C>T (p.Pro22Ser) c.79G>C (p.D27H) c.94C>T (p.P32S) c.109C>T (p.Gln37Ter) c.220C>T (p.Arg74Ter) c.271_283del (p.Ala91fs) c.294dup (p.Glu99Ter) c.365_366dup (p.Phe123fs) c.458_459del (p.Thr153fs) c.458C>T (p.Thr153fle) c.553C>T (p.Arg185Cys) c.611_614del (p.Glu204fs) c.640+1G>A c.656T>C (p.lle219Thr) c.721C>T(p.P241S); c.728_746delinsACAGA (p.Arg243fs) c.790C>T (p.Gln264Ter) c.859C>T (p.Arg287Trp) c.1034T>C(p.V345A)	[190] [191] [192]

### **Therapeutic Considerations**

Dystroglycanopathies are very heterogeneous diseases at a clinical and genetic level in which numerous genes are involved. Phenotypic identification and genetic characterization have made it possible to advance in the development of innovative therapies, currently in the preclinical phases, which could be implemented in the coming years [81]. Gene therapy is considered a simple treatment strategy. But the expression of glycosyltransferases may also be strictly controlled, and it must consider the glycosylation status in muscle progenitor cells and changes during differentiation.

# **S**ne Therapy

Several preclinical studies have investigated the use of recombinant adeno-associated virus (AAV) to deliver functional FKRP, as well as other genes involved in glycosylation, such as LARGE1 and Beta-1,4 N-acetylgalactosaminyltransferase 2 (B4GALNT2 previously GALGT2) [144–145].

The use of CRISPR-Cas9 technology in combination with patient-specific iPS cells for the future development of autologous cell transplantation for FKRP has a big potential, because this approach uses functional  $\alpha$ -DG( $\alpha$ -DG) glycosylation in geneedited WWS iPS cell-derived myotubes [146].

The expression of B4GALNT2 was shown to have a therapeutic effect on various types of muscular dystrophy models, such as dystrophin-deficient and laminin deficient mice [147].

Overexpression of the LARGE gene increases matriglycan modification and enhances laminin-binding activity [80,119,129]. LARGE1 gene therapy has called attention as a treatment method that does not depend on the type of causative gene [43,148–149].

# armacological Therapy

Corticosteroids are anti-inflammatory drugs used as palliatives in Duchenne muscular dystrophy (DMD), as they improve muscle strength and function. Bisphosphonates prevent loss of bone density [150]. Its combined use in FKRP mouse models has been shown to achieve a decrease in muscle degeneration. The use of selective estrogen receptor modulators, tamoxifen and raloxifene, have shown to inhibit fibrosis and improve muscle strength and respiratory function [151]. Using zebrafish as a model, it has been shown that pentic acid can rescue dystrophic pathology in DMD models [144].

The effect of coenzyme nicotinamide adenine dinucleotide (NAD+) was tested in FKRP zebrafish morphants. The study reported decreased muscle degeneration and improved muscle organization and function when treatment occurred at gastrulation [152].

Many FCMD patients have a transposon insertion in the fukutin gene [127] that results in abnormal splicing of fukutin [153]. Administration of antisense nucleotides capable of correcting this splicing abnormality restores the normal function of fukutin in both fukutin KI mice and human patient-derived cells [43].

### **Ribitol Supplementation Therapy**

Defects in the pathway for the incorporation of ribitol 5-phosphate into Core 3 of  $\alpha$ -DG, which is carried out by or in the FCMD, FKRP and ISPD genes, bring together different pathologies associated with DMC [145]. Previous studies demonstrated that CDP-Rbo supplementation rescued the compromised O-glycosylation enzymatic pathway in an ISPD deficient cell line [154–155]. Ribitol supplementation [156–157], gene therapy targeting the LARGE1 gene [148], and the use of clinicalgrade induced pluripotent cells (iPSCs) [158] appear to be promising experimental models for studying the pathogenesis of α-dystroglycanopathies and for testing potential drugs or break throughs in the development of autologous therapies [155,159–160]. The application of the generation of embryoid bodies from human induced pluripotent stem cells that model the basal lamina to evaluate an experimental ribitol supplement therapy has been reported [161].

## **Cell Therapy**

Since skeletal muscle is a highly regenerative tissue, cell-based therapeutic approaches focusing on the delivery of muscle stem cells/early progenitor cells to replace diseased muscle tissue with healthy myofibers and satellite cells are highly attractive. To date, two studies have been reported on the use of cell transplantation for FKRP-associated dystroglycanopathies [162–163].

### **Conclusions**

The knowledge acquired about the dystroglycanopathies in recent years was of great importance. The elucidation of the genes involved in the formation of sugar chain structure of  $\alpha$ -DG allowed opening steps to new research and new alternative therapies.

It will be the joint effort of different fields of biology that will enable us to assume research and strategies that will allow us to implement effective therapies and precise diagnoses for these pathologies.

# eclaration of Conflicting Interests

The author declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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