

Structural Mechanism of Laforin Function in Glycogen Dephosphorylation and Lafora Disease

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SUMMARY

Glycogen is the major mammalian glucose storage cache and is critical for energy homeostasis. Glycogen synthesis in neurons must be tightly controlled due to neuronal sensitivity to perturbations in glycogen metabolism. Lafora disease (LD) is a fatal, congenital, neurodegenerative epilepsy. Mutations in the gene encoding the glycogen phosphatase laforin result in hyperphosphorylated glycogen that forms water-insoluble inclusions called Lafora bodies (LBs). LBs induce neuronal apoptosis and are the causative agent of LD. The mechanism of glycogen dephosphorylation by laforin and dysfunction in LD is unknown. We report the crystal structure of laforin bound to phosphoglucan product, revealing its unique integrated tertiary and quaternary structure. Structure-guided mutagenesis combined with biophysical and biochemical analyses reveal the basis for normal function of laforin in glycogen metabolism. Analyses of LD patient mutations define the mechanism by which subsets of mutations disrupt laforin function. These data provide fundamental insights connecting glycogen metabolism to neurodegenerative disease.

INTRODUCTION

Glycogen, the major glucose storage molecule in animals, plays an essential role in energy metabolism throughout the body. The brain is the organ most susceptible to decreases in glucose availability (Dinuzzo et al., 2014; Fryer and Brown, 2014). During the past 20 years, the perceived role of brain glycogen has shifted from an emergency energy supply to a dynamic participant in brain metabolism (Dinuzzo et al., 2014; Fryer and Brown, 2014; Swanson, 1992). While neuronal glycogen was thought to be limited to embryonic neurons, adult neurons express both

glycogen synthase and glycogen phosphorylase, and they produce low levels of glycogen (Duran et al., 2014; Lovatt et al., 2007; Pfeiffer-Guglielmi et al., 2003; Saez et al., 2014; Vilchez et al., 2007). However, glycogen synthesis in neurons must be tightly controlled because both overaccumulation and aberrant accumulation induce neuronal apoptosis (DePaoli-Roach et al., 2010; Duran et al., 2014; Turnbull et al., 2011; Valles-Ortega et al., 2011; Vilchez et al., 2007).

Glycogen is a branched polymer of glucose units joined by α -1,4-glycosidic linkages, formed by glycogen synthase, and branches occurring every 12–14 units via α -1,6-glycosidic branches, created by branching enzyme (Roach et al., 2012). Branches within glycogen are evenly distributed, resulting in a spherical structure with exposed nonreducing chain ends. This unique organization allows cells to store up to \approx 55,000 glucose units in a water-soluble form that can be rapidly released during bursts of metabolic energetics. Mutations in the *Epilepsy, Progressive Myoclonus 2A (EPM2A)* gene encoding the glycogen phosphatase laforin cause Lafora disease (LD) (Minassian et al., 1998; Serratosa et al., 1999). Laforin mutations result in hyperphosphorylated glycogen with decreased branching and increased glucose chain length that eventually forms an insoluble glucan inclusion called a Lafora body (LB) (Nitschke et al., 2013; Tagliabracci et al., 2011). Notably, it was recognized that LBs share a similar morphology with plant amylopectin, the major constituent of starch (Yokoi et al., 1968). Starch also contains branched polymers of glucose, but it is water insoluble due to its longer glucose chains of 20–25 glucose units that are branched in clusters and form helices that exclude water (Blennow and Engelsen, 2010; Gallant et al., 1997). Recent studies demonstrated that LBs possess longer glucose chains than glycogen and they contain monoester-bound phosphate, both molecular features that resemble plant starch rather than human glycogen (Nitschke et al., 2013; Tagliabracci et al., 2011). Indeed, LD is one of a family of glycogen storage diseases and offers a unique window into glycogen metabolism and associated diseases.

LD is a fatal, autosomal recessive, neurodegenerative disorder and one of the five major progressive myoclonic epilepsies (Berkovic et al., 1991; Minassian et al., 2001). LD patients typically present with an epileptic event in the second decade of life

that invariably progresses to intractable and fatal seizures within 5–10 years (Berkovic et al., 1991; Minassian et al., 2001). Disease symptoms result from neuronal apoptosis driven by the accumulation of the cytoplasmic, hyperphosphorylated, water-insoluble LB inclusions, which are normally prevented by the action of laforin (Duran et al., 2014; Turnbull et al., 2011; Valles-Ortega et al., 2011; Vilchez et al., 2007). These inclusions occur throughout the body, but neurons are acutely sensitive to polyglucosan-mediated toxicity (DePaoli-Roach et al., 2010; Duran et al., 2014; Valles-Ortega et al., 2011; Vilchez et al., 2007).

Laforin is the founding member of the glucan phosphatase family and dephosphorylates glycogen both in vitro and in vivo (Gentry et al., 2007; Tagliabracci et al., 2007, 2008; Worby et al., 2006). Laforin possesses a carbohydrate binding module (CBM) family 20 domain followed by a dual specificity phosphatase (DSP) domain (Minassian et al., 1998; Serratosa et al., 1999; Wang et al., 2002). Highlighting its basic biological importance, laforin orthologs are found in all vertebrates as well as several unicellular eukaryotes (Gentry et al., 2007; Gentry and Pace, 2009). While having a critical role in normal glycogen metabolism and aberrant LB formation, the mechanism of glycogen dephosphorylation by laforin is not known. Further, LD mutations are distributed throughout the primary sequence of laforin, leaving open the critical question of the mechanism(s) by which mutations in laforin lead to LB formation.

Plants utilize a cyclic process of reversible phosphorylation by glucan dikinases and glucan phosphatases for efficient starch degradation (Silver et al., 2014; Streb and Zeeman, 2012). Phosphorylation of starch outer glucans results in solubilization, thereby allowing degradation by starch hydrolyzing amylases, and subsequent dephosphorylation by the glucan phosphatases starch excess 4 (SEX4) and like sex four2 (LSF2). All known glucan phosphatases are members of the protein tyrosine phosphatase (PTP) superfamily within the dual specificity phosphatases (DSPs) clade (Gentry et al., 2009; Gentry et al., 2007; Tonks, 2006). The DSP domain is an ~150 amino acid domain that is <10% identical among the 65 human DSPs. The heterogeneous DSPs all utilize a cysteine residue at the base of the active site within the conserved C_xR catalytic motif to perform nucleophilic attack on the phosphorus atom of the substrate (Tonks, 2006). Differences within the DSP domain and active site allow for different members of the DSP family to specifically dephosphorylate proteinaceous substrates, glucans, lipids, or nucleic acids (Moorhead et al., 2009; Tonks, 2006).

Each glucan phosphatase possesses unique features that enable it to bind and dephosphorylate phosphorylated glucans. We recently demonstrated the molecular basis for plant glucan phosphatase function (Meekins et al., 2013, 2014; Vander Kooi et al., 2010). SEX4 possesses a DSP and CBM domain followed by a carboxy-terminal (CT) motif. In SEX4, the DSP and CBM share an extensive interdomain interface that forms a continuous binding pocket to engage a hexasaccharide. Conversely, LSF2 possesses only a DSP domain and CT motif and utilizes two secondary binding sites (SBSs) to engage glucan substrates. Laforin possesses CBM and DSP domains in the reverse orientation as SEX4, it lacks a CT motif, and it possesses a unique interdomain linker, suggesting a distinct physical basis for the specific activity of laforin versus glycogen.

Despite glycogen metabolism being a mainstay in textbooks, work on Lafora disease has yielded a number of recent breakthroughs regarding glycogen biology, including the discovery of glycogen dephosphorylation as well as putative therapies for glycogen storage diseases. Laforin is critical for regulating the phosphorylation state of glycogen, which is necessary to maintain glycogen homeostasis. Understanding the mechanism(s) underlying human disease mutations in laforin is critical for understanding the molecular basis for LD and normal glycogen metabolism. Herein, we report the structure of laforin bound to phosphoglucan product. The structure reveals the unique tertiary and quaternary architecture of laforin, providing a platform for understanding its causative role in LD. We define the basis for subsets of mutations that selectively disrupt laforin substrate engagement and specific activity. These data provide a key connection between this critical fundamental biological process and human neurodegenerative disease.

RESULTS

Crystal Structure of Laforin Bound to Maltotetraose

The structure of human laforin bound to maltotetraose and phosphate was determined using tungstate single-wavelength anomalous diffraction (SAD) phasing and refined to a resolution of 2.4 Å (Figure 1A and Table 1). There are two laforin molecules in the asymmetric unit that are superimposable, and each possesses a CBM and DSP domain. Laforin is the only known phosphatase in the human proteome that contains CBM and DSP domains, and the structure displays unique tertiary and quaternary structure. The N-terminal CBM domain is intimately associated with the DSP domain, forming an integrated tertiary structure. Furthermore, the two laforin molecules form an antiparallel dimer mediated entirely by the DSP domain (Figure 1A). Multiple maltotetraose molecules are bound to the laforin CBM and DSP domains, along with a single phosphate at the DSP active site. Maltotetraose is a soluble oligosaccharide composed of six glucose moieties with α -1,4-glycosidic linkages and serves as a model for the linear portions of glycogen.

Laforin is the only known human phosphatase that can dephosphorylate glycogen substrates (Gentry et al., 2007; Sherwood et al., 2013; Worby et al., 2006). To confirm the function of the crystallized laforin construct, we tested the ability of the catalytically active protein to both bind and dephosphorylate glycogen. Laforin readily bound to glycogen, whereas the prototypical human DSP protein phosphatase vaccinia H1-related (VHR/ DSP3) did not (Figure 1B). Additionally, laforin readily dephosphorylated glycogen (Figure 1C). This activity was specific, since mutation of the catalytic nucleophile (C266S) lacked specific activity. Additionally, VHR also lacked glycogen phosphatase activity. Thus, the crystallized laforin construct represents an active specific glycogen phosphatase.

This structure opens unique avenues for understanding laforin function in glycogen dephosphorylation and the effect of LD disease mutations. A significant number of LD mutations are localized to regions of the protein involved in physical interactions, including the DSP active site, CBM binding site, CBM-DSP interface, buried regions within individual domains, and the DSP-DSP dimer interface. Thus, the laforin structure allows us to probe the

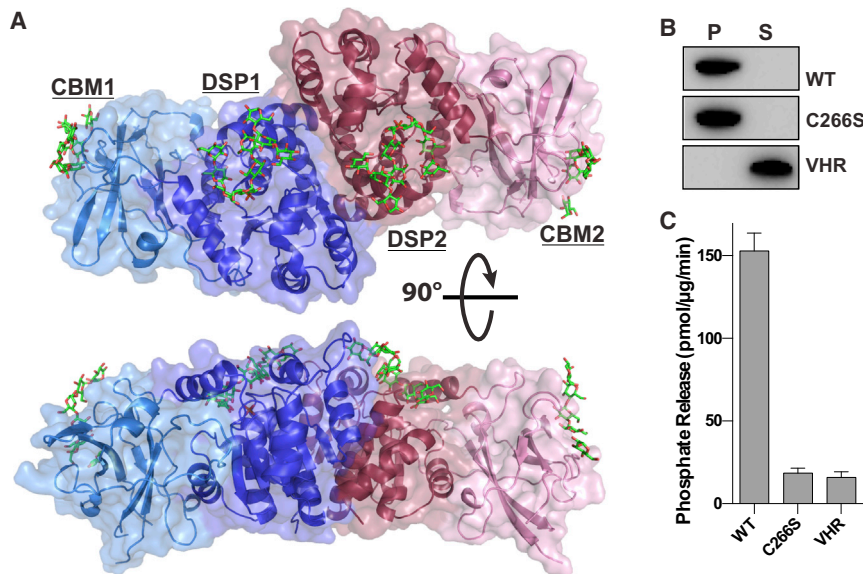


Figure 1. Crystal Structure of Human Laforin Bound to Maltotetraose and Phosphate

(A) Structure of human laforin bound to maltotetraose (green) and phosphate (orange). The laforin structure is an antiparallel dimer with one molecule in blue and the other in red. Each molecule contains a CBM and DSP domain yielding a tetramodular CBM1-DSP1-DSP2-CBM2 structure. Maltotetraose chains are bound at the CBM and DSP domains with a single phosphate molecule located at the base of the catalytic site.

(B) Glycogen cosedimentation assay to measure glycogen binding. Recombinant laforin wild-type (WT), laforin C266S, and VHR were incubated with glycogen bound to Con A agarose beads, glycogen was pelleted by centrifugation, and proteins in the pellet (P) and supernatant (S) were separated by SDS-PAGE and visualized by western analysis. Glycogen-bound proteins are in the pellet, and unbound proteins are in the supernatant.

(C) Specific activity of laforin WT, inactive mutant laforin C266S, and human VHR against glycogen. A total of 100 ng of protein was incubated with 45 µg glycogen for 20 min. Each bar is the mean ± SEM of six replicates; $p < 0.05$.

mechanism(s) by which LD patient mutations affect laforin function.

Maltotetraose and Phosphate Bound at the Laforin Active Site

A bound maltotetraose chain and phosphate are tightly integrated into the laforin DSP domain active site, with a contact area of 655 Å². The active site is broad, with a cross section ~17 Å wide and 10 Å deep, and is formed by residues from different portions of the DSP domain whose variability among DSP family members yields unique substrate specificities (Figure 2A). The laforin DSP domain is structurally most similar to the plant glucan phosphatases LSF2 and SEX4, with a root-mean-square deviation (rmsd) of 2.0 and 2.1 Å (Figure S1, available online), respectively. The electron density of maltotetraose in the laforin active site allows unambiguous assignment of glucan chain orientation, labeled Glc1–Glc6 from the nonreducing to reducing end (Figure 2B). Seven conserved laforin residues make extensive interactions with the maltotetraose glucose moieties (Figure 2B). M236 fully engages the concave surface of the maltotetraose chain. T142, Y146, T194, D197, R241, N267, and Y304 interact with the convex surface, which involves many hydrogen bonds as well as van der Waals interactions. Each of these interacting residues is strictly conserved in laforin vertebrate orthologs (Figure S2A). At the base of the active site is the PTP-loop C_X5R motif (HCNAGVGR, residues 265–272). The cysteine within this motif sits at the base of the catalytic cleft and functions as a nucleophile during catalysis. The O3 and O2 of Glc3 are positioned 2.7 Å and 3.8 Å from the phosphate, respectively, with the O6 at a distance of 7.7 Å pointed toward solvent (Figure S3).

To investigate the role of these residues in laforin function, we generated alanine mutations of M236, T142, and D197. Additionally, we tested multiple LD mutations in the DSP domain that are localized to the DSP active site, including R171H and G240S (Figure 2B). We first tested the effect of these mutations on la-

forin stability and folding by measuring protein stability using differential scanning fluorimetry (DSF). None of the mutations showed a significant decrease in stability (Table S1). After

Table 1. Data Collection and Refinement Statistics

Data Collection	Native	Tungstate
Beamline	APS 22-ID	APS 22-ID
Wavelength	1.0000	1.2148
Space group	P4 ₃ 2 ₁ 2	P4 ₃ 2 ₁ 2
Unit cell parameters (a, b, c)	78.71, 78.71, 321.12	78.68, 78.68, 321.10
Unique reflections	37,585	38,389
Completeness (%)	92.3 (92.0)	99.8 (100.0)
Resolution (Å)	20.0–2.4 (2.49–2.40)	20.0–3.0 (3.11–3.00)
R _{merge} (%)	9.7 (51.6)	10.5 (26.4)
Redundancy	3.6 (2.8)	3.8 (3.6)
1/σ(I)	11.6 (2.5)	12.3 (5.7)
Refinement		
Resolution limits (Å)	20.00–2.40	
No. reflections/no. to compute R _{free}	33,896/3,582	
R/R _{free}	16.6/21.1	
No. protein residues	624	
No. glucose/phosphate	30/2	
No. of water	143	
Rmsd bond (Å)	0.008	
Rmsd angle (°)	1.54	
Ramachandran favored/outlier (%)	98.24/0.0	
MolProbity score	0.92 (100 th percentile)	

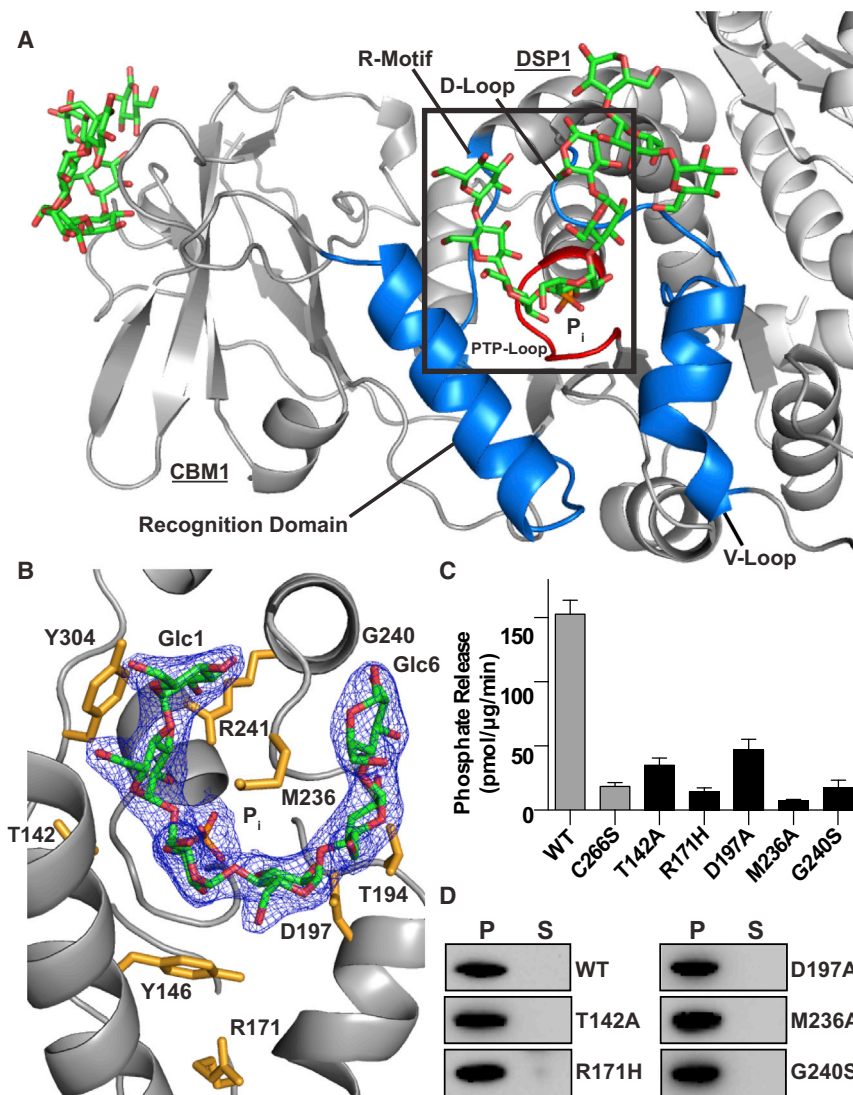


Figure 2. Maltohexaose Bound at the Laforin DSP Domain Active Site

(A) Ribbon diagram of one laforin molecule with maltotetraose (green), phosphate (orange), backbone (gray), DSP domain motifs (blue), active site loop (red), and the second molecule (light gray). (B) Inset from (A), showing a 2Fo-Fc electron density map (1.3 σ) with maltotetraose (green) and phosphate (orange). Glucose moieties (Glc) are numbered from nonreducing to reducing end. Side chains of interacting residues are colored yellow. See also Figure S3. (C) Specific activity of laforin WT and C266S (from Figure 1) compared to DSP active site mutants. Each bar is the mean \pm SEM of six replicates; $p < 0.05$. (D) Glycogen cosedimentation assay of WT and DSP active site mutants.

Laforin CBM-Glucan Interactions

A separate maltotetraose chain is bound to the canonical carbohydrate binding site in the laforin CBM domain (Figure 3A). Aromatic and hydrophilic residues within the CBM engage the maltotetraose, with a contact area of 327 \AA^2 , via a network of stacking interactions and hydrophilic residues including the side chains of W32, K87, W99, and D107 and backbone of G103. The tryptophan residues interact with one face of the glucose units. W32, K87, and W99 are strictly conserved, and D107 is highly conserved in all laforin orthologs (Figure S2B).

To define the contribution of the CBM to laforin activity, we mutated K87 and W99 to alanine, as well as assessed the LD patient mutation W32G. W32G showed reduced stability with the T_m decreased by

5.1 $^\circ\text{C}$, indicating that it contributes to protein stability (Table S1). Indeed, the structure identifies a number of key core hydrophobic residues including F5, Y86, and F88 that are mutated in LD and likely destabilize the protein. In contrast, we found the other mutant proteins had stability equivalent to that of WT protein. To determine how these mutations impact laforin binding to glycogen, we employed the Con A-glycogen experiment. We found that W32G was the most deleterious, being found entirely in the supernatant fraction. K87A and W99A also exhibited decreased glycogen binding by $\sim 40\%$ (Figure 3B). The CBM mutations resulted in a 57%–96% decrease in glycogen phosphatase activity (Figure 3C). Thus, LD and CBM mutations that affect CBM domain stability and/or direct glucan engagement lead to loss of enzymatic activity. These data indicate the importance of glycogen engagement by the CBM domain for laforin activity.

establishing that the laforin point mutants did not destabilize the protein, we tested the mutants' phosphatase activity against glycogen. In all cases, we found a significant decrease in specific activity, ranging from a 69% to 95% decrease compared to wild-type (WT) (Figure 2C). M236A was the most deleterious to laforin activity, consistent with its observed extended interaction surface with the bound carbohydrate. Since the laforin DSP independently engages a glucan chain, a loss of glucan phosphatase activity may indicate a decrease in glucan binding. To assess binding to glycogen, we incubated laforin with glycogen coupled to Concanavalin A (Con A) Agarose (GE Healthcare), washed the agarose beads, performed a low-speed spin, and used western analysis to determine if laforin was bound to glycogen in the pellet or remained in the supernatant. We found that the laforin DSP mutants pelleted with glycogen similarly to WT (Figure 2D). Thus, mutations in the DSP domain active site channel do not dramatically decrease glycogen binding but specifically impact the ability of laforin to dephosphorylate glycogen.

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Laforin Architecture and Dynamics

While each domain is clearly important for laforin function, the structure reveals a unique tertiary architecture with tightly

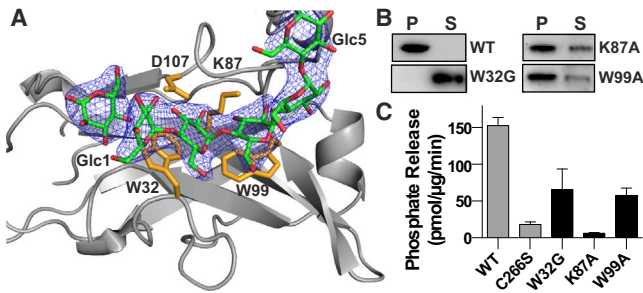


Figure 3. Maltose Bound at the Laforin CBM Domain

(A) Close up of the laforin CBM domain with bound maltose (green) 2Fo-Fc electron density map (1.0 σ) with CBM residues (yellow). Glucose moieties (Glc) are numbered from nonreducing to reducing end.

(B) Glycogen cosedimentation assay of WT and CBM domain mutants.

(C) Specific activity of laforin WT and C266S (from Figure 1) compared to CBM domain mutants. Each bar is the mean \pm SEM of six replicates; $p < 0.05$.

integrated CBM and DSP domains, with a contact area of 1,154 \AA^2 . The relative orientation of the CBM and DSP domains is dramatically different than that observed in the plant glucan phosphatase SEX4 (Figure S1). Most strikingly, the two carbohydrate binding sites are not proximal, as was seen in SEX4 (Figure S1). Thus, we next probed the relationship and coupling between the protein domains.

We employed deuterium exchange mass spectrometry (DXMS) to elucidate the conformational dynamics of laforin in solution and the effect of binding to its physiological substrate glycogen in terms of changes in tertiary and quaternary structure. Optimization of the assay allowed 100% coverage of C-terminally His₆-tagged laforin WT and allowed us to assess the change in peptide mass with increasing time (Figure S4A). Strong differences in protection were observed for different regions of the protein (Figures 4A and 4B). Among the most protected regions, defined as <20% deuteration from 10 s to 1,000 s, are residues 122–129 in the CBM domain and 290–296 in the DSP domain. These are residues in the complementary surfaces of the CBM-DSP domain interface, indicating that the observed tertiary structure is stable in solution and suggesting that there is minimal structural rearrangement of this region. Additionally, residues 218–227 and 252–256, both located at the dimer interface in the DSP domain, exhibit <20% deuteration throughout the experiment, consistent with stable dimer association in solution.

Regions that are highly accessible to solvent, defined as greater than or equal to 50% deuterated after 300 s in D₂O, are primarily within surface-exposed regions (Figure 4B). The region that is the most highly solvent accessible, exhibiting >90% deuteration at all time points, is residues 66–84 of the CBM domain, which corresponds to the disordered loop not modeled in the crystal structure (Figure 4A). The CBM-DSP domain linker and DSP recognition motif also displayed >70% deuteration at 300 s (Figures 4A and 4B), yet other peptides did not display high levels of deuteration, suggestive of local dynamics, but not global conformational changes.

Next, we utilized DXMS to evaluate structural changes in laforin in the presence of its endogenous substrate glycogen. In

the presence of glycogen, we observed significant substrate-induced decreases in the exchange rate within both the CBM and DSP domain (Figures 4C and S4B). Three peptides within the CBM exhibited 13%–27% decreases of deuteration in the presence of glycogen: residues 21–52, residues 60–65, and residues 98–111. The identified oligosaccharide-binding residues W32, W99, and D107 lay within these regions (Figure 4A). Likewise, regions within the DSP domain surrounding the active site also showed significant protection, decreases of 13%–45%, including within the recognition domain (residues 130–145, 146–155), V loop (residues 193–217), D loop (residues 228–236), PTP loop (267–281), and R motif (308–317). Changes were restricted to the surface of the laforin structure encompassing the active site cleft, with little to no change on the opposite face (Figure 4D). Peptides involved in the CBM-DSP domain interface and the dimerization interface did not exhibit significant changes in the presence of glycogen. Indeed, there were no significant increases in deuteration in any region of laforin, suggesting that there is not an increase in protein dynamics associated with substrate binding. Intriguingly, the CBM- and DSP-bound maltose chains are positioned in a parallel orientation, i.e., the nonreducing and reducing ends are oriented in the same direction. When residue conservation is mapped onto the surface of the structure, very high levels of conservation are observed for the active site and intervening region to the CBM binding site (Figure 4E). These regions closely resemble the less-deuterated regions in the presence of glycogen, suggestive of a physical path for cooperative binding of glycogen.

The CBM and DSP Domain Interaction Is Critical for Function and Impacts Lafora Disease

Given the extensive interdomain interface, stable dynamics, and conservation, we hypothesized that the observed CBM/DSP orientation exhibited in the laforin tertiary structure is critical for its function (Figure 5A). Indeed, many of the residues that participate in interdomain interactions are mutated in LD, including I126T, Q293L, Y294N, and P301L. We generated an alanine mutation of V8 as well as analyzed the LD mutants I126T, Y294N, and P301L, all of which are located at the interdomain interface. P301L displayed a decreased T_m , while all other mutant proteins were stable and well folded with no significant difference in T_m compared to WT protein (Table S1). We investigated the glycogen binding of these mutations using the Con A-glycogen assay and found that each mutant pellets with glycogen, indicating that glycogen binding is not dramatically affected (Figure 5B). Next, we tested the glycogen phosphatase activity of each mutant and found that V8A, I126T, Y294N, and P301L significantly decreased the glycogen phosphatase activity of laforin by 47%–93% (Figure 5C). Thus, mutations within the CBM-DSP interface significantly decrease glycogen phosphatase activity without diminishing glycogen binding.

To further explore the structural dynamics of the CBM-DSP interface, we performed DXMS analysis of the LD patient mutation laforin Y294N. Strikingly, the Y294N mutation significantly increased the solvent accessibility of the CBM/DSP interface (Figures 5D, 5E, and S4C). Specifically, residues 290–296 and 297–307 were 95% and 45% more deuterated in laforin Y294N than in WT, respectively. These residues surround Y294 and

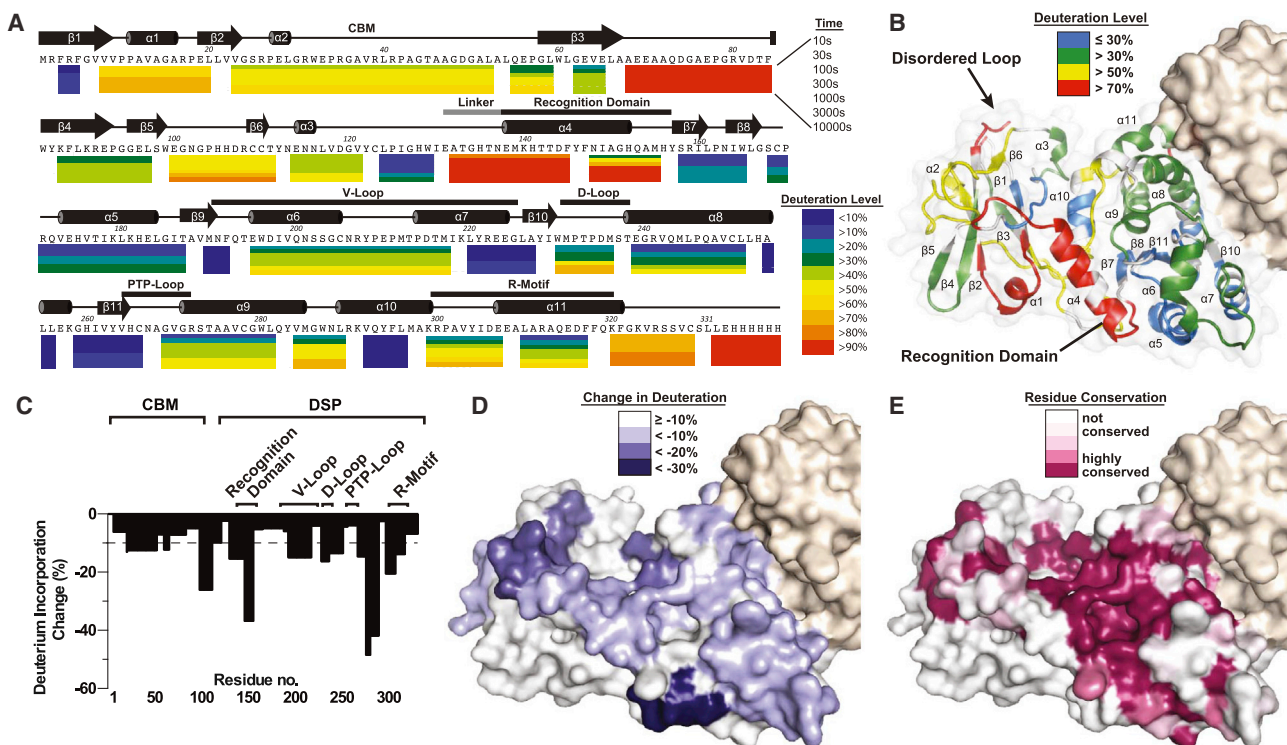


Figure 4. Deuterium Exchange Mass Spectrometry Analysis of Laforin Architectural Dynamics

(A) Laforin dynamics from DXMS expressed as percent deuteriation of laforin peptides with the lowest deuteriation in blue to highest in red, as indicated. Each colored bar below the primary sequence represents percent deuteriation at one of seven time points. Secondary structure elements are depicted above the primary sequence with the CBM domain and DSP motifs labeled.

(B) Surface mapping of the percent of deuterium exchange of each peptide at 300 s was mapped onto the laforin structure. Residues 69–79 were 100% deuteriated throughout the DXMS experiment, and these residues were disordered and not modeled in the structure (labeled disordered loop).

(C) Maximal percent change in deuteriation of laforin ± glycogen versus the laforin primary sequence. The dotted line designates 10% decrease. See also Figure S4B.

(D) Surface mapping depicting the regions that exhibit decreased deuteriation in the presence of glycogen.

(E) Surface model of laforin illustrating amino acid conservation using the sequences from Figures S2A and S2B.

include the DSP domain residues that comprise the CBM-DSP interface (Figure 5E). Further, CBM domain residues comprising the opposite side of the interface also exhibit increased deuteriation compared to WT with residues 113–121 and 122–129 increasing 50% and 39%, respectively. While the CBM/DSP interface exhibited increased deuteriation, there was no significant change in deuteriation of peptides comprising the dimer interface. These results indicate that the unique tertiary structure with specific CBM/DSP orientation is required for laforin activity and provides an explanation for the role that mutations of these interfacial residues play in LD.

Laforin Dimerization

The physical and functional role of laforin oligomerization has been controversial (Dukhande et al., 2011; Koksai and Cingolani, 2011; Liu et al., 2006, 2009; Sánchez-Martín et al., 2013). In situ and in vivo experiments have demonstrated that laforin is able to both dimerize and multimerize, but different models have been proposed for the extent to which oligomers form and which domains contribute. The structure reveals that laforin is an anti-parallel dimer mediated exclusively by the DSP domains, with an

extended 906 Å² interface and high shape complementarity ($Sc = 0.81$) (Figure 1A).

To examine the structure of laforin in solution, we employed inline size exclusion chromatography-small angle X-ray scattering (SEC-SAXS). Laforin eluted from SEC as a single species (Figure S5A) with molecular size and shape parameters $R_g = 33.9 \pm 0.2$ Å and $D_{max} = 119$ Å. R_g was invariant across a 25-fold concentration range as the protein eluted from the column (33.7 ± 0.7 Å for the entire range), indicating that the protein was not exchanging between oligomeric states. The experimental scattering curve demonstrated strong agreement with the theoretical profile calculated for the laforin dimer (predicted $R_g = 31.2$ Å and $D_{max} = 121$ Å) observed in the crystal structure ($\chi^2 = 1.1$) (Figures 6A, S5B, and S5C). In contrast, the theoretical scattering profile for a single copy of laforin monomer (predicted $R_g = 21.2$ Å and $D_{max} = 78$ Å) showed a marked deviation from the experimental data ($\chi^2 = 27.0$). An ab initio reconstruction of the average molecular envelope revealed an extended bilobal shape that closely matched the crystal structure of laforin (Figure 6B). The excluded volume indicated that the molecular weight of the protein was 72 kDa, matching

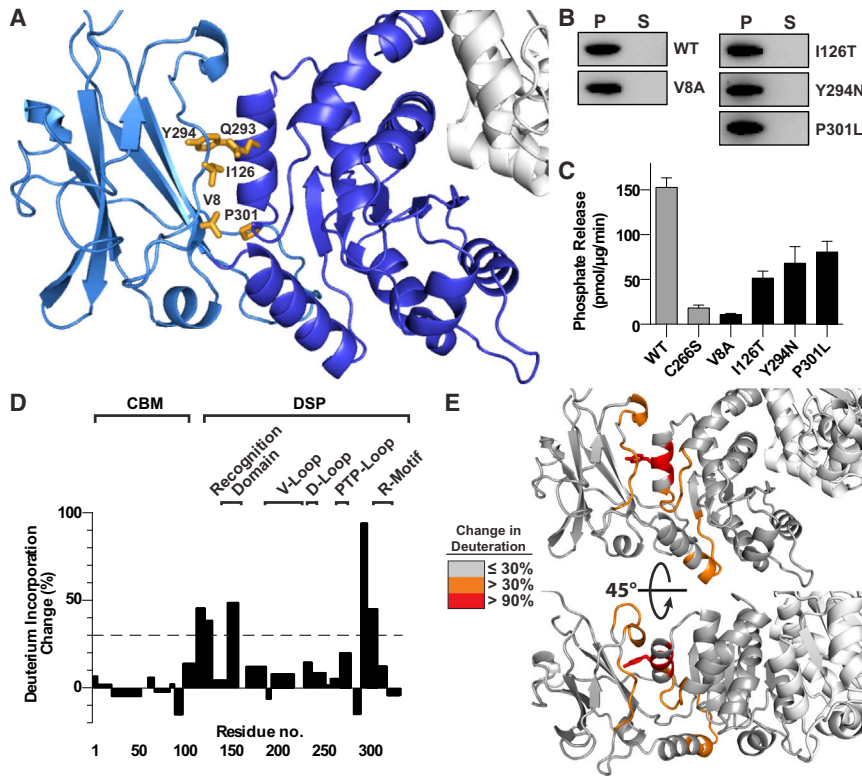


Figure 5. The Laforin CBM-DSP Interface Is Necessary for Laforin Function

(A) Close up of the CBM (light blue) DSP (dark blue) domain interface. Residues that comprise the CBM-DSP interface are yellow.

(B) Glycogen cosedimentation assay of WT and CBM-DSP interface mutants.

(C) Specific activity of laforin WT and C266S (from Figure 1) compared to CBM-DSP interface mutants. Each bar is the mean \pm SEM of six replicates; $p < 0.05$.

(D) Structural dynamics of the laforin mutant Y294N with a bar graph depicts maximal percent change in deuteration between WT and Y294N; positive values depict increased deuteration in Y294N, and negative values depict decreases. The dotted line designates 30% change in deuteration.

(E) Surface mapping of regions that exhibit changes in deuteration between WT and Y294N.

the theoretical molecular weight of the laforin dimer (74 kDa) and further confirming that laforin forms a stable dimer in solution. Thus, the solution scattering data demonstrate that the tertiary and quaternary structure of laforin observed in the crystal structure represent the native conformation of the protein in solution.

Closer examination of the dimerization interface demonstrates that it contains a core hydrophobic region comprised of I229, M231, L251, C250, and F321 and outer hydrophilic regions containing D215, R222, E239, Q243, and K320 (Figure 6C). In particular, F321 is buried at the interface and is mutated in the LD patient mutation F321S.

To determine the effect of mutations at the laforin dimeric interface on function, we next tested the effect of F321S or L251A mutations. Both mutations significantly affected laforin stability, resulting in $>5^{\circ}\text{C}$ decreases in protein stability (Table S1). Mutation at the dimer interface also significantly decreased glycogen phosphatase activity, with observed activity decreases of 56%–62%, but without global effects on glycogen binding (Figures 6D and 6E).

Laforin Substrate Binding Cooperativity

To examine the oligomeric state of the F321S LD mutation, we employed analytical ultracentrifugation (AU). Consistent with SEC and SAXS solution measurements, WT laforin exists as a stable dimer in solution with a single peak observed at 79 ± 6 kDa (Figures 7A and S6A). Strikingly, the F321S mutation disrupts dimerization, existing as a single species of 41 ± 6 kDa, consistent with the monomeric molecular weight of laforin (Figures 7B and S6B).

presence of malto-oligosaccharides from maltose of degree of polymerization 2 (DP2) through DP24. We saw little effect with DP2–DP4 oligosaccharides and a minimal length-dependent increase in T_m for DP5–DP8, with each additional glucose moiety yielding an $\sim 0.3^{\circ}\text{C}$ ΔT_m (Figure 7C). Strikingly, there was a dramatic increase in stabilization when using longer oligosaccharides, with a 24-mer yielding a ΔT_m increase of $\sim 14^{\circ}\text{C}$. Further, when the concentration dependence was measured, a shorter oligosaccharide showed a broad transition with two binding events, whereas the longer oligosaccharide showed a single high-affinity binding with a Hill coefficient of 1.41, demonstrating striking cooperativity in binding (Figure 7D). We next assessed the effect of laforin mutant protein for cooperative binding. F321S exhibited a significantly decreased degree of stabilization by glucan substrate and, notably, loss of cooperativity, with a Hill coefficient of 0.95 (Figure 7E). Overall, these data provide a mechanism for laforin dimerization in its function and provide a basis for understanding the effect of LD mutations that disrupt dimerization.

DISCUSSION

The role of glycogen dephosphorylation is an emerging area of research that directly impacts our understanding of normal glycogen metabolism as well as dysfunction in the fatal human epilepsy Lafora disease. The data presented here provide fundamental insights into how laforin functions in glycogen dephosphorylation and uncover features of the laforin protein that provide a basis to understand disease mechanisms. The domain orientation in laforin is unique and distinct from the other known

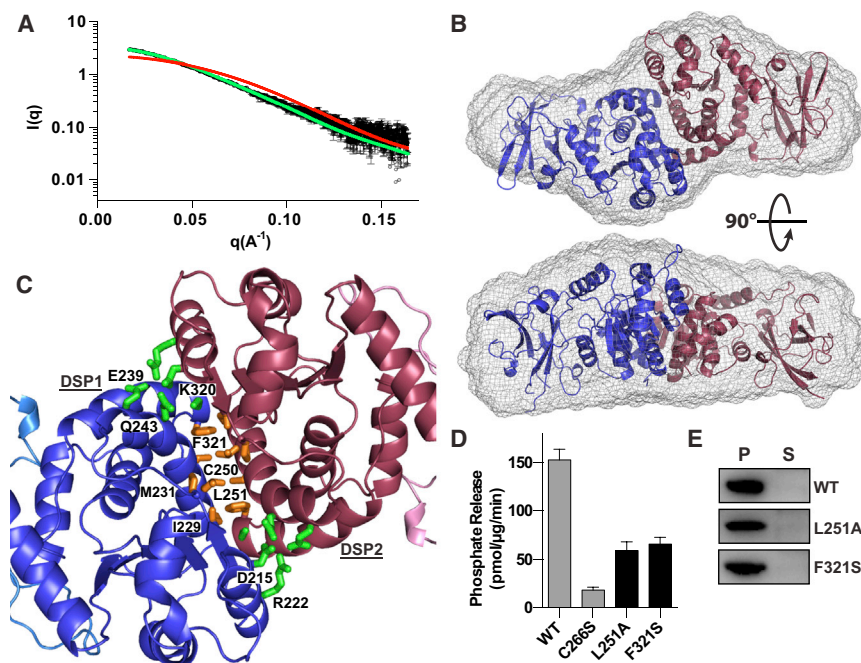


Figure 6. Laforin Quaternary Structure Is Critical for Its Function

(A) Scattering curve of laforin C266S (black) with curves computed from the crystallographic structure for monomer (red) and dimer (green).

(B) Ab initio model of laforin C266S with the computed surface (gray mesh) overlaid with the crystal structure.

(C) The laforin DSP domain dimer interface with interfacial hydrophilic (green) and hydrophobic interacting residues (gold). Residues from DSP1 are labeled.

(D) Specific activity of WT and C266S (from Figure 1) compared to dimer interface mutants. Each bar is the mean \pm SEM of six replicates; $p < 0.05$.

(E) Glycogen cosedimentation assay of WT and dimer interface mutants.

glucan phosphatases with the carbohydrate binding regions of the CBM and DSP domains distant from each other. In contrast, SEX4 employs a continuous binding pocket that spans both the CBM binding site and DSP active site, allowing it to bind a single maltohexaose chain (Meekins et al., 2014). The laforin dimer produces a unique tetramodular architecture with CBM-DSP-DSP-CBM domains aligned and engaging multiple maltohexaose chains. Structure-guided and patient mutations are described that result in loss of functional interactions in the CBM binding site, DSP active site, CBM/DSP interface, and dimerization interface. Our data demonstrate that this unique tertiary and quaternary structure allows for engagement of glycogen by the CBM domain, cooperative substrate binding by the dimer, engagement of the phosphorylated glucan at the DSP active site, and finally catalysis.

This model has direct implications for our understanding of the source and position-specific function of glycogen phosphorylation on both glycogen superstructure and LB formation, which is currently being explored. Some propose that phosphate is introduced by a glycogen synthase side reaction, while others suggest specific incorporation by a yet-unidentified human glycogen kinase (Nitschke et al., 2013; Tagliabracci et al., 2011). In plants, regulated phosphorylation of starch is better characterized. Glucan, water dikinase (GWD) specifically phosphorylates the C6 position that targets phosphoglucan, water dikinase (PWD) to phosphorylate C3 hydroxyls. These phosphorylation events induce steric strain and disrupt the helical structure of starch and allow hydrolytic enzymes to processively degrade starch (Blennow and Engelsen, 2010; Silver et al., 2014; Streb and Zeeman, 2012). Complementary to the dikinases, the plant glucan phosphatases possess distinct preference for dephosphorylating glucose positions. SEX4 preferentially dephosphorylates the C6 position, while LSF2 exclusively

and C6-hydroxyls for SEX4 (Meekins et al., 2013, 2014). Glycogen can be phosphorylated at the C2, C3, and C6 positions (Nitschke et al., 2013). The laforin structure shows preferential binding of the glucan with C3/C2 oriented toward the catalytic center, suggesting that it will preferentially dephosphorylate the C3 or C2 position (Figure S3). Indeed, the C6 hydroxyls are pointed away from the laforin active site into the solvent. As with SEX4, laforin will likely be able to act on all positions to some extent but would require the opposite glucan orientation to accommodate dephosphorylation of the C6 position and thus is expected to have significantly reduced activity level for the C6 position.

Currently, the position of phosphorylation with respect to α -1,6 branch points is unresolved. It has been reported that C6 monophosphates may be selectively elevated in LBs and predominate near the center of glycogen molecules, thereby affecting branching (Nitschke et al., 2013). Intriguingly, an additional portion of an oligosaccharide chain is observed in the crystal structure of laforin near the DSP active site, and a single-ordered glucose moiety is observed near the CBM binding site. This is likely a consequence of the high concentration of carbohydrate used in crystallization, but it may suggest the possibility of preferential binding to specific glycogen regions, particularly branch points. The laforin structure demonstrates that a shallow and wide binding pocket (10 Å \times 17 Å) is a conserved feature of glucan phosphatases from plants to animals and an architectural feature that would accommodate a branched glycogen chain. The ability of laforin to bind and localize to specific subregions within glycogen is an area of ongoing research that will require temporal and quantitative comparisons of wild-type and mutant proteins in the context of physiological substrates. These results will further define the role of phosphorylation/dephosphorylation in LB formation and disease progression.

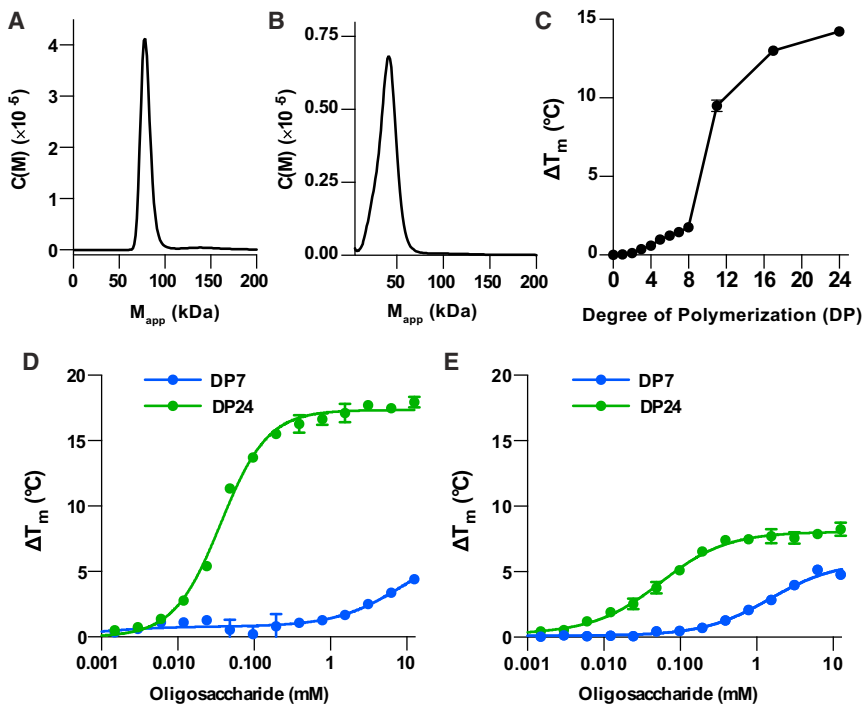


Figure 7. Cooperative Binding to Glucan Substrates

(A) Sedimentation velocity experiments reveal that laforin WT exists as a dimer in solution. (B) Sedimentation velocity experiments reveal that laforin F321S exists primarily as a monomer in solution. (C) Thermal shift assay with 1 mM oligosaccharides reveals that laforin C266S is more stabilized in the presence of an increasing degree of polymerization (DP). (D) Concentration-dependent increase in laforin C266S melting temperature (ΔT_m) with maltoheptaose (DP7) and maltodextrins (DP24). (E) Concentration-dependent increase in laforin F321S melting temperature (ΔT_m) with maltoheptaose (DP7) and maltodextrins (DP24).

The physical and physiological relevance of laforin dimerization has been a point of debate in the field (Dukhande et al., 2011; Liu et al., 2006, 2009; Sánchez-Martín et al., 2013). The crystal structure, along with solution-based measurements using complementary techniques, establishes that laforin exists as an antiparallel dimer mediated by the DSP domain. Notably, laforin does not possess a CT motif, which is required for stability and function in plant glucan phosphatases. Instead, the laforin dimer interface partially overlaps with the region where the CT motif binds to the DSP domain (Figure S1). Furthermore, we demonstrate that the LD mutation F321S disrupts laforin dimerization and diminishes glycogen phosphatase activity and cooperative substrate binding. Dimerization is a common theme regulating the activity of transmembrane receptor-like PTPs (RPTPs). A helix-turn-helix wedge motif of one RPTP obstructs the active site of the dimer partner, possibly allowing a ligand-dependent manner to modulate phosphatase activity (Tonks, 2006, 2013). Conversely, dimerization of cytoplasmic PTPs is much less common, and DSPs are typically regarded as monomeric enzymes. The human phosphoinositide lipid phosphatase myotubularin related protein 2 (MTMR2) and the *Vaccinia* virus gene H1 (VH1) are both DSPs that dimerize. The laforin DSP domain forms its dimer interface, while both MTMR2 and VH1 form dimers via ancillary domains. MTMR2 forms a dimer via its C-terminal coiled-coil domain, while the VH1 dimer interface is formed by a swapped N-terminal α helix (Berger et al., 2003; Koksai and Cingolani, 2011; Phan et al., 2007).

LD offers a unique window into both normal cellular glycogen metabolism and disease implications when the process is perturbed. Recently, a study using a novel glycogen detection method demonstrated that neurons contain low levels of glycogen, and this glycogen protects neurons from hypoxic stress in vitro and in vivo (Saez et al., 2014). Yet, glycogen

based aggregates that form in brain, eye, and peripheral nervous system neurons of individuals 30 years old and greater (Cavanagh, 1999; Sinadinos et al., 2014). The function of corpora amylacea, as well as their impact on cellular homeostasis and involvement of laforin, is unknown, yet recent evidence suggests that corpora amylacea may constitute an important factor in age-related neurological decline (Sinadinos et al., 2014).

In addition to impacting neuronal processes, perturbations in glycogen metabolism are a feature of glycogen storage diseases, diabetes, cardiac pre-excitation syndromes, and cancer (Light, 2006; Roach, 2002; Roach et al., 2012; Ros and Schulze, 2012). Given the central importance of glycogen to cellular homeostasis, glycogen (de)phosphorylation may impact other aspects of human disease that are yet unexplored. The role of regulated carbohydrate phosphorylation in diverse biological systems by glucan phosphatases, glucan dikinases, and yet to be discovered enzymes opens unique avenues for understanding unexplored aspects of glycogen metabolism as well as human diseases.

EXPERIMENTAL PROCEDURES

Please refer to Supplemental Experimental Procedures for additional experimental details.

Cloning, Protein Expression, and Protein Purification

H. sapiens (Hs) laforin residues 1–328 were expressed from pET28b (Novagen) as an N-terminal His₆ tag fusion. Wild-type and mutant proteins were expressed in BL21-CodonPlus *E. coli* cells and purified using IMAC and SEC.

Crystal Structure Determination

High-quality single native and tungstate cocrystals were obtained and data collected on the 22-ID beamline of SER-CAT at the Advanced Photon Source, Argonne National Laboratory to 2.4 Å and 3.0 Å, respectively (Table 1).

Phosphatase Assays

We utilized a phosphatase assay that employs malachite green to quantify inorganic phosphate release from rabbit muscle glycogen; assays were performed six times or more (Sherwood et al., 2013; Tagliabracci et al., 2007, 2008).

Glycogen Binding Assays

Glycogen binding assays were performed using Concanavalin A (Con A) Agarose beads (Sigma) as described in the [Supplemental Experimental Procedures](#).

Hydrogen Deuterium Exchange Mass Spectrometry

DXMS experiments were performed as previously described (Hsu et al., 2009; Pantazatos et al., 2004). A total of 134 high-quality peptides with 100% coverage of laforin were obtained. Exchange of laforin with and without glycogen was performed three independent times, with all samples prepared in triplicate.

Size Exclusion Chromatography Small-Angle X-Ray Scattering

SEC-SAXS experiments were performed at the 18-ID beamline of BioCAT, Advanced Photon Source, Argonne National Laboratory using an inline Superdex 75 10/300 column (GE). Guinier analysis demonstrated that the system is monodisperse with particles that have a radius of gyration (R_g) = 33.9 ± 0.2 Å (Figure S5B). Pair-distance distribution function, $P(r)$, demonstrated a well-defined D_{max} = 119 Å and real space R_g (35.0 Å) consistent with Guinier analysis (Figure S5C).

Analytical Ultracentrifugation

Sedimentation velocity experiments were conducted using a Beckman XL-I analytical ultracentrifuge (Beckman Coulter). Protein was concentrated to 5 μM in size exclusion buffer.

Differential Scanning Fluorimetry

Thermal stability experiments were carried out using a CFX96 Real-Time PCR system (BioRad). A total of 2 μM of protein was combined with 5X SYPRO Orange Protein Gel Stain (Invitrogen) with or without maltoligosaccharides, and stability was assessed from 20°C to 90°C at 1°C/50 s in triplicate.

ACCESSION NUMBERS

The atomic coordinates and structure factors are available in the Protein Data Bank under ID code 4RKK.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, and one table and can be found with this article online at <http://dx.doi.org/10.1016/j.molcel.2014.11.020>.

AUTHOR CONTRIBUTIONS

M.S.G. and C.W.V.K. conceived the experiments, analyzed data, and wrote the paper. C.W.V.K. collected X-ray diffraction data and determined the crystal structure. M.R. conceived experiments, purified protein, generated and optimized crystals, generated mutants, performed DSF and phosphatase assays, analyzed data, and generated figures. M.K.B. generated mutants, performed assays, analyzed DXMS data, and generated figures. B.K.W. and H.S. performed DXMS experiments under the supervision of S.L. and V.L.W. A.R.S. generated constructs, purified proteins, performed assays, analyzed DXMS data, and generated figures. M.W.P., S.C., and C.W.V.K. performed SAXS experiments and analysis. L.M.H. performed and analyzed AUC data. S.H. generated mutants and performed binding experiments. P.S. provided constructs and analyzed data. T.M.B., B.C.P., K.D.A., D.A.M., B.D.T., V.V.D., and A.O.T. generated constructs, purified proteins, performed crystallization screens, and performed assays.

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